

Reporting Limit Outliers

Lab Reporting Batch ID: PH111

Laboratory: LL

EDD Filename: PH111

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-576-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	3.44	4.91	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.748	4.91	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.104	4.91	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0940	4.91	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.156	4.91	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.377	4.91	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.149	4.91	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.397	4.91	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.294	4.91	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.0919	4.91	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.366	4.91	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.160	4.91	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.475	4.91	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.278	0.982	PQL	ng/Kg	
OCDF	JB	1.34	9.82	PQL	ng/Kg		
SL-577-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.148	4.98	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.0872	4.98	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0451	4.98	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0610	4.98	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.0751	4.98	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.0773	4.98	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.0551	4.98	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JBQ	0.0378	4.98	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.0745	4.98	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.0627	4.98	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.107	4.98	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.0553	4.98	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0442	4.98	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.0714	0.996	PQL	ng/Kg	
OCDD	JB	0.620	9.96	PQL	ng/Kg		
OCDF	JBQ	0.181	9.96	PQL	ng/Kg		
SL-578-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	2.40	5.08	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.597	5.08	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0881	5.08	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0716	5.08	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.113	5.08	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.320	5.08	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.143	5.08	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.348	5.08	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.504	5.08	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.0832	5.08	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.234	5.08	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.150	5.08	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.364	5.08	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.175	1.02	PQL	ng/Kg	
OCDF	JB	1.32	10.2	PQL	ng/Kg		

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eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-579-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	1.36	4.98	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.422	4.98	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0917	4.98	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0742	4.98	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.125	4.98	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.316	4.98	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.121	4.98	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.390	4.98	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.506	4.98	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.226	4.98	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.139	4.98	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.235	4.98	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.113	0.996	PQL	ng/Kg	
	OCDF	JB	0.803	9.96	PQL	ng/Kg	
SL-879-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	1.05	4.90	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JBQ	0.170	4.90	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.121	4.90	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.212	4.90	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.697	4.90	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.220	4.90	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.882	4.90	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.745	4.90	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.166	4.90	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.588	4.90	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.208	4.90	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.664	4.90	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.0995	0.980	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.307	0.980	PQL	ng/Kg	
	OCDF	JB	2.82	9.80	PQL	ng/Kg	

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-577-SA5D-SB-0.0-0.5	ARSENIC	J	1.81	3.93	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.544	0.982	PQL	mg/Kg	
	CADMIUM	J	0.799	0.982	PQL	mg/Kg	
	TIN	J	2.37	9.82	PQL	mg/Kg	

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-577-SA5D-SB-0.0-0.5	SELENIUM	J	0.253	0.393	PQL	mg/Kg	J (all detects)
	THALLIUM	J	0.187	0.196	PQL	mg/Kg	

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Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-576-SA5D-SB-3.5-4.5	EFH (C21-C30)	J	5.1	5.3	PQL	mg/Kg	J (all detects)
SL-578-SA5D-SB-0.0-0.5	EFH (C15-C20)	J	2.6	5.3	PQL	mg/Kg	J (all detects)

Method: 8081B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-576-SA5D-SB-0.0-0.5	4,4'-DDT	J	0.40	1.7	PQL	ug/Kg	J (all detects)
SL-579-SA5D-SB-0.0-0.5	4,4'-DDE	J	0.86	1.7	PQL	ug/Kg	J (all detects)
	4,4'-DDT	J	0.98	1.7	PQL	ug/Kg	
SL-879-SA5D-SB-0.0-0.5	4,4'-DDE	J	0.82	1.7	PQL	ug/Kg	J (all detects)
	4,4'-DDT	J	1.0	1.7	PQL	ug/Kg	

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-578-SA5D-SB-0.0-0.5	1-METHYLNAPHTHALENE	J	1.4	1.8	PQL	ug/Kg	J (all detects)
	BENZO(A)ANTHRACENE	J	1.2	1.8	PQL	ug/Kg	
	BENZO(A)PYRENE	J	1.2	1.8	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	1.1	1.8	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	0.89	1.8	PQL	ug/Kg	
	INDENO(1,2,3-CD)PYRENE	J	0.86	1.8	PQL	ug/Kg	
SL-581-SA5D-SB-0.0-0.5	2-METHYLNAPHTHALENE	J	1.6	1.7	PQL	ug/Kg	J (all detects)
	BENZO(B)FLUORANTHENE	J	0.98	1.7	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	12	18	PQL	ug/Kg	
	CHRYSENE	J	1.1	1.7	PQL	ug/Kg	
	FLUORANTHENE	J	1.1	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	0.87	1.7	PQL	ug/Kg	
	PYRENE	J	0.90	1.7	PQL	ug/Kg	
SL-581-SA5D-SB-2.5-3.5	NAPHTHALENE	J	0.74	1.8	PQL	ug/Kg	J (all detects)

LDC #: 30695N4

VALIDATION COMPLETENESS WORKSHEET

Date: 11/11/13

SDG #: PH111

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: al

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	-	Sampling dates: 9/19/13
II.	ICP/MS Tune	-	
III.	Calibration	-	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	N	CS
VII.	Duplicate Sample Analysis	N	
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	N	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	N	
XV.	Field Blanks	SW	EB = EB-091813 FB = FB-011113 01/10/13 32

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

(PH111)
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

(PH029)
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Validated Samples: soil

1	SL-577-SA5D-SB-0.0-0.5	11		21		31	
2		12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/16/13 **Soil factor applied:** 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ **Associated Samples:** All Soil

Analyte	Blank ID	Sample Identification															
	FB-041613 (SDG: PH032)	Action Limit	No Qualifiers														
Sn	0.0029	1.45															
Mo	0.0132	6.6															

Sampling date: 9/18/13 **Soil factor applied:** 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ **Associated Samples:** All Soil

Analyte	Blank ID	Sample Identification															
	EB1-091813 (SDG: PH110)	Action Limit	No Qualifiers														
Ca	0.0464	23.2															
Mo	0.0083	4.15															

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH112

Prepared for

CDM Smith
555 17th Street, Suite 1100
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Prepared by

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Carlsbad, California 92010

December 11, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on September 23, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by EPA SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Pesticides by EPA SW 846 Method 8081A
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A and 7471B
Herbicides by EPA SW 846 Method 8151A
Perchlorate by EPA Method 6850
Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M
TPH as Extractables by EPA SW 846 Method 8015M
Dioxins and Dibenzofurans by EPA Method 1613B

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks and field duplicates. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits with the exception of several samples for PCBs. The associated sample results were qualified as detected estimated (J) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pairs for SVOCs, metals and TPH as extractables. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits with the exception of one DUP for metals. The associated sample results were qualified as detected estimated (J) as applicable. The details regarding the qualification of data are provided in Enclosure I.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one LCS/LCSD pair for PCBs. The associated sample results were qualified as detected estimated (J) as applicable. The details regarding the qualification of data are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met with the following exceptions:

Diluted Sample	Analyte	%D (Limits)	Associated Samples	Flag	A or P
SL-568-SA5D-SB-2.5-3.5	Zinc	16 (≤ 10)	All soil samples in SDG PH112	J (all detects) UJ (all non-detects)	A

The associated sample results were qualified as detected estimated (J).

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH112	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pair was collected and analyzed for SVOCS, pesticides, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables and dioxins. All RPDs were within QC limits with the exception of several SVOCS, PCBs, metals, TPH as extractables. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No volatile contaminants were found in the trip blank.

One equipment blank (from SDG PH114) was collected and analyzed for SVOCS, pesticides, PCBs, metals, herbicides, perchlorate, TPH as gasoline, TPH as extractables and dioxins. The

equipment blank had detections for SVOCs, metals and dioxins. The associated sample results were not detected or were significantly greater than the concentrations found in the equipment blank, therefore no data were qualified.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, perchlorate, TPH as gasoline, TPH as extractables and dioxins. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1

Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
23-Sep-2013	SL-568-SA5D-SB-0.0-0.5	7208981	N	3050B	6010C	III
23-Sep-2013	SL-568-SA5D-SB-0.0-0.5	7208981	N	3050B	6020A	III
23-Sep-2013	SL-568-SA5D-SB-0.0-0.5	7208981	N	3546	8015M	III
23-Sep-2013	SL-568-SA5D-SB-0.0-0.5	7208981	N	3546	8081B	III
23-Sep-2013	SL-568-SA5D-SB-0.0-0.5	7208981	N	3546	8082A	III
23-Sep-2013	SL-568-SA5D-SB-0.0-0.5	7208981	N	3546	8270D SIM	III
23-Sep-2013	SL-568-SA5D-SB-0.0-0.5	7208981	N	3550B	8151A	III
23-Sep-2013	SL-568-SA5D-SB-0.0-0.5	7208981	N	METHOD	1613B	III
23-Sep-2013	SL-568-SA5D-SB-0.0-0.5	7208981	N	METHOD	7471B	III
23-Sep-2013	TB-092313	7208980	TB	5030B	8015M	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5	7208982	N	3050B	6010C	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5	7208982	N	3050B	6020A	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5	7208982	N	3546	8015M	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5	7208982	N	3546	8081B	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5	7208982	N	3546	8082A	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5	7208982	N	3546	8270D SIM	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5	7208982	N	3550B	8151A	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5	7208982	N	5035A	8015M	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5	7208982	N	METHOD	7471B	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MS	7208983	MS	3050B	6010C	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MS	7208983	MS	3050B	6020A	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MS	7208983	MS	3546	8015M	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MS	7208983	MS	3546	8081B	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MS	7208983	MS	3546	8082A	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MS	7208983	MS	3546	8270D SIM	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MS	7208983	MS	3550B	8151A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MS	7208983	MS	5035A	8015M	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MS	7208983	MS	METHOD	7471B	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MSD	7208984	MSD	3050B	6010C	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MSD	7208984	MSD	3050B	6020A	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MSD	7208984	MSD	3546	8015M	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MSD	7208984	MSD	3546	8081B	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MSD	7208984	MSD	3546	8082A	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MSD	7208984	MSD	3546	8270D SIM	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MSD	7208984	MSD	3550B	8151A	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MSD	7208984	MSD	5035A	8015M	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5MSD	7208984	MSD	METHOD	7471B	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5DUP	7208985	DUP	3050B	6010C	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5DUP	7208985	DUP	3050B	6020A	III
23-Sep-2013	SL-568-SA5D-SB-2.5-3.5DUP	7208985	DUP	METHOD	7471B	III
23-Sep-2013	SL-868-SA5D-SB-2.5-3.5	7208986	FD	3050B	6010C	III
23-Sep-2013	SL-868-SA5D-SB-2.5-3.5	7208986	FD	3050B	6020A	III
23-Sep-2013	SL-868-SA5D-SB-2.5-3.5	7208986	FD	3546	8015M	III
23-Sep-2013	SL-868-SA5D-SB-2.5-3.5	7208986	FD	3546	8081B	III
23-Sep-2013	SL-868-SA5D-SB-2.5-3.5	7208986	FD	3546	8082A	III
23-Sep-2013	SL-868-SA5D-SB-2.5-3.5	7208986	FD	3546	8270D SIM	III
23-Sep-2013	SL-868-SA5D-SB-2.5-3.5	7208986	FD	3550B	8151A	III
23-Sep-2013	SL-868-SA5D-SB-2.5-3.5	7208986	FD	5035A	8015M	III
23-Sep-2013	SL-868-SA5D-SB-2.5-3.5	7208986	FD	METHOD	7471B	III
23-Sep-2013	SL-574-SA5D-SB-0.0-0.5	7208989	N	3546	8081B	III
23-Sep-2013	SL-574-SA5D-SB-0.0-0.5	7208989	N	3546	8270D SIM	III
23-Sep-2013	SL-574-SA5D-SB-0.0-0.5	7208989	N	3550B	8151A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
23-Sep-2013	SL-574-SA5D-SB-0.0-0.5	7208989	N	METHOD	1613B	III
23-Sep-2013	SL-572-SA5D-SB-0.0-0.5	7208988	N	3546	8081B	III
23-Sep-2013	SL-572-SA5D-SB-0.0-0.5	7208988	N	3550B	8151A	III
23-Sep-2013	SL-572-SA5D-SB-0.0-0.5	7208988	N	METHOD	1613B	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5	7208987	N	3050B	6010C	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5	7208987	N	3050B	6020A	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5	7208987	N	3546	8015M	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5	7208987	N	3546	8081B	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5	7208987	N	3546	8082A	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5	7208987	N	3546	8270D SIM	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5	7208987	N	3550B	8151A	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5	7208987	N	METHOD	1613B	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5	7208987	N	METHOD	6850	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5	7208987	N	METHOD	7471B	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5MSD	P208987M241750A	MSD	METHOD	6850	III
23-Sep-2013	SL-571-SA5D-SB-0.0-0.5MS	P208987R241737A	MS	METHOD	6850	III
23-Sep-2013	SL-583-SA5D-SB-0.0-0.5	7208990	N	3050B	6010C	III
23-Sep-2013	SL-583-SA5D-SB-0.0-0.5	7208990	N	3050B	6020A	III
23-Sep-2013	SL-583-SA5D-SB-0.0-0.5	7208990	N	3546	8015M	III
23-Sep-2013	SL-583-SA5D-SB-0.0-0.5	7208990	N	3546	8082A	III
23-Sep-2013	SL-583-SA5D-SB-0.0-0.5	7208990	N	3546	8270D SIM	III
23-Sep-2013	SL-583-SA5D-SB-0.0-0.5	7208990	N	METHOD	1613B	III
23-Sep-2013	SL-583-SA5D-SB-0.0-0.5	7208990	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS								
Method:	6010C			Matrix: SO					

Sample ID: SL-568-SA5D-SB-0.0-0.5		Collected: 9/23/2013 7:30:00			Analysis Type: REA			Dilution: 1		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
VANADIUM	50.1		0.131	MDL	1.01	PQL	mg/Kg	J	E	
TITANIUM	790		0.171	MDL	1.01	PQL	mg/Kg	J	E	

Sample ID: SL-568-SA5D-SB-0.0-0.5		Collected: 9/23/2013 7:30:00			Analysis Type: RES			Dilution: 1		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
ALUMINUM	14800		7.27	MDL	40.3	PQL	mg/Kg	J	E	
ANTIMONY	4.03	U	0.746	MDL	4.03	PQL	mg/Kg	UJ	Q	
BERYLLIUM	0.458	J	0.0676	MDL	1.01	PQL	mg/Kg	J	Z	
BORON	7.90	J	0.847	MDL	10.1	PQL	mg/Kg	J	Z	
CADMIUM	0.868	J	0.0767	MDL	1.01	PQL	mg/Kg	J	Z	
COBALT	7.00		0.0999	MDL	1.01	PQL	mg/Kg	J	E	
IRON	22500		3.65	MDL	40.3	PQL	mg/Kg	J	E	
LITHIUM	16.0		0.34	MDL	4.0	PQL	mg/Kg	J	E	
MAGNESIUM	6990		1.68	MDL	10.1	PQL	mg/Kg	J	E	
MANGANESE	287		0.0837	MDL	1.01	PQL	mg/Kg	J	E	
MOLYBDENUM	1.44	J	0.171	MDL	2.02	PQL	mg/Kg	U	F	
POTASSIUM	3520		8.41	MDL	101	PQL	mg/Kg	J	Q, E	
TIN	2.54	J	0.222	MDL	10.1	PQL	mg/Kg	U	B	
ZINC	68.8		0.202	MDL	4.03	PQL	mg/Kg	J	Q, E, A	

Sample ID: SL-568-SA5D-SB-2.5-3.5		Collected: 9/23/2013 8:00:00			Analysis Type: REA			Dilution: 1		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
VANADIUM	60.1		0.139	MDL	1.07	PQL	mg/Kg	J	E	

Sample ID: SL-568-SA5D-SB-2.5-3.5		Collected: 9/23/2013 8:00:00			Analysis Type: REA2			Dilution: 2		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
TITANIUM	1170		0.362	MDL	2.13	PQL	mg/Kg	J	E	

Sample ID: SL-568-SA5D-SB-2.5-3.5		Collected: 9/23/2013 8:00:00			Analysis Type: RES			Dilution: 1		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
ALUMINUM	21900		7.69	MDL	42.6	PQL	mg/Kg	J	E	

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-568-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.26	U	0.789	MDL	4.26	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.602	J	0.0714	MDL	1.07	PQL	mg/Kg	J	Z
BORON	10.1	J	0.896	MDL	10.7	PQL	mg/Kg	J	Z
CADMIUM	0.342	J	0.0810	MDL	1.07	PQL	mg/Kg	J	Z
COBALT	8.49		0.106	MDL	1.07	PQL	mg/Kg	J	E
IRON	27700		3.86	MDL	42.6	PQL	mg/Kg	J	E
LITHIUM	35.1		0.36	MDL	4.3	PQL	mg/Kg	J	E
MAGNESIUM	7660		1.78	MDL	10.7	PQL	mg/Kg	J	E
MANGANESE	283		0.0885	MDL	1.07	PQL	mg/Kg	J	E
POTASSIUM	3590		8.89	MDL	107	PQL	mg/Kg	J	Q, E
TIN	3.30	J	0.235	MDL	10.7	PQL	mg/Kg	U	B
ZINC	83.4		0.213	MDL	4.26	PQL	mg/Kg	J	Q, E, A
Zirconium	4.53	J	0.896	MDL	5.33	PQL	mg/Kg	J	Z

Sample ID: SL-571-SA5D-SB-0.0-0.5 Collected: 9/23/2013 10:25:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
VANADIUM	58.0		0.132	MDL	1.01	PQL	mg/Kg	J	E

Sample ID: SL-571-SA5D-SB-0.0-0.5 Collected: 9/23/2013 10:25:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	35500		7.34	MDL	81.1	PQL	mg/Kg	J	E
TITANIUM	1090		0.345	MDL	2.03	PQL	mg/Kg	J	E

Sample ID: SL-571-SA5D-SB-0.0-0.5 Collected: 9/23/2013 10:25:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	18900		7.31	MDL	40.6	PQL	mg/Kg	J	E
ANTIMONY	4.06	U	0.750	MDL	4.06	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.611	J	0.0679	MDL	1.01	PQL	mg/Kg	J	Z
BORON	6.99	J	0.852	MDL	10.1	PQL	mg/Kg	J	Z
CADMIUM	0.251	J	0.0771	MDL	1.01	PQL	mg/Kg	J	Z
COBALT	7.78		0.100	MDL	1.01	PQL	mg/Kg	J	E
LITHIUM	38.6		0.34	MDL	4.1	PQL	mg/Kg	J	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-571-SA5D-SB-0.0-0.5 Collected: 9/23/2013 10:25:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MAGNESIUM	8190		1.69	MDL	10.1	PQL	mg/Kg	J	E
MANGANESE	270		0.0841	MDL	1.01	PQL	mg/Kg	J	E
POTASSIUM	4370		8.46	MDL	101	PQL	mg/Kg	J	Q, E
TIN	2.83	J	0.223	MDL	10.1	PQL	mg/Kg	U	B
ZINC	76.7		0.203	MDL	4.06	PQL	mg/Kg	J	Q, E, A
Zirconium	2.62	J	0.852	MDL	5.07	PQL	mg/Kg	J	Z

Sample ID: SL-583-SA5D-SB-0.0-0.5 Collected: 9/23/2013 12:40:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
VANADIUM	61.0		0.135	MDL	1.04	PQL	mg/Kg	J	E

Sample ID: SL-583-SA5D-SB-0.0-0.5 Collected: 9/23/2013 12:40:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	31200		7.53	MDL	83.2	PQL	mg/Kg	J	E
TITANIUM	1010		0.354	MDL	2.08	PQL	mg/Kg	J	E

Sample ID: SL-583-SA5D-SB-0.0-0.5 Collected: 9/23/2013 12:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	22600		7.50	MDL	41.6	PQL	mg/Kg	J	E
ANTIMONY	4.16	U	0.770	MDL	4.16	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.758	J	0.0697	MDL	1.04	PQL	mg/Kg	J	Z
CADMIUM	0.341	J	0.0791	MDL	1.04	PQL	mg/Kg	J	Z
COBALT	8.40		0.103	MDL	1.04	PQL	mg/Kg	J	E
LITHIUM	25.4		0.35	MDL	4.2	PQL	mg/Kg	J	E
MAGNESIUM	7570		1.74	MDL	10.4	PQL	mg/Kg	J	E
MANGANESE	408		0.0864	MDL	1.04	PQL	mg/Kg	J	E
POTASSIUM	7160		8.68	MDL	104	PQL	mg/Kg	J	Q, E
SODIUM	84.7	J	17.4	MDL	104	PQL	mg/Kg	J	Z
TIN	3.21	J	0.229	MDL	10.4	PQL	mg/Kg	U	B
ZINC	80.0		0.208	MDL	4.16	PQL	mg/Kg	J	Q, E, A
Zirconium	5.17	J	0.874	MDL	5.20	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-868-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:10:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
TITANIUM	891		0.169	MDL	0.995	PQL	mg/Kg	J	E
VANADIUM	53.7		0.129	MDL	0.995	PQL	mg/Kg	J	E

Sample ID: SL-868-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	18100		7.18	MDL	39.8	PQL	mg/Kg	J	E
ANTIMONY	3.98	U	0.737	MDL	3.98	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.534	J	0.0667	MDL	0.995	PQL	mg/Kg	J	Z
BORON	8.73	J	0.836	MDL	9.95	PQL	mg/Kg	J	Z
CADMIUM	0.390	J	0.0756	MDL	0.995	PQL	mg/Kg	J	Z
COBALT	7.63		0.0985	MDL	0.995	PQL	mg/Kg	J	E
IRON	23800		3.60	MDL	39.8	PQL	mg/Kg	J	E
LITHIUM	28.5		0.34	MDL	4.0	PQL	mg/Kg	J	E
MAGNESIUM	6710		1.66	MDL	9.95	PQL	mg/Kg	J	E
MANGANESE	307		0.0826	MDL	0.995	PQL	mg/Kg	J	E
POTASSIUM	3020		8.30	MDL	99.5	PQL	mg/Kg	J	Q, E
TIN	2.66	J	0.219	MDL	9.95	PQL	mg/Kg	U	B
ZINC	66.1		0.199	MDL	3.98	PQL	mg/Kg	J	Q, E, A
Zirconium	3.82	J	0.836	MDL	4.98	PQL	mg/Kg	J	Z

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-568-SA5D-SB-0.0-0.5 Collected: 9/23/2013 7:30:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.324	J	0.101	MDL	0.403	PQL	mg/Kg	J	Z

Sample ID: SL-568-SA5D-SB-0.0-0.5 Collected: 9/23/2013 7:30:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.110	J	0.0262	MDL	0.202	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-568-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:00:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.195	J	0.107	MDL	0.426	PQL	mg/Kg	J	Z

Sample ID: SL-568-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:00:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0348	J	0.0277	MDL	0.213	PQL	mg/Kg	J	Z

Sample ID: SL-571-SA5D-SB-0.0-0.5 Collected: 9/23/2013 10:25:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.178	J	0.101	MDL	0.406	PQL	mg/Kg	J	Z

Sample ID: SL-583-SA5D-SB-0.0-0.5 Collected: 9/23/2013 12:40:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.332	J	0.104	MDL	0.416	PQL	mg/Kg	J	Z

Sample ID: SL-583-SA5D-SB-0.0-0.5 Collected: 9/23/2013 12:40:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0487	J	0.0271	MDL	0.208	PQL	mg/Kg	J	Z

Sample ID: SL-868-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:10:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.168	J	0.0995	MDL	0.398	PQL	mg/Kg	J	Z

Sample ID: SL-868-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:10:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0294	J	0.0259	MDL	0.199	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 7471B **Matrix:** SO

Sample ID: SL-568-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0123	J	0.0099	MDL	0.0165	PQL	mg/Kg	J	Z, FD

Sample ID: SL-571-SA5D-SB-0.0-0.5 Collected: 9/23/2013 10:25:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0160	J	0.0098	MDL	0.0163	PQL	mg/Kg	J	Z

Sample ID: SL-868-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0158	U	0.0095	MDL	0.0158	PQL	mg/Kg	UJ	FD

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-568-SA5D-SB-0.0-0.5 Collected: 9/23/2013 7:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8,9-HPCDF	0.444	JB	0.0280	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.713	JB	0.0476	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDF	0.679	JB	0.0396	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	1.53	JB	0.0515	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.649	JB	0.0392	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	1.25	JB	0.0492	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDF	0.566	JB	0.0402	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	2.73	JB	0.0580	MDL	5.00	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HxCDF	0.653	JB	0.0376	MDL	5.00	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	1.34	JB	0.0531	MDL	5.00	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.102	JQ	0.0378	MDL	1.00	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.702	JQ	0.114	MDL	1.00	PQL	ng/Kg	J	Z

Sample ID: SL-571-SA5D-SB-0.0-0.5 Collected: 9/23/2013 10:25:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.275	JB	0.0353	MDL	5.04	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-571-SA5D-SB-0.0-0.5 Collected: 9/23/2013 10:25:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	0.0964	JB	0.0111	MDL	5.04	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0500	JBQ	0.0185	MDL	5.04	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0351	JBQ	0.0205	MDL	5.04	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0551	JBQ	0.0235	MDL	5.04	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.0837	JBQ	0.0216	MDL	5.04	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0686	JB	0.0203	MDL	5.04	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0616	JBQ	0.0212	MDL	5.04	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.111	JB	0.0251	MDL	5.04	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0867	JBQ	0.0329	MDL	5.04	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.138	JBQ	0.0176	MDL	5.04	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0734	JBQ	0.0221	MDL	5.04	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0575	JBQ	0.0186	MDL	5.04	PQL	ng/Kg	U	B
OCDD	1.19	JB	0.0512	MDL	10.1	PQL	ng/Kg	U	B
OCDF	0.305	JBQ	0.0540	MDL	10.1	PQL	ng/Kg	U	B

Sample ID: SL-572-SA5D-SB-0.0-0.5 Collected: 9/23/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	3.11	JB	0.0343	MDL	4.88	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.744	JB	0.0148	MDL	4.88	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.143	JBQ	0.0234	MDL	4.88	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.152	JB	0.0311	MDL	4.88	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.249	JBQ	0.0286	MDL	4.88	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.539	JB	0.0340	MDL	4.88	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.274	JB	0.0253	MDL	4.88	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.659	JB	0.0308	MDL	4.88	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.534	JB	0.0303	MDL	4.88	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.322	JBQ	0.0377	MDL	4.88	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.511	JB	0.0307	MDL	4.88	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.207	JB	0.0274	MDL	4.88	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.460	JBQ	0.0311	MDL	4.88	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.111	JQ	0.0538	MDL	0.975	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.288	JQ	0.0709	MDL	0.975	PQL	ng/Kg	J	Z
OCDF	1.52	JB	0.0376	MDL	9.75	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-574-SA5D-SB-0.0-0.5		Collected: 9/23/2013 9:20:00		Analysis Type: RES		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.175	JB	0.0260	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0910	JBQ	0.0121	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0767	JBQ	0.0188	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0708	JBQ	0.0254	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.144	JBQ	0.0232	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.138	JB	0.0269	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.110	JBQ	0.0210	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.150	JB	0.0261	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.138	JB	0.0257	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.127	JBQ	0.0338	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.163	JBQ	0.0193	MDL	5.00	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0790	JBQ	0.0223	MDL	5.00	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.146	JB	0.0192	MDL	5.00	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0625	JQ	0.0572	MDL	1.00	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0615	JQ	0.0431	MDL	1.00	PQL	ng/Kg	J	Z
OCDD	0.428	JB	0.0306	MDL	10.0	PQL	ng/Kg	U	B
OCDF	0.246	JBQ	0.0495	MDL	10.0	PQL	ng/Kg	U	B

Sample ID: SL-583-SA5D-SB-0.0-0.5		Collected: 9/23/2013 12:40:00		Analysis Type: RES		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	4.83	JB	0.0754	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	1.07	JB	0.0541	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.285	JBQ	0.0966	MDL	5.18	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.300	JBQ	0.0699	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.693	JB	0.0656	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.470	JB	0.0744	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.491	JB	0.0588	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.588	JB	0.0737	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.615	JB	0.0892	MDL	5.18	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.303	JBQ	0.0599	MDL	5.18	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.614	JBQ	0.0580	MDL	5.18	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0878	JQ	0.0682	MDL	1.04	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.582	JQ	0.119	MDL	1.04	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-583-SA5D-SB-0.0-0.5 Collected: 9/23/2013 12:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
OCDF	2.79	JBQ	0.222	MDL	10.4	PQL	ng/Kg	J	Z

Method Category:	SVOA	
Method:	8015M	Matrix: SO

Sample ID: SL-568-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:00:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	22		2.1	MDL	5.3	PQL	mg/Kg	J	Q, Q, Q, FD
EFH (C30-C40)	26		4.3	MDL	11	PQL	mg/Kg	J	Q, Q, FD

Sample ID: SL-583-SA5D-SB-0.0-0.5 Collected: 9/23/2013 12:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	3.6	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z

Sample ID: SL-868-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:10:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	12		2.0	MDL	5.1	PQL	mg/Kg	J	FD
EFH (C30-C40)	45		4.1	MDL	10	PQL	mg/Kg	J	FD

Method Category:	SVOA	
Method:	8081B	Matrix: SO

Sample ID: SL-568-SA5D-SB-0.0-0.5 Collected: 9/23/2013 7:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DELTA-BHC	0.51	J	0.46	MDL	0.85	PQL	ug/Kg	J	Z
METHOXYCHLOR	4.1	J	1.7	MDL	6.8	PQL	ug/Kg	J	Z

Sample ID: SL-572-SA5D-SB-0.0-0.5 Collected: 9/23/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
4,4'-DDE	0.60	J	0.33	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8081B **Matrix:** SO

Sample ID: SL-572-SA5D-SB-0.0-0.5 Collected: 9/23/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
4,4'-DDT	1.4	J	0.35	MDL	1.7	PQL	ug/Kg	J	Z
gamma-BHC (Lindane)	0.24	J	0.17	MDL	0.83	PQL	ug/Kg	J	Z

Method Category: SVOA
Method: 8082A **Matrix:** SO

Sample ID: SL-568-SA5D-SB-0.0-0.5 Collected: 9/23/2013 7:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
Aroclor 5460	15	J	10	MDL	34	PQL	ug/Kg	J	Z, L

Sample ID: SL-568-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	7.3	J	4.7	MDL	18	PQL	ug/Kg	J	Z, S, FD

Sample ID: SL-571-SA5D-SB-0.0-0.5 Collected: 9/23/2013 10:25:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	12	J	4.5	MDL	17	PQL	ug/Kg	J	Z, S

Sample ID: SL-868-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	17	U	4.5	MDL	17	PQL	ug/Kg	UJ	FD

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-568-SA5D-SB-0.0-0.5 Collected: 9/23/2013 7:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(A)ANTHRACENE	16	J	6.8	MDL	17	PQL	ug/Kg	J	Z
BENZO(E)PYRENE	39	J	34	MDL	170	PQL	ug/Kg	J	Z
Butylbenzylphthalate	110	J	61	MDL	180	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-568-SA5D-SB-0.0-0.5 Collected: 9/23/2013 7:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
INDENO(1,2,3-CD)PYRENE	11	J	6.8	MDL	17	PQL	ug/Kg	J	Z

Sample ID: SL-568-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(A)ANTHRACENE	0.94	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	1.2	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	0.87	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	1.2	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z, FD
Butylbenzylphthalate	19	U	6.3	MDL	19	PQL	ug/Kg	UJ	FD
CHRYSENE	1.5	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z
Diethylphthalate	19	U	6.3	MDL	19	PQL	ug/Kg	UJ	Q
INDENO(1,2,3-CD)PYRENE	1.8	U	0.70	MDL	1.8	PQL	ug/Kg	UJ	FD
PHENANTHRENE	0.93	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-571-SA5D-SB-0.0-0.5 Collected: 9/23/2013 10:25:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	0.79	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-574-SA5D-SB-0.0-0.5 Collected: 9/23/2013 9:20:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	0.69	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-583-SA5D-SB-0.0-0.5 Collected: 9/23/2013 12:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	0.92	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-583-SA5D-SB-0.0-0.5 Collected: 9/23/2013 12:40:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	1.5	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(A)ANTHRACENE	0.95	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-583-SA5D-SB-0.0-0.5 Collected: 9/23/2013 12:40:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(A)PYRENE	1.0	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	1.2	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	0.84	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	12	J	6.2	MDL	19	PQL	ug/Kg	J	Z
INDENO(1,2,3-CD)PYRENE	0.89	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-868-SA5D-SB-2.5-3.5 Collected: 9/23/2013 8:10:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(A)ANTHRACENE	0.82	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	0.98	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(B)FLUORANTHENE	1.4	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	1.2	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	1.7	U	0.68	MDL	1.7	PQL	ug/Kg	UJ	FD
Butylbenzylphthalate	7.5	J	6.1	MDL	18	PQL	ug/Kg	J	Z, FD
CHRYSENE	1.1	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
FLUORANTHENE	1.4	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
INDENO(1,2,3-CD)PYRENE	0.76	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z, FD
PHENANTHRENE	0.86	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	1.4	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PrepPH112

eQAPP Name: CDM_SSFL_131101_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
*#	Professional Judgment
A	ICP Serial Dilution
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Field Blank Contamination
FD	Field Duplicate Precision
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH112

Method Blank Outlier Report

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2730B371140	10/2/2013 11:40:00 AM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF OCDD OCDF	0.0965 ng/Kg 0.0595 ng/Kg 0.0657 ng/Kg 0.0334 ng/Kg 0.0515 ng/Kg 0.0256 ng/Kg 0.0453 ng/Kg 0.0369 ng/Kg 0.0754 ng/Kg 0.0738 ng/Kg 0.0876 ng/Kg 0.0332 ng/Kg 0.0646 ng/Kg 0.249 ng/Kg 0.137 ng/Kg	SL-568-SA5D-SB-0.0-0.5 SL-571-SA5D-SB-0.0-0.5 SL-572-SA5D-SB-0.0-0.5 SL-574-SA5D-SB-0.0-0.5 SL-583-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.275 ng/Kg	0.275U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0964 ng/Kg	0.0964U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0500 ng/Kg	0.0500U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0351 ng/Kg	0.0351U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.0551 ng/Kg	0.0551U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.0837 ng/Kg	0.0837U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0686 ng/Kg	0.0686U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.0616 ng/Kg	0.0616U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.111 ng/Kg	0.111U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.0867 ng/Kg	0.0867U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.138 ng/Kg	0.138U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0734 ng/Kg	0.0734U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.0575 ng/Kg	0.0575U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	OCDD	1.19 ng/Kg	1.19U ng/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	OCDF	0.305 ng/Kg	0.305U ng/Kg
SL-572-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.143 ng/Kg	0.143U ng/Kg
SL-572-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.152 ng/Kg	0.152U ng/Kg
SL-572-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.249 ng/Kg	0.249U ng/Kg
SL-572-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.322 ng/Kg	0.322U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.175 ng/Kg	0.175U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0910 ng/Kg	0.0910U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0767 ng/Kg	0.0767U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0708 ng/Kg	0.0708U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.144 ng/Kg	0.144U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.110 ng/Kg	0.110U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.150 ng/Kg	0.150U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.138 ng/Kg	0.138U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.127 ng/Kg	0.127U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.163 ng/Kg	0.163U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-574-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0790 ng/Kg	0.0790U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.146 ng/Kg	0.146U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	OCDD	0.428 ng/Kg	0.428U ng/Kg
SL-574-SA5D-SB-0.0-0.5(RES)	OCDF	0.246 ng/Kg	0.246U ng/Kg
SL-583-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.285 ng/Kg	0.285U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P27437AB221412	10/3/2013 2:12:00 PM	CALCIUM TIN ZINC	9.48 mg/Kg 1.46 mg/Kg 0.850 mg/Kg	SL-568-SA5D-SB-0.0-0.5 SL-568-SA5D-SB-2.5-3.5 SL-571-SA5D-SB-0.0-0.5 SL-583-SA5D-SB-0.0-0.5 SL-868-SA5D-SB-2.5-3.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-568-SA5D-SB-0.0-0.5(RES)	TIN	2.54 mg/Kg	2.54U mg/Kg
SL-568-SA5D-SB-2.5-3.5(RES)	TIN	3.30 mg/Kg	3.30U mg/Kg
SL-571-SA5D-SB-0.0-0.5(RES)	TIN	2.83 mg/Kg	2.83U mg/Kg
SL-583-SA5D-SB-0.0-0.5(RES)	TIN	3.21 mg/Kg	3.21U mg/Kg
SL-868-SA5D-SB-2.5-3.5(RES)	TIN	2.66 mg/Kg	2.66U mg/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041613(REA2/TOT)	4/16/2013 3:15:00 PM	MOLYBDENUM TIN	0.0132 mg/L 0.0029 mg/L	SL-568-SA5D-SB-0.0-0.5 SL-568-SA5D-SB-2.5-3.5 SL-571-SA5D-SB-0.0-0.5 SL-572-SA5D-SB-0.0-0.5 SL-574-SA5D-SB-0.0-0.5 SL-583-SA5D-SB-0.0-0.5 SL-868-SA5D-SB-2.5-3.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-568-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	1.44 mg/Kg	1.44U mg/Kg

Surrogate Outlier Report

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8082A

Matrix: SO

Sample ID (Analysis Type)	Surrogate	Sample % Recovery	% Recovery Limits	Affected Compounds	Flag
SL-568-SA5D-SB- 2.5-3.5	DECACHLOROBIPHENYL	138	45.00-120.00	All Target Analytes	J (all detects)
	TETRACHLORO-M-XYLENE	145	45.00-120.00		
SL-571-SA5D-SB- 0.0-0.5	DECACHLOROBIPHENYL	137	45.00-120.00	All Target Analytes	J(all detects)
	TETRACHLORO-M-XYLENE	132	45.00-120.00		
SL-868-SA5D-SB- 2.5-3.5	DECACHLOROBIPHENYL	130	45.00-120.00	All Target Analytes	J(all detects)
	TETRACHLORO-M-XYLENE	121	45.00-120.00		

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8015M

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-568-SA5D-SB-2.5-3.5MS SL-568-SA5D-SB-2.5-3.5MSD (SL-568-SA5D-SB-2.5-3.5)	EFH (C30-C40)	144	526	49.00-123.00	76 (20.00)	EFH (C30-C40)	J (all detects)
SL-568-SA5D-SB-2.5-3.5MS SL-568-SA5D-SB-2.5-3.5MSD (SL-568-SA5D-SB-2.5-3.5)	EFH (C21-C30)	29	193	49.00-123.00	59 (20.00)	EFH (C21-C30)	J(all detects) UJ(all non-detects)

Method: 8270D SIM

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-568-SA5D-SB-2.5-3.5MS (SL-568-SA5D-SB-2.5-3.5)	Diethylphthalate	73	-	76.00-127.00	-	Diethylphthalate	J(all detects) UJ(all non-detects)

Method: 6010C

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-568-SA5D-SB-2.5-3.5MS (TOT) SL-568-SA5D-SB-2.5-3.5MSD (TOT) (SL-568-SA5D-SB-0.0-0.5 SL-568-SA5D-SB-2.5-3.5 SL-571-SA5D-SB-0.0-0.5 SL-583-SA5D-SB-0.0-0.5 SL-868-SA5D-SB-2.5-3.5)	CALCIUM MANGANESE PHOSPHORUS TITANIUM	1949 230 173 177	2450 342 168 236	75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00	- - - -	CALCIUM MANGANESE PHOSPHORUS TITANIUM	No Qual, >4x
SL-568-SA5D-SB-2.5-3.5MS (TOT) SL-568-SA5D-SB-2.5-3.5MSD (TOT) (SL-568-SA5D-SB-0.0-0.5 SL-568-SA5D-SB-2.5-3.5 SL-571-SA5D-SB-0.0-0.5 SL-583-SA5D-SB-0.0-0.5 SL-868-SA5D-SB-2.5-3.5)	ALUMINUM IRON MAGNESIUM	-136 -5142 -455	-131 -6500 -524	75.00-125.00 75.00-125.00 75.00-125.00	- - -	ALUMINUM IRON MAGNESIUM	No Qual, >4x
SL-568-SA5D-SB-2.5-3.5MS (TOT) SL-568-SA5D-SB-2.5-3.5MSD (TOT) (SL-568-SA5D-SB-0.0-0.5 SL-568-SA5D-SB-2.5-3.5 SL-571-SA5D-SB-0.0-0.5 SL-583-SA5D-SB-0.0-0.5 SL-868-SA5D-SB-2.5-3.5)	ANTIMONY POTASSIUM ZINC	44 48 57	42 64 66	75.00-125.00 75.00-125.00 75.00-125.00	- - -	ANTIMONY POTASSIUM ZINC	J(all detects) UJ(all non-detects)

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6020A

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-568-SA5D-SB-2.5-3.5MS (TOT) (SL-568-SA5D-SB-0.0-0.5 SL-568-SA5D-SB-2.5-3.5 SL-571-SA5D-SB-0.0-0.5 SL-583-SA5D-SB-0.0-0.5 SL-868-SA5D-SB-2.5-3.5)	STRONTIUM	145	-	75.00-125.00	-	STRONTIUM	No Qual, >4x

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-568-SA5D-SB-2.5-3.5DUP (TOT)	ALUMINUM	43	20.00	J (all detects) UJ (all non-detects) As, Be, B, Cd, Mo, Na, Zr No Qual, OK by Difference
(SL-568-SA5D-SB-0.0-0.5)	ARSENIC	21	20.00	
SL -568-SA5D-SB-2.5-3.5	BERYLLIUM	31	20.00	
SL -571-SA5D-SB-0.0-0.5	BORON	30	20.00	
SL -583-SA5D-SB-0.0-0.5	CADMIUM	32	20.00	
SL -868-SA5D-SB-2.5-3.5)	COBALT	30	20.00	
	IRON	39	20.00	
	LITHIUM	56	20.00	
	MAGNESIUM	33	20.00	
	MANGANESE	29	20.00	
	MOLYBDENUM	200	20.00	
	POTASSIUM	31	20.00	
	SODIUM	25	20.00	
	TITANIUM	51	20.00	
	VANADIUM	30	20.00	
	ZINC	35	20.00	
	Zirconium	28	20.00	

Method: 6020A
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-568-SA5D-SB-2.5-3.5DUP (TOT)	SELENIUM	26	20.00	No Qual, OK by Difference
(SL-568-SA5D-SB-0.0-0.5)				
SL -568-SA5D-SB-2.5-3.5				
SL -571-SA5D-SB-0.0-0.5				
SL -583-SA5D-SB-0.0-0.5				
SL -868-SA5D-SB-2.5-3.5)				

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8082A

Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P32699AQ241313A P32699AY241254A (SL-568-SA5D-SB-0.0-0.5 SL-568-SA5D-SB-2.5-3.5 SL-571-SA5D-SB-0.0-0.5 SL-583-SA5D-SB-0.0-0.5 SL-868-SA5D-SB-2.5-3.5)	Aroclor 5442	96	94	62.00-87.00	-	Aroclor 5432, Aroclor 5442, Aroclor 5460,	J (all detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 160.3M
Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-568-SA5D-SB-2.5-3.5	SL-868-SA5D-SB-2.5-3.5			
MOISTURE	6.2	1.5	122		No Qualifiers Applied

Method: 6010C
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-568-SA5D-SB-2.5-3.5 (TOT)	SL-868-SA5D-SB-2.5-3.5 (TOT)			
ALUMINUM	21900	18100	19	50.00	No Qualifiers Applied
ARSENIC	4.32	4.83	11	50.00	
BARIUM	102	91.9	10	50.00	
BERYLLIUM	0.602	0.534	12	50.00	
BORON	10.1	8.73	15	50.00	
CADMIUM	0.342	0.390	13	50.00	
CALCIUM	39600	43500	9	50.00	
CHROMIUM	33.3	32.7	2	50.00	
COBALT	8.49	7.63	11	50.00	
COPPER	17.1	15.8	8	50.00	
IRON	27700	23800	15	50.00	
LEAD	7.55	6.79	11	50.00	
LITHIUM	35.1	28.5	21	50.00	
MAGNESIUM	7660	6710	13	50.00	
MANGANESE	283	307	8	50.00	
NICKEL	18.6	16.6	11	50.00	
PHOSPHORUS	468	487	4	50.00	
POTASSIUM	3590	3020	17	50.00	
SODIUM	200	176	13	50.00	
TIN	3.30	2.66	21	50.00	
TITANIUM	1170	891	27	50.00	
VANADIUM	60.1	53.7	11	50.00	
ZINC	83.4	66.1	23	50.00	
Zirconium	4.53	3.82	17	50.00	

Method: 6020A
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-568-SA5D-SB-2.5-3.5 (TOT)	SL-868-SA5D-SB-2.5-3.5 (TOT)			
SELENIUM	0.195	0.168	15	50.00	No Qualifiers Applied
SILVER	0.0348	0.0294	17	50.00	
STRONTIUM	95.6	97.0	1	50.00	
THALLIUM	0.302	0.269	12	50.00	

Method: 7471B
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-568-SA5D-SB-2.5-3.5 (TOT)	SL-868-SA5D-SB-2.5-3.5 (TOT)			
MERCURY	0.0123	0.0158 U	200	50.00	J(all detects) UJ(all non-detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8015M

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-568-SA5D-SB-2.5-3.5	SL-868-SA5D-SB-2.5-3.5			
EFH (C21-C30)	22	12	59	50.00	J(all detects)
EFH (C30-C40)	26	45	54	50.00	

Method: 8082A

Matrix: SO

Analyte	Concentration (ug/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-568-SA5D-SB-2.5-3.5	SL-868-SA5D-SB-2.5-3.5			
AROCLOR 1254	7.3	17 U	200	50.00	J(all detects) UJ(all non-detects)

Method: 8270D SIM

Matrix: SO

Analyte	Concentration (ug/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-568-SA5D-SB-2.5-3.5	SL-868-SA5D-SB-2.5-3.5			
BENZO(A)ANTHRACENE	0.94	0.82	14	50.00	No Qualifiers Applied
BENZO(A)PYRENE	1.2	0.98	20	50.00	
BENZO(B)FLUORANTHENE	2.1	1.4	40	50.00	
BENZO(G,H,I)PERYLENE	0.87	1.2	32	50.00	
CHRYSENE	1.5	1.1	31	50.00	
FLUORANTHENE	2.1	1.4	40	50.00	
PHENANTHRENE	0.93	0.86	8	50.00	
PYRENE	1.9	1.4	30	50.00	
BENZO(K)FLUORANTHENE	1.2	1.7 U	200	50.00	J(all detects) UJ(all non-detects)
Butylbenzylphthalate	19 U	7.5	200	50.00	
INDENO(1,2,3-CD)PYRENE	1.8 U	0.76	200	50.00	

Method: 9045M

Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-568-SA5D-SB-2.5-3.5	SL-868-SA5D-SB-2.5-3.5			
PH	7.99	8.14	2	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-568-SA5D-SB-0.0-0.5	1,2,3,4,7,8,9-HPCDF	JB	0.444	5.00	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8-HxCDD	JB	0.713	5.00	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.679	5.00	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	1.53	5.00	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.649	5.00	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	1.25	5.00	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.566	5.00	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	2.73	5.00	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.653	5.00	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	1.34	5.00	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.102	1.00	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.702	1.00	PQL	ng/Kg	
	SL-571-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	0.275	5.04	PQL	
1,2,3,4,6,7,8-HPCDF		JB	0.0964	5.04	PQL	ng/Kg	
1,2,3,4,7,8,9-HPCDF		JBQ	0.0500	5.04	PQL	ng/Kg	
1,2,3,4,7,8-HxCDD		JBQ	0.0351	5.04	PQL	ng/Kg	
1,2,3,4,7,8-HXCDF		JBQ	0.0551	5.04	PQL	ng/Kg	
1,2,3,6,7,8-HXCDD		JBQ	0.0837	5.04	PQL	ng/Kg	
1,2,3,6,7,8-HXCDF		JB	0.0686	5.04	PQL	ng/Kg	
1,2,3,7,8,9-HXCDD		JBQ	0.0616	5.04	PQL	ng/Kg	
1,2,3,7,8,9-HXCDF		JB	0.111	5.04	PQL	ng/Kg	
1,2,3,7,8-PECDD		JBQ	0.0867	5.04	PQL	ng/Kg	
1,2,3,7,8-PECDF		JBQ	0.138	5.04	PQL	ng/Kg	
2,3,4,6,7,8-HXCDF		JBQ	0.0734	5.04	PQL	ng/Kg	
2,3,4,7,8-PECDF		JBQ	0.0575	5.04	PQL	ng/Kg	
OCDD		JB	1.19	10.1	PQL	ng/Kg	
OCDF	JBQ	0.305	10.1	PQL	ng/Kg		
SL-572-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	3.11	4.88	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.744	4.88	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.143	4.88	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.152	4.88	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.249	4.88	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.539	4.88	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.274	4.88	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.659	4.88	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.534	4.88	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.322	4.88	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.511	4.88	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.207	4.88	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.460	4.88	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.111	0.975	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.288	0.975	PQL	ng/Kg	
OCDF	JB	1.52	9.75	PQL	ng/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-574-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	0.175	5.00	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0910	5.00	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0767	5.00	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0708	5.00	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.144	5.00	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.138	5.00	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.110	5.00	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.150	5.00	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.138	5.00	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.127	5.00	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.163	5.00	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.0790	5.00	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.146	5.00	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0625	1.00	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0615	1.00	PQL	ng/Kg	
	OCDD	JB	0.428	10.0	PQL	ng/Kg	
	OCDF	JBQ	0.246	10.0	PQL	ng/Kg	
SL-583-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	4.83	5.18	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	1.07	5.18	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.285	5.18	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.300	5.18	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.693	5.18	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.470	5.18	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.491	5.18	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.588	5.18	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.615	5.18	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.303	5.18	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.614	5.18	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0878	1.04	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.582	1.04	PQL	ng/Kg	
OCDF	JBQ	2.79	10.4	PQL	ng/Kg		

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-568-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.458	1.01	PQL	mg/Kg	J (all detects)
	BORON	J	7.90	10.1	PQL	mg/Kg	
	CADMIUM	J	0.868	1.01	PQL	mg/Kg	
	MOLYBDENUM	J	1.44	2.02	PQL	mg/Kg	
SL-568-SA5D-SB-2.5-3.5	TIN	J	2.54	10.1	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.602	1.07	PQL	mg/Kg	
	BORON	J	10.1	10.7	PQL	mg/Kg	
	CADMIUM	J	0.342	1.07	PQL	mg/Kg	
	TIN	J	3.30	10.7	PQL	mg/Kg	
SL-571-SA5D-SB-0.0-0.5	Zirconium	J	4.53	5.33	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.611	1.01	PQL	mg/Kg	
	BORON	J	6.99	10.1	PQL	mg/Kg	
	CADMIUM	J	0.251	1.01	PQL	mg/Kg	
	TIN	J	2.83	10.1	PQL	mg/Kg	
Zirconium	J	2.62	5.07	PQL	mg/Kg		

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Reporting Limit Outliers

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-583-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.758	1.04	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.341	1.04	PQL	mg/Kg	
	SODIUM	J	84.7	104	PQL	mg/Kg	
	TIN	J	3.21	10.4	PQL	mg/Kg	
	Zirconium	J	5.17	5.20	PQL	mg/Kg	
SL-868-SA5D-SB-2.5-3.5	BERYLLIUM	J	0.534	0.995	PQL	mg/Kg	J (all detects)
	BORON	J	8.73	9.95	PQL	mg/Kg	
	CADMIUM	J	0.390	0.995	PQL	mg/Kg	
	TIN	J	2.66	9.95	PQL	mg/Kg	
	Zirconium	J	3.82	4.98	PQL	mg/Kg	

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-568-SA5D-SB-0.0-0.5	SELENIUM	J	0.324	0.403	PQL	mg/Kg	J (all detects)
	SILVER	J	0.110	0.202	PQL	mg/Kg	
SL-568-SA5D-SB-2.5-3.5	SELENIUM	J	0.195	0.426	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0348	0.213	PQL	mg/Kg	
SL-571-SA5D-SB-0.0-0.5	SELENIUM	J	0.178	0.406	PQL	mg/Kg	J (all detects)
SL-583-SA5D-SB-0.0-0.5	SELENIUM	J	0.332	0.416	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0487	0.208	PQL	mg/Kg	
SL-868-SA5D-SB-2.5-3.5	SELENIUM	J	0.168	0.398	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0294	0.199	PQL	mg/Kg	

Method: 7471B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-568-SA5D-SB-2.5-3.5	MERCURY	J	0.0123	0.0165	PQL	mg/Kg	J (all detects)
SL-571-SA5D-SB-0.0-0.5	MERCURY	J	0.0160	0.0163	PQL	mg/Kg	J (all detects)

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-583-SA5D-SB-0.0-0.5	EFH (C15-C20)	J	3.6	5.2	PQL	mg/Kg	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8081B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-568-SA5D-SB-0.0-0.5	DELTA-BHC	J	0.51	0.85	PQL	ug/Kg	J (all detects)
	METHOXYCHLOR	J	4.1	6.8	PQL	ug/Kg	
SL-572-SA5D-SB-0.0-0.5	4,4'-DDE	J	0.60	1.7	PQL	ug/Kg	J (all detects)
	4,4'-DDT	J	1.4	1.7	PQL	ug/Kg	
	gamma-BHC (Lindane)	J	0.24	0.83	PQL	ug/Kg	

Method: 8082A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-568-SA5D-SB-0.0-0.5	Aroclor 5460	J	15	34	PQL	ug/Kg	J (all detects)
SL-568-SA5D-SB-2.5-3.5	AROCLOR 1254	J	7.3	18	PQL	ug/Kg	J (all detects)
SL-571-SA5D-SB-0.0-0.5	AROCLOR 1254	J	12	17	PQL	ug/Kg	J (all detects)

Method: 8270D SIM
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-568-SA5D-SB-0.0-0.5	BENZO(A)ANTHRACENE	J	16	17	PQL	ug/Kg	J (all detects)
	BENZO(E)PYRENE	J	39	170	PQL	ug/Kg	
	Butylbenzylphthalate	J	110	180	PQL	ug/Kg	
	INDENO(1,2,3-CD)PYRENE	J	11	17	PQL	ug/Kg	
SL-568-SA5D-SB-2.5-3.5	BENZO(A)ANTHRACENE	J	0.94	1.8	PQL	ug/Kg	J (all detects)
	BENZO(A)PYRENE	J	1.2	1.8	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	0.87	1.8	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	1.2	1.8	PQL	ug/Kg	
	CHRYSENE	J	1.5	1.8	PQL	ug/Kg	
	PHENANTHRENE	J	0.93	1.8	PQL	ug/Kg	
SL-571-SA5D-SB-0.0-0.5	NAPHTHALENE	J	0.79	1.7	PQL	ug/Kg	J (all detects)
SL-574-SA5D-SB-0.0-0.5	NAPHTHALENE	J	0.69	1.7	PQL	ug/Kg	J (all detects)
SL-583-SA5D-SB-0.0-0.5	1-METHYLNAPHTHALENE	J	0.92	1.7	PQL	ug/Kg	J (all detects)
	2-METHYLNAPHTHALENE	J	1.5	1.7	PQL	ug/Kg	
	BENZO(A)ANTHRACENE	J	0.95	1.7	PQL	ug/Kg	
	BENZO(A)PYRENE	J	1.0	1.7	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	1.2	1.7	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	0.84	1.7	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	12	19	PQL	ug/Kg	
	INDENO(1,2,3-CD)PYRENE	J	0.89	1.7	PQL	ug/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH112

Laboratory: LL

EDD Filename: PH112

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-868-SA5D-SB-2.5-3.5	BENZO(A)ANTHRACENE	J	0.82	1.7	PQL	ug/Kg	J (all detects)
	BENZO(A)PYRENE	J	0.98	1.7	PQL	ug/Kg	
	BENZO(B)FLUORANTHENE	J	1.4	1.7	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	1.2	1.7	PQL	ug/Kg	
	Butylbenzylphthalate	J	7.5	18	PQL	ug/Kg	
	CHRYSENE	J	1.1	1.7	PQL	ug/Kg	
	FLUORANTHENE	J	1.4	1.7	PQL	ug/Kg	
	INDENO(1,2,3-CD)PYRENE	J	0.76	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	0.86	1.7	PQL	ug/Kg	
	PYRENE	J	1.4	1.7	PQL	ug/Kg	

LDC #: 3069504

VALIDATION COMPLETENESS WORKSHEET

Date: 11/11/13

SDG #: PH112

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: [Signature]

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	—	Sampling dates: 9/23/13
II.	ICP/MS Tune	—	
III.	Calibration	—	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	—	
VI.	Matrix Spike Analysis	SW	MS/D
VII.	Duplicate Sample Analysis	SW	DUP
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	SW	Zn: 16% (All Soil - J/W/ACA)
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	—	(2,3)
XV.	Field Blanks	SW	EB = EB10912513 TB = FB011613

Note: A = Acceptable ND = No compounds detected D = Duplicate (PH114)
 N = Not provided/applicable R = Rinsate TB = Trip blank (PH032)
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples:

1	SL-568-SA5D-SB-0.0-0.5	11		21		31	
2	SL-568-SA5D-SB-2.5-3.5	12		22		32	
3	SL-868-SA5D-SB-2.5-3.5	13		23		33	
4	SL-571-SA5D-SB-0.0-0.5	14		24		34	
5	SL-583-SA5D-SB-0.0-0.5	15		25		35	
6	SL-568-SA5D-SB-2.5-3.5MS	16		26		36	
7	SL-568-SA5D-SB-2.5-3.5MSD	17		27		37	
8	SL-568-SA5D-SB-2.5-3.5DUP	18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F
Sampling date: 4/11/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification														
	FB-041613 (SDG: PH032)	Action Limit	1	2 ^{gm}												
Mo	0.0132	6.60	1.44	0.388												
Sn	0.0029	1.45														

Sampling date: 9/25/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification														
	EB1-092513 (SDG: PH114)	Action Limit	No Qualifiers													
Ca	0.0498	24.9														

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".



QUALITY ASSURANCE SUMMARY

FORM 5A (MS/MSD)

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

SDG No.: PH112

Matrix: SOIL

Level

(low/med):

LOW

Background Lab Sample ID: 7208982BKG Matrix Spike Lab Sample ID: 7208983MS Matrix Spike Duplicate Lab Sample ID: 7208984MSD
Batch Id(s): P27437A, P27438A

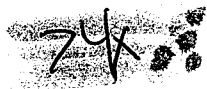
Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Control Limit		M
		Result	C	Result	C	Result	C				%R	Q	%R	Q		%R	RPD	
Aluminum		20503.4170		20234.2723		20248.9573		198.0198	194.1748	MG/KG	-136		-131		0		20	P
Antimony		0.7400	U	21.5891		20.3621		49.5050	48.5437	MG/KG	44	N	42	N	6	75 - 125	20	P
Arsenic		4.0560		18.3941		19.2252		14.8515	14.5631	MG/KG	97		104		4	75 - 125	20	P
Barium		95.6450		281.0584		273.8534		198.0198	194.1748	MG/KG	94		92		3	75 - 125	20	P
Beryllium		0.5650	B	5.3426		5.3204		4.9505	4.8544	MG/KG	97		98		0	75 - 125	20	P
Boron		9.4400	B	200.6564		203.6835		198.0198	194.1748	MG/KG	97		100		1	75 - 125	20	P
Cadmium		0.3210	B	4.9574		5.0456		4.9505	4.8544	MG/KG	94		97		2	75 - 125	20	P
Calcium		37146.4410		44866.6911		46661.0204		396.0396	388.3495	MG/KG	1949		2450		4		20	P
Chromium		31.2820		55.6653		52.4748		19.8020	19.4175	MG/KG	123		109		6	75 - 125	20	P
Cobalt		7.9630		51.7772		51.8155		49.5050	48.5437	MG/KG	89		90		0	75 - 125	20	P
Copper		16.0030		40.0436		39.8757		24.7525	24.2718	MG/KG	97		98		0	75 - 125	20	P
Iron		25969.1730		20878.0931		19658.7447		99.0099	97.0874	MG/KG	-5142		-6500		6		20	P
Lead		7.0820		20.2119		20.3718		14.8515	14.5631	MG/KG	88		91		1	75 - 125	20	P
Lithium		32.9190		122.9356		122.3932		99.0099	97.0874	MG/KG	91		92		0	75 - 125	20	P
Magnesium		7184.0330		6282.9455		6167.3728		198.0198	194.1748	MG/KG	-455		-524		2		20	P
Manganese		265.7460		379.4307		431.6019		49.5050	48.5437	MG/KG	230		342		13		20	P
Mercury		0.0115	B	0.1903		0.1711		0.1555	0.1554	MG/KG	115		103		11	65 - 135	20	CV
Molybdenum		0.1700	U	184.2960		182.8087		198.0198	194.1748	MG/KG	93		94		1	75 - 125	20	P
Nickel		17.4050		61.6040		59.7816		49.5050	48.5437	MG/KG	89		87		3	75 - 125	20	P
Phosphorus		439.1830		610.4188		602.5252		99.0099	97.0874	MG/KG	173		168		1		20	P
Potassium		3366.5940		3843.9644		3983.5971		990.0990	970.8738	MG/KG	48	N	64	N	4	75 - 125	20	P
Selenium	78	0.1832	B	2.2752		2.3786		1.9802	1.9417	MG/KG	106		113		4	75 - 125	20	MS
Silver	107	0.0327	B	11.0396		11.0718		9.9010	9.7087	MG/KG	111		114		0	75 - 125	20	MS
Sodium		187.3580		1115.4762		1097.0592		990.0990	970.8738	MG/KG	94		94		2	75 - 125	20	P
Strontium	88	89.7000		101.1881		97.2816		7.9208	7.7670	MG/KG	145		98		4		20	MS
Thallium	203	0.2830		0.7689		0.7363		0.3960	0.3883	MG/KG	123		117		4	75 - 125	20	MS

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor</p> <p>MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U= Below MDL, B= Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>
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Lancaster
Laboratories



QUALITY ASSURANCE SUMMARY

FORM 5A (MS/MSD)

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

SDG No.: PH112

Matrix: SOIL

Level

(low/med):

LOW

Background Lab Sample ID: 7208982BKG Matrix Spike Lab Sample ID: 7208983MS Matrix Spike Duplicate Lab Sample ID: 7208984MSD

Batch Id(s): P27437A, P27438A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Control Limit			
		Result	C	Result	C	Result	C				%R	Q	%R	Q		%R	RPD	M	
Tin		3.0930	B	335.6079		337.5825		396.0396	388.3495	MG/KG	84		86		1	75 - 125	20	P	
*Titanium		1095.0880		1270.0455		1323.9398		99.0099	97.0874	MG/KG	177		236		4			20	P
Vanadium		56.3670		107.3485		110.4146		49.5050	48.5437	MG/KG	103		111		3	75 - 125	20	P	
Zinc		78.1960		106.6188		110.4631		49.5050	48.5437	MG/KG	57	N	66	N	4	75 - 125	20	P	
Zirconium		4.2450	B	98.3792		100.4874		99.0099	97.0874	MG/KG	95		99		2	75 - 125	20	P	

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor</p> <p>MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U= Below MDL, B= Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>
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Background Lab Sample ID: 7208982BKG
 Batch ID(s): P27437A, P27438A
 Concentration Units: MG/KG

Duplicate Lab Sample ID: 7208985DUP

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			20503.4170		13205.6703		43	*	P
Antimony			0.7400	U	0.7327	U			P
Arsenic		4.0	4.0560		3.2822	B	21		P
Barium			95.6450		88.6614		8		P
Beryllium			0.5650	B	0.4119	B	31		P
Boron			9.4400	B	6.9733	B	30		P
Cadmium			0.3210	B	0.4416	B	32		P
Calcium			37146.4410		42777.6000		14		P
Chromium			31.2820		27.8752		12		P
Cobalt			7.9630		5.8970		30	*	P
Copper			16.0030		13.1168		20		P
Iron			25969.1730		17476.5436		39	*	P
Lead		3.0	7.0820		6.8614		3		P
Lithium		4.0	32.9190		18.4238		56	*	P
Magnesium			7184.0330		5124.0703		33	*	P
Manganese			265.7460		354.7842		29	*	P
Mercury			0.0115	B	0.0111	B	4		CV
Molybdenum			0.1700	U	0.3881	B	200		P
Nickel			17.4050		14.8426		16		P
Phosphorus			439.1830		503.4861		14		P
Potassium			3366.5940		2468.9218		31	*	P
Selenium	78		0.1832	B	0.2380	B	26		MS
Silver	107		0.0327	B	0.0359	B	9		MS
Sodium		100.0	187.3580		146.4515		25		P
Strontium	88		89.7000		84.5050		6		MS
Thallium	203	0.2	0.2830		0.2509		12		MS
Tin			3.0930	B	2.5505	B	19		P
Titanium			1095.0880		652.4723		51	*	P
Vanadium			56.3670		41.7030		30	*	P
Zinc			78.1960		54.7109		35	*	P
Zirconium		5.0	4.2450	B	5.6347		28		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).
 The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

ok by difference

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: Duplicate Out of Spec
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**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH113

Prepared for

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Prepared by

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December 11, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on September 24, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by EPA SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A and 7471B
Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M
TPH as Extractables by EPA SW 846 Method 8015M
Dioxins and Dibenzofurans by EPA Method 1613B

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks, and field duplicates. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed (in SDG PH109) at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pairs for metals and TPH as extractables. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits with the exception of one DUP for metals. The associated sample results were qualified as detected estimated (J). The details regarding the qualification of data are provided in Enclosure I.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met with the following exceptions:

Diluted Sample	Analyte	%D (Limits)	Associated Samples	Flag	A or P
SL-565-SA5D-SB-0.0-0.5	Strontium	24 (≤ 10)	All soil samples in SDG PH113	J (all detects) UJ (all non-detects)	A

The associated sample results were qualified as detected estimated (J).

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH113	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pairs were collected and analyzed for SVOC, PCBs, metals, TPH as extractables, and dioxins. All RPDs were within QC limits except for several SVOCs, metals, TPH as extractables, and dioxins. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No volatile contaminants were found in the trip blank.

One equipment blank (from SDG PH114) was collected and analyzed for SVOCs, PCBs, metals, TPH as gasoline, TPH as extractables and dioxins. The equipment blank had detections for SVOCs, metals and dioxins. The associated sample results were not detected or were

significantly greater than the concentrations found in the equipment blanks, therefore no data were qualified.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, PCBs, metals, TPH as gasoline, TPH as extractables and dioxins. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1

Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5	7211497	N	3050B	6010C	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5	7211497	N	3050B	6020A	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5	7211497	N	3546	8015M	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5	7211497	N	3546	8082A	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5	7211497	N	3546	8270D SIM	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5	7211497	N	METHOD	1613B	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5	7211497	N	METHOD	7471B	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MS	7211498	MS	3050B	6010C	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MS	7211498	MS	3050B	6020A	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MS	7211498	MS	3546	8015M	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MS	7211498	MS	3546	8082A	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MS	7211498	MS	3546	8270D SIM	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MS	7211498	MS	METHOD	1613B	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MS	7211498	MS	METHOD	7471B	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MSD	7211499	MSD	3050B	6010C	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MSD	7211499	MSD	3050B	6020A	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MSD	7211499	MSD	3546	8015M	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MSD	7211499	MSD	3546	8082A	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MSD	7211499	MSD	3546	8270D SIM	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MSD	7211499	MSD	METHOD	1613B	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5MSD	7211499	MSD	METHOD	7471B	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5DUP	7211500	DUP	3050B	6010C	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5DUP	7211500	DUP	3050B	6020A	III
24-Sep-2013	SL-565-SA5D-SB-0.0-0.5DUP	7211500	DUP	METHOD	7471B	III
24-Sep-2013	SL-865-SA5D-SB-0.0-0.5	7211501	FD	3050B	6010C	III
24-Sep-2013	SL-865-SA5D-SB-0.0-0.5	7211501	FD	3050B	6020A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
24-Sep-2013	SL-865-SA5D-SB-0.0-0.5	7211501	FD	3546	8015M	III
24-Sep-2013	SL-865-SA5D-SB-0.0-0.5	7211501	FD	3546	8082A	III
24-Sep-2013	SL-865-SA5D-SB-0.0-0.5	7211501	FD	3546	8270D SIM	III
24-Sep-2013	SL-865-SA5D-SB-0.0-0.5	7211501	FD	METHOD	1613B	III
24-Sep-2013	SL-865-SA5D-SB-0.0-0.5	7211501	FD	METHOD	7471B	III
24-Sep-2013	TB-092413	7211494	TB	5030B	8015M	III
24-Sep-2013	SL-558-SA5D-SB-0.0-0.5	7211495	N	3050B	6010C	III
24-Sep-2013	SL-558-SA5D-SB-0.0-0.5	7211495	N	3050B	6020A	III
24-Sep-2013	SL-558-SA5D-SB-0.0-0.5	7211495	N	3546	8015M	III
24-Sep-2013	SL-558-SA5D-SB-0.0-0.5	7211495	N	3546	8082A	III
24-Sep-2013	SL-558-SA5D-SB-0.0-0.5	7211495	N	3546	8270D SIM	III
24-Sep-2013	SL-558-SA5D-SB-0.0-0.5	7211495	N	METHOD	1613B	III
24-Sep-2013	SL-558-SA5D-SB-0.0-0.5	7211495	N	METHOD	7471B	III
24-Sep-2013	SL-558-SA5D-SB-4.0-5.0	7211496	N	3050B	6010C	III
24-Sep-2013	SL-558-SA5D-SB-4.0-5.0	7211496	N	3050B	6020A	III
24-Sep-2013	SL-558-SA5D-SB-4.0-5.0	7211496	N	3546	8015M	III
24-Sep-2013	SL-558-SA5D-SB-4.0-5.0	7211496	N	3546	8082A	III
24-Sep-2013	SL-558-SA5D-SB-4.0-5.0	7211496	N	3546	8270D SIM	III
24-Sep-2013	SL-558-SA5D-SB-4.0-5.0	7211496	N	5035A	8015M	III
24-Sep-2013	SL-558-SA5D-SB-4.0-5.0	7211496	N	METHOD	7471B	III
24-Sep-2013	SL-557-SA5D-SB-0.0-0.5	7211502	N	3050B	6010C	III
24-Sep-2013	SL-557-SA5D-SB-0.0-0.5	7211502	N	3050B	6020A	III
24-Sep-2013	SL-557-SA5D-SB-0.0-0.5	7211502	N	3546	8015M	III
24-Sep-2013	SL-557-SA5D-SB-0.0-0.5	7211502	N	3546	8082A	III
24-Sep-2013	SL-557-SA5D-SB-0.0-0.5	7211502	N	3546	8270D SIM	III
24-Sep-2013	SL-557-SA5D-SB-0.0-0.5	7211502	N	METHOD	1613B	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
24-Sep-2013	SL-557-SA5D-SB-0.0-0.5	7211502	N	METHOD	7471B	III
24-Sep-2013	SL-566-SA5D-SB-0.0-0.5	7211503	N	3050B	6010C	III
24-Sep-2013	SL-566-SA5D-SB-0.0-0.5	7211503	N	3050B	6020A	III
24-Sep-2013	SL-566-SA5D-SB-0.0-0.5	7211503	N	3546	8015M	III
24-Sep-2013	SL-566-SA5D-SB-0.0-0.5	7211503	N	3546	8082A	III
24-Sep-2013	SL-566-SA5D-SB-0.0-0.5	7211503	N	3546	8270D SIM	III
24-Sep-2013	SL-566-SA5D-SB-0.0-0.5	7211503	N	METHOD	1613B	III
24-Sep-2013	SL-566-SA5D-SB-0.0-0.5	7211503	N	METHOD	7471B	III
24-Sep-2013	SL-566-SA5D-SB-4.0-5.0	7211504	N	3050B	6010C	III
24-Sep-2013	SL-566-SA5D-SB-4.0-5.0	7211504	N	3050B	6020A	III
24-Sep-2013	SL-566-SA5D-SB-4.0-5.0	7211504	N	3546	8015M	III
24-Sep-2013	SL-566-SA5D-SB-4.0-5.0	7211504	N	3546	8082A	III
24-Sep-2013	SL-566-SA5D-SB-4.0-5.0	7211504	N	3546	8270D SIM	III
24-Sep-2013	SL-566-SA5D-SB-4.0-5.0	7211504	N	5035A	8015M	III
24-Sep-2013	SL-566-SA5D-SB-4.0-5.0	7211504	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-557-SA5D-SB-0.0-0.5	Collected: 9/24/2013 9:35:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.17	U	0.771	MDL	4.17	PQL	mg/Kg	UJ	Q
ARSENIC	2.79	J	0.729	MDL	4.17	PQL	mg/Kg	J	Z
BERYLLIUM	0.843	J	0.0698	MDL	1.04	PQL	mg/Kg	J	Z
CADMIUM	0.965	J	0.0792	MDL	1.04	PQL	mg/Kg	J	Z
COBALT	10.3		0.103	MDL	1.04	PQL	mg/Kg	J	E
TIN	2.91	J	0.229	MDL	10.4	PQL	mg/Kg	U	B
POTASSIUM	4870		8.69	MDL	104	PQL	mg/Kg	J	Q

Sample ID: SL-558-SA5D-SB-0.0-0.5	Collected: 9/24/2013 8:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.17	U	0.772	MDL	4.17	PQL	mg/Kg	UJ	Q
ARSENIC	3.27	J	0.730	MDL	4.17	PQL	mg/Kg	J	Z
BERYLLIUM	0.686	J	0.0699	MDL	1.04	PQL	mg/Kg	J	Z
COBALT	8.29		0.103	MDL	1.04	PQL	mg/Kg	J	E
MOLYBDENUM	0.198	J	0.177	MDL	2.09	PQL	mg/Kg	U	F
POTASSIUM	6280		8.70	MDL	104	PQL	mg/Kg	J	Q
SODIUM	92.0	J	17.4	MDL	104	PQL	mg/Kg	J	Z
TIN	2.34	J	0.230	MDL	10.4	PQL	mg/Kg	U	B
Zirconium	5.19	J	0.876	MDL	5.22	PQL	mg/Kg	J	Z

Sample ID: SL-558-SA5D-SB-4.0-5.0	Collected: 9/24/2013 8:45:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.27	U	0.790	MDL	4.27	PQL	mg/Kg	UJ	Q
ARSENIC	1.72	J	0.747	MDL	4.27	PQL	mg/Kg	J	Z
BERYLLIUM	0.476	J	0.0715	MDL	1.07	PQL	mg/Kg	J	Z
CADMIUM	0.847	J	0.0811	MDL	1.07	PQL	mg/Kg	J	Z
COBALT	4.95		0.106	MDL	1.07	PQL	mg/Kg	J	E
POTASSIUM	2740		8.90	MDL	107	PQL	mg/Kg	J	Q
SODIUM	102	J	17.8	MDL	107	PQL	mg/Kg	J	Z
TIN	2.08	J	0.235	MDL	10.7	PQL	mg/Kg	U	B
Zirconium	4.44	J	0.896	MDL	5.34	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-565-SA5D-SB-0.0-0.5	Collected: 9/24/2013 7:40:00	Analysis Type: REA	Dilution: 5						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	55000		19.0	MDL	210	PQL	mg/Kg	J	FD

Sample ID: SL-565-SA5D-SB-0.0-0.5	Collected: 9/24/2013 7:40:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.21	U	0.779	MDL	4.21	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.858	J	0.0705	MDL	1.05	PQL	mg/Kg	J	Z
BORON	3.09	J	0.884	MDL	10.5	PQL	mg/Kg	J	Z, FD
CADMIUM	0.840	J	0.0800	MDL	1.05	PQL	mg/Kg	J	Z
COBALT	9.18		0.104	MDL	1.05	PQL	mg/Kg	J	E
POTASSIUM	4120		8.78	MDL	105	PQL	mg/Kg	J	Q
TIN	2.63	J	0.231	MDL	10.5	PQL	mg/Kg	U	B

Sample ID: SL-566-SA5D-SB-0.0-0.5	Collected: 9/24/2013 12:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.08	U	0.755	MDL	4.08	PQL	mg/Kg	UJ	Q
ARSENIC	2.79	J	0.714	MDL	4.08	PQL	mg/Kg	J	Z
BERYLLIUM	0.913	J	0.0683	MDL	1.02	PQL	mg/Kg	J	Z
COBALT	10.5		0.101	MDL	1.02	PQL	mg/Kg	J	E
POTASSIUM	4780		8.50	MDL	102	PQL	mg/Kg	J	Q
TIN	2.88	J	0.224	MDL	10.2	PQL	mg/Kg	U	B

Sample ID: SL-566-SA5D-SB-4.0-5.0	Collected: 9/24/2013 1:15:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.21	U	0.779	MDL	4.21	PQL	mg/Kg	UJ	Q
ARSENIC	3.14	J	0.737	MDL	4.21	PQL	mg/Kg	J	Z
BERYLLIUM	0.642	J	0.0706	MDL	1.05	PQL	mg/Kg	J	Z
CADMIUM	0.947	J	0.0800	MDL	1.05	PQL	mg/Kg	J	Z
COBALT	8.80		0.104	MDL	1.05	PQL	mg/Kg	J	E
POTASSIUM	4610		8.78	MDL	105	PQL	mg/Kg	J	Q
TIN	3.07	J	0.232	MDL	10.5	PQL	mg/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-865-SA5D-SB-0.0-0.5 Collected: 9/24/2013 7:50:00 Analysis Type: REA2 Dilution: 5

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	32700		18.9	MDL	209	PQL	mg/Kg	J	FD

Sample ID: SL-865-SA5D-SB-0.0-0.5 Collected: 9/24/2013 7:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.18	U	0.773	MDL	4.18	PQL	mg/Kg	UJ	Q
ARSENIC	4.16	J	0.731	MDL	4.18	PQL	mg/Kg	J	Z
BERYLLIUM	0.692	J	0.0700	MDL	1.04	PQL	mg/Kg	J	Z
BORON	23.9		0.877	MDL	10.4	PQL	mg/Kg	J	FD
COBALT	7.91		0.103	MDL	1.04	PQL	mg/Kg	J	E
POTASSIUM	4490		8.71	MDL	104	PQL	mg/Kg	J	Q
TIN	2.63	J	0.230	MDL	10.4	PQL	mg/Kg	U	B

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-557-SA5D-SB-0.0-0.5 Collected: 9/24/2013 9:35:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.261	J	0.104	MDL	0.417	PQL	mg/Kg	J	Z

Sample ID: SL-557-SA5D-SB-0.0-0.5 Collected: 9/24/2013 9:35:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0358	J	0.0271	MDL	0.208	PQL	mg/Kg	J	Z
STRONTIUM	40.8		0.0708	MDL	0.417	PQL	mg/Kg	J	E, A

Sample ID: SL-558-SA5D-SB-0.0-0.5 Collected: 9/24/2013 8:30:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.354	J	0.104	MDL	0.417	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-558-SA5D-SB-0.0-0.5			Collected: 9/24/2013 8:30:00			Analysis Type: RES		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0392	J	0.0271	MDL	0.209	PQL	mg/Kg	J	Z
STRONTIUM	75.9		0.0709	MDL	0.417	PQL	mg/Kg	J	E, A

Sample ID: SL-558-SA5D-SB-4.0-5.0			Collected: 9/24/2013 8:45:00			Analysis Type: REA		Dilution: 5	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	98.5		0.181	MDL	1.07	PQL	mg/Kg	J	E, A

Sample ID: SL-558-SA5D-SB-4.0-5.0			Collected: 9/24/2013 8:45:00			Analysis Type: REA2		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.251	J	0.107	MDL	0.427	PQL	mg/Kg	J	Z

Sample ID: SL-558-SA5D-SB-4.0-5.0			Collected: 9/24/2013 8:45:00			Analysis Type: RES		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
THALLIUM	0.183	J	0.0320	MDL	0.213	PQL	mg/Kg	J	Z

Sample ID: SL-565-SA5D-SB-0.0-0.5			Collected: 9/24/2013 7:40:00			Analysis Type: REA		Dilution: 5	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	114		0.179	MDL	1.05	PQL	mg/Kg	J	E, A

Sample ID: SL-565-SA5D-SB-0.0-0.5			Collected: 9/24/2013 7:40:00			Analysis Type: REA2		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.166	J	0.105	MDL	0.421	PQL	mg/Kg	J	Z

Sample ID: SL-566-SA5D-SB-0.0-0.5			Collected: 9/24/2013 12:30:00			Analysis Type: REA2		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.386	J	0.102	MDL	0.408	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-566-SA5D-SB-0.0-0.5 Collected: 9/24/2013 12:30:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0384	J	0.0265	MDL	0.204	PQL	mg/Kg	J	Z
STRONTIUM	38.9		0.0693	MDL	0.408	PQL	mg/Kg	J	E, A

Sample ID: SL-566-SA5D-SB-4.0-5.0 Collected: 9/24/2013 1:15:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.116	J	0.105	MDL	0.421	PQL	mg/Kg	J	Z

Sample ID: SL-566-SA5D-SB-4.0-5.0 Collected: 9/24/2013 1:15:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0311	J	0.0274	MDL	0.211	PQL	mg/Kg	J	Z
STRONTIUM	30.0		0.0716	MDL	0.421	PQL	mg/Kg	J	E, A

Sample ID: SL-865-SA5D-SB-0.0-0.5 Collected: 9/24/2013 7:50:00 Analysis Type: REA Dilution: 5

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	169		0.178	MDL	1.04	PQL	mg/Kg	J	E, A

Sample ID: SL-865-SA5D-SB-0.0-0.5 Collected: 9/24/2013 7:50:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.266	J	0.104	MDL	0.418	PQL	mg/Kg	J	Z

Method Category:	METALS	
Method:	7471B	Matrix: SO

Sample ID: SL-565-SA5D-SB-0.0-0.5 Collected: 9/24/2013 7:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0140	J	0.0103	MDL	0.0171	PQL	mg/Kg	J	Z, FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	7471B	Matrix: SO

Sample ID: SL-566-SA5D-SB-4.0-5.0	Collected: 9/24/2013 1:15:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0132	J	0.0104	MDL	0.0174	PQL	mg/Kg	J	Z

Sample ID: SL-865-SA5D-SB-0.0-0.5	Collected: 9/24/2013 7:50:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0292		0.0101	MDL	0.0168	PQL	mg/Kg	J	FD

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-557-SA5D-SB-0.0-0.5	Collected: 9/24/2013 9:35:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	0.844	JB	0.0142	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.120	JBQ	0.0227	MDL	5.11	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.155	JB	0.0377	MDL	5.11	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.180	JBQ	0.0345	MDL	5.11	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.464	JBQ	0.0396	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.192	JB	0.0327	MDL	5.11	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.505	JBQ	0.0356	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.345	JB	0.0392	MDL	5.11	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.166	JBQ	0.0408	MDL	5.11	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.413	JB	0.0352	MDL	5.11	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.181	JB	0.0333	MDL	5.11	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.461	JBQ	0.0365	MDL	5.11	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.442	J	0.0831	MDL	1.02	PQL	ng/Kg	J	Z
OCDF	1.86	JB	0.0413	MDL	10.2	PQL	ng/Kg	J	Z

Sample ID: SL-558-SA5D-SB-0.0-0.5	Collected: 9/24/2013 8:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	2.62	JB	0.0235	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.239	JBQ	0.0341	MDL	5.20	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.267	JBQ	0.0483	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.261	JBQ	0.0352	MDL	5.20	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-558-SA5D-SB-0.0-0.5 Collected: 9/24/2013 8:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,6,7,8-HXCDD	0.555	JB	0.0501	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.195	JBQ	0.0333	MDL	5.20	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.324	JB	0.0470	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.178	JB	0.0376	MDL	5.20	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.248	JBQ	0.0381	MDL	5.20	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.320	JB	0.0290	MDL	5.20	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.250	JB	0.0334	MDL	5.20	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.224	JBQ	0.0274	MDL	5.20	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0701	JQ	0.0554	MDL	1.04	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.197	JQ	0.0501	MDL	1.04	PQL	ng/Kg	J	Z

Sample ID: SL-565-SA5D-SB-0.0-0.5 Collected: 9/24/2013 7:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.477	JBQ	0.0292	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.219	JBQ	0.0136	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.124	JBQ	0.0236	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.136	JBQ	0.0264	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.212	JB	0.0226	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.114	JB	0.0277	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.206	JB	0.0203	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.134	JBQ	0.0255	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.153	JBQ	0.0243	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.254	JBQ	0.0335	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.379	JBQ	0.0202	MDL	5.25	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.155	JBQ	0.0200	MDL	5.25	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.223	JB	0.0200	MDL	5.25	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0976	J	0.0523	MDL	1.05	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0682	JQ	0.0439	MDL	1.05	PQL	ng/Kg	J	Z
OCDD	2.22	JB	0.0373	MDL	10.5	PQL	ng/Kg	J	Z, FD
OCDF	0.385	JB	0.0465	MDL	10.5	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-566-SA5D-SB-0.0-0.5 Collected: 9/24/2013 12:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	2.02	JB	0.0236	MDL	5.06	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.360	JBQ	0.0117	MDL	5.06	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.102	JBQ	0.0240	MDL	5.06	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0826	JBQ	0.0260	MDL	5.06	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.128	JB	0.0220	MDL	5.06	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.276	JBQ	0.0277	MDL	5.06	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.100	JBQ	0.0187	MDL	5.06	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.306	JB	0.0271	MDL	5.06	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDF	0.356	JB	0.0252	MDL	5.06	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0695	JBQ	0.0303	MDL	5.06	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.256	JB	0.0185	MDL	5.06	PQL	ng/Kg	U	B
2,3,4,6,7,8-HxCDF	0.135	JBQ	0.0214	MDL	5.06	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.218	JBQ	0.0183	MDL	5.06	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.116	JQ	0.0417	MDL	1.01	PQL	ng/Kg	J	Z
OCDF	0.857	JB	0.0367	MDL	10.1	PQL	ng/Kg	J	Z

Sample ID: SL-865-SA5D-SB-0.0-0.5 Collected: 9/24/2013 7:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.722	JB	0.0295	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.267	JB	0.0137	MDL	5.03	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.147	JB	0.0197	MDL	5.03	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0869	JBQ	0.0280	MDL	5.03	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.146	JB	0.0237	MDL	5.03	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.144	JBQ	0.0304	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.143	JB	0.0223	MDL	5.03	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.146	JBQ	0.0284	MDL	5.03	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDF	0.175	JB	0.0235	MDL	5.03	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.157	JBQ	0.0371	MDL	5.03	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.275	JBQ	0.0235	MDL	5.03	PQL	ng/Kg	U	B
2,3,4,6,7,8-HxCDF	0.159	JB	0.0237	MDL	5.03	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.241	JB	0.0229	MDL	5.03	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.105	JQ	0.0633	MDL	1.01	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0816	JQ	0.0522	MDL	1.01	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-865-SA5D-SB-0.0-0.5	Collected: 9/24/2013 7:50:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
OCDD	4.69	JB	0.0276	MDL	10.1	PQL	ng/Kg	J	Z, FD
OCDF	0.582	JBQ	0.0391	MDL	10.1	PQL	ng/Kg	U	B

Method Category:	SVOA	
Method:	8015M	Matrix: SO

Sample ID: SL-565-SA5D-SB-0.0-0.5	Collected: 9/24/2013 7:40:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	4.3	J	2.1	MDL	5.3	PQL	mg/Kg	J	Z, Q, Q, Q, FD
EFH (C30-C40)	13		4.2	MDL	11	PQL	mg/Kg	J	Q, Q, Q, FD

Sample ID: SL-566-SA5D-SB-0.0-0.5	Collected: 9/24/2013 12:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	4.9	J	2.1	MDL	5.1	PQL	mg/Kg	J	Z

Sample ID: SL-865-SA5D-SB-0.0-0.5	Collected: 9/24/2013 7:50:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	8.7		2.1	MDL	5.2	PQL	mg/Kg	J	FD
EFH (C30-C40)	29		4.2	MDL	10	PQL	mg/Kg	J	FD

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-557-SA5D-SB-0.0-0.5	Collected: 9/24/2013 9:35:00	Analysis Type: RES-BASE/NEUTRAL	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.73	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
CHRYSENE	1.5	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
FLUORANTHENE	1.3	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
NAPHTHALENE	1.1	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
PHENANTHRENE	1.1	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	1.3	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-558-SA5D-SB-0.0-0.5 Collected: 9/24/2013 8:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	1.2	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(B)FLUORANTHENE	1.7	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	0.86	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	0.79	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	7.1	J	6.3	MDL	19	PQL	ug/Kg	J	Z
Butylbenzylphthalate	10	J	6.3	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	1.7	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z
FLUORANTHENE	1.4	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	1.6	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
PHENANTHRENE	1.2	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
PYRENE	1.3	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-565-SA5D-SB-0.0-0.5 Collected: 9/24/2013 7:40:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BIS(2-ETHYLHEXYL)PHTHALATE	19	U	6.3	MDL	19	PQL	ug/Kg	UJ	FD
NAPHTHALENE	1.7	U	0.70	MDL	1.7	PQL	ug/Kg	UJ	FD

Sample ID: SL-566-SA5D-SB-0.0-0.5 Collected: 9/24/2013 12:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	1.1	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
2-METHYLNAPHTHALENE	1.5	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
ACENAPHTHYLENE	0.45	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
ANTHRACENE	0.70	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(B)FLUORANTHENE	1.3	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	9.1	J	6.1	MDL	18	PQL	ug/Kg	J	Z
Butylbenzylphthalate	10	J	6.1	MDL	18	PQL	ug/Kg	J	Z
CHRYSENE	1.1	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
FLUORANTHENE	1.1	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
FLUORENE	0.70	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
PHENANTHRENE	1.2	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	1.1	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-566-SA5D-SB-4.0-5.0 Collected: 9/24/2013 1:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	0.91	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-865-SA5D-SB-0.0-0.5 Collected: 9/24/2013 7:50:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BIS(2-ETHYLHEXYL)PHTHALATE	11	J	6.3	MDL	19	PQL	ug/Kg	J	Z, FD
NAPHTHALENE	0.71	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z, FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
*#	Professional Judgment
A	ICP Serial Dilution
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Field Blank Contamination
FD	Field Duplicate Precision
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I

EPA Level III ADR Outliers

(Including Manual Review Outliers)

Quality Control Outlier Reports

PH113

Method Blank Outlier Report

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2730B371140	10/2/2013 11:40:00 AM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDD 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDD 1,2,3,7,8,9-HxCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HxCDF 2,3,4,7,8-PECDF OCDD OCDF	0.0965 ng/Kg 0.0595 ng/Kg 0.0657 ng/Kg 0.0334 ng/Kg 0.0515 ng/Kg 0.0256 ng/Kg 0.0453 ng/Kg 0.0369 ng/Kg 0.0754 ng/Kg 0.0738 ng/Kg 0.0876 ng/Kg 0.0332 ng/Kg 0.0646 ng/Kg 0.249 ng/Kg 0.137 ng/Kg	SL-557-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-0.0-0.5 SL-565-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-0.0-0.5 SL-865-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-557-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.120 ng/Kg	0.120U ng/Kg
SL-557-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.155 ng/Kg	0.155U ng/Kg
SL-557-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.180 ng/Kg	0.180U ng/Kg
SL-557-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HxCDF	0.192 ng/Kg	0.192U ng/Kg
SL-557-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HxCDF	0.345 ng/Kg	0.345U ng/Kg
SL-557-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.166 ng/Kg	0.166U ng/Kg
SL-557-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.413 ng/Kg	0.413U ng/Kg
SL-558-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.239 ng/Kg	0.239U ng/Kg
SL-558-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HxCDF	0.195 ng/Kg	0.195U ng/Kg
SL-558-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HxCDF	0.178 ng/Kg	0.178U ng/Kg
SL-558-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.248 ng/Kg	0.248U ng/Kg
SL-558-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.320 ng/Kg	0.320U ng/Kg
SL-558-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.224 ng/Kg	0.224U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.477 ng/Kg	0.477U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.219 ng/Kg	0.219U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.124 ng/Kg	0.124U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.136 ng/Kg	0.136U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.212 ng/Kg	0.212U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HxCDD	0.114 ng/Kg	0.114U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HxCDF	0.206 ng/Kg	0.206U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HxCDD	0.134 ng/Kg	0.134U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HxCDF	0.153 ng/Kg	0.153U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.254 ng/Kg	0.254U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.379 ng/Kg	0.379U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HxCDF	0.155 ng/Kg	0.155U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.223 ng/Kg	0.223U ng/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	OCDF	0.385 ng/Kg	0.385U ng/Kg
SL-566-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.102 ng/Kg	0.102U ng/Kg
SL-566-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0826 ng/Kg	0.0826U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-566-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.128 ng/Kg	0.128U ng/Kg
SL-566-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.100 ng/Kg	0.100U ng/Kg
SL-566-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.356 ng/Kg	0.356U ng/Kg
SL-566-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.0695 ng/Kg	0.0695U ng/Kg
SL-566-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.256 ng/Kg	0.256U ng/Kg
SL-566-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.135 ng/Kg	0.135U ng/Kg
SL-566-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.218 ng/Kg	0.218U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.267 ng/Kg	0.267U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.147 ng/Kg	0.147U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0869 ng/Kg	0.0869U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.146 ng/Kg	0.146U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.143 ng/Kg	0.143U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.146 ng/Kg	0.146U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.175 ng/Kg	0.175U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.157 ng/Kg	0.157U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.275 ng/Kg	0.275U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.159 ng/Kg	0.159U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.241 ng/Kg	0.241U ng/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	OCDF	0.582 ng/Kg	0.582U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P26837AB221826	9/27/2013 6:26:00 PM	CALCIUM TIN ZINC	7.61 mg/Kg 1.45 mg/Kg 0.570 mg/Kg	SL-557-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-4.0-5.0 SL-565-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-4.0-5.0 SL-865-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-557-SA5D-SB-0.0-0.5(RES)	TIN	2.91 mg/Kg	2.91U mg/Kg
SL-558-SA5D-SB-0.0-0.5(RES)	TIN	2.34 mg/Kg	2.34U mg/Kg
SL-558-SA5D-SB-4.0-5.0(RES)	TIN	2.08 mg/Kg	2.08U mg/Kg
SL-565-SA5D-SB-0.0-0.5(RES)	TIN	2.63 mg/Kg	2.63U mg/Kg
SL-566-SA5D-SB-0.0-0.5(RES)	TIN	2.88 mg/Kg	2.88U mg/Kg
SL-566-SA5D-SB-4.0-5.0(RES)	TIN	3.07 mg/Kg	3.07U mg/Kg
SL-865-SA5D-SB-0.0-0.5(RES)	TIN	2.63 mg/Kg	2.63U mg/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Field Blank Outlier Report

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041613(REA2/TOT)	4/16/2013 3:15:00 PM	MOLYBDENUM TIN	0.0132 mg/L 0.0029 mg/L	SL-557-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-4.0-5.0 SL-565-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-4.0-5.0 SL-865-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-558-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	0.198 mg/Kg	0.198U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8015M
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-565-SA5D-SB-0.0-0.5MSD (SL-565-SA5D-SB-0.0-0.5)	EFH (C12-C14) EFH (C15-C20) EFH (C8-C11)	- - -	- 124 -	49.00-123.00 49.00-123.00 49.00-123.00	30 (20.00) 86 (20.00) 32 (20.00)	EFH (C12-C14) EFH (C15-C20) EFH (C8-C11)	J (all detects)
SL-565-SA5D-SB-0.0-0.5MS SL-565-SA5D-SB-0.0-0.5MSD (SL-565-SA5D-SB-0.0-0.5)	EFH (C30-C40)	-39	354	49.00-123.00	167 (20.00)	EFH (C30-C40)	J(all detects) R(all non-detects)
SL-565-SA5D-SB-0.0-0.5MS SL-565-SA5D-SB-0.0-0.5MSD (SL-565-SA5D-SB-0.0-0.5)	EFH (C21-C30)	11	170	49.00-123.00	130 (20.00)	EFH (C21-C30)	J(all detects) UJ(all non-detects)

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-565-SA5D-SB-0.0-0.5MS (TOT) SL-565-SA5D-SB-0.0-0.5MSD (TOT) (SL-557-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-4.0-5.0 SL-565-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-4.0-5.0 SL-865-SA5D-SB-0.0-0.5)	ALUMINUM CALCIUM MAGNESIUM PHOSPHORUS POTASSIUM TITANIUM	3790 5761 266 165 207 790	4255 2417 312 173 193 747	75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00	- - - - - -	ALUMINUM CALCIUM MAGNESIUM PHOSPHORUS POTASSIUM TITANIUM	J(all detects) Al, Ca, Mg, P, Ti, No Qual, >4x
SL-565-SA5D-SB-0.0-0.5MS (TOT) SL-565-SA5D-SB-0.0-0.5MSD (TOT) (SL-557-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-4.0-5.0 SL-565-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-4.0-5.0 SL-865-SA5D-SB-0.0-0.5)	ANTIMONY IRON	26 -20245	28 -21138	75.00-125.00 75.00-125.00	- -	ANTIMONY IRON	J(all detects) UJ(all non-detects) Fe, No Qual, >4x Sb, Post Spike = 95%
SL-565-SA5D-SB-0.0-0.5MS (TOT) SL-565-SA5D-SB-0.0-0.5MSD (TOT) (SL-557-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-4.0-5.0 SL-565-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-4.0-5.0 SL-865-SA5D-SB-0.0-0.5)	MANGANESE	150	68	75.00-125.00	-	MANGANESE	No Qual, >4x

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6020A

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-565-SA5D-SB-0.0-0.5MS (TOT) SL-565-SA5D-SB-0.0-0.5MSD (TOT) (SL-557-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-0.0-0.5 SL-558-SA5D-SB-4.0-5.0 SL-565-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-0.0-0.5 SL-566-SA5D-SB-4.0-5.0 SL-865-SA5D-SB-0.0-0.5)	STRONTIUM	1026	638	75.00-125.00	-	STRONTIUM	No Qual, >4x

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-565-SA5D-SB-0.0-0.5DUP (TOT)	ARSENIC	26	20.00	J (all detects) UJ (all non-detects)
(SL-557-SA5D-SB-0.0-0.5	BORON	116	20.00	
SL-558-SA5D-SB-0.0-0.5	COBALT	23	20.00	As, B, No Qual, OK by Difference
SL-558-SA5D-SB-4.0-5.0				
SL-565-SA5D-SB-0.0-0.5				
SL-566-SA5D-SB-0.0-0.5				
SL-566-SA5D-SB-4.0-5.0				
SL-865-SA5D-SB-0.0-0.5)				

Method: 6020A
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-565-SA5D-SB-0.0-0.5DUP (TOT)	SELENIUM	26	20.00	J(all detects) UJ(all non-detects)
(SL-557-SA5D-SB-0.0-0.5	STRONTIUM	23	20.00	
SL-558-SA5D-SB-0.0-0.5				Se, No Qual, OK by Difference
SL-558-SA5D-SB-4.0-5.0				
SL-565-SA5D-SB-0.0-0.5				
SL-566-SA5D-SB-0.0-0.5				
SL-566-SA5D-SB-4.0-5.0				
SL-865-SA5D-SB-0.0-0.5)				

Field Duplicate RPD Report

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 160.3M

Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-565-SA5D-SB-0.0-0.5	SL-865-SA5D-SB-0.0-0.5			
MOISTURE	5.9	5.2	13		No Qualifiers Applied

Method: 1613B

Matrix: SO

Analyte	Concentration (ng/Kg)		Sample RPD	eQAPP RPD	Flag	
	SL-565-SA5D-SB-0.0-0.5	SL-865-SA5D-SB-0.0-0.5				
1,2,3,4,6,7,8-HPCDD	0.477	0.722	41	50.00	No Qualifiers Applied	
1,2,3,4,6,7,8-HPCDF	0.219	0.267	20	50.00		
1,2,3,4,7,8,9-HPCDF	0.124	0.147	17	50.00		
1,2,3,4,7,8-HxCDD	0.136	0.0869	44	50.00		
1,2,3,4,7,8-HxCDF	0.212	0.146	37	50.00		
1,2,3,6,7,8-HxCDD	0.114	0.144	23	50.00		
1,2,3,6,7,8-HxCDF	0.206	0.143	36	50.00		
1,2,3,7,8,9-HxCDD	0.134	0.146	9	50.00		
1,2,3,7,8,9-HxCDF	0.153	0.175	13	50.00		
1,2,3,7,8-PECDD	0.254	0.157	47	50.00		
1,2,3,7,8-PECDF	0.379	0.275	32	50.00		
2,3,4,6,7,8-HxCDF	0.155	0.159	3	50.00		
2,3,4,7,8-PECDF	0.223	0.241	8	50.00		
2,3,7,8-TCDD	0.0976	0.105	7	50.00		
2,3,7,8-TCDF	0.0682	0.0816	18	50.00		
OCDF	0.385	0.582	41	50.00		
OCDD	2.22	4.69	71	50.00		J(all detects)

Method: 6010C

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-565-SA5D-SB-0.0-0.5 (TOT)	SL-865-SA5D-SB-0.0-0.5 (TOT)			
ALUMINUM	22700	20400	11	50.00	No Qualifiers Applied
ARSENIC	4.92	4.16	17	50.00	
BARIUM	149	151	1	50.00	
BERYLLIUM	0.858	0.692	21	50.00	
CADMIUM	0.840	1.19	34	50.00	
CALCIUM	52200	61000	16	50.00	
CHROMIUM	30.9	29.6	4	50.00	
COBALT	9.18	7.91	15	50.00	
COPPER	30.5	25.7	17	50.00	
LEAD	11.2	16.3	37	50.00	
LITHIUM	25.5	22.5	12	50.00	
MAGNESIUM	8530	7520	13	50.00	
MANGANESE	275	263	4	50.00	
NICKEL	22.2	18.8	17	50.00	
PHOSPHORUS	427	533	22	50.00	
POTASSIUM	4120	4490	9	50.00	
SODIUM	134	131	2	50.00	
TIN	2.63	2.63	0	50.00	
TITANIUM	924	914	1	50.00	
VANADIUM	65.0	59.1	10	50.00	
ZINC	85.3	83.0	3	50.00	
Zirconium	6.59	6.15	7	50.00	
BORON	3.09	23.9	154	50.00	J(all detects)
IRON	55000	32700	51	50.00	

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Field Duplicate RPD Report

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6020A
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-565-SA5D-SB-0.0-0.5 (TOT)	SL-865-SA5D-SB-0.0-0.5 (TOT)			
SELENIUM	0.166	0.266	46	50.00	No Qualifiers Applied
STRONTIUM	114	169	39	50.00	
THALLIUM	0.341	0.319	7	50.00	

Method: 7471B
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-565-SA5D-SB-0.0-0.5 (TOT)	SL-865-SA5D-SB-0.0-0.5 (TOT)			
MERCURY	0.0140	0.0292	70	50.00	J(all detects)

Method: 8015M
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-565-SA5D-SB-0.0-0.5	SL-865-SA5D-SB-0.0-0.5			
EFH (C21-C30)	4.3	8.7	68	50.00	J(all detects)
EFH (C30-C40)	13	29	76	50.00	

Method: 8270D SIM
Matrix: SO

Analyte	Concentration (ug/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-565-SA5D-SB-0.0-0.5	SL-865-SA5D-SB-0.0-0.5			
BIS(2-ETHYLHEXYL)PHTHALATE	19 U	11	200	50.00	J(all detects) UJ(all non-detects)
NAPHTHALENE	1.7 U	0.71	200	50.00	

Method: 9045M
Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-565-SA5D-SB-0.0-0.5	SL-865-SA5D-SB-0.0-0.5			
PH	7.54	7.72	2	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-557-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	0.844	5.11	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JBQ	0.120	5.11	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.155	5.11	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.180	5.11	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.464	5.11	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.192	5.11	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.505	5.11	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.345	5.11	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.166	5.11	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.413	5.11	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.181	5.11	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.461	5.11	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.442	1.02	PQL	ng/Kg	
	OCDF	JB	1.86	10.2	PQL	ng/Kg	
SL-558-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	2.62	5.20	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JBQ	0.239	5.20	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.267	5.20	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.261	5.20	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.555	5.20	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.195	5.20	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.324	5.20	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.178	5.20	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.248	5.20	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.320	5.20	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.250	5.20	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.224	5.20	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0701	1.04	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.197	1.04	PQL	ng/Kg	
SL-565-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.477	5.25	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.219	5.25	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.124	5.25	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.136	5.25	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.212	5.25	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.114	5.25	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.206	5.25	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.134	5.25	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.153	5.25	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.254	5.25	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.379	5.25	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.155	5.25	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.223	5.25	PQL	ng/Kg	
	2,3,7,8-TCDD	J	0.0976	1.05	PQL	ng/Kg	
2,3,7,8-TCDF	JQ	0.0682	1.05	PQL	ng/Kg		
OCDD	JB	2.22	10.5	PQL	ng/Kg		
OCDF	JB	0.385	10.5	PQL	ng/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-566-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	2.02	5.06	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.360	5.06	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.102	5.06	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0826	5.06	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.128	5.06	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.276	5.06	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.100	5.06	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.306	5.06	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.356	5.06	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.0695	5.06	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.256	5.06	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.135	5.06	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.218	5.06	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.116	1.01	PQL	ng/Kg	
	OCDF	JB	0.857	10.1	PQL	ng/Kg	
SL-865-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	0.722	5.03	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.267	5.03	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.147	5.03	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0869	5.03	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.146	5.03	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.144	5.03	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.143	5.03	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JBQ	0.146	5.03	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.175	5.03	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.157	5.03	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.275	5.03	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.159	5.03	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.241	5.03	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.105	1.01	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0816	1.01	PQL	ng/Kg	
OCDD	JB	4.69	10.1	PQL	ng/Kg		
OCDF	JBQ	0.582	10.1	PQL	ng/Kg		

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-557-SA5D-SB-0.0-0.5	ARSENIC	J	2.79	4.17	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.843	1.04	PQL	mg/Kg	
	CADMIUM	J	0.965	1.04	PQL	mg/Kg	
	TIN	J	2.91	10.4	PQL	mg/Kg	
SL-558-SA5D-SB-0.0-0.5	ARSENIC	J	3.27	4.17	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.686	1.04	PQL	mg/Kg	
	MOLYBDENUM	J	0.198	2.09	PQL	mg/Kg	
	SODIUM	J	92.0	104	PQL	mg/Kg	
	TIN	J	2.34	10.4	PQL	mg/Kg	
Zirconium	J	5.19	5.22	PQL	mg/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-558-SA5D-SB-4.0-5.0	ARSENIC	J	1.72	4.27	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.476	1.07	PQL	mg/Kg	
	CADMIUM	J	0.847	1.07	PQL	mg/Kg	
	SODIUM	J	102	107	PQL	mg/Kg	
	TIN	J	2.08	10.7	PQL	mg/Kg	
SL-565-SA5D-SB-0.0-0.5	Zirconium	J	4.44	5.34	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.858	1.05	PQL	mg/Kg	
SL-565-SA5D-SB-0.0-0.5	BORON	J	3.09	10.5	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.840	1.05	PQL	mg/Kg	
	TIN	J	2.63	10.5	PQL	mg/Kg	
SL-566-SA5D-SB-0.0-0.5	ARSENIC	J	2.79	4.08	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.913	1.02	PQL	mg/Kg	
	TIN	J	2.88	10.2	PQL	mg/Kg	
SL-566-SA5D-SB-4.0-5.0	ARSENIC	J	3.14	4.21	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.642	1.05	PQL	mg/Kg	
	CADMIUM	J	0.947	1.05	PQL	mg/Kg	
	TIN	J	3.07	10.5	PQL	mg/Kg	
SL-865-SA5D-SB-0.0-0.5	ARSENIC	J	4.16	4.18	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.692	1.04	PQL	mg/Kg	
	TIN	J	2.63	10.4	PQL	mg/Kg	

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-557-SA5D-SB-0.0-0.5	SELENIUM	J	0.261	0.417	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0358	0.208	PQL	mg/Kg	
SL-558-SA5D-SB-0.0-0.5	SELENIUM	J	0.354	0.417	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0392	0.209	PQL	mg/Kg	
SL-558-SA5D-SB-4.0-5.0	SELENIUM	J	0.251	0.427	PQL	mg/Kg	J (all detects)
	THALLIUM	J	0.183	0.213	PQL	mg/Kg	
SL-565-SA5D-SB-0.0-0.5	SELENIUM	J	0.166	0.421	PQL	mg/Kg	J (all detects)
SL-566-SA5D-SB-0.0-0.5	SELENIUM	J	0.386	0.408	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0384	0.204	PQL	mg/Kg	
SL-566-SA5D-SB-4.0-5.0	SELENIUM	J	0.116	0.421	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0311	0.211	PQL	mg/Kg	
SL-865-SA5D-SB-0.0-0.5	SELENIUM	J	0.266	0.418	PQL	mg/Kg	J (all detects)

Method: 7471B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-565-SA5D-SB-0.0-0.5	MERCURY	J	0.0140	0.0171	PQL	mg/Kg	J (all detects)
SL-566-SA5D-SB-4.0-5.0	MERCURY	J	0.0132	0.0174	PQL	mg/Kg	J (all detects)

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

12/4/2013 8:00:17 AM

ADR version 1.7.0.207

Page 3 of 4

Reporting Limit Outliers

Lab Reporting Batch ID: PH113

Laboratory: LL

EDD Filename: PH113

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-565-SA5D-SB-0.0-0.5	EFH (C21-C30)	J	4.3	5.3	PQL	mg/Kg	J (all detects)
SL-566-SA5D-SB-0.0-0.5	EFH (C15-C20)	J	4.9	5.1	PQL	mg/Kg	J (all detects)

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-557-SA5D-SB-0.0-0.5	2-METHYLNAPHTHALENE	J	0.73	1.7	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	1.5	1.7	PQL	ug/Kg	
	FLUORANTHENE	J	1.3	1.7	PQL	ug/Kg	
	NAPHTHALENE	J	1.1	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	1.1	1.7	PQL	ug/Kg	
SL-558-SA5D-SB-0.0-0.5	PYRENE	J	1.3	1.7	PQL	ug/Kg	J (all detects)
	2-METHYLNAPHTHALENE	J	1.2	1.8	PQL	ug/Kg	
	BENZO(B)FLUORANTHENE	J	1.7	1.8	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	0.86	1.8	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	0.79	1.8	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	7.1	19	PQL	ug/Kg	
	Butylbenzylphthalate	J	10	19	PQL	ug/Kg	
	CHRYSENE	J	1.7	1.8	PQL	ug/Kg	
	FLUORANTHENE	J	1.4	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	1.6	1.8	PQL	ug/Kg	
PHENANTHRENE	J	1.2	1.8	PQL	ug/Kg		
SL-566-SA5D-SB-0.0-0.5	PYRENE	J	1.3	1.8	PQL	ug/Kg	J (all detects)
	1-METHYLNAPHTHALENE	J	1.1	1.7	PQL	ug/Kg	
	2-METHYLNAPHTHALENE	J	1.5	1.7	PQL	ug/Kg	
	ACENAPHTHYLENE	J	0.45	1.7	PQL	ug/Kg	
	ANTHRACENE	J	0.70	1.7	PQL	ug/Kg	
	BENZO(B)FLUORANTHENE	J	1.3	1.7	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	9.1	18	PQL	ug/Kg	
	Butylbenzylphthalate	J	10	18	PQL	ug/Kg	
	CHRYSENE	J	1.1	1.7	PQL	ug/Kg	
	FLUORANTHENE	J	1.1	1.7	PQL	ug/Kg	
	FLUORENE	J	0.70	1.7	PQL	ug/Kg	
SL-566-SA5D-SB-4.0-5.0	PHENANTHRENE	J	1.2	1.7	PQL	ug/Kg	J (all detects)
	PYRENE	J	1.1	1.7	PQL	ug/Kg	
SL-865-SA5D-SB-0.0-0.5	NAPHTHALENE	J	0.91	1.8	PQL	ug/Kg	J (all detects)
SL-865-SA5D-SB-0.0-0.5	BIS(2-ETHYLHEXYL)PHTHALATE	J	11	19	PQL	ug/Kg	J (all detects)
	NAPHTHALENE	J	0.71	1.7	PQL	ug/Kg	

LDC #: 30695P4

VALIDATION COMPLETENESS WORKSHEET

Date: 11/11/13

SDG #: PH113

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: [Signature]
2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	=	Sampling dates: 9/24/13
II.	ICP/MS Tune	=	
III.	Calibration	-	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	SW	MS/D
VII.	Duplicate Sample Analysis	SW	DUP
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	SW	Sc: 24%: J/U/A(A): All
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	A	(3,4)
XV.	Field Blanks	SW	EB = EB31-0012S13 FB = FB-041613

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

D = Duplicate
TB = Trip blank
EB = Equipment blank

(PH114)

(PH032)

Validated Samples:

1	SL-558-SA5D-SB-0.0-0.5	11		21		31	
2	SL-558-SA5D-SB-4.0-5.0	12		22		32	
3	SL-565-SA5D-SB-0.0-0.5	13		23		33	
4	SL-865-SA5D-SB-0.0-0.5	14		24		34	
5	SL-557-SA5D-SB-0.0-0.5	15		25		35	
6	SL-566-SA5D-SB-0.0-0.5	16		26		36	
7	SL-566-SA5D-SB-4.0-5.0	17		27		37	
8	SL-565-SA5D-SB-0.0-0.5MS	18		28		38	
9	SL-565-SA5D-SB-0.0-0.5MSD	19		29		39	
10	SL-565-SA5D-SB-0.0-0.5DUP	20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F
Sampling date: 4/11/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification											
	FB-041613 (SDG: PH032)	Action Limit	1										
Mo	0.0132	6.60	0.198										
Sn	0.0029	1.45											

Sampling date: 9/25/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification											
	EB1-092513 (SDG: PH114)	Action Limit	No Qualifiers										
Ca	0.0498	24.9											

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

30.05 = 95% (J/U/A)

Background Lab Sample ID: 7211497BKG

Matrix Spike Lab Sample ID: 7211498MS

Matrix Spike Duplicate Lab Sample ID: 7211499MSD

Batch Id(s): P26837A, P26838A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit		M	
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q		%R
Aluminum		21361.9673		28720.2845		29871.7210		194.1748	200.0000	MG/KG	3790		4255*		4		20	P
Antimony		0.7327	U	12.7068		14.1710		48.5437	50.0000	MG/KG	26	N	28	N	11	75 - 125	20	P
Arsenic		4.6297		17.8670		19.0360		14.5631	15.0000	MG/KG	91		96		6	75 - 125	20	P
Barium		140.4020		318.7709		312.3290		194.1748	200.0000	MG/KG	92		86		2	75 - 125	20	P
Beryllium		0.8069	B	5.3981		5.3760		4.8544	5.0000	MG/KG	95		91		0	75 - 125	20	P
Boron		2.9069	B	196.4427		195.4540		194.1748	200.0000	MG/KG	100		96		1	75 - 125	20	P
Cadmium		0.7901	B	5.1738		5.2120		4.8544	5.0000	MG/KG	90		88		1	75 - 125	20	P
Calcium		49111.1168		71483.1961		58778.1980		388.3495	400.0000	MG/KG	5761		2417		20		20	P
Chromium		29.0634		52.3447		53.5080		19.4175	20.0000	MG/KG	120		122		2	75 - 125	20	P
Cobalt		8.6347		51.0437		49.1410		48.5437	50.0000	MG/KG	87		81		4	75 - 125	20	P
Copper		28.7158		50.3903		52.0280		24.2718	25.0000	MG/KG	89		93		3	75 - 125	20	P
Iron		51717.3416		32061.9922		30579.6320		97.0874	100.0000	MG/KG	-20245		-21138		5		20	P
Lead		10.5119		23.1388		24.1080		14.5631	15.0000	MG/KG	87		91		4	75 - 125	20	P
Lithium		23.9812		118.8757		121.8050		97.0874	100.0000	MG/KG	98		98		2	75 - 125	20	P
Magnesium		8024.4119		8541.3621		8648.7740		194.1748	200.0000	MG/KG	266		312		1		20	P
Manganese		258.9208		331.9068		293.0770		48.5437	50.0000	MG/KG	150		68*		12		20	P
Mercury		0.0131	B	0.2093		0.2027		0.1657	0.1650	MG/KG	118		115		3	65 - 135	20	CV
Molybdenum		0.1683	U	168.6126		171.2500		194.1748	200.0000	MG/KG	87		86		2	75 - 125	20	P
Nickel		20.8743		59.4194		58.8690		48.5437	50.0000	MG/KG	79		76		1	75 - 125	20	P
Phosphorus		401.7644		561.7699		574.5650		97.0874	100.0000	MG/KG	165		173		2		20	P
Potassium		3878.0040		5889.9835		5806.5610		970.8738	1000.0000	MG/KG	207	N	193	N	1	75 - 125	20	P
Selenium	78	0.1566	B	2.0951		2.2600		1.9417	2.0000	MG/KG	100		105		8	75 - 125	20	MS
Silver	107	0.0257	U	10.0039		9.5780		9.7087	10.0000	MG/KG	103		96		4	75 - 125	20	MS
Sodium		125.8812		1038.5990		1057.5930		970.8738	1000.0000	MG/KG	94		93		2	75 - 125	20	P
Strontium	88	106.8317		186.5534		157.8500		7.7670	8.0000	MG/KG	1026		638		17		20	MS
Thallium	203	0.3210		0.6586		0.7008		0.3883	0.4000	MG/KG	87		95		6	75 - 125	20	MS

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor</p> <p>MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U = Below MDL, B = Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>
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Lancaster
Laboratories



QUALITY ASSURANCE SUMMARY

FORM 5A (MS/MSD)

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

SDG No.: PH113

Matrix: SOIL

Level

(low/med):

LOW

Background Lab Sample ID: 7211497BKG Matrix Spike Lab Sample ID: 7211498MS Matrix Spike Duplicate Lab Sample ID: 7211499MSD
Batch Id(s): P26837A, P26838A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit					
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q	%R	RPD	M	
Tin		2.4772	B	296.3835		301.9100		388.3495	400.0000	MG/KG	76		75		2		75 - 125	20	P	
Titanium		869.4851		1636.4621		1616.3240		97.0874	100.0000	MG/KG	790		747*		1				20	P
Vanadium		61.2079		121.1320		122.3520		48.5437	50.0000	MG/KG	123		122		1		75 - 125	20	P	
Zinc		80.2406		120.0718		123.4260		48.5437	50.0000	MG/KG	82		86		3		75 - 125	20	P	
Zirconium		6.2020		92.9175		94.6960		97.0874	100.0000	MG/KG	89		88		2		75 - 125	20	P	

Note: Results shown are reported on an as-received basis.

<p>METHODS: P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS: U= Below MDL, B= Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS</p>
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Background Lab Sample ID: 7211497BKG
 Batch ID(s): P26837A, P26838A
 Concentration Units: MG/KG

Duplicate Lab Sample ID: 7211500DUP

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			21361.9673		21564.8456		1		P
Antimony			-3.6020	B	-2.8456	B	-23		P
Arsenic		4.0	4.6297		3.5641	B	26		P
Barium			140.4020		147.6029		5		P
Beryllium			0.8069	B	0.7728	B	4		P
Boron		9.9	2.9069	B	10.9786		116		P
Cadmium			0.7901	B	0.9194	B	15		P
Calcium			49111.1168		49676.3602		1		P
Chromium			29.0634		29.7068		2		P
Cobalt			8.6347		10.8262		23	*	P
Copper			28.7158		26.8039		7		P
Iron			51717.3416		42604.5534		19		P
Lead		3.0	10.5119		9.7718		7		P
Lithium			23.9812		24.1932		1		P
Magnesium			8024.4119		8193.6563		2		P
Manganese			258.9208		307.5728		17		P
Mercury			0.0131	B	0.0160	B	20		CV
Molybdenum			0.1683	U	0.1650	U			P
Nickel			20.8743		22.1631		6		P
Phosphorus			401.7644		405.6204		1		P
Potassium			3878.0040		3971.0660		2		P
Selenium	78		0.1566	B	0.1202	B	26		MS
Silver	107		0.0257	U	0.0252	U			MS
Sodium		99.0	125.8812		127.4194		1		P
Strontium	88		106.8317		134.0777		23	*	MS
Thallium	203	0.2	0.3210		0.3412		6		MS
Tin			2.4772	B	2.5087	B	1		P
Titanium			869.4851		892.2961		3		P
Vanadium			61.2079		61.6272		1		P
Zinc			80.2406		77.2447		4		P
Zirconium		5.0	6.2020		6.1835		0		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).
 The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

ok by difference

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: Duplicate Out of Spec
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**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH114

Prepared for

CDM Smith
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
2701 Loker Ave West, Suite 220
Carlsbad, California 92010

December 11, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on September 25, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by EPA SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Pesticides by EPA SW 846 Method 8081A
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A, 7470A and 7471B
Herbicides by EPA SW 846 Method 8151A
Perchlorate by Method 6850
Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M
TPH as Extractables by EPA SW 846 Method 8015M
Dioxins and Dibenzofurans by EPA Method 1613B

Wet Chemistry:

Fluoride by EPA Method 300.0

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks, and field duplicates. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of several blanks for metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits with the exception of one sample for TPH as extractables. No data were qualified due to high %Rs since the associated results were non-detected.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pairs for metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of several LCS/LCSD pairs for SVOCs, pesticides and TPH as extractables. The associated sample results were qualified as non-detected estimated (UJ). The details regarding the qualification of data are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH114	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

Four field duplicate pairs were collected and analyzed for SVOCs, PCBs, metals, TPH as gasoline and TPH as extractables. All RPDs were within QC limits with the exception of several SVOCs and TPH as extractables. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No volatile contaminants were found in the trip blank.

One equipment blank was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, perchlorate, TPH as gasoline, TPH as extractables, dioxins and fluoride. The equipment blank had detections for SVOCs, metals and dioxins. The associated sample results were not detected or were significantly greater than the concentrations found in the equipment blanks, therefore no data were qualified. The equipment blank outlier reports are presented in Enclosure I.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, perchlorate, TPH as gasoline, TPH as extractables, dioxins and fluoride. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were

qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1

Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
25-Sep-2013	SL-573-SA5D-SB-0.0-0.5	7213407	N	3546	8081B	III
25-Sep-2013	SL-573-SA5D-SB-0.0-0.5	7213407	N	3546	8270D SIM	III
25-Sep-2013	TB-092513	7213405	TB	5030B	8015M	III
25-Sep-2013	SL-573-SA5D-SB-4.0-5.0	7213408	N	3546	8081B	III
25-Sep-2013	SL-569-SA5D-SB-0.0-0.5	7213406	N	3050B	6010C	III
25-Sep-2013	SL-569-SA5D-SB-0.0-0.5	7213406	N	3050B	6020A	III
25-Sep-2013	SL-569-SA5D-SB-0.0-0.5	7213406	N	3546	8015M	III
25-Sep-2013	SL-569-SA5D-SB-0.0-0.5	7213406	N	3546	8082A	III
25-Sep-2013	SL-569-SA5D-SB-0.0-0.5	7213406	N	3546	8270D SIM	III
25-Sep-2013	SL-569-SA5D-SB-0.0-0.5	7213406	N	METHOD	1613B	III
25-Sep-2013	SL-569-SA5D-SB-0.0-0.5	7213406	N	METHOD	7471B	III
25-Sep-2013	SL-563-SA5D-SB-0.0-0.5	7213411	N	3050B	6010C	III
25-Sep-2013	SL-563-SA5D-SB-0.0-0.5	7213411	N	3050B	6020A	III
25-Sep-2013	SL-563-SA5D-SB-0.0-0.5	7213411	N	3546	8015M	III
25-Sep-2013	SL-563-SA5D-SB-0.0-0.5	7213411	N	3546	8082A	III
25-Sep-2013	SL-563-SA5D-SB-0.0-0.5	7213411	N	3546	8270D SIM	III
25-Sep-2013	SL-563-SA5D-SB-0.0-0.5	7213411	N	METHOD	1613B	III
25-Sep-2013	SL-563-SA5D-SB-0.0-0.5	7213411	N	METHOD	7471B	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0	7213412	N	3050B	6010C	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0	7213412	N	3050B	6020A	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0	7213412	N	3546	8015M	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0	7213412	N	3546	8082A	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0	7213412	N	3546	8270D SIM	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0	7213412	N	5035A	8015M	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0	7213412	N	METHOD	7471B	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MS	7213413	MS	3050B	6010C	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MS	7213413	MS	3050B	6020A	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MS	7213413	MS	3546	8015M	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MS	7213413	MS	3546	8082A	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MS	7213413	MS	3546	8270D SIM	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MS	7213413	MS	5035A	8015M	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MS	7213413	MS	METHOD	7471B	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MSD	7213414	MSD	3050B	6010C	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MSD	7213414	MSD	3050B	6020A	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MSD	7213414	MSD	3546	8015M	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MSD	7213414	MSD	3546	8082A	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MSD	7213414	MSD	3546	8270D SIM	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MSD	7213414	MSD	5035A	8015M	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0MSD	7213414	MSD	METHOD	7471B	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0DUP	7213415	DUP	3050B	6010C	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0DUP	7213415	DUP	3050B	6020A	III
25-Sep-2013	SL-563-SA5D-SB-4.0-5.0DUP	7213415	DUP	METHOD	7471B	III
25-Sep-2013	SL-863-SA5D-SB-4.0-5.0	7213416	FD	3050B	6010C	III
25-Sep-2013	SL-863-SA5D-SB-4.0-5.0	7213416	FD	3050B	6020A	III
25-Sep-2013	SL-863-SA5D-SB-4.0-5.0	7213416	FD	3546	8015M	III
25-Sep-2013	SL-863-SA5D-SB-4.0-5.0	7213416	FD	3546	8082A	III
25-Sep-2013	SL-863-SA5D-SB-4.0-5.0	7213416	FD	3546	8270D SIM	III
25-Sep-2013	SL-863-SA5D-SB-4.0-5.0	7213416	FD	5035A	8015M	III
25-Sep-2013	SL-863-SA5D-SB-4.0-5.0	7213416	FD	METHOD	7471B	III
25-Sep-2013	SL-562-SA5D-SB-0.0-0.5	7213409	N	3050B	6010C	III
25-Sep-2013	SL-562-SA5D-SB-0.0-0.5	7213409	N	3050B	6020A	III
25-Sep-2013	SL-562-SA5D-SB-0.0-0.5	7213409	N	3546	8015M	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
25-Sep-2013	SL-562-SA5D-SB-0.0-0.5	7213409	N	3546	8082A	III
25-Sep-2013	SL-562-SA5D-SB-0.0-0.5	7213409	N	3546	8270D SIM	III
25-Sep-2013	SL-562-SA5D-SB-0.0-0.5	7213409	N	METHOD	1613B	III
25-Sep-2013	SL-562-SA5D-SB-0.0-0.5	7213409	N	METHOD	7471B	III
25-Sep-2013	SL-562-SA5D-SB-4.0-5.0	7213410	N	3050B	6010C	III
25-Sep-2013	SL-562-SA5D-SB-4.0-5.0	7213410	N	3050B	6020A	III
25-Sep-2013	SL-562-SA5D-SB-4.0-5.0	7213410	N	3546	8015M	III
25-Sep-2013	SL-562-SA5D-SB-4.0-5.0	7213410	N	3546	8082A	III
25-Sep-2013	SL-562-SA5D-SB-4.0-5.0	7213410	N	3546	8270D SIM	III
25-Sep-2013	SL-562-SA5D-SB-4.0-5.0	7213410	N	5035A	8015M	III
25-Sep-2013	SL-562-SA5D-SB-4.0-5.0	7213410	N	METHOD	7471B	III
25-Sep-2013	EB1-092513	7213403	EB	3005A	6010C	III
25-Sep-2013	EB1-092513	7213403	EB	3510C	8015M	III
25-Sep-2013	EB1-092513	7213403	EB	3510C	8081B	III
25-Sep-2013	EB1-092513	7213403	EB	3510C	8082A	III
25-Sep-2013	EB1-092513	7213403	EB	3510C	8270D SIM	III
25-Sep-2013	EB1-092513	7213403	EB	5030B	8015M	III
25-Sep-2013	EB1-092513	7213403	EB	Gen Prep	300.0	III
25-Sep-2013	EB1-092513	7213403	EB	Gen Prep	6850	III
25-Sep-2013	EB1-092513	7213403	EB	M3010A	6020A	III
25-Sep-2013	EB1-092513	7213403	EB	METHOD	1613B	III
25-Sep-2013	EB1-092513	7213403	EB	METHOD	7470A	III
25-Sep-2013	EB1-092513	7213403	EB	METHOD	8151A	III
25-Sep-2013	EB1-092513MSD	P213403M241621A	MSD	Gen Prep	6850	III
25-Sep-2013	EB1-092513MS	P213403R241608A	MS	Gen Prep	6850	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6010C **Matrix: AQ**

Sample ID: EB1-092513 Collected: 9/25/2013 3:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	0.0498	J	0.0334	MDL	0.400	PQL	mg/L	U	B

Method Category: METALS
Method: 6010C **Matrix: SO**

Sample ID: SL-562-SA5D-SB-0.0-0.5 Collected: 9/25/2013 1:35:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	25300		7.47	MDL	41.5	PQL	mg/Kg	J	E
ANTIMONY	4.15	U	0.767	MDL	4.15	PQL	mg/Kg	UJ	Q
ARSENIC	2.93	J	0.725	MDL	4.15	PQL	mg/Kg	J	Z
BERYLLIUM	0.886	J	0.0694	MDL	1.04	PQL	mg/Kg	J	Z
CADMIUM	0.315	J	0.0788	MDL	1.04	PQL	mg/Kg	J	Z
CALCIUM	18600		3.46	MDL	20.7	PQL	mg/Kg	J	E
MAGNESIUM	8040		1.73	MDL	10.4	PQL	mg/Kg	J	E
MANGANESE	417		0.0860	MDL	1.04	PQL	mg/Kg	J	E
MOLYBDENUM	0.184	J	0.176	MDL	2.07	PQL	mg/Kg	U	F
PHOSPHORUS	661		3.00	MDL	10.4	PQL	mg/Kg	J	E
POTASSIUM	6840		8.64	MDL	104	PQL	mg/Kg	J	Q
SODIUM	102	J	17.3	MDL	104	PQL	mg/Kg	J	Z
TIN	2.69	J	0.228	MDL	10.4	PQL	mg/Kg	U	B
VANADIUM	62.4		0.135	MDL	1.04	PQL	mg/Kg	J	Q
ZINC	83.7		0.207	MDL	4.15	PQL	mg/Kg	J	Q

Sample ID: SL-562-SA5D-SB-0.0-0.5 Collected: 9/25/2013 1:35:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	32900		7.50	MDL	82.9	PQL	mg/Kg	J	E

Sample ID: SL-562-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:50:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	28500		8.05	MDL	44.6	PQL	mg/Kg	J	E
ANTIMONY	4.46	U	0.826	MDL	4.46	PQL	mg/Kg	UJ	Q
ARSENIC	3.59	J	0.781	MDL	4.46	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS

Method: 6010C

Matrix: SO

Sample ID: SL-562-SA5D-SB-4.0-5.0

Collected: 9/25/2013 1:50:00

Analysis Type: REA

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.955	J	0.0748	MDL	1.12	PQL	mg/Kg	J	Z
BORON	7.21	J	0.938	MDL	11.2	PQL	mg/Kg	J	Z
CADMIUM	0.105	J	0.0848	MDL	1.12	PQL	mg/Kg	J	Z
CALCIUM	20500		3.73	MDL	22.3	PQL	mg/Kg	J	E
MAGNESIUM	6590		1.86	MDL	11.2	PQL	mg/Kg	J	E
MANGANESE	435		0.0926	MDL	1.12	PQL	mg/Kg	J	E
MOLYBDENUM	0.213	J	0.190	MDL	2.23	PQL	mg/Kg	U	F
PHOSPHORUS	160		3.23	MDL	11.2	PQL	mg/Kg	J	E
POTASSIUM	3370		9.31	MDL	112	PQL	mg/Kg	J	Q
TIN	2.84	J	0.246	MDL	11.2	PQL	mg/Kg	U	B
VANADIUM	66.0		0.145	MDL	1.12	PQL	mg/Kg	J	Q
ZINC	58.3		0.223	MDL	4.46	PQL	mg/Kg	J	Q

Sample ID: SL-562-SA5D-SB-4.0-5.0

Collected: 9/25/2013 1:50:00

Analysis Type: REA2

Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	34300		8.08	MDL	89.3	PQL	mg/Kg	J	E

Sample ID: SL-563-SA5D-SB-0.0-5.0

Collected: 9/25/2013 12:15:00

Analysis Type: REA

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	24600		7.39	MDL	41.0	PQL	mg/Kg	J	E
ANTIMONY	4.10	U	0.759	MDL	4.10	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.766	J	0.0687	MDL	1.03	PQL	mg/Kg	J	Z
CADMIUM	0.372	J	0.0779	MDL	1.03	PQL	mg/Kg	J	Z
MAGNESIUM	7730		1.71	MDL	10.3	PQL	mg/Kg	J	E
MANGANESE	390		0.0851	MDL	1.03	PQL	mg/Kg	J	E
PHOSPHORUS	550		2.96	MDL	10.3	PQL	mg/Kg	J	E
POTASSIUM	4470		8.55	MDL	103	PQL	mg/Kg	J	Q
TIN	2.44	J	0.226	MDL	10.3	PQL	mg/Kg	U	B
VANADIUM	60.8		0.133	MDL	1.03	PQL	mg/Kg	J	Q
ZINC	71.8		0.205	MDL	4.10	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS

Method: 6010C

Matrix: SO

Sample ID: SL-563-SA5D-SB-0.0-0.5 Collected: 9/25/2013 12:15:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	71700		6.85	MDL	41.0	PQL	mg/Kg	J	E
IRON	29800		7.42	MDL	82.0	PQL	mg/Kg	J	E

Sample ID: SL-563-SA5D-SB-4.0-5.0 Collected: 9/25/2013 12:45:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	19300		7.56	MDL	41.9	PQL	mg/Kg	J	E
ANTIMONY	4.19	U	0.776	MDL	4.19	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.655	J	0.0702	MDL	1.05	PQL	mg/Kg	J	Z
CADMIUM	0.439	J	0.0797	MDL	1.05	PQL	mg/Kg	J	Z
IRON	26900		3.80	MDL	41.9	PQL	mg/Kg	J	E
MAGNESIUM	6980		1.75	MDL	10.5	PQL	mg/Kg	J	E
MANGANESE	303		0.0870	MDL	1.05	PQL	mg/Kg	J	E
PHOSPHORUS	521		3.03	MDL	10.5	PQL	mg/Kg	J	E
POTASSIUM	3280		8.74	MDL	105	PQL	mg/Kg	J	Q
TIN	2.39	J	0.231	MDL	10.5	PQL	mg/Kg	U	B
VANADIUM	45.0		0.136	MDL	1.05	PQL	mg/Kg	J	Q
ZINC	70.3		0.210	MDL	4.19	PQL	mg/Kg	J	Q

Sample ID: SL-563-SA5D-SB-4.0-5.0 Collected: 9/25/2013 12:45:00 Analysis Type: REA2 Dilution: 5

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	112000		17.5	MDL	105	PQL	mg/Kg	J	E

Sample ID: SL-569-SA5D-SB-0.0-0.5 Collected: 9/25/2013 10:00:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	18000		7.33	MDL	40.7	PQL	mg/Kg	J	E
ANTIMONY	4.07	U	0.752	MDL	4.07	PQL	mg/Kg	UJ	Q
ARSENIC	3.80	J	0.712	MDL	4.07	PQL	mg/Kg	J	Z
BERYLLIUM	0.593	J	0.0681	MDL	1.02	PQL	mg/Kg	J	Z
CADMIUM	0.114	J	0.0773	MDL	1.02	PQL	mg/Kg	J	Z
CALCIUM	5800		3.40	MDL	20.3	PQL	mg/Kg	J	E
MAGNESIUM	7310		1.70	MDL	10.2	PQL	mg/Kg	J	E
MANGANESE	241		0.0844	MDL	1.02	PQL	mg/Kg	J	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS

Method: 6010C

Matrix: SO

Sample ID: SL-569-SA5D-SB-0.0-0.5 Collected: 9/25/2013 10:00:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
PHOSPHORUS	632		2.94	MDL	10.2	PQL	mg/Kg	J	E
POTASSIUM	3830		8.48	MDL	102	PQL	mg/Kg	J	Q
TIN	2.56	J	0.224	MDL	10.2	PQL	mg/Kg	U	B
VANADIUM	63.7		0.132	MDL	1.02	PQL	mg/Kg	J	Q
ZINC	57.5		0.203	MDL	4.07	PQL	mg/Kg	J	Q

Sample ID: SL-569-SA5D-SB-0.0-0.5 Collected: 9/25/2013 10:00:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	25800		7.36	MDL	81.3	PQL	mg/Kg	J	E

Sample ID: SL-863-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:00:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	16800		7.63	MDL	42.4	PQL	mg/Kg	J	E
ANTIMONY	4.24	U	0.784	MDL	4.24	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.560	J	0.0709	MDL	1.06	PQL	mg/Kg	J	Z
BORON	8.83	J	0.890	MDL	10.6	PQL	mg/Kg	J	Z
CADMIUM	0.402	J	0.0805	MDL	1.06	PQL	mg/Kg	J	Z
IRON	25000		3.83	MDL	42.4	PQL	mg/Kg	J	E
MAGNESIUM	6440		1.77	MDL	10.6	PQL	mg/Kg	J	E
MANGANESE	296		0.0879	MDL	1.06	PQL	mg/Kg	J	E
PHOSPHORUS	505		3.06	MDL	10.6	PQL	mg/Kg	J	E
POTASSIUM	2860		8.83	MDL	106	PQL	mg/Kg	J	Q
TIN	2.35	J	0.233	MDL	10.6	PQL	mg/Kg	U	B
VANADIUM	40.1		0.138	MDL	1.06	PQL	mg/Kg	J	Q
ZINC	63.0		0.212	MDL	4.24	PQL	mg/Kg	J	Q

Sample ID: SL-863-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:00:00 Analysis Type: REA2 Dilution: 5

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	127000		17.7	MDL	106	PQL	mg/Kg	J	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS

Method: 6020A

Matrix: SO

Sample ID: SL-562-SA5D-SB-0.0-0.5 Collected: 9/25/2013 1:35:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0479	J	0.0269	MDL	0.207	PQL	mg/Kg	J	Z, Q, E
STRONTIUM	69.6		0.0705	MDL	0.415	PQL	mg/Kg	J	E
THALLIUM	0.417		0.0311	MDL	0.207	PQL	mg/Kg	J	Q, E

Sample ID: SL-562-SA5D-SB-0.0-0.5 Collected: 9/25/2013 1:35:00 Analysis Type: REA4 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.296	J	0.104	MDL	0.415	PQL	mg/Kg	J	Z

Sample ID: SL-562-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:50:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0655	J	0.0290	MDL	0.223	PQL	mg/Kg	J	Z, Q, E
STRONTIUM	82.7		0.0759	MDL	0.446	PQL	mg/Kg	J	E
THALLIUM	0.466		0.0335	MDL	0.223	PQL	mg/Kg	J	Q, E

Sample ID: SL-562-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:50:00 Analysis Type: REA4 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.222	J	0.112	MDL	0.446	PQL	mg/Kg	J	Z

Sample ID: SL-563-SA5D-SB-0.0-0.5 Collected: 9/25/2013 12:15:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0487	J	0.0267	MDL	0.205	PQL	mg/Kg	J	Z, Q, E
THALLIUM	0.378		0.0308	MDL	0.205	PQL	mg/Kg	J	Q, E

Sample ID: SL-563-SA5D-SB-0.0-0.5 Collected: 9/25/2013 12:15:00 Analysis Type: REA3 Dilution: 5

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	128		0.174	MDL	1.03	PQL	mg/Kg	J	E

Sample ID: SL-563-SA5D-SB-0.0-0.5 Collected: 9/25/2013 12:15:00 Analysis Type: REA4 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.327	J	0.103	MDL	0.410	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-563-SA5D-SB-4.0-5.0			Collected: 9/25/2013 12:45:00			Analysis Type: REA2			Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
SILVER	0.0378	J	0.0273	MDL	0.210	PQL	mg/Kg	J	Z, Q, E	
THALLIUM	0.289		0.0315	MDL	0.210	PQL	mg/Kg	J	Q, E	

Sample ID: SL-563-SA5D-SB-4.0-5.0			Collected: 9/25/2013 12:45:00			Analysis Type: REA3			Dilution: 10	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
STRONTIUM	206		0.356	MDL	2.10	PQL	mg/Kg	J	E	

Sample ID: SL-563-SA5D-SB-4.0-5.0			Collected: 9/25/2013 12:45:00			Analysis Type: REA4			Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
SELENIUM	0.239	J	0.105	MDL	0.419	PQL	mg/Kg	J	Z	

Sample ID: SL-569-SA5D-SB-0.0-0.5			Collected: 9/25/2013 10:00:00			Analysis Type: REA2			Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
SILVER	0.203	U	0.0264	MDL	0.203	PQL	mg/Kg	UJ	E	
STRONTIUM	23.7		0.0691	MDL	0.407	PQL	mg/Kg	J	E	
THALLIUM	0.223		0.0305	MDL	0.203	PQL	mg/Kg	J	Q, E	

Sample ID: SL-569-SA5D-SB-0.0-0.5			Collected: 9/25/2013 10:00:00			Analysis Type: REA4			Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
SELENIUM	0.257	J	0.102	MDL	0.407	PQL	mg/Kg	J	Z	

Sample ID: SL-863-SA5D-SB-4.0-5.0			Collected: 9/25/2013 1:00:00			Analysis Type: REA2			Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
SILVER	0.0392	J	0.0275	MDL	0.212	PQL	mg/Kg	J	Z, Q, E	
THALLIUM	0.332		0.0318	MDL	0.212	PQL	mg/Kg	J	Q, E	

Sample ID: SL-863-SA5D-SB-4.0-5.0			Collected: 9/25/2013 1:00:00			Analysis Type: REA3			Dilution: 10	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code	
STRONTIUM	235		0.360	MDL	2.12	PQL	mg/Kg	J	E	

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6020A **Matrix: SO**

Sample ID: SL-863-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:00:00 Analysis Type: REA4 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.207	J	0.106	MDL	0.424	PQL	mg/Kg	J	Z

Method Category: METALS
Method: 7471B **Matrix: SO**

Sample ID: SL-562-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0159	J	0.0105	MDL	0.0175	PQL	mg/Kg	J	Z

Sample ID: SL-563-SA5D-SB-4.0-5.0 Collected: 9/25/2013 12:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0130	J	0.0103	MDL	0.0172	PQL	mg/Kg	J	Z

Sample ID: SL-863-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0126	J	0.0106	MDL	0.0176	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 1613B **Matrix: AQ**

Sample ID: EB1-092513 Collected: 9/25/2013 3:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.426	JBQ	0.151	MDL	9.78	PQL	pg/L	U	B
1,2,3,4,6,7,8-HPCDF	0.207	JB	0.0772	MDL	9.78	PQL	pg/L	U	B
1,2,3,4,7,8,9-HPCDF	0.276	JB	0.0916	MDL	9.78	PQL	pg/L	U	B
1,2,3,4,7,8-HxCDD	0.202	JB	0.139	MDL	9.78	PQL	pg/L	U	B
1,2,3,4,7,8-HxCDF	0.196	JB	0.0719	MDL	9.78	PQL	pg/L	U	B
1,2,3,6,7,8-HxCDD	0.353	JBQ	0.148	MDL	9.78	PQL	pg/L	U	B
1,2,3,7,8,9-HxCDD	0.256	JBQ	0.141	MDL	9.78	PQL	pg/L	U	B
1,2,3,7,8,9-HxCDF	0.217	JBQ	0.0732	MDL	9.78	PQL	pg/L	U	B
1,2,3,7,8-PECDD	0.293	JBQ	0.194	MDL	9.78	PQL	pg/L	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: AQ

Sample ID: EB1-092513 Collected: 9/25/2013 3:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,7,8-PECDF	0.638	JBQ	0.0873	MDL	9.78	PQL	pg/L	U	B
2,3,4,6,7,8-HXCDF	0.174	JBQ	0.0691	MDL	9.78	PQL	pg/L	U	B
2,3,4,7,8-PECDF	0.274	JBQ	0.0782	MDL	9.78	PQL	pg/L	U	B
OCDD	2.39	JB	0.200	MDL	19.6	PQL	pg/L	U	B
OCDF	1.30	JBQ	0.188	MDL	19.6	PQL	pg/L	U	B

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-562-SA5D-SB-0.0-0.5 Collected: 9/25/2013 1:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.95	JB	0.0284	MDL	5.08	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.495	JB	0.0171	MDL	5.08	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0976	JBQ	0.0245	MDL	5.08	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0832	JBQ	0.0294	MDL	5.08	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.149	JBQ	0.0242	MDL	5.08	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.138	JB	0.0315	MDL	5.08	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.130	JBQ	0.0226	MDL	5.08	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.194	JB	0.0283	MDL	5.08	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.0658	JBQ	0.0233	MDL	5.08	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.117	JBQ	0.0355	MDL	5.08	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.276	JB	0.0242	MDL	5.08	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.162	JBQ	0.0216	MDL	5.08	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.259	JB	0.0238	MDL	5.08	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.159	JQ	0.0502	MDL	1.02	PQL	ng/Kg	J	Z
OCDF	0.770	JB	0.0299	MDL	10.2	PQL	ng/Kg	J	Z

Sample ID: SL-563-SA5D-SB-0.0-0.5 Collected: 9/25/2013 12:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.283	JBQ	0.0216	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0983	JB	0.00682	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0571	JBQ	0.00991	MDL	5.31	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	SVOA		
Method:	1613B	Matrix:	SO

Sample ID: SL-563-SA5D-SB-0.0-0.5 Collected: 9/25/2013 12:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HxCDD	0.0678	JBQ	0.0187	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.0797	JBQ	0.0148	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.0823	JBQ	0.0203	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDF	0.101	JB	0.0138	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0651	JBQ	0.0181	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDF	0.0760	JB	0.0155	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.140	JB	0.0265	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.173	JBQ	0.0154	MDL	5.31	PQL	ng/Kg	U	B
2,3,4,6,7,8-HxCDF	0.0911	JB	0.0133	MDL	5.31	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.126	JBQ	0.0144	MDL	5.31	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.0419	JQ	0.0307	MDL	1.06	PQL	ng/Kg	J	Z
OCDD	1.10	JB	0.0180	MDL	10.6	PQL	ng/Kg	U	B
OCDF	0.253	JBQ	0.0278	MDL	10.6	PQL	ng/Kg	U	B

Sample ID: SL-569-SA5D-SB-0.0-0.5 Collected: 9/25/2013 10:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.70	JB	0.0246	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.491	JB	0.0155	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0876	JBQ	0.0240	MDL	5.05	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0701	JBQ	0.0229	MDL	5.05	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.104	JB	0.0234	MDL	5.05	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.128	JBQ	0.0241	MDL	5.05	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDF	0.0850	JBQ	0.0216	MDL	5.05	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.108	JBQ	0.0231	MDL	5.05	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDF	0.0751	JB	0.0243	MDL	5.05	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.309	JB	0.0221	MDL	5.05	PQL	ng/Kg	U	B
2,3,4,6,7,8-HxCDF	0.104	JBQ	0.0226	MDL	5.05	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.112	J	0.0487	MDL	1.01	PQL	ng/Kg	J	Z
OCDF	1.04	JB	0.0298	MDL	10.1	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	SVOA	
Method:	8015M	Matrix: AQ

Sample ID: EB1-092513 Collected: 9/25/2013 3:00:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C30-C40)	0.49	U	0.097	MDL	0.49	PQL	mg/L	UJ	L
EFH (C8-C11)	0.097	U	0.049	MDL	0.097	PQL	mg/L	UJ	L

Method Category:	SVOA	
Method:	8015M	Matrix: SO

Sample ID: SL-562-SA5D-SB-0.0-0.5 Collected: 9/25/2013 1:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	4.1	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z

Sample ID: SL-562-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:50:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	2.2	J	2.2	MDL	5.5	PQL	mg/Kg	J	Z

Sample ID: SL-563-SA5D-SB-0.0-0.5 Collected: 9/25/2013 12:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	2.9	J	2.1	MDL	5.3	PQL	mg/Kg	J	Z

Sample ID: SL-563-SA5D-SB-4.0-5.0 Collected: 9/25/2013 12:45:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	5.4	U	2.2	MDL	5.4	PQL	mg/Kg	UJ	FD
EFH (C30-C40)	11	U	4.3	MDL	11	PQL	mg/Kg	UJ	FD

Sample ID: SL-569-SA5D-SB-0.0-0.5 Collected: 9/25/2013 10:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C8-C11)	3.8	J	2.0	MDL	5.1	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8015M **Matrix:** SO

Sample ID: SL-863-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:00:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	5.0	J	2.1	MDL	5.3	PQL	mg/Kg	J	Z, FD
EFH (C30-C40)	8.5	J	4.3	MDL	11	PQL	mg/Kg	J	Z, FD

Method Category: SVOA
Method: 8081B **Matrix:** AQ

Sample ID: EB1-092513 Collected: 9/25/2013 3:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ENDRIN	0.016	U	0.0066	MDL	0.016	PQL	ug/L	UJ	L, E
ENDRIN KETONE	0.016	U	0.0041	MDL	0.016	PQL	ug/L	UJ	E

Method Category: SVOA
Method: 8081B **Matrix:** SO

Sample ID: SL-573-SA5D-SB-0.0-0.5 Collected: 9/25/2013 7:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALPHA-BHC	0.28	J	0.17	MDL	0.84	PQL	ug/Kg	J	Z
HEPTACHLOR	0.19	J	0.17	MDL	0.84	PQL	ug/Kg	J	Z
HEPTACHLOR EPOXIDE	0.22	J	0.17	MDL	0.84	PQL	ug/Kg	J	Z

Method Category: SVOA
Method: 8270D SIM **Matrix:** AQ

Sample ID: EB1-092513 Collected: 9/25/2013 3:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BIS(2-ETHYLHEXYL)PHTHALATE	1.0	U	0.052	MDL	1.0	PQL	ug/L	UJ	E
Diethylphthalate	0.83	J	0.052	MDL	1.0	PQL	ug/L	J	Z
Di-n-butylphthalate	0.22	J	0.052	MDL	1.0	PQL	ug/L	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

12/4/2013 7:42:00 AM

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Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-562-SA5D-SB-0.0-0.5 Collected: 9/25/2013 1:35:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	0.92	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
2-METHYLNAPHTHALENE	1.6	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	0.95	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	8.1	J	6.3	MDL	19	PQL	ug/Kg	J	Z
FLUORANTHENE	1.5	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
INDENO(1,2,3-CD)PYRENE	0.71	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
PYRENE	1.7	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-562-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:50:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(A)ANTHRACENE	1.0	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	1.5	J	0.37	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-563-SA5D-SB-0.0-0.5 Collected: 9/25/2013 12:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	1.2	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-563-SA5D-SB-4.0-5.0 Collected: 9/25/2013 12:45:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	5.6		0.72	MDL	1.8	PQL	ug/Kg	J	FD
2-METHYLNAPHTHALENE	12		0.72	MDL	1.8	PQL	ug/Kg	J	FD
NAPHTHALENE	7.8		0.72	MDL	1.8	PQL	ug/Kg	J	FD

Sample ID: SL-569-SA5D-SB-0.0-0.5 Collected: 9/25/2013 10:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	0.99	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
NAPHTHALENE	0.79	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-573-SA5D-SB-0.0-0.5 Collected: 9/25/2013 7:45:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.77	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(B)FLUORANTHENE	1.3	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	0.69	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z
CHRYSENE	1.2	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
FLUORANTHENE	1.1	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z
PHENANTHRENE	1.1	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	0.98	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-863-SA5D-SB-4.0-5.0 Collected: 9/25/2013 1:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	1.8	U	0.71	MDL	1.8	PQL	ug/Kg	UJ	FD
2-METHYLNAPHTHALENE	1.8	U	0.71	MDL	1.8	PQL	ug/Kg	UJ	FD
NAPHTHALENE	1.8	U	0.71	MDL	1.8	PQL	ug/Kg	UJ	FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

12/4/2013 7:42:00 AM

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Data Qualifier Summary

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Method Blank Contamination
E	Laboratory Control Precision
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Field Blank Contamination
FD	Field Duplicate Precision
L	Laboratory Control Precision
L	Laboratory Control Spike Lower Estimation
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

12/4/2013 7:42:00 AM

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH114

Method Blank Outlier Report

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2760B371524	10/6/2013 3:24:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF 2,3,7,8-TCDD OCDD OCDF	0.518 pg/L 0.528 pg/L 0.306 pg/L 0.207 pg/L 0.225 pg/L 0.126 pg/L 0.268 pg/L 0.369 pg/L 0.383 pg/L 0.345 pg/L 0.494 pg/L 0.369 pg/L 0.155 pg/L 0.346 pg/L 2.28 pg/L 0.989 pg/L	EB1-092513

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB1-092513(RES)	1,2,3,4,6,7,8-HPCDD	0.426 pg/L	0.426U pg/L
EB1-092513(RES)	1,2,3,4,6,7,8-HPCDF	0.207 pg/L	0.207U pg/L
EB1-092513(RES)	1,2,3,4,7,8,9-HPCDF	0.276 pg/L	0.276U pg/L
EB1-092513(RES)	1,2,3,4,7,8-HxCDD	0.202 pg/L	0.202U pg/L
EB1-092513(RES)	1,2,3,4,7,8-HXCDF	0.196 pg/L	0.196U pg/L
EB1-092513(RES)	1,2,3,6,7,8-HXCDD	0.353 pg/L	0.353U pg/L
EB1-092513(RES)	1,2,3,7,8,9-HXCDD	0.256 pg/L	0.256U pg/L
EB1-092513(RES)	1,2,3,7,8,9-HXCDF	0.217 pg/L	0.217U pg/L
EB1-092513(RES)	1,2,3,7,8-PECDD	0.293 pg/L	0.293U pg/L
EB1-092513(RES)	1,2,3,7,8-PECDF	0.638 pg/L	0.638U pg/L
EB1-092513(RES)	2,3,4,6,7,8-HXCDF	0.174 pg/L	0.174U pg/L
EB1-092513(RES)	2,3,4,7,8-PECDF	0.274 pg/L	0.274U pg/L
EB1-092513(RES)	OCDD	2.39 pg/L	2.39U pg/L
EB1-092513(RES)	OCDF	1.30 pg/L	1.30U pg/L

Method Blank Outlier Report

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2730B371140	10/2/2013 11:40:00 AM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF OCDD OCDF	0.0965 ng/Kg 0.0595 ng/Kg 0.0657 ng/Kg 0.0334 ng/Kg 0.0515 ng/Kg 0.0256 ng/Kg 0.0453 ng/Kg 0.0369 ng/Kg 0.0754 ng/Kg 0.0738 ng/Kg 0.0876 ng/Kg 0.0332 ng/Kg 0.0646 ng/Kg 0.249 ng/Kg 0.137 ng/Kg	SL-562-SA5D-SB-0.0-0.5 SL-563-SA5D-SB-0.0-0.5 SL-569-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-562-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0976 ng/Kg	0.0976U ng/Kg
SL-562-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0832 ng/Kg	0.0832U ng/Kg
SL-562-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.149 ng/Kg	0.149U ng/Kg
SL-562-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.130 ng/Kg	0.130U ng/Kg
SL-562-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.0658 ng/Kg	0.0658U ng/Kg
SL-562-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.117 ng/Kg	0.117U ng/Kg
SL-562-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.276 ng/Kg	0.276U ng/Kg
SL-562-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.162 ng/Kg	0.162U ng/Kg
SL-562-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.259 ng/Kg	0.259U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.283 ng/Kg	0.283U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0983 ng/Kg	0.0983U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0571 ng/Kg	0.0571U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0678 ng/Kg	0.0678U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0797 ng/Kg	0.0797U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.0823 ng/Kg	0.0823U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.101 ng/Kg	0.101U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.0651 ng/Kg	0.0651U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.0760 ng/Kg	0.0760U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.140 ng/Kg	0.140U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.173 ng/Kg	0.173U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0911 ng/Kg	0.0911U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.126 ng/Kg	0.126U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	OCDD	1.10 ng/Kg	1.10U ng/Kg
SL-563-SA5D-SB-0.0-0.5(RES)	OCDF	0.253 ng/Kg	0.253U ng/Kg
SL-569-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0876 ng/Kg	0.0876U ng/Kg
SL-569-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0701 ng/Kg	0.0701U ng/Kg
SL-569-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.104 ng/Kg	0.104U ng/Kg
SL-569-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.128 ng/Kg	0.128U ng/Kg
SL-569-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0850 ng/Kg	0.0850U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-569-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.108 ng/Kg	0.108U ng/Kg
SL-569-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.0751 ng/Kg	0.0751U ng/Kg
SL-569-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.309 ng/Kg	0.309U ng/Kg
SL-569-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.104 ng/Kg	0.104U ng/Kg

Method: 6010C
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P27035BB221457	9/30/2013 2:57:00 PM	CALCIUM	0.0618 mg/L	EB1-092513

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB1-092513(RES)	CALCIUM	0.0498 mg/L	0.0498U mg/L

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P27437BB220610	10/4/2013 6:10:00 AM	CALCIUM LITHIUM TIN ZINC	8.09 mg/Kg 0.37 mg/Kg 1.38 mg/Kg 0.535 mg/Kg	SL-562-SA5D-SB-0.0-0.5 SL-562-SA5D-SB-4.0-5.0 SL-563-SA5D-SB-0.0-0.5 SL-563-SA5D-SB-4.0-5.0 SL-569-SA5D-SB-0.0-0.5 SL-863-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-562-SA5D-SB-0.0-0.5(REA)	TIN	2.69 mg/Kg	2.69U mg/Kg
SL-562-SA5D-SB-4.0-5.0(REA)	TIN	2.84 mg/Kg	2.84U mg/Kg
SL-563-SA5D-SB-0.0-0.5(REA)	TIN	2.44 mg/Kg	2.44U mg/Kg
SL-563-SA5D-SB-4.0-5.0(REA)	TIN	2.39 mg/Kg	2.39U mg/Kg
SL-569-SA5D-SB-0.0-0.5(REA)	TIN	2.56 mg/Kg	2.56U mg/Kg
SL-863-SA5D-SB-4.0-5.0(REA)	TIN	2.35 mg/Kg	2.35U mg/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041613(REA2/TOT)	4/16/2013 3:15:00 PM	MOLYBDENUM TIN	0.0132 mg/L 0.0029 mg/L	SL-562-SA5D-SB-0.0-0.5 SL-562-SA5D-SB-4.0-5.0 SL-563-SA5D-SB-0.0-0.5 SL-563-SA5D-SB-4.0-5.0 SL-569-SA5D-SB-0.0-0.5 SL-573-SA5D-SB-0.0-0.5 SL-573-SA5D-SB-4.0-5.0 SL-863-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-562-SA5D-SB-0.0-0.5(REA)	MOLYBDENUM	0.184 mg/Kg	0.184U mg/Kg
SL-562-SA5D-SB-4.0-5.0(REA)	MOLYBDENUM	0.213 mg/Kg	0.213U mg/Kg

Surrogate Outlier Report

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8015M

Matrix: AQ

<i>Sample ID (Analysis Type)</i>	<i>Surrogate</i>	<i>Sample % Recovery</i>	<i>% Recovery Limits</i>	<i>Affected Compounds</i>	<i>Flag</i>
EB1-092513 (REA)	CHLOROBENZENE	126	37.00-125.00	All Target Analytes	J (all detects)

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-563-SA5D-SB-4.0-5.0MS (TOT) SL-563-SA5D-SB-4.0-5.0MSD (TOT) (SL-562-SA5D-SB-0.0-0.5 SL-562-SA5D-SB-4.0-5.0 SL-563-SA5D-SB-0.0-0.5 SL-563-SA5D-SB-4.0-5.0 SL-569-SA5D-SB-0.0-0.5 SL-863-SA5D-SB-4.0-5.0)	POTASSIUM TITANIUM VANADIUM	192 652 127	128 412 -	75.00-125.00 75.00-125.00 75.00-125.00	- - -	POTASSIUM TITANIUM VANADIUM	J (all detects) Ti, No Qual, >4x
SL-563-SA5D-SB-4.0-5.0MS (TOT) SL-563-SA5D-SB-4.0-5.0MSD (TOT) (SL-562-SA5D-SB-0.0-0.5 SL-562-SA5D-SB-4.0-5.0 SL-563-SA5D-SB-0.0-0.5 SL-563-SA5D-SB-4.0-5.0 SL-569-SA5D-SB-0.0-0.5 SL-863-SA5D-SB-4.0-5.0)	CALCIUM IRON MAGNESIUM	-1269 2744 726	12670 -6195 -89	75.00-125.00 75.00-125.00 75.00-125.00	44 (20.00) 38 (20.00) 23 (20.00)	CALCIUM IRON MAGNESIUM	J(all detects) UJ(all non-detects) No Qual %R, >4x
SL-563-SA5D-SB-4.0-5.0MS (TOT) SL-563-SA5D-SB-4.0-5.0MSD (TOT) (SL-562-SA5D-SB-0.0-0.5 SL-562-SA5D-SB-4.0-5.0 SL-563-SA5D-SB-0.0-0.5 SL-563-SA5D-SB-4.0-5.0 SL-569-SA5D-SB-0.0-0.5 SL-863-SA5D-SB-4.0-5.0)	ALUMINUM ANTIMONY MANGANESE PHOSPHORUS ZINC	4770 47 56 - -	2132 48 219 293 69	75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00	21 (20.00) - 24 (20.00) 26 (20.00) -	ALUMINUM ANTIMONY MANGANESE PHOSPHORUS ZINC	J(all detects) UJ(all non-detects) Al, Mn, P, No Qual %R, >4x

Method: 6020A

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-563-SA5D-SB-4.0-5.0MS (TOT) SL-563-SA5D-SB-4.0-5.0MSD (TOT) (SL-562-SA5D-SB-0.0-0.5 SL-562-SA5D-SB-4.0-5.0 SL-563-SA5D-SB-0.0-0.5 SL-563-SA5D-SB-4.0-5.0 SL-569-SA5D-SB-0.0-0.5 SL-863-SA5D-SB-4.0-5.0)	SILVER STRONTIUM THALLIUM	145 318 174	- 950 -	75.00-125.00 75.00-125.00 75.00-125.00	27 (20.00) 21 (20.00) 33 (20.00)	SILVER STRONTIUM THALLIUM	J(all detects) UJ(all non-detects) Sr, No Qual %R, >4x

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6020A
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-563-SA5D-SB-4.0-5.0DUP (TOT)	SELENIUM	41	20.00	No Qual, OK by Difference
(SL-562-SA5D-SB-0.0-0.5)	SILVER	28	20.00	
SL-562-SA5D-SB-4.0-5.0	THALLIUM	31	20.00	
SL-563-SA5D-SB-0.0-0.5				
SL-563-SA5D-SB-4.0-5.0				
SL-569-SA5D-SB-0.0-0.5				
SL-863-SA5D-SB-4.0-5.0				

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8081B

Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P32730AQ241157A P32730AY241212A (EB1-092513)	ENDRIN ENDRIN KETONE	24 136	- -	43.00-139.00 51.00-133.00	107 (30.00) 38 (30.00)	ENDRIN ENDRIN KETONE	J (all detects) UJ (all non-detects)

Method: 8015M

Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P32732AQ320750A P32732AY320811A (EB1-092513)	EFH (C30-C40) EFH (C8-C11)	67 64	69 61	70.00-130.00 70.00-130.00	- -	EFH (C30-C40) EFH (C8-C11)	J (all detects) UJ (all non-detects)

Method: 8270D SIM

Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P4WKLCSQ261241 P4WKLCSY261310 (EB1-092513)	BENZO(B)FLUORANTHENE	136	133	74.00-130.00	-	BENZO(B)FLUORANTHENE	J (all detects)
P4WKLCSQ261241 P4WKLCSY261310 (EB1-092513)	BIS(2-ETHYLHEXYL)PHTHALAT	185	-	70.00-143.00	87 (30.00)	BIS(2-ETHYLHEXYL)PHTHALA	J (all detects) UJ (all non-detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 160.3M

Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-563-SA5D-SB-4.0-5.0	SL-863-SA5D-SB-4.0-5.0			
MOISTURE	7.4	6.5	13		No Qualifiers Applied

Method: 6010C

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-563-SA5D-SB-4.0-5.0 (TOT)	SL-863-SA5D-SB-4.0-5.0 (TOT)			
ALUMINUM	19300	16800	14	50.00	No Qualifiers Applied
ARSENIC	4.38	4.49	2	50.00	
BARIUM	75.0	67.1	11	50.00	
BERYLLIUM	0.655	0.560	16	50.00	
BORON	10.5	8.83	17	50.00	
CADMIUM	0.439	0.402	9	50.00	
CALCIUM	112000	127000	13	50.00	
CHROMIUM	24.7	23.0	7	50.00	
COBALT	9.99	7.40	30	50.00	
COPPER	14.7	12.6	15	50.00	
IRON	26900	25000	7	50.00	
LEAD	10.0	8.55	16	50.00	
LITHIUM	26.2	24.0	9	50.00	
MAGNESIUM	6980	6440	8	50.00	
MANGANESE	303	296	2	50.00	
NICKEL	15.9	14.5	9	50.00	
PHOSPHORUS	521	505	3	50.00	
POTASSIUM	3280	2860	14	50.00	
SODIUM	146	144	1	50.00	
TIN	2.39	2.35	2	50.00	
TITANIUM	873	734	17	50.00	
VANADIUM	45.0	40.1	12	50.00	
ZINC	70.3	63.0	11	50.00	
Zirconium	7.13	6.31	12	50.00	

Method: 6020A

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-563-SA5D-SB-4.0-5.0 (TOT)	SL-863-SA5D-SB-4.0-5.0 (TOT)			
SELENIUM	0.239	0.207	14	50.00	No Qualifiers Applied
SILVER	0.0378	0.0392	4	50.00	
STRONTIUM	206	235	13	50.00	
THALLIUM	0.289	0.332	14	50.00	

Method: 7471B

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-563-SA5D-SB-4.0-5.0 (TOT)	SL-863-SA5D-SB-4.0-5.0 (TOT)			
MERCURY	0.0130	0.0126	3	50.00	No Qualifiers Applied

Field Duplicate RPD Report

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8015M

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-563-SA5D-SB-4.0-5.0	SL-863-SA5D-SB-4.0-5.0			
EFH (C21-C30)	5.4 U	5.0	200	50.00	J(all detects)
EFH (C30-C40)	11 U	8.5	200	50.00	UJ(all non-detects)

Method: 8270D SIM

Matrix: SO

Analyte	Concentration (ug/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-563-SA5D-SB-4.0-5.0	SL-863-SA5D-SB-4.0-5.0			
1-METHYLNAPHTHALENE	5.6	1.8 U	200	50.00	J(all detects) UJ(all non-detects)
2-METHYLNAPHTHALENE	12	1.8 U	200	50.00	
NAPHTHALENE	7.8	1.8 U	200	50.00	

Method: 9045M

Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-563-SA5D-SB-4.0-5.0	SL-863-SA5D-SB-4.0-5.0			
PH	8.29	8.35	1	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-092513	1,2,3,4,6,7,8-HPCDD	JBQ	0.426	9.78	PQL	pg/L	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.207	9.78	PQL	pg/L	
	1,2,3,4,7,8,9-HPCDF	JB	0.276	9.78	PQL	pg/L	
	1,2,3,4,7,8-HxCDD	JB	0.202	9.78	PQL	pg/L	
	1,2,3,4,7,8-HxCDF	JB	0.196	9.78	PQL	pg/L	
	1,2,3,6,7,8-HxCDD	JBQ	0.353	9.78	PQL	pg/L	
	1,2,3,7,8,9-HxCDD	JBQ	0.256	9.78	PQL	pg/L	
	1,2,3,7,8,9-HxCDF	JBQ	0.217	9.78	PQL	pg/L	
	1,2,3,7,8-PECDD	JBQ	0.293	9.78	PQL	pg/L	
	1,2,3,7,8-PECDF	JBQ	0.638	9.78	PQL	pg/L	
	2,3,4,6,7,8-HxCDF	JBQ	0.174	9.78	PQL	pg/L	
	2,3,4,7,8-PECDF	JBQ	0.274	9.78	PQL	pg/L	
	OCDD	JB	2.39	19.6	PQL	pg/L	
	OCDF	JBQ	1.30	19.6	PQL	pg/L	

Method: 6010C

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-092513	CALCIUM	J	0.0498	0.400	PQL	mg/L	J (all detects)

Method: 8270D SIM

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-092513	Diethylphthalate	J	0.83	1.0	PQL	ug/L	J (all detects)
	Di-n-butylphthalate	J	0.22	1.0	PQL	ug/L	

Reporting Limit Outliers

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-562-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	1.95	5.08	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.495	5.08	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0976	5.08	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0832	5.08	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.149	5.08	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.138	5.08	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.130	5.08	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.194	5.08	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JBQ	0.0658	5.08	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.117	5.08	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.276	5.08	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.162	5.08	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.259	5.08	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.159	1.02	PQL	ng/Kg	
	OCDF	JB	0.770	10.2	PQL	ng/Kg	
SL-563-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.283	5.31	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.0983	5.31	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0571	5.31	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0678	5.31	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.0797	5.31	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.0823	5.31	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.101	5.31	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JBQ	0.0651	5.31	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.0760	5.31	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.140	5.31	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.173	5.31	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.0911	5.31	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.126	5.31	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0419	1.06	PQL	ng/Kg	
	OCDD	JB	1.10	10.6	PQL	ng/Kg	
OCDF	JBQ	0.253	10.6	PQL	ng/Kg		
SL-569-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	1.70	5.05	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.491	5.05	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0876	5.05	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0701	5.05	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.104	5.05	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.128	5.05	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.0850	5.05	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JBQ	0.108	5.05	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.0751	5.05	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.309	5.05	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.104	5.05	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.112	1.01	PQL	ng/Kg	
	OCDF	JB	1.04	10.1	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-562-SA5D-SB-0.0-0.5	ARSENIC	J	2.93	4.15	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.886	1.04	PQL	mg/Kg	
	CADMIUM	J	0.315	1.04	PQL	mg/Kg	
	MOLYBDENUM	J	0.184	2.07	PQL	mg/Kg	
	SODIUM	J	102	104	PQL	mg/Kg	
SL-562-SA5D-SB-4.0-5.0	TIN	J	2.69	10.4	PQL	mg/Kg	J (all detects)
	ARSENIC	J	3.59	4.46	PQL	mg/Kg	
	BERYLLIUM	J	0.955	1.12	PQL	mg/Kg	
	BORON	J	7.21	11.2	PQL	mg/Kg	
	CADMIUM	J	0.105	1.12	PQL	mg/Kg	
SL-563-SA5D-SB-0.0-0.5	MOLYBDENUM	J	0.213	2.23	PQL	mg/Kg	J (all detects)
	TIN	J	2.84	11.2	PQL	mg/Kg	
	BERYLLIUM	J	0.766	1.03	PQL	mg/Kg	
SL-563-SA5D-SB-4.0-5.0	CADMIUM	J	0.372	1.03	PQL	mg/Kg	J (all detects)
	TIN	J	2.44	10.3	PQL	mg/Kg	
	BERYLLIUM	J	0.655	1.05	PQL	mg/Kg	
SL-569-SA5D-SB-0.0-0.5	CADMIUM	J	0.439	1.05	PQL	mg/Kg	J (all detects)
	TIN	J	2.39	10.5	PQL	mg/Kg	
	ARSENIC	J	3.80	4.07	PQL	mg/Kg	
	BERYLLIUM	J	0.593	1.02	PQL	mg/Kg	
SL-863-SA5D-SB-4.0-5.0	CADMIUM	J	0.114	1.02	PQL	mg/Kg	J (all detects)
	TIN	J	2.56	10.2	PQL	mg/Kg	
	BORON	J	8.83	10.6	PQL	mg/Kg	
	BERYLLIUM	J	0.560	1.06	PQL	mg/Kg	
SL-863-SA5D-SB-4.0-5.0	CADMIUM	J	0.402	1.06	PQL	mg/Kg	J (all detects)
	TIN	J	2.35	10.6	PQL	mg/Kg	
	BORON	J	8.83	10.6	PQL	mg/Kg	

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-562-SA5D-SB-0.0-0.5	SELENIUM	J	0.296	0.415	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0479	0.207	PQL	mg/Kg	
SL-562-SA5D-SB-4.0-5.0	SELENIUM	J	0.222	0.446	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0655	0.223	PQL	mg/Kg	
SL-563-SA5D-SB-0.0-0.5	SELENIUM	J	0.327	0.410	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0487	0.205	PQL	mg/Kg	
SL-563-SA5D-SB-4.0-5.0	SELENIUM	J	0.239	0.419	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0378	0.210	PQL	mg/Kg	
SL-569-SA5D-SB-0.0-0.5	SELENIUM	J	0.257	0.407	PQL	mg/Kg	J (all detects)
SL-863-SA5D-SB-4.0-5.0	SELENIUM	J	0.207	0.424	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0392	0.212	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 7471B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-562-SA5D-SB-4.0-5.0	MERCURY	J	0.0159	0.0175	PQL	mg/Kg	J (all detects)
SL-563-SA5D-SB-4.0-5.0	MERCURY	J	0.0130	0.0172	PQL	mg/Kg	J (all detects)
SL-863-SA5D-SB-4.0-5.0	MERCURY	J	0.0126	0.0176	PQL	mg/Kg	J (all detects)

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-562-SA5D-SB-0.0-0.5	EFH (C15-C20)	J	4.1	5.2	PQL	mg/Kg	J (all detects)
SL-562-SA5D-SB-4.0-5.0	EFH (C15-C20)	J	2.2	5.5	PQL	mg/Kg	J (all detects)
SL-563-SA5D-SB-0.0-0.5	EFH (C15-C20)	J	2.9	5.3	PQL	mg/Kg	J (all detects)
SL-569-SA5D-SB-0.0-0.5	EFH (C8-C11)	J	3.8	5.1	PQL	mg/Kg	J (all detects)
SL-863-SA5D-SB-4.0-5.0	EFH (C21-C30)	J	5.0	5.3	PQL	mg/Kg	J (all detects)
	EFH (C30-C40)	J	8.5	11	PQL	mg/Kg	J (all detects)

Method: 8081B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-573-SA5D-SB-0.0-0.5	ALPHA-BHC	J	0.28	0.84	PQL	ug/Kg	J (all detects)
	HEPTACHLOR	J	0.19	0.84	PQL	ug/Kg	
	HEPTACHLOR EPOXIDE	J	0.22	0.84	PQL	ug/Kg	

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-562-SA5D-SB-0.0-0.5	1-METHYLNAPHTHALENE	J	0.92	1.8	PQL	ug/Kg	J (all detects)
	2-METHYLNAPHTHALENE	J	1.6	1.8	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	0.95	1.8	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	8.1	19	PQL	ug/Kg	
	FLUORANTHENE	J	1.5	1.8	PQL	ug/Kg	
	INDENO(1,2,3-CD)PYRENE	J	0.71	1.8	PQL	ug/Kg	
	PYRENE	J	1.7	1.8	PQL	ug/Kg	
SL-562-SA5D-SB-4.0-5.0	BENZO(A)ANTHRACENE	J	1.0	1.8	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	1.5	1.8	PQL	ug/Kg	
SL-563-SA5D-SB-0.0-0.5	NAPHTHALENE	J	1.2	1.8	PQL	ug/Kg	J (all detects)
SL-569-SA5D-SB-0.0-0.5	CHRYSENE	J	0.99	1.7	PQL	ug/Kg	J (all detects)
	NAPHTHALENE	J	0.79	1.7	PQL	ug/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH114

Laboratory: LL

EDD Filename: PH114

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-573-SA5D-SB-0.0-0.5	2-METHYLNAPHTHALENE	J	0.77	1.7	PQL	ug/Kg	J (all detects)
	BENZO(B)FLUORANTHENE	J	1.3	1.7	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	0.69	1.7	PQL	ug/Kg	
	CHRYSENE	J	1.2	1.7	PQL	ug/Kg	
	FLUORANTHENE	J	1.1	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	1.1	1.7	PQL	ug/Kg	
	PYRENE	J	0.98	1.7	PQL	ug/Kg	

LDC #: 30695Q4

VALIDATION COMPLETENESS WORKSHEET

Date: 11/11/13

SDG #: PH114

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: OR

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	-	Sampling dates: 9/25/13
II.	ICP/MS Tune	-	
III.	Calibration	-	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	SW MS/D	
VII.	Duplicate Sample Analysis	SW Dup	
VIII.	Laboratory Control Samples (LCS)	A LCS	
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	SA	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	N	
XV.	Field Blanks	SW EB=1 FB=PH11613	

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

(PH1032)

Validated Samples:

1	EB1-092513	11		21		31	
2	SL-569-SA5D-SB-0.0-0.5	12		22		32	
3	SL-562-SA5D-SB-0.0-0.5	13		23		33	
4	SL-562-SA5D-SB-4.0-5.0	14		24		34	
5	SL-563-SA5D-SB-0.0-0.5	15		25		35	
6	SL-563-SA5D-SB-4.0-5.0	16		26		36	
7	SL-863-SA5D-SB-4.0-5.0	17		27		37	
8	SL-563-SA5D-SB-4.0-5.0MS	18		28		38	
9	SL-563-SA5D-SB-4.0-5.0MSD	19		29		39	
10	SL-563-SA5D-SB-4.0-5.0DUP	20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All Soil

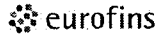
Analyte	Blank ID	Sample Identification												
	FB-041613 (SDG: PH032)	Action Limit	3	4										
Mo	0.0132	6.60	0.184	0.213										
Sn	0.0029	1.45												

Sampling date: 9/25/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All Soil

Analyte	Blank ID	Sample Identification												
	EB1-092513 (SDG: PH114)	Action Limit	No Qualifiers											
Ca	0.0498	24.9												

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".



Lancaster Laboratories

Redout qual J/05/1A

QUALITY ASSURANCE SUMMARY

FORM 5A (MS/MSD)

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

SDG No.: PH114

Matrix: SOIL

Level

(low/med):

LOW

Background Lab Sample ID: 7213412BKG Matrix Spike Lab Sample ID: 7213413MS Matrix Spike Duplicate Lab Sample ID: 7213414MSD
Batch Id(s): P27437B, P27438B

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit					
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q	%R	RPD	M	
Aluminum		17878.5447		27231.1314		22142.2410		196.0784	200.0000	MG/KG	4770		2132				20	P		
Antimony		0.7184	U	23.2598		24.0140		49.0196	50.0000	MG/KG	47	N	48	N	3		75 - 125	20	P	
Arsenic		4.0563		18.1157		20.2310		14.7059	15.0000	MG/KG	96		108		11		75 - 125	20	P	
Barium		69.4544		274.0500		262.8070		196.0784	200.0000	MG/KG	104		97		4		75 - 125	20	P	
Beryllium		0.6068	B	5.4039		5.2930		4.9020	5.0000	MG/KG	98		94		2		75 - 125	20	P	
Boron		9.7126		202.7608		209.2510		196.0784	200.0000	MG/KG	98		100		3		75 - 125	20	P	
Cadmium		0.4068	B	4.7637		4.9160		4.9020	5.0000	MG/KG	89		90		3		75 - 125	20	P	
Calcium		103698.6214		98722.0608		154379.6500		392.1569	400.0000	MG/KG	-1269		12670						20	P
Chromium		22.8583		46.0098		46.2520		19.6078	20.0000	MG/KG	118		117		1		75 - 125	20	P	
Cobalt		9.2485		52.5167		52.3060		49.0196	50.0000	MG/KG	88		86		0		75 - 125	20	P	
Copper		13.6282		39.8294		34.2550		24.5098	25.0000	MG/KG	107		83		15		75 - 125	20	P	
Iron		24877.7913		27568.1686		18683.2090		98.0392	100.0000	MG/KG	2744		-6195						20	P
Lead		9.2612		22.7373		20.7950		14.7059	15.0000	MG/KG	92		77		9		75 - 125	20	P	
Lithium		24.2932		127.4765		124.0170		98.0392	100.0000	MG/KG	105		100		3		75 - 125	20	P	
Magnesium		6459.1845		7882.1588		6281.0860		196.0784	200.0000	MG/KG	726		-89						20	P
Manganese		280.4961		307.9814		390.0880		49.0196	50.0000	MG/KG	56		219						20	P
Mercury		0.0120	B	0.1861		0.1780		0.1582	0.1579	MG/KG	110		105		4		65 - 135	20	CV	
Molybdenum		0.1650	U	183.9118		192.3370		196.0784	200.0000	MG/KG	94		96		4		75 - 125	20	P	
Nickel		14.7039		60.1461		57.0660		49.0196	50.0000	MG/KG	93		85		5		75 - 125	20	P	
Phosphorus		482.6728		595.0990		775.3080		98.0392	100.0000	MG/KG	115		293						20	P
Potassium		3039.9311		4926.9382		4317.3800		980.3922	1000.0000	MG/KG	192	N	128	N	13		75 - 125	20	P	
Selenium	78	0.2210	B	2.5706		2.2160		1.9608	2.0000	MG/KG	120		100		15		75 - 125	20	MS	
Silver	107	0.0350	B	14.2294		10.8400		9.8039	10.0000	MG/KG	145	N	108		27	*	75 - 125	20	MS	
Sodium		135.6126		1104.5265		1159.6700		980.3922	1000.0000	MG/KG	99		102		5		75 - 125	20	P	
Strontium	88	190.3883		215.2941		266.4000		7.8431	8.0000	MG/KG	318		950						20	MS
Thallium	203	0.2680		0.9514		0.6850		0.3922	0.4000	MG/KG	174	N	104		33	*	75 - 125	20	MS	

Note: Results shown are reported on an as-received basis.

METHODS:

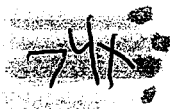
P = ICP Atomic Emission Spectrometer CV = Cold Vapor
MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence

CONCENTRATION QUALIFIERS:

U = Below MDL, B = Below LOQ

FLAGS:

N = Matrix Spike OOS, * = Duplicate OOS



Background Lab Sample ID: 7213412BKG Matrix Spike Lab Sample ID: 7213413MS Matrix Spike Duplicate Lab Sample ID: 7213414MSD
 Batch Id(s): P27437B, P27438B

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit				
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q	%R	RPD	M
Tin		2.2155	B	350.3735		363.5780		392.1569	400.0000	MG/KG	89		90		4	75 - 125	20	P	
Titanium		808.8485		1447.6000		1220.7000		98.0392	100.0000	MG/KG	652		412		17			20	P
Vanadium		41.7117		103.8686		99.4470		49.0196	50.0000	MG/KG	127	N	115		4	75 - 125	20	P	
Zinc		65.0524		115.5784		99.4290		49.0196	50.0000	MG/KG	103		69	N	15	75 - 125	20	P	
Zirconium		6.6010		100.9382		103.2720		98.0392	100.0000	MG/KG	96		97		2	75 - 125	20	P	

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U = Below MDL, B = Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS
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Background Lab Sample ID: 7213412BKG
 Batch ID(s): P27437B, P27438B
 Concentration Units: MG/KG

Duplicate Lab Sample ID: 7213415DUP

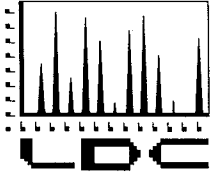
Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			17878.5447		19147.7243		7		P
Antimony			0.7184	U	0.7184	U			P
Arsenic		3.9	4.0563		3.7330	B	8		P
Barium			69.4544		68.7670		1		P
Beryllium			0.6068	B	0.6252	B	3		P
Boron		9.7	9.7126		10.4825		8		P
Cadmium			0.4068	B	0.3951	B	3		P
Calcium			103698.6214		106029.2816		2		P
Chromium			22.8583		23.9330		5		P
Cobalt			9.2485		8.1777		12		P
Copper			13.6282		13.8417		2		P
Iron			24877.7913		26053.1476		5		P
Lead		2.9	9.2612		9.3175		1		P
Lithium			24.2932		24.9107		3		P
Magnesium			6459.1845		6621.6670		2		P
Manganese			280.4961		274.6301		2		P
Mercury			0.0120	B	0.0104	B	14		CV
Molybdenum			0.1650	U	0.1650	U			P
Nickel			14.7039		15.0544		2		P
Phosphorus			482.6728		481.7631		0		P
Potassium			3039.9311		3135.3010		3		P
Selenium	78		0.2210	B	0.3340	B	41		MS
Silver	107		0.0350	B	0.0466	B	28		MS
Sodium		97.1	135.6126		138.8379		2		P
Strontium	88		190.3883		202.2330		6		MS
Thallium	203	0.2	0.2680		0.3656		31		MS
Tin			2.2155	B	2.3301	B	5		P
Titanium			808.8485		969.5320		18		P
Vanadium			41.7117		45.0845		8		P
Zinc			65.0524		65.7515		1		P
Zirconium		4.9	6.6010		6.9291		5		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).
 The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

ok by difference

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: * = Duplicate Out of Spec
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LABORATORY DATA CONSULTANTS, INC.

2701 Loker Ave. West, Suite 220, Carlsbad, CA 92010 Bus: 760-827-1100 Fax: 760-827-1099

CDM
555 17th Street, Suite 1100
Denver, CO 80202
ATTN: Mrs. Cherie Zakowski

December 23, 2013

SUBJECT: Santa Susana Field Laboratory, Subarea 5D Data Validation

Dear Mrs. Zakowski,

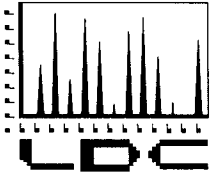
Enclosed are the final validation reports for the fractions listed below. These SDGs were received on November 13, 2013. Attachment 1 is a summary of the samples that were reviewed for each analysis.

LDC Project # 30842:

<u>SDG #</u>	<u>Fraction</u>
PH116, PH118 PH119, PH120 PH122, PH123	Semivolatiles, Chlorinated Pesticides, Polychlorinated Biphenyls, Metals, Herbicides, Wet Chemistry, Total Petroleum Hydrocarbons as Gasoline, Total Petroleum Hydrocarbons as Extractables, Dioxins/Dibenzofurans, Hexavalent Chromium, Fluoride

The data validation was performed under Level III & IV guidelines. The analyses were validated using the following documents, as applicable to each method:

- Quality Assurance Project Plan for Santa Susana Field Laboratory, RCRA Facility Investigation, Surficial Media Operable Unit, March 2009, Revision 4
- USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review, June 2008
- USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review, January 2010
- Polychlorinated Dioxins/Dibenzofurans Data Review, September 2005
- EPA SW 846, Third Edition, Test Methods for Evaluating Solid Waste, update 1, July 1992; update IIA, August 1993; update II, September 1994; update IIB, January 1995; update III, December 1996; update IIIA, April 1998; IIIB, November 2004; Update IV, February 2007



Please feel free to contact us if you have any questions.

Sincerely,

A handwritten signature in black ink, appearing to read 'S. McKellar', written in a cursive style.

Shauna McKellar
Project Manager/Chemist

90/10 ADR/IV LDC #30842 (CDM Federal Programs-Chantilly VA / Santa Susana Field Laboratory, Subarea 5D)

LDC	SDG#	DATE REC'D	(4) DATE DUE	SVOA (8270D -SIM)		Pest. (8081B)		PCBs (8082A)		Metals & Hg (SW846)		Herbs. (8151A)		TPH-G (8015M)		TPH-E (8015M)		Dioxins (1613B)		Cr(VI) (7199)		F (300.0)															
				W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S
A	PH116	11/13/13	12/13/13	0	2	0	2	0	2	0	2	-	-	1	1	0	2	0	1	-	-	-	-														
B	PH118	11/13/13	12/13/13	0	4	-	-	0	4	0	4	0	4	1	3	0	4	0	1	-	-	0	13														
C	PH119	11/13/13	12/13/13	0	6	0	2	0	6	0	6	0	2	1	3	0	6	0	3	-	-	0	5														
D	PH120	11/13/13	12/13/13	1	2	1	2	1	2	1	2	1	0	2	1	1	2	1	1	1	0	1	9														
E	PH122	11/13/13	12/13/13	0	13	-	-	0	13	0	13	-	-	1	9	0	13	0	4	-	-	-	-														
F	PH123	11/13/13	12/13/13	0	4	0	2	0	2	0	2	0	4	1	1	0	2	0	1	-	-	0	2														
Total	T/SM			1	31	1	8	1	29	1	29	1	10	7	18	1	29	1	11	1	0	1	29	0	0	0	0	0	0	0	0	0	0	0	0	0	210

Shaded cells indicate Level IV validation (all other cells are ADR review). These sample counts do not include MS/MSD, and DUPs

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH116

Prepared for

CDM Smith
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
2701 Loker Ave West, Suite 220
Carlsbad, California 92010

December 19, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on September 26, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by Environmental Protection Agency (EPA) SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)

Pesticides by EPA SW 846 8081B

Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A

Metals by EPA SW 846 Method 6010C, 6020A and 7471B

Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M

TPH as Extractables by EPA SW 846 Method 8015M

Dioxins and Dibenzofurans by EPA Method 1613B

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), method blanks, trip blanks, equipment blanks and field blanks. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VIII. Laboratory Duplicates Sample

The laboratory has indicated that there were no laboratory duplicate (DUP) analyses specified for the samples in this SDG, and therefore laboratory duplicate analyses were not performed for this SDG.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution was not performed for this SDG.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH116	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

No field duplicates were identified in this SDG.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No volatile contaminants were found in the trip blank.

One equipment blank (from SDG PH114) was collected and analyzed for SVOCs, pesticides, PCBs, metals, TPH as gasoline, TPH as extractables and dioxins. The equipment blank had detections for SVOCs, metals and dioxins. The associated sample results were not detected or were significantly greater than the concentrations found in the equipment blank, therefore no data were qualified.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, pesticides, PCBs, metals, TPH as gasoline, TPH as extractables and dioxins. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were not detected or were significantly greater than the concentrations found in the field blank, therefore no data were qualified.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1

Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
26-Sep-2013	SL-512-SA5D-SB-0.0-0.5	7215511	N	3050B	6010C	III
26-Sep-2013	SL-512-SA5D-SB-0.0-0.5	7215511	N	3050B	6020A	III
26-Sep-2013	SL-512-SA5D-SB-0.0-0.5	7215511	N	3546	8015M	III
26-Sep-2013	SL-512-SA5D-SB-0.0-0.5	7215511	N	3546	8081B	III
26-Sep-2013	SL-512-SA5D-SB-0.0-0.5	7215511	N	3546	8082A	III
26-Sep-2013	SL-512-SA5D-SB-0.0-0.5	7215511	N	3546	8270D SIM	III
26-Sep-2013	SL-512-SA5D-SB-0.0-0.5	7215511	N	METHOD	1613B	III
26-Sep-2013	SL-512-SA5D-SB-0.0-0.5	7215511	N	METHOD	7471B	III
26-Sep-2013	SL-512-SA5D-SB-0.0-0.5MSD	P215511M241040A	MSD	3546	8081B	III
26-Sep-2013	SL-512-SA5D-SB-0.0-0.5MS	P215511R241025A	MS	3546	8081B	III
26-Sep-2013	TB1-092613	7215510	TB	5030B	8015M	III
26-Sep-2013	SL-512-SA5D-SB-4.0-5.0	7215512	N	3050B	6010C	III
26-Sep-2013	SL-512-SA5D-SB-4.0-5.0	7215512	N	3050B	6020A	III
26-Sep-2013	SL-512-SA5D-SB-4.0-5.0	7215512	N	3546	8015M	III
26-Sep-2013	SL-512-SA5D-SB-4.0-5.0	7215512	N	3546	8081B	III
26-Sep-2013	SL-512-SA5D-SB-4.0-5.0	7215512	N	3546	8082A	III
26-Sep-2013	SL-512-SA5D-SB-4.0-5.0	7215512	N	3546	8270D SIM	III
26-Sep-2013	SL-512-SA5D-SB-4.0-5.0	7215512	N	5035A	8015M	III
26-Sep-2013	SL-512-SA5D-SB-4.0-5.0	7215512	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH116

Laboratory: LL

EDD Filename: PrepPH116_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-512-SA5D-SB-0.0-0.5 Collected: 9/26/2013 7:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.948	J	0.0689	MDL	1.03	PQL	mg/Kg	J	Z
BORON	5.74	J	0.864	MDL	10.3	PQL	mg/Kg	J	Z
CADMIUM	0.347	J	0.0782	MDL	1.03	PQL	mg/Kg	J	Z
Zirconium	3.68	J	0.864	MDL	5.14	PQL	mg/Kg	J	Z
TIN	2.97	J	0.226	MDL	10.3	PQL	mg/Kg	U	B

Sample ID: SL-512-SA5D-SB-4.0-5.0 Collected: 9/26/2013 8:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	1.03	J	0.0750	MDL	1.12	PQL	mg/Kg	J	Z
BORON	7.63	J	0.940	MDL	11.2	PQL	mg/Kg	J	Z
CADMIUM	0.270	J	0.0851	MDL	1.12	PQL	mg/Kg	J	Z
TIN	3.61	J	0.246	MDL	11.2	PQL	mg/Kg	U	B
Zirconium	4.72	J	0.940	MDL	5.60	PQL	mg/Kg	J	Z

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-512-SA5D-SB-0.0-0.5 Collected: 9/26/2013 7:35:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.198	J	0.103	MDL	0.411	PQL	mg/Kg	J	Z

Sample ID: SL-512-SA5D-SB-0.0-0.5 Collected: 9/26/2013 7:35:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0482	J	0.0267	MDL	0.206	PQL	mg/Kg	J	Z

Sample ID: SL-512-SA5D-SB-4.0-5.0 Collected: 9/26/2013 8:00:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.148	J	0.112	MDL	0.448	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH116

Laboratory: LL

EDD Filename: PrepPH116_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6020A **Matrix: SO**

Sample ID: SL-512-SA5D-SB-4.0-5.0 Collected: 9/26/2013 8:00:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0436	J	0.0291	MDL	0.224	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 1613B **Matrix: SO**

Sample ID: SL-512-SA5D-SB-0.0-0.5 Collected: 9/26/2013 7:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.273	JB	0.0203	MDL	5.12	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0996	JBQ	0.00731	MDL	5.12	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0440	JB	0.0106	MDL	5.12	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0321	JBQ	0.0213	MDL	5.12	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.0538	JBQ	0.0142	MDL	5.12	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.143	JB	0.0230	MDL	5.12	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.0675	JBQ	0.0128	MDL	5.12	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.307	JBQ	0.0206	MDL	5.12	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDF	0.367	JB	0.0133	MDL	5.12	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0551	JB	0.0288	MDL	5.12	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.215	JBQ	0.0163	MDL	5.12	PQL	ng/Kg	U	B
2,3,4,6,7,8-HxCDF	0.0367	JBQ	0.0127	MDL	5.12	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0925	JBQ	0.0161	MDL	5.12	PQL	ng/Kg	U	B
OCDD	1.70	JB	0.0195	MDL	10.2	PQL	ng/Kg	J	Z
OCDF	0.201	JB	0.0273	MDL	10.2	PQL	ng/Kg	U	B

Method Category: SVOA
Method: 8270D SIM **Matrix: SO**

Sample ID: SL-512-SA5D-SB-0.0-0.5 Collected: 9/26/2013 7:35:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.76	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(A)ANTHRACENE	0.76	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH116

Laboratory: LL

EDD Filename: PrepPH116_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-512-SA5D-SB-0.0-0.5 Collected: 9/26/2013 7:35:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	1.2	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	0.99	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z
FLUORANTHENE	1.2	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	0.84	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
PYRENE	1.2	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-512-SA5D-SB-4.0-5.0 Collected: 9/26/2013 8:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	1.0	J	0.76	MDL	1.9	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH116

Laboratory: LL

EDD Filename: PrepPH116_v1

eQAPP Name: CDM_SSFL_131101_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Method Blank Contamination
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

12/18/2013 2:29:13 PM

ADR version 1.7.0.207

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH116

Method Blank Outlier Report

Lab Reporting Batch ID: PH116

Laboratory: LL

EDD Filename: PrepPH116_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2730B371140	10/2/2013 11:40:00 AM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF OCDD OCDF	0.0965 ng/Kg 0.0595 ng/Kg 0.0657 ng/Kg 0.0334 ng/Kg 0.0515 ng/Kg 0.0256 ng/Kg 0.0453 ng/Kg 0.0369 ng/Kg 0.0754 ng/Kg 0.0738 ng/Kg 0.0876 ng/Kg 0.0332 ng/Kg 0.0646 ng/Kg 0.249 ng/Kg 0.137 ng/Kg	SL-512-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-512-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.273 ng/Kg	0.273U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0996 ng/Kg	0.0996U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0440 ng/Kg	0.0440U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0321 ng/Kg	0.0321U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.0538 ng/Kg	0.0538U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0675 ng/Kg	0.0675U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.367 ng/Kg	0.367U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.0551 ng/Kg	0.0551U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.215 ng/Kg	0.215U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0367 ng/Kg	0.0367U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.0925 ng/Kg	0.0925U ng/Kg
SL-512-SA5D-SB-0.0-0.5(RES)	OCDF	0.201 ng/Kg	0.201U ng/Kg

Method: 6010C

Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P27437AB221412	10/3/2013 2:12:00 PM	CALCIUM TIN ZINC	9.48 mg/Kg 1.46 mg/Kg 0.850 mg/Kg	SL-512-SA5D-SB-0.0-0.5 SL-512-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-512-SA5D-SB-0.0-0.5(RES)	TIN	2.97 mg/Kg	2.97U mg/Kg
SL-512-SA5D-SB-4.0-5.0(RES)	TIN	3.61 mg/Kg	3.61U mg/Kg

Reporting Limit Outliers

Lab Reporting Batch ID: PH116

Laboratory: LL

EDD Filename: PrepPH116_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-512-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	0.273	5.12	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0996	5.12	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0440	5.12	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0321	5.12	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0538	5.12	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.143	5.12	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0675	5.12	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.307	5.12	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.367	5.12	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.0551	5.12	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.215	5.12	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0367	5.12	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0925	5.12	PQL	ng/Kg	
	OCDD	JB	1.70	10.2	PQL	ng/Kg	
OCDF	JB	0.201	10.2	PQL	ng/Kg		

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-512-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.948	1.03	PQL	mg/Kg	J (all detects)
	BORON	J	5.74	10.3	PQL	mg/Kg	
	CADMIUM	J	0.347	1.03	PQL	mg/Kg	
	TIN	J	2.97	10.3	PQL	mg/Kg	
	Zirconium	J	3.68	5.14	PQL	mg/Kg	
SL-512-SA5D-SB-4.0-5.0	BERYLLIUM	J	1.03	1.12	PQL	mg/Kg	J (all detects)
	BORON	J	7.63	11.2	PQL	mg/Kg	
	CADMIUM	J	0.270	1.12	PQL	mg/Kg	
	TIN	J	3.61	11.2	PQL	mg/Kg	
	Zirconium	J	4.72	5.60	PQL	mg/Kg	

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-512-SA5D-SB-0.0-0.5	SELENIUM	J	0.198	0.411	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0482	0.206	PQL	mg/Kg	
SL-512-SA5D-SB-4.0-5.0	SELENIUM	J	0.148	0.448	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0436	0.224	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH116

Laboratory: LL

EDD Filename: PrepPH116_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-512-SA5D-SB-0.0-0.5	2-METHYLNAPHTHALENE	J	0.76	1.8	PQL	ug/Kg	J (all detects)
	BENZO(A)ANTHRACENE	J	0.76	1.8	PQL	ug/Kg	
	BENZO(B)FLUORANTHENE	J	1.2	1.8	PQL	ug/Kg	
	CHRYSENE	J	0.99	1.8	PQL	ug/Kg	
	FLUORANTHENE	J	1.2	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	0.84	1.8	PQL	ug/Kg	
	PYRENE	J	1.2	1.8	PQL	ug/Kg	
SL-512-SA5D-SB-4.0-5.0	NAPHTHALENE	J	1.0	1.9	PQL	ug/Kg	J (all detects)

LDC #: 30842A4

VALIDATION COMPLETENESS WORKSHEET

Date: 12/4/13

SDG #: PH116

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: AL

2nd Reviewer: W

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	—	Sampling dates: 9/26/13
II.	ICP/MS Tune	—	
III.	Calibration	—	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	—	
VI.	Matrix Spike Analysis	N	CS
VII.	Duplicate Sample Analysis	N	↓
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	N	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	—	
XV.	Field Blanks	SW	FB = FB-011613 EB = EB1-092513

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

(PH032)

(PH114)

Validated Samples:

so.1

1	SL-512-SA5D-SB-0.0-0.5	11		21		31	
2	SL-512-SA5D-SB-4.0-5.0	12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: Associated Samples: All

Analyte	Blank ID	Sample Identification														
	FB-041613 (SDG: PH032)	Action Limit	No Qualifiers													
Mo	0.0132	6.60														
Sn	0.0029	1.45														

Sampling date: 9/25/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: Associated Samples: All

Analyte	Blank ID	Sample Identification														
	EB1-092513 (SDG: PH114)	Action Limit	No Qualifiers													
Ca	0.0498	24.9														

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH118

Prepared for

CDM Smith
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
2701 Loker Ave West, Suite 220
Carlsbad, California 92010

December 23, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on September 30, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by Environmental Protection Agency (EPA) SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A and 7471B
Herbicides by EPA SW 846 Method 8151A
Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M
TPH as Extractables by EPA SW 846 Method 8015M
Dioxins and Dibenzofurans by EPA Method 1613B

Wet Chemistry:

Fluoride by EPA Method 300.0

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks and field duplicates. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met with the exception of one sample for TPH as gasoline. The associated sample result was qualified as non-detected estimated (UJ). The details regarding the qualification of data are provided in Enclosure I.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of several MS/MSD pairs for metals and fluoride. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one LCS/LCSD pair for herbicides. The dinoseb result in samples SL-582-SA5D-SB-0.0-0.5, SL-582-SA5D-SB-10.0-11.0, SL-582-SA5D-SB-4.0-5.0 and SL-882-SA5D-SB-4.0-5.0 were qualified as rejected (R) due to LCS/LCSD %Rs grossly outside QC limits (i.e., $\leq 10\%$). The details regarding the qualification of data are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH118	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pair was collected and analyzed for SVOCs, PCBs, metals, herbicides, TPH as gasoline and TPH as extractables. All RPDs were within QC limits. The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No volatile contaminants were found in the trip blank.

One equipment blank (from SDG PH120) was collected and analyzed for SVOCs, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins and fluoride. The equipment blank had detections for SVOCs, metals and dioxins. The associated sample results were not detected or were significantly greater than the concentrations found in the equipment blank, therefore no data were qualified.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins and fluoride. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were not detected or were significantly greater than the concentrations found in the field blank, therefore no data were qualified.

XV. Overall Assessment of Data

The dinoseb results in four samples were rejected due to LCS/LCSD %Rs grossly outside of QC limits. These results are not useable for all purposes.

The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. The remainder of the data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
30-Sep-2013	TB-093013	7218893	TB	5030B	8015M	III
30-Sep-2013	SL-582-SA5D-SB-0.0-0.5	7218894	N	3050B	6010C	III
30-Sep-2013	SL-582-SA5D-SB-0.0-0.5	7218894	N	3050B	6020A	III
30-Sep-2013	SL-582-SA5D-SB-0.0-0.5	7218894	N	3546	8015M	III
30-Sep-2013	SL-582-SA5D-SB-0.0-0.5	7218894	N	3546	8082A	III
30-Sep-2013	SL-582-SA5D-SB-0.0-0.5	7218894	N	3546	8270D SIM	III
30-Sep-2013	SL-582-SA5D-SB-0.0-0.5	7218894	N	3550B	8151A	III
30-Sep-2013	SL-582-SA5D-SB-0.0-0.5	7218894	N	METHOD	1613B	III
30-Sep-2013	SL-582-SA5D-SB-0.0-0.5	7218894	N	METHOD	7471B	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0	7218895	N	3050B	6010C	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0	7218895	N	3050B	6020A	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0	7218895	N	3546	8015M	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0	7218895	N	3546	8082A	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0	7218895	N	3546	8270D SIM	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0	7218895	N	3550B	8151A	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0	7218895	N	5035A	8015M	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0	7218895	N	METHOD	7471B	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MS	7218896	MS	3050B	6010C	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MS	7218896	MS	3050B	6020A	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MS	7218896	MS	3546	8015M	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MS	7218896	MS	3546	8082A	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MS	7218896	MS	3546	8270D SIM	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MS	7218896	MS	3550B	8151A	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MS	7218896	MS	5035A	8015M	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MS	7218896	MS	METHOD	7471B	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MSD	7218897	MSD	3050B	6010C	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MSD	7218897	MSD	3050B	6020A	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MSD	7218897	MSD	3546	8015M	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MSD	7218897	MSD	3546	8082A	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MSD	7218897	MSD	3546	8270D SIM	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MSD	7218897	MSD	3550B	8151A	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MSD	7218897	MSD	5035A	8015M	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0MSD	7218897	MSD	METHOD	7471B	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0DUP	7218898	DUP	3050B	6010C	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0DUP	7218898	DUP	3050B	6020A	III
30-Sep-2013	SL-582-SA5D-SB-4.0-5.0DUP	7218898	DUP	METHOD	7471B	III
30-Sep-2013	SL-882-SA5D-SB-4.0-5.0	7218899	FD	3050B	6010C	III
30-Sep-2013	SL-882-SA5D-SB-4.0-5.0	7218899	FD	3050B	6020A	III
30-Sep-2013	SL-882-SA5D-SB-4.0-5.0	7218899	FD	3546	8015M	III
30-Sep-2013	SL-882-SA5D-SB-4.0-5.0	7218899	FD	3546	8082A	III
30-Sep-2013	SL-882-SA5D-SB-4.0-5.0	7218899	FD	3546	8270D SIM	III
30-Sep-2013	SL-882-SA5D-SB-4.0-5.0	7218899	FD	3550B	8151A	III
30-Sep-2013	SL-882-SA5D-SB-4.0-5.0	7218899	FD	5035A	8015M	III
30-Sep-2013	SL-882-SA5D-SB-4.0-5.0	7218899	FD	METHOD	7471B	III
30-Sep-2013	SL-582-SA5D-SB-10.0-11.0	7218900	N	3050B	6010C	III
30-Sep-2013	SL-582-SA5D-SB-10.0-11.0	7218900	N	3050B	6020A	III
30-Sep-2013	SL-582-SA5D-SB-10.0-11.0	7218900	N	3546	8015M	III
30-Sep-2013	SL-582-SA5D-SB-10.0-11.0	7218900	N	3546	8082A	III
30-Sep-2013	SL-582-SA5D-SB-10.0-11.0	7218900	N	3546	8270D SIM	III
30-Sep-2013	SL-582-SA5D-SB-10.0-11.0	7218900	N	3550B	8151A	III
30-Sep-2013	SL-582-SA5D-SB-10.0-11.0	7218900	N	5035A	8015M	III
30-Sep-2013	SL-582-SA5D-SB-10.0-11.0	7218900	N	METHOD	7471B	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
30-Sep-2013	SL-595-SA5D-SB-0.0-0.5	7218901	N	METHOD	300.0	III
30-Sep-2013	SL-595-SA5D-SB-0.0-0.5DUP	P218901D271803A	DUP	METHOD	300.0	III
30-Sep-2013	SL-595-SA5D-SB-0.0-0.5MS	P218901R272125A	MS	METHOD	300.0	III
30-Sep-2013	SL-595-SA5D-SB-4.0-5.0	7218902	N	METHOD	300.0	III
30-Sep-2013	SL-595-SA5D-SB-9.0-10.0	7218903	N	METHOD	300.0	III
30-Sep-2013	SL-595-SA5D-SB-14.0-15.0	7218904	N	METHOD	300.0	III
30-Sep-2013	SL-595-SA5D-SB-17.0-18.0	7218905	N	METHOD	300.0	III
30-Sep-2013	SL-596-SA5D-SB-0.0-0.5	7218906	N	METHOD	300.0	III
30-Sep-2013	SL-596-SA5D-SB-4.0-5.0	7218907	N	METHOD	300.0	III
30-Sep-2013	SL-596-SA5D-SB-11.5-12.5	7218908	N	METHOD	300.0	III
30-Sep-2013	SL-597-SA5D-SB-0.0-0.5	7218909	N	METHOD	300.0	III
30-Sep-2013	SL-597-SA5D-SB-4.0-5.0	7218910	N	METHOD	300.0	III
30-Sep-2013	SL-597-SA5D-SB-9.0-10.0	7218911	N	METHOD	300.0	III
30-Sep-2013	SL-597-SA5D-SB-9.0-10.0DU	P218911D272045B	DUP	METHOD	300.0	III
30-Sep-2013	SL-597-SA5D-SB-9.0-10.0MS	P218911R270121B	MS	METHOD	300.0	III
30-Sep-2013	SL-597-SA5D-SB-14.0-15.0	7218912	N	METHOD	300.0	III
30-Sep-2013	SL-597-SA5D-SB-17.0-18.0	7218913	N	METHOD	300.0	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: GENCHEM
Method: 300.0 **Matrix:** SO

Sample ID: SL-595-SA5D-SB-0.0-0.5 Collected: 9/30/2013 9:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.0		0.44	MDL	1.1	PQL	mg/Kg	J	Q

Sample ID: SL-595-SA5D-SB-14.0-15.0 Collected: 9/30/2013 10:20:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	8.9		0.46	MDL	1.2	PQL	mg/Kg	J	Q

Sample ID: SL-595-SA5D-SB-17.0-18.0 Collected: 9/30/2013 10:30:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	5.3		0.92	MDL	2.3	PQL	mg/Kg	J	Q

Sample ID: SL-595-SA5D-SB-4.0-5.0 Collected: 9/30/2013 10:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.2		0.44	MDL	1.1	PQL	mg/Kg	J	Q

Sample ID: SL-595-SA5D-SB-9.0-10.0 Collected: 9/30/2013 10:10:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	6.1		0.93	MDL	2.3	PQL	mg/Kg	J	Q

Sample ID: SL-596-SA5D-SB-0.0-0.5 Collected: 9/30/2013 12:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.1		0.43	MDL	1.1	PQL	mg/Kg	J	Q

Sample ID: SL-596-SA5D-SB-11.5-12.5 Collected: 9/30/2013 12:55:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.5		0.90	MDL	2.2	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: GENCHEM
Method: 300.0 **Matrix:** SO

Sample ID: SL-596-SA5D-SB-4.0-5.0		Collected: 9/30/2013 12:45:00		Analysis Type: RES				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	1.9	J	0.89	MDL	2.2	PQL	mg/Kg	J	Z, Q

Sample ID: SL-597-SA5D-SB-0.0-0.5		Collected: 9/30/2013 1:40:00		Analysis Type: RES				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.7		0.88	MDL	2.2	PQL	mg/Kg	J	Q

Sample ID: SL-597-SA5D-SB-14.0-15.0		Collected: 9/30/2013 2:10:00		Analysis Type: RES				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.9		0.89	MDL	2.2	PQL	mg/Kg	J	Q

Sample ID: SL-597-SA5D-SB-17.0-18.0		Collected: 9/30/2013 2:15:00		Analysis Type: RES				Dilution: 1	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	3.5		0.44	MDL	1.1	PQL	mg/Kg	J	Q

Sample ID: SL-597-SA5D-SB-4.0-5.0		Collected: 9/30/2013 1:50:00		Analysis Type: RES				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.8		0.89	MDL	2.2	PQL	mg/Kg	J	Q

Sample ID: SL-597-SA5D-SB-9.0-10.0		Collected: 9/30/2013 2:00:00		Analysis Type: RES				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	4.2		0.90	MDL	2.3	PQL	mg/Kg	J	Q

* denotes a non-reportable result

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Data Qualifier Summary

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-582-SA5D-SB-0.0-0.5 Collected: 9/30/2013 8:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.15	U	0.767	MDL	4.15	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.816	J	0.0694	MDL	1.04	PQL	mg/Kg	J	Z
BORON	6.31	J	0.871	MDL	10.4	PQL	mg/Kg	J	Z
POTASSIUM	5110		8.64	MDL	104	PQL	mg/Kg	J	Q
SODIUM	98.2	J	17.3	MDL	104	PQL	mg/Kg	J	Z
TIN	2.86	J	0.228	MDL	10.4	PQL	mg/Kg	U	B

Sample ID: SL-582-SA5D-SB-10.0-11.0 Collected: 9/30/2013 9:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.43	U	0.820	MDL	4.43	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.765	J	0.0742	MDL	1.11	PQL	mg/Kg	J	Z
BORON	5.51	J	0.930	MDL	11.1	PQL	mg/Kg	J	Z
POTASSIUM	3380		9.24	MDL	111	PQL	mg/Kg	J	Q
TIN	2.93	J	0.244	MDL	11.1	PQL	mg/Kg	U	B

Sample ID: SL-582-SA5D-SB-4.0-5.0 Collected: 9/30/2013 8:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.49	U	0.830	MDL	4.49	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.771	J	0.0751	MDL	1.12	PQL	mg/Kg	J	Z
BORON	5.70	J	0.942	MDL	11.2	PQL	mg/Kg	J	Z
POTASSIUM	3880		9.35	MDL	112	PQL	mg/Kg	J	Q
TIN	2.98	J	0.247	MDL	11.2	PQL	mg/Kg	U	B

Sample ID: SL-882-SA5D-SB-4.0-5.0 Collected: 9/30/2013 9:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.39	U	0.813	MDL	4.39	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.789	J	0.0736	MDL	1.10	PQL	mg/Kg	J	Z
BORON	6.43	J	0.923	MDL	11.0	PQL	mg/Kg	J	Z
POTASSIUM	4170		9.16	MDL	110	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-882-SA5D-SB-4.0-5.0	Collected: 9/30/2013 9:00:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
TIN	2.85	J	0.242	MDL	11.0	PQL	mg/Kg	U	B

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-582-SA5D-SB-0.0-0.5	Collected: 9/30/2013 8:45:00	Analysis Type: REA	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.277	J	0.104	MDL	0.415	PQL	mg/Kg	J	Z

Sample ID: SL-582-SA5D-SB-0.0-0.5	Collected: 9/30/2013 8:45:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0311	J	0.0269	MDL	0.207	PQL	mg/Kg	J	Z

Sample ID: SL-582-SA5D-SB-10.0-11.0	Collected: 9/30/2013 9:10:00	Analysis Type: REA	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.150	J	0.111	MDL	0.443	PQL	mg/Kg	J	Z

Sample ID: SL-582-SA5D-SB-4.0-5.0	Collected: 9/30/2013 8:55:00	Analysis Type: REA	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.220	J	0.112	MDL	0.449	PQL	mg/Kg	J	Z

Sample ID: SL-882-SA5D-SB-4.0-5.0	Collected: 9/30/2013 9:00:00	Analysis Type: REA	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.184	J	0.110	MDL	0.439	PQL	mg/Kg	J	Z

* denotes a non-reportable result

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Data Qualifier Summary

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 1613B Matrix: SO

Sample ID: SL-582-SA5D-SB-0.0-0.5 Collected: 9/30/2013 8:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	0.0334	JBQ	0.0167	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0673	JB	0.0375	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.0836	JB	0.0349	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.0554	JQ	0.0392	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.0566	JBQ	0.0325	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.118	JB	0.0379	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDF	0.141	JQ	0.0320	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.0935	JBQ	0.0370	MDL	5.29	PQL	ng/Kg	U	B
2,3,4,6,7,8-HxCDF	0.0497	JBQ	0.0333	MDL	5.29	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0515	J	0.0364	MDL	5.29	PQL	ng/Kg	J	Z
OCDD	0.839	JB	0.0376	MDL	10.6	PQL	ng/Kg	U	B
OCDF	0.203	JBQ	0.0611	MDL	10.6	PQL	ng/Kg	U	B

Method Category: SVOA
Method: 8015M Matrix: SO

Sample ID: SL-582-SA5D-SB-4.0-5.0 Collected: 9/30/2013 8:55:00 Analysis Type: REA Dilution: 23.28

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
GASOLINE RANGE ORGANICS (C5-C12)	1.1	U	0.2	MDL	1.1	PQL	mg/Kg	UJ	H

Method Category: SVOA
Method: 8151A Matrix: SO

Sample ID: SL-582-SA5D-SB-0.0-0.5 Collected: 9/30/2013 8:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	25	U	9.4	MDL	25	PQL	ug/Kg	R	L

* denotes a non-reportable result

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Data Qualifier Summary

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8151A **Matrix:** SO

Sample ID: SL-582-SA5D-SB-10.0-11.0 Collected: 9/30/2013 9:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	27	U	10	MDL	27	PQL	ug/Kg	R	L

Sample ID: SL-582-SA5D-SB-4.0-5.0 Collected: 9/30/2013 8:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	27	U	10	MDL	27	PQL	ug/Kg	R	L

Sample ID: SL-882-SA5D-SB-4.0-5.0 Collected: 9/30/2013 9:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	27	U	10	MDL	27	PQL	ug/Kg	R	L

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-582-SA5D-SB-0.0-0.5 Collected: 9/30/2013 8:45:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	0.91	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-582-SA5D-SB-10.0-11.0 Collected: 9/30/2013 9:10:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	0.82	J	0.74	MDL	1.9	PQL	ug/Kg	J	Z
2-METHYLNAPHTHALENE	1.0	J	0.74	MDL	1.9	PQL	ug/Kg	J	Z
NAPHTHALENE	0.99	J	0.74	MDL	1.9	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Method Blank Contamination
H	Extraction to Analysis Estimation
L	Laboratory Control Spike Lower Rejection
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH118

QC Outlier Report: HoldingTimes

Lab Reporting Batch ID: PH118
EDD Filename: PrepPH118_v1

Laboratory: LL
eQAPP Name: CDM_SSFL_131101_Lan

Method: 8015M Preparation Method: 3546
Matrix: SO

Sample ID	Type	Actual	Criteria	Units	Flag
SL-582-SA5D-SB-4.0-5.0 (REA)	Extraction To Analysis	8.00	7.00	DAYS	J(all detects)
SL-582-SA5D-SB-4.0-5.0MS (REA)		8.00	7.00	DAYS	UJ(all non-detects)
SL-582-SA5D-SB-4.0-5.0MSD (REA)		8.00	7.00	DAYS	

Method Blank Outlier Report

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2770B371443	10/9/2013 2:43:00 PM	1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HxCDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF OCDD OCDF	0.0524 ng/Kg 0.0495 ng/Kg 0.0584 ng/Kg 0.0979 ng/Kg 0.0632 ng/Kg 0.0594 ng/Kg 0.0547 ng/Kg 0.104 ng/Kg 0.386 ng/Kg 0.187 ng/Kg	SL-582-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-582-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0334 ng/Kg	0.0334U ng/Kg
SL-582-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0673 ng/Kg	0.0673U ng/Kg
SL-582-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0836 ng/Kg	0.0836U ng/Kg
SL-582-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0566 ng/Kg	0.0566U ng/Kg
SL-582-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HxCDD	0.118 ng/Kg	0.118U ng/Kg
SL-582-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.0935 ng/Kg	0.0935U ng/Kg
SL-582-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0497 ng/Kg	0.0497U ng/Kg
SL-582-SA5D-SB-0.0-0.5(RES)	OCDD	0.839 ng/Kg	0.839U ng/Kg
SL-582-SA5D-SB-0.0-0.5(RES)	OCDF	0.203 ng/Kg	0.203U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P27537CB221359	10/10/2013 1:59:00 PM	ALUMINUM CALCIUM TIN ZINC	8.51 mg/Kg 15.2 mg/Kg 1.39 mg/Kg 1.38 mg/Kg	SL-582-SA5D-SB-0.0-0.5 SL-582-SA5D-SB-10.0-11.0 SL-582-SA5D-SB-4.0-5.0 SL-882-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-582-SA5D-SB-0.0-0.5(RES)	TIN	2.86 mg/Kg	2.86U mg/Kg
SL-582-SA5D-SB-10.0-11.0(RES)	TIN	2.93 mg/Kg	2.93U mg/Kg
SL-582-SA5D-SB-4.0-5.0(RES)	TIN	2.98 mg/Kg	2.98U mg/Kg
SL-882-SA5D-SB-4.0-5.0(RES)	TIN	2.85 mg/Kg	2.85U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PH118_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-582-SA5D-SB-4.0-5.0MS (TOT) SL-582-SA5D-SB-4.0-5.0MSD (TOT) (SL-582-SA5D-SB-0.0-0.5 SL-582-SA5D-SB-10.0-11.0 SL-582-SA5D-SB-4.0-5.0 SL-882-SA5D-SB-4.0-5.0)	ALUMINUM IRON MAGNESIUM MANGANESE POTASSIUM TITANIUM	3336 1811 396 133 173 499	3044 975 332 148 167 469	75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00	- - - - - -	ALUMINUM IRON MAGNESIUM MANGANESE POTASSIUM TITANIUM	J (all detects) Al, Fe, Mg, Mn, Ti No Qual, >4X
SL-582-SA5D-SB-4.0-5.0MS (TOT) SL-582-SA5D-SB-4.0-5.0MSD (TOT) (SL-582-SA5D-SB-0.0-0.5 SL-582-SA5D-SB-10.0-11.0 SL-582-SA5D-SB-4.0-5.0 SL-882-SA5D-SB-4.0-5.0)	CALCIUM	-261	57	75.00-125.00	-	CALCIUM	No Qual, >4X
SL-582-SA5D-SB-4.0-5.0MS (TOT) SL-582-SA5D-SB-4.0-5.0MSD (TOT) (SL-582-SA5D-SB-0.0-0.5 SL-582-SA5D-SB-10.0-11.0 SL-582-SA5D-SB-4.0-5.0 SL-882-SA5D-SB-4.0-5.0)	ANTIMONY	41	43	75.00-125.00	-	ANTIMONY	J(all detects) UJ(all non-detects)

Method: 6020A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-582-SA5D-SB-4.0-5.0MSD (TOT) (SL-582-SA5D-SB-0.0-0.5 SL-582-SA5D-SB-10.0-11.0 SL-582-SA5D-SB-4.0-5.0 SL-882-SA5D-SB-4.0-5.0)	STRONTIUM	-	145	75.00-125.00	-	STRONTIUM	No Qual, >4X

Method: 300.0
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-595-SA5D-SB-0.0-0.5MS (SL-595-SA5D-SB-0.0-0.5 SL-595-SA5D-SB-14.0-15.0 SL-595-SA5D-SB-17.0-18.0 SL-595-SA5D-SB-4.0-5.0 SL-595-SA5D-SB-9.0-10.0 SL-596-SA5D-SB-0.0-0.5 SL-596-SA5D-SB-11.5-12.5 SL-596-SA5D-SB-4.0-5.0 SL-597-SA5D-SB-0.0-0.5 SL-597-SA5D-SB-4.0-5.0)	FLUORIDE	65	-	80.00-120.00	-	FLUORIDE	J(all detects) UJ(all non-detects)
SL-597-SA5D-SB-9.0-10.0MS (SL-597-SA5D-SB-14.0-15.0 SL-597-SA5D-SB-17.0-18.0 SL-597-SA5D-SB-9.0-10.0)	FLUORIDE	23	-	80.00-120.00	-	FLUORIDE	J(all detects) R(all non-detects)

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8151A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P32767AQ241601A (SL-582-SA5D-SB-0.0-0.5 SL-582-SA5D-SB-10.0-11.0 SL-582-SA5D-SB-4.0-5.0 SL-882-SA5D-SB-4.0-5.0)	DINOSEB	9	-	10.00-56.00	-	DINOSEB	J (all detects) R (all non-detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 160.3M

Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-582-SA5D-SB-4.0-5.0	SL-882-SA5D-SB-4.0-5.0			
MOISTURE	11.7	11.6	1		No Qualifiers Applied

Method: 6010C

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-582-SA5D-SB-4.0-5.0 (TOT)	SL-882-SA5D-SB-4.0-5.0 (TOT)			
ALUMINUM	23800	25200	6	50.00	No Qualifiers Applied
ARSENIC	6.58	6.42	2	50.00	
BARIUM	95.3	108	12	50.00	
BERYLLIUM	0.771	0.789	2	50.00	
BORON	5.70	6.43	12	50.00	
CALCIUM	26200	30200	14	50.00	
CHROMIUM	34.1	34.8	2	50.00	
COBALT	8.11	8.59	6	50.00	
IRON	30200	31200	3	50.00	
LEAD	5.23	5.50	5	50.00	
LITHIUM	29.5	30.8	4	50.00	
MAGNESIUM	7620	7760	2	50.00	
MANGANESE	326	396	19	50.00	
NICKEL	14.0	15.2	8	50.00	
PHOSPHORUS	433	462	6	50.00	
POTASSIUM	3880	4170	7	50.00	
SODIUM	146	144	1	50.00	
TIN	2.98	2.85	4	50.00	
TITANIUM	1360	1460	7	50.00	
VANADIUM	67.8	69.2	2	50.00	
ZINC	69.4	69.5	0	50.00	
Zirconium	8.23	7.78	6	50.00	

Method: 6020A

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-582-SA5D-SB-4.0-5.0 (TOT)	SL-882-SA5D-SB-4.0-5.0 (TOT)			
SELENIUM	0.220	0.184	18	50.00	No Qualifiers Applied
STRONTIUM	61.4	62.7	2	50.00	
THALLIUM	0.297	0.318	7	50.00	

Method: 9045M

Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-582-SA5D-SB-4.0-5.0	SL-882-SA5D-SB-4.0-5.0			
PH	8.18	8.26	1	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-582-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JBQ	0.0334	5.29	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8-HxCDD	JB	0.0673	5.29	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.0836	5.29	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JQ	0.0554	5.29	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.0566	5.29	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.118	5.29	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JQ	0.141	5.29	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0935	5.29	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.0497	5.29	PQL	ng/Kg	
	2,3,4,7,8-PECDF	J	0.0515	5.29	PQL	ng/Kg	
	OCDD	JB	0.839	10.6	PQL	ng/Kg	
	OCDF	JBQ	0.203	10.6	PQL	ng/Kg	

Method: 300.0

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-596-SA5D-SB-4.0-5.0	FLUORIDE	J	1.9	2.2	PQL	mg/Kg	J (all detects)

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-582-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.816	1.04	PQL	mg/Kg	J (all detects)
	BORON	J	6.31	10.4	PQL	mg/Kg	
	SODIUM	J	98.2	104	PQL	mg/Kg	
	TIN	J	2.86	10.4	PQL	mg/Kg	
SL-582-SA5D-SB-10.0-11.0	BERYLLIUM	J	0.765	1.11	PQL	mg/Kg	J (all detects)
	BORON	J	5.51	11.1	PQL	mg/Kg	
	TIN	J	2.93	11.1	PQL	mg/Kg	
SL-582-SA5D-SB-4.0-5.0	BERYLLIUM	J	0.771	1.12	PQL	mg/Kg	J (all detects)
	BORON	J	5.70	11.2	PQL	mg/Kg	
	TIN	J	2.98	11.2	PQL	mg/Kg	
SL-882-SA5D-SB-4.0-5.0	BERYLLIUM	J	0.789	1.10	PQL	mg/Kg	J (all detects)
	BORON	J	6.43	11.0	PQL	mg/Kg	
	TIN	J	2.85	11.0	PQL	mg/Kg	

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-582-SA5D-SB-0.0-0.5	SELENIUM	J	0.277	0.415	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0311	0.207	PQL	mg/Kg	
SL-582-SA5D-SB-10.0-11.0	SELENIUM	J	0.150	0.443	PQL	mg/Kg	J (all detects)

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Reporting Limit Outliers

Lab Reporting Batch ID: PH118

Laboratory: LL

EDD Filename: PrepPH118_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-582-SA5D-SB-4.0-5.0	SELENIUM	J	0.220	0.449	PQL	mg/Kg	J (all detects)
SL-882-SA5D-SB-4.0-5.0	SELENIUM	J	0.184	0.439	PQL	mg/Kg	J (all detects)

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-582-SA5D-SB-0.0-0.5	NAPHTHALENE	J	0.91	1.7	PQL	ug/Kg	J (all detects)
SL-582-SA5D-SB-10.0-11.0	1-METHYLNAPHTHALENE	J	0.82	1.9	PQL	ug/Kg	J (all detects)
	2-METHYLNAPHTHALENE	J	1.0	1.9	PQL	ug/Kg	
	NAPHTHALENE	J	0.99	1.9	PQL	ug/Kg	

LDC #: 30842B4

VALIDATION COMPLETENESS WORKSHEET

Date: 12/4/13

SDG #: PH118

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: CRZ

2nd Reviewer: ✓

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	—	Sampling dates: 9/30/13
II.	ICP/MS Tune	—	
III.	Calibration	—	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	—	
VI.	Matrix Spike Analysis	SW	MS/D
VII.	Duplicate Sample Analysis	A	DUP
VIII.	Laboratory Control Samples (LCS)	N	
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	A	DUP
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	—	(2,3)
XV.	Field Blanks	SW	FB=FB-041613 EB=EB1-100213

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

(PH032)
D = Duplicate
TB = Trip blank
EB = Equipment blank

(PH120)

Validated Samples:

Soil

1	SL-582-SA5D-SB-0.0-0.5	11		21		31	
2	SL-582-SA5D-SB-4.0-5.0	12		22		32	
3	SL-882-SA5D-SB-4.0-5.0	13		23		33	
4	SL-582-SA5D-SB-10.0-11.0	14		24		34	
5	SL-582-SA5D-SB-4.0-5.0MS	15		25		35	
6	SL-582-SA5D-SB-4.0-5.0MSD	16		26		36	
7	SL-582-SA5D-SB-4.0-5.0DUP	17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg **Reason:** F
Sampling date: 4/11/13 **Soil factor applied** 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ **Associated Samples:** All

Analyte	Blank ID	Sample Identification														
	FB-041613 (SDG: PH032)	Action Limit	No Qualifiers													
Mo	0.0132	6.60														
Sn	0.0029	1.45														

Sampling date: 10/2/13 **Soil factor applied** 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ **Associated Samples:** All

Analyte	Blank ID	Sample Identification														
	EB1-100213 (SDG: PH120)	Action Limit	No Qualifiers													
Ba	0.00058	0.29														
B	0.0098	4.9														
Ca	0.0427	21.35														
Cu	0.0028	1.4														
Mo	0.0086	4.3														

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

Background Lab Sample ID: 7218895BKG Matrix Spike Lab Sample ID: 7218896MS Matrix Spike Duplicate Lab Sample ID: 7218897MSD
Batch Id(s): P27537C, P27538C

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Control Limit		
		Result	C	Result	C	Result	C				%R	Q	%R	Q		%R	RPD	M
Aluminum		21028.9604		27635.5040		27056.3208		198.0198	198.0198	MG/KG	3336		3044		2		20	P
Antimony		0.7327	U	20.3356		21.3129		49.5050	49.5050	MG/KG	41	N	43	N	5	75 - 125	20	P
Arsenic		5.8129		20.9980		22.0158		14.8515	14.8515	MG/KG	102		109		5	75 - 125	20	P
Barium		84.1475		286.5891		284.4525		198.0198	198.0198	MG/KG	102		101		1	75 - 125	20	P
Beryllium		0.6812	B	5.5525		5.5356		4.9505	4.9505	MG/KG	98		98		0	75 - 125	20	P
Boron		5.0366	B	197.8356		196.5158		198.0198	198.0198	MG/KG	97		97		1	75 - 125	20	P
Cadmium		0.0752	U	4.6901		4.6911		4.9505	4.9505	MG/KG	95		95		0	75 - 125	20	P
Calcium		23163.8347		22131.7158		23390.9634		396.0396	396.0396	MG/KG	-261		57		6		20	P
Chromium		30.1436		52.9337		53.0931		19.8020	19.8020	MG/KG	115		116		0	75 - 125	20	P
Cobalt		7.1653		53.6525		55.1604		49.5050	49.5050	MG/KG	94		97		3	75 - 125	20	P
Copper		1.4356	U	19.0248		19.0792		24.7525	24.7525	MG/KG	77		77		0	75 - 125	20	P
Iron		26645.2277		28438.2729		27610.3663		99.0099	99.0099	MG/KG	1811		975		3		20	P
Lead		4.6198		18.7762		19.6535		14.8515	14.8515	MG/KG	95		101		5	75 - 125	20	P
Lithium		26.0406		127.3188		126.4257		99.0099	99.0099	MG/KG	102		101		1	75 - 125	20	P
Magnesium		6724.4861		7508.5713		7382.1129		198.0198	198.0198	MG/KG	396		332		2		20	P
Manganese		287.7614		353.4455		360.8505		49.5050	49.5050	MG/KG	133		148		2		20	P
Mercury		0.0099	U	0.1915		0.1855		0.1626	0.1621	MG/KG	118		114		3	65 - 135	20	CV
Molybdenum		0.1683	U	182.0455		181.1455		198.0198	198.0198	MG/KG	92		91		0	75 - 125	20	P
Nickel		12.3673		59.3059		59.5733		49.5050	49.5050	MG/KG	95		95		0	75 - 125	20	P
Phosphorus		382.6109		491.6386		483.5663		99.0099	99.0099	MG/KG	110		102		2	75 - 125	20	P
Potassium		3426.0297		5140.4109		5078.1644		990.0990	990.0990	MG/KG	173	N	167	N	1	75 - 125	20	P
Selenium	78	0.1939	B	2.0382		2.0457		1.9802	1.9802	MG/KG	93		94		0	75 - 125	20	MS
Silver	107	0.0257	U	10.0069		10.3459		9.9010	9.9010	MG/KG	101		104		3	75 - 125	20	MS
Sodium		128.6416		1117.9545		1103.8050		990.0990	990.0990	MG/KG	100		98		1	75 - 125	20	P
Strontium	88	54.2519		62.7556		65.7139		7.9208	7.9208	MG/KG	107		145		5		20	MS
Thallium	203	0.2624		0.6840		0.6626		0.3960	0.3960	MG/KG	106		101		3	75 - 125	20	MS

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U = Below MDL, B = Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>
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QUALITY ASSURANCE SUMMARY
 FORM 5A (MS/MSD)
 MATRIX SPIKE/MATRIX SPIKE DUPLICATE
 SDG No.: PH118
 Matrix: SOIL Level: LOW
 (low/med):

Background Lab Sample ID: 7218895BKG Matrix Spike Lab Sample ID: 7218896MS Matrix Spike Duplicate Lab Sample ID: 7218897MSD
 Batch Id(s): P27537C, P27538C

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit				
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q	%R	RPD	M
Tin		2.6356	B	339.9950		340.1089		396.0396	396.0396	MG/KG	85		85		0	75 - 125	20	P	
Titanium		1201.8515	B	1695.5218		1666.4851		99.0099	99.0099	MG/KG	499		469		2			20	P
Vanadium		59.8436		117.2040		117.4287		49.5050	49.5050	MG/KG	116		116		0	75 - 125	20	P	
Zinc		61.2614		112.4723		112.5584		49.5050	49.5050	MG/KG	103		104		0	75 - 125	20	P	
Zirconium		7.2713		102.9337		101.8772		99.0099	99.0099	MG/KG	97		96		1	75 - 125	20	P	

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL, B= Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS
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**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH119

Prepared for

CDM Smith
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Prepared by

Laboratory Data Consultants, Inc
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Carlsbad, California 92010

December 23, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level IV data validation results for samples collected on October 1, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by Environmental Protection Agency (EPA) SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)

Pesticides by EPA SW 846 Method 8081B

Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A

Metals by EPA SW 846 Method 6010C, 6020A, and 7471B

Herbicides by EPA SW 846 Method 8151A

Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M

TPH as Extractables by EPA SW 846 Method 8015M

Dioxins and Dibenzofurans by EPA Method 1613B

Wet Chemistry:

Fluoride by EPA Method 300.0

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Automated Data Review outliers are presented in Enclosure I. Method specific Level IV DVRs are presented in Enclosure II.

All sample results were subjected to Level IV data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibrations, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards, interference check (ICSA and ICSAB) samples, matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), method blanks, trip blanks, equipment blanks, field blanks, and the raw data to confirm sample quantitation and identification.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the initial and continuing calibrations, ICB/CCBs, interference check samples and internal standards (except dioxins) which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

All criteria for the initial calibration of each method were met.

III. Continuing Calibration

All criteria for the initial calibration verifications and continuing calibration of each method were met with the exception of herbicides. The associated sample results were qualified as non-detected estimated (UJ). The details regarding the qualification of data are provided in Enclosures II.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for dioxins and metals. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosures I and II.

No contaminant concentrations were detected in the initial or continuing calibration blanks with the exception of several metals. The associated sample results were qualified as non-detected (U) due to initial or continuing calibration blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure II.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

The frequency of analysis was met.

The criteria for analysis were met.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of two MS/MSD pairs for metals and fluoride. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosures I and II.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one LCS/LCSD pairs for PCBs. No data were qualified due to high %Rs since the associated results were non-detected.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH119	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

No field duplicates were identified in this SDG.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No volatile contaminants were found in the trip blank.

One equipment blank (from SDG PH120) was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins and fluoride. The equipment blank had detections for SVOCs, metals and dioxins. The associated sample results were not detected or were significantly greater than the concentrations found in the equipment blank, therefore no data were qualified.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins and fluoride. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to equipment blank contamination as applicable. The sample results that

were not detected or were significantly greater than the concentrations found in the equipment blanks were not qualified. The equipment blank outlier reports are presented in Enclosures I and II.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
01-Oct-2013	TB-100113	7220652	TB	5030B	8015M	IV
01-Oct-2013	SL-598-SA5D-SB-0.0-0.5	7220655	N	METHOD	300.0	IV
01-Oct-2013	SL-598-SA5D-SB-0.0-0.5DUP	P220655D272153A	DUP	METHOD	300.0	IV
01-Oct-2013	SL-598-SA5D-SB-0.0-0.5MS	P220655R272209A	MS	METHOD	300.0	IV
01-Oct-2013	SL-598-SA5D-SB-4.0-5.0	7220656	N	METHOD	300.0	IV
01-Oct-2013	SL-598-SA5D-SB-9.0-10.0	7220657	N	METHOD	300.0	IV
01-Oct-2013	SL-598-SA5D-SB-14.0-15.0	7220658	N	METHOD	300.0	IV
01-Oct-2013	SL-598-SA5D-SB-18.0-19.0	7220659	N	METHOD	300.0	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5	7220653	N	3050B	6010C	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5	7220653	N	3050B	6020A	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5	7220653	N	3546	8015M	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5	7220653	N	3546	8081B	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5	7220653	N	3546	8082A	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5	7220653	N	3546	8270D SIM	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5	7220653	N	3550B	8151A	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5	7220653	N	METHOD	1613B	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5	7220653	N	METHOD	7471B	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5MSD	P220653M241917A	MSD	3550B	8151A	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5MSD	P220653M322257A	MSD	3546	8015M	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5MS	P220653R241850A	MS	3550B	8151A	IV
01-Oct-2013	SL-570-SA5D-SB-0.0-0.5MS	P220653R322236A	MS	3546	8015M	IV
01-Oct-2013	SL-570-SA5D-SB-4.0-5.0	7220654	N	3050B	6010C	IV
01-Oct-2013	SL-570-SA5D-SB-4.0-5.0	7220654	N	3050B	6020A	IV
01-Oct-2013	SL-570-SA5D-SB-4.0-5.0	7220654	N	3546	8015M	IV
01-Oct-2013	SL-570-SA5D-SB-4.0-5.0	7220654	N	3546	8081B	IV
01-Oct-2013	SL-570-SA5D-SB-4.0-5.0	7220654	N	3546	8082A	IV

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
01-Oct-2013	SL-570-SA5D-SB-4.0-5.0	7220654	N	3546	8270D SIM	IV
01-Oct-2013	SL-570-SA5D-SB-4.0-5.0	7220654	N	3550B	8151A	IV
01-Oct-2013	SL-570-SA5D-SB-4.0-5.0	7220654	N	5035A	8015M	IV
01-Oct-2013	SL-570-SA5D-SB-4.0-5.0	7220654	N	METHOD	7471B	IV
01-Oct-2013	SL-564-SA5D-SB-0.0-0.5	7220660	N	3050B	6010C	IV
01-Oct-2013	SL-564-SA5D-SB-0.0-0.5	7220660	N	3050B	6020A	IV
01-Oct-2013	SL-564-SA5D-SB-0.0-0.5	7220660	N	3546	8015M	IV
01-Oct-2013	SL-564-SA5D-SB-0.0-0.5	7220660	N	3546	8082A	IV
01-Oct-2013	SL-564-SA5D-SB-0.0-0.5	7220660	N	3546	8270D SIM	IV
01-Oct-2013	SL-564-SA5D-SB-0.0-0.5	7220660	N	METHOD	1613B	IV
01-Oct-2013	SL-564-SA5D-SB-0.0-0.5	7220660	N	METHOD	7471B	IV
01-Oct-2013	SL-564-SA5D-SB-4.0-5.0	7220661	N	3050B	6010C	IV
01-Oct-2013	SL-564-SA5D-SB-4.0-5.0	7220661	N	3050B	6020A	IV
01-Oct-2013	SL-564-SA5D-SB-4.0-5.0	7220661	N	3546	8015M	IV
01-Oct-2013	SL-564-SA5D-SB-4.0-5.0	7220661	N	3546	8082A	IV
01-Oct-2013	SL-564-SA5D-SB-4.0-5.0	7220661	N	3546	8270D SIM	IV
01-Oct-2013	SL-564-SA5D-SB-4.0-5.0	7220661	N	5035A	8015M	IV
01-Oct-2013	SL-564-SA5D-SB-4.0-5.0	7220661	N	METHOD	7471B	IV
01-Oct-2013	SL-607-SA5D-SB-0.0-0.5	7220662	N	3050B	6010C	IV
01-Oct-2013	SL-607-SA5D-SB-0.0-0.5	7220662	N	3050B	6020A	IV
01-Oct-2013	SL-607-SA5D-SB-0.0-0.5	7220662	N	3546	8015M	IV
01-Oct-2013	SL-607-SA5D-SB-0.0-0.5	7220662	N	3546	8082A	IV
01-Oct-2013	SL-607-SA5D-SB-0.0-0.5	7220662	N	3546	8270D SIM	IV
01-Oct-2013	SL-607-SA5D-SB-0.0-0.5	7220662	N	METHOD	1613B	IV
01-Oct-2013	SL-607-SA5D-SB-0.0-0.5	7220662	N	METHOD	7471B	IV
01-Oct-2013	SL-607-SA5D-SB-4.0-5.0	7220663	N	3050B	6010C	IV

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
01-Oct-2013	SL-607-SA5D-SB-4.0-5.0	7220663	N	3050B	6020A	IV
01-Oct-2013	SL-607-SA5D-SB-4.0-5.0	7220663	N	3546	8015M	IV
01-Oct-2013	SL-607-SA5D-SB-4.0-5.0	7220663	N	3546	8082A	IV
01-Oct-2013	SL-607-SA5D-SB-4.0-5.0	7220663	N	3546	8270D SIM	IV
01-Oct-2013	SL-607-SA5D-SB-4.0-5.0	7220663	N	5035A	8015M	IV
01-Oct-2013	SL-607-SA5D-SB-4.0-5.0	7220663	N	METHOD	7471B	IV

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: GENCHEM
Method: 300.0 **Matrix:** SO

Sample ID: SL-598-SA5D-SB-0.0-0.5 Collected: 10/1/2013 8:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.2		0.42	MDL	1.1	PQL	mg/Kg	J	Q

Sample ID: SL-598-SA5D-SB-14.0-15.0 Collected: 10/1/2013 9:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	3.1		0.47	MDL	1.2	PQL	mg/Kg	J	Q

Sample ID: SL-598-SA5D-SB-18.0-19.0 Collected: 10/1/2013 9:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	3.8		0.47	MDL	1.2	PQL	mg/Kg	J	Q

Sample ID: SL-598-SA5D-SB-4.0-5.0 Collected: 10/1/2013 8:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.9		0.45	MDL	1.1	PQL	mg/Kg	J	Q

Sample ID: SL-598-SA5D-SB-9.0-10.0 Collected: 10/1/2013 8:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.5		0.46	MDL	1.1	PQL	mg/Kg	J	Q

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-564-SA5D-SB-0.0-0.5 Collected: 10/1/2013 12:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.27	U	0.790	MDL	4.27	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.974	J	0.0715	MDL	1.07	PQL	mg/Kg	J	Z
BORON	7.13	J	0.896	MDL	10.7	PQL	mg/Kg	J	Z
POTASSIUM	3920		8.90	MDL	107	PQL	mg/Kg	J	Q
TIN	2.75	J	0.235	MDL	10.7	PQL	mg/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS

Method: 6010C

Matrix: SO

Sample ID: SL-564-SA5D-SB-4.0-5.0

Collected: 10/1/2013 1:00:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.42	U	0.818	MDL	4.42	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.864	J	0.0740	MDL	1.11	PQL	mg/Kg	J	Z
BORON	9.38	J	0.928	MDL	11.1	PQL	mg/Kg	J	Z
POTASSIUM	4150		9.22	MDL	111	PQL	mg/Kg	J	Q
TIN	2.73	J	0.243	MDL	11.1	PQL	mg/Kg	U	B

Sample ID: SL-570-SA5D-SB-0.0-5.0

Collected: 10/1/2013 10:10:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.28	U	0.791	MDL	4.28	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.667	J	0.0716	MDL	1.07	PQL	mg/Kg	J	Z
BORON	6.97	J	0.898	MDL	10.7	PQL	mg/Kg	J	Z
POTASSIUM	4350		8.92	MDL	107	PQL	mg/Kg	J	Q
TIN	2.90	J	0.235	MDL	10.7	PQL	mg/Kg	U	B

Sample ID: SL-570-SA5D-SB-4.0-5.0

Collected: 10/1/2013 10:20:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.43	U	0.819	MDL	4.43	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.556	J	0.0741	MDL	1.11	PQL	mg/Kg	J	Z
BORON	6.60	J	0.929	MDL	11.1	PQL	mg/Kg	J	Z
CADMIUM	0.218	J	0.0841	MDL	1.11	PQL	mg/Kg	J	Z
LEAD	3.05	J	0.553	MDL	3.32	PQL	mg/Kg	J	Z
POTASSIUM	3520		9.23	MDL	111	PQL	mg/Kg	J	Q
TIN	0.977	J	0.243	MDL	11.1	PQL	mg/Kg	U	B, F

Sample ID: SL-607-SA5D-SB-0.0-5.0

Collected: 10/1/2013 1:55:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.25	U	0.787	MDL	4.25	PQL	mg/Kg	UJ	Q
BORON	4.64	J	0.893	MDL	10.6	PQL	mg/Kg	J	Z
MOLYBDENUM	0.190	J	0.181	MDL	2.13	PQL	mg/Kg	J	Z
POTASSIUM	4870		8.87	MDL	106	PQL	mg/Kg	J	Q
TIN	3.04	J	0.234	MDL	10.6	PQL	mg/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

12/23/2013 8:19:53 AM

ADR version 1.7.0.207

Page 2 of 8

Data Qualifier Summary

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-607-SA5D-SB-4.0-5.0 Collected: 10/1/2013 2:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.61	U	0.852	MDL	4.61	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.937	J	0.0771	MDL	1.15	PQL	mg/Kg	J	Z
BORON	4.47	J	0.967	MDL	11.5	PQL	mg/Kg	J	Z
CADMIUM	0.0979	J	0.0875	MDL	1.15	PQL	mg/Kg	J	Z
POTASSIUM	3590		9.60	MDL	115	PQL	mg/Kg	J	Q
TIN	3.11	J	0.253	MDL	11.5	PQL	mg/Kg	U	B

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-564-SA5D-SB-0.0-0.5 Collected: 10/1/2013 12:50:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.227	J	0.107	MDL	0.427	PQL	mg/Kg	J	Z

Sample ID: SL-564-SA5D-SB-0.0-0.5 Collected: 10/1/2013 12:50:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0504	J	0.0277	MDL	0.213	PQL	mg/Kg	J	Z

Sample ID: SL-564-SA5D-SB-4.0-5.0 Collected: 10/1/2013 1:00:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.195	J	0.111	MDL	0.442	PQL	mg/Kg	J	Z

Sample ID: SL-570-SA5D-SB-0.0-0.5 Collected: 10/1/2013 10:10:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.177	J	0.107	MDL	0.428	PQL	mg/Kg	J	Z

Sample ID: SL-570-SA5D-SB-4.0-5.0 Collected: 10/1/2013 10:20:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.177	J	0.111	MDL	0.443	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-607-SA5D-SB-0.0-0.5 Collected: 10/1/2013 1:55:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.177	J	0.106	MDL	0.425	PQL	mg/Kg	J	Z

Sample ID: SL-607-SA5D-SB-0.0-0.5 Collected: 10/1/2013 1:55:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0545	J	0.0277	MDL	0.213	PQL	mg/Kg	J	Z

Sample ID: SL-607-SA5D-SB-4.0-5.0 Collected: 10/1/2013 2:05:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0456	J	0.0299	MDL	0.230	PQL	mg/Kg	J	Z

Method Category: METALS
Method: 7471B **Matrix:** SO

Sample ID: SL-607-SA5D-SB-0.0-0.5 Collected: 10/1/2013 1:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0121	J	0.0107	MDL	0.0178	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-564-SA5D-SB-0.0-0.5 Collected: 10/1/2013 12:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0954	JBQ	0.0175	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0856	JBQ	0.00744	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0361	JBQ	0.0115	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0308	JQ	0.0164	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDF	0.0209	JBQ	0.0152	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDF	0.0273	JBQ	0.0137	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0569	JBQ	0.0156	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDF	0.0890	JBQ	0.0154	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0728	JB	0.0158	MDL	5.22	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-564-SA5D-SB-0.0-0.5 Collected: 10/1/2013 12:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,3,4,6,7,8-HXCDF	0.0550	JBQ	0.0134	MDL	5.22	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0515	JBQ	0.0164	MDL	5.22	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0495	JBQ	0.0383	MDL	1.04	PQL	ng/Kg	U	B
OCDD	0.289	JBQ	0.0230	MDL	10.4	PQL	ng/Kg	U	B
OCDF	0.188	JB	0.0263	MDL	10.4	PQL	ng/Kg	U	B

Sample ID: SL-570-SA5D-SB-0.0-0.5 Collected: 10/1/2013 10:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.235	JBQ	0.0215	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0877	JBQ	0.00698	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0742	JBQ	0.00992	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0355	JQ	0.0180	MDL	5.32	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0512	JBQ	0.0113	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0568	JBQ	0.0192	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0321	JBQ	0.0105	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0504	JB	0.0178	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0565	JBQ	0.0121	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0532	JB	0.0253	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0599	JBQ	0.0133	MDL	5.32	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0544	JB	0.0112	MDL	5.32	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0432	JBQ	0.0131	MDL	5.32	PQL	ng/Kg	U	B
OCDD	1.53	JB	0.0222	MDL	10.6	PQL	ng/Kg	U	B
OCDF	0.167	JBQ	0.0230	MDL	10.6	PQL	ng/Kg	U	B

Sample ID: SL-607-SA5D-SB-0.0-0.5 Collected: 10/1/2013 1:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.239	JB	0.0205	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0307	JB	0.00698	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0229	JBQ	0.00934	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0561	J	0.0230	MDL	5.25	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0311	JBQ	0.0131	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.108	JBQ	0.0242	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0334	JBQ	0.0122	MDL	5.25	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-607-SA5D-SB-0.0-0.5 Collected: 10/1/2013 1:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,7,8,9-HXCDD	0.256	JBQ	0.0219	MDL	5.25	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.411	JB	0.0134	MDL	5.25	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0276	JBQ	0.0228	MDL	5.25	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0885	JBQ	0.0131	MDL	5.25	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0348	JBQ	0.0119	MDL	5.25	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0442	JB	0.0129	MDL	5.25	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.0277	JQ	0.0252	MDL	1.05	PQL	ng/Kg	J	Z
OCDD	0.701	JBQ	0.0226	MDL	10.5	PQL	ng/Kg	U	B
OCDF	0.125	JBQ	0.0214	MDL	10.5	PQL	ng/Kg	U	B

Method Category: SVOA
Method: 8151A **Matrix:** SO

Sample ID: SL-570-SA5D-SB-0.0-0.5 Collected: 10/1/2013 10:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	26	U	9.8	MDL	26	PQL	ug/Kg	UJ	C

Sample ID: SL-570-SA5D-SB-4.0-5.0 Collected: 10/1/2013 10:20:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	27	U	10	MDL	27	PQL	ug/Kg	UJ	C

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-570-SA5D-SB-4.0-5.0 Collected: 10/1/2013 10:20:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	1.1	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z
ANTHRACENE	0.41	J	0.37	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	1.5	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	1.0	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	1.1	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-607-SA5D-SB-0.0-0.5 Collected: 10/1/2013 1:55:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	0.82	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-607-SA5D-SB-4.0-5.0 Collected: 10/1/2013 2:05:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	1.2	J	0.77	MDL	1.9	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Method Blank Contamination
C	Continuing Calibration Verification Percent Difference Lower Estimation
F	Field Blank Contamination
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH119

Method Blank Outlier Report

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2810B372320	10/10/2013 11:20:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF 2,3,7,8-TCDD OCDD OCDF	0.0871 ng/Kg 0.0654 ng/Kg 0.0445 ng/Kg 0.0397 ng/Kg 0.0350 ng/Kg 0.0378 ng/Kg 0.0212 ng/Kg 0.0565 ng/Kg 0.0514 ng/Kg 0.0743 ng/Kg 0.0495 ng/Kg 0.0383 ng/Kg 0.0417 ng/Kg 0.306 ng/Kg 0.161 ng/Kg	SL-564-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-564-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.0954 ng/Kg	0.0954U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0856 ng/Kg	0.0856U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0361 ng/Kg	0.0361U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0209 ng/Kg	0.0209U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0273 ng/Kg	0.0273U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.0569 ng/Kg	0.0569U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.0890 ng/Kg	0.0890U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.0728 ng/Kg	0.0728U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0550 ng/Kg	0.0550U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.0515 ng/Kg	0.0515U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	2,3,7,8-TCDD	0.0495 ng/Kg	0.0495U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	OCDD	0.289 ng/Kg	0.289U ng/Kg
SL-564-SA5D-SB-0.0-0.5(RES)	OCDF	0.188 ng/Kg	0.188U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.235 ng/Kg	0.235U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0877 ng/Kg	0.0877U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0742 ng/Kg	0.0742U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0512 ng/Kg	0.0512U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.0568 ng/Kg	0.0568U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0321 ng/Kg	0.0321U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.0504 ng/Kg	0.0504U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.0565 ng/Kg	0.0565U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.0532 ng/Kg	0.0532U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.0599 ng/Kg	0.0599U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0544 ng/Kg	0.0544U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.0432 ng/Kg	0.0432U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	OCDD	1.53 ng/Kg	1.53U ng/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	OCDF	0.167 ng/Kg	0.167U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.239 ng/Kg	0.239U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0307 ng/Kg	0.0307U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-607-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0229 ng/Kg	0.0229U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0311 ng/Kg	0.0311U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.108 ng/Kg	0.108U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0334 ng/Kg	0.0334U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.0276 ng/Kg	0.0276U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.0885 ng/Kg	0.0885U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0348 ng/Kg	0.0348U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.0442 ng/Kg	0.0442U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	OCDD	0.701 ng/Kg	0.701U ng/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	OCDF	0.125 ng/Kg	0.125U ng/Kg

Method: 6010C

Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P27537CB221359	10/10/2013 1:59:00 PM	ALUMINUM CALCIUM TIN ZINC	8.51 mg/Kg 15.2 mg/Kg 1.39 mg/Kg 1.38 mg/Kg	SL-564-SA5D-SB-0.0-0.5 SL-564-SA5D-SB-4.0-5.0 SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-607-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-564-SA5D-SB-0.0-0.5(RES)	TIN	2.75 mg/Kg	2.75U mg/Kg
SL-564-SA5D-SB-4.0-5.0(RES)	TIN	2.73 mg/Kg	2.73U mg/Kg
SL-570-SA5D-SB-0.0-0.5(RES)	TIN	2.90 mg/Kg	2.90U mg/Kg
SL-570-SA5D-SB-4.0-5.0(RES)	TIN	0.977 mg/Kg	0.977U mg/Kg
SL-607-SA5D-SB-0.0-0.5(RES)	TIN	3.04 mg/Kg	3.04U mg/Kg
SL-607-SA5D-SB-4.0-5.0(RES)	TIN	3.11 mg/Kg	3.11U mg/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C

Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041613(REA2/TOT)	4/16/2013 3:15:00 PM	MOLYBDENUM TIN	0.0132 mg/L 0.0029 mg/L	SL-564-SA5D-SB-0.0-0.5 SL-564-SA5D-SB-4.0-5.0 SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-598-SA5D-SB-0.0-0.5 SL-598-SA5D-SB-14.0-15.0 SL-598-SA5D-SB-18.0-19.0 SL-598-SA5D-SB-4.0-5.0 SL-598-SA5D-SB-9.0-10.0 SL-607-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-570-SA5D-SB-4.0-5.0(RES)	TIN	0.977 mg/Kg	0.977U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 300.0

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-598-SA5D-SB-0.0-0.5MS (SL-598-SA5D-SB-0.0-0.5 SL-598-SA5D-SB-14.0-15.0 SL-598-SA5D-SB-18.0-19.0 SL-598-SA5D-SB-4.0-5.0 SL-598-SA5D-SB-9.0-10.0)	FLUORIDE	49	-	80.00-120.00	-	FLUORIDE	J (all detects) UJ (all non-detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-564-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.0954	5.22	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0856	5.22	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0361	5.22	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0308	5.22	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0209	5.22	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0273	5.22	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.0569	5.22	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.0890	5.22	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.0728	5.22	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0550	5.22	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0515	5.22	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.0495	1.04	PQL	ng/Kg	
	OCDD	JBQ	0.289	10.4	PQL	ng/Kg	
	OCDF	JB	0.188	10.4	PQL	ng/Kg	
SL-570-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.235	5.32	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0877	5.32	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0742	5.32	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0355	5.32	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0512	5.32	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.0568	5.32	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0321	5.32	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.0504	5.32	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.0565	5.32	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.0532	5.32	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0599	5.32	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.0544	5.32	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0432	5.32	PQL	ng/Kg	
	OCDD	JB	1.53	10.6	PQL	ng/Kg	
OCDF	JBQ	0.167	10.6	PQL	ng/Kg		
SL-607-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	0.239	5.25	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.0307	5.25	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0229	5.25	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	J	0.0561	5.25	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0311	5.25	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.108	5.25	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0334	5.25	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.256	5.25	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.411	5.25	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.0276	5.25	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0885	5.25	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0348	5.25	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.0442	5.25	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0277	1.05	PQL	ng/Kg	
OCDD	JBQ	0.701	10.5	PQL	ng/Kg		
OCDF	JBQ	0.125	10.5	PQL	ng/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-564-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.974	1.07	PQL	mg/Kg	J (all detects)
	BORON	J	7.13	10.7	PQL	mg/Kg	
	TIN	J	2.75	10.7	PQL	mg/Kg	
SL-564-SA5D-SB-4.0-5.0	BERYLLIUM	J	0.864	1.11	PQL	mg/Kg	J (all detects)
	BORON	J	9.38	11.1	PQL	mg/Kg	
	TIN	J	2.73	11.1	PQL	mg/Kg	
SL-570-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.667	1.07	PQL	mg/Kg	J (all detects)
	BORON	J	6.97	10.7	PQL	mg/Kg	
	TIN	J	2.90	10.7	PQL	mg/Kg	
SL-570-SA5D-SB-4.0-5.0	BERYLLIUM	J	0.556	1.11	PQL	mg/Kg	J (all detects)
	BORON	J	6.60	11.1	PQL	mg/Kg	
	CADMIUM	J	0.218	1.11	PQL	mg/Kg	
	LEAD	J	3.05	3.32	PQL	mg/Kg	
	TIN	J	0.977	11.1	PQL	mg/Kg	
SL-607-SA5D-SB-0.0-0.5	BORON	J	4.64	10.6	PQL	mg/Kg	J (all detects)
	MOLYBDENUM	J	0.190	2.13	PQL	mg/Kg	
	TIN	J	3.04	10.6	PQL	mg/Kg	
SL-607-SA5D-SB-4.0-5.0	BERYLLIUM	J	0.937	1.15	PQL	mg/Kg	J (all detects)
	BORON	J	4.47	11.5	PQL	mg/Kg	
	CADMIUM	J	0.0979	1.15	PQL	mg/Kg	
	TIN	J	3.11	11.5	PQL	mg/Kg	

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-564-SA5D-SB-0.0-0.5	SELENIUM	J	0.227	0.427	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0504	0.213	PQL	mg/Kg	
SL-564-SA5D-SB-4.0-5.0	SELENIUM	J	0.195	0.442	PQL	mg/Kg	J (all detects)
SL-570-SA5D-SB-0.0-0.5	SELENIUM	J	0.177	0.428	PQL	mg/Kg	J (all detects)
SL-570-SA5D-SB-4.0-5.0	SELENIUM	J	0.177	0.443	PQL	mg/Kg	J (all detects)
SL-607-SA5D-SB-0.0-0.5	SELENIUM	J	0.177	0.425	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0545	0.213	PQL	mg/Kg	
SL-607-SA5D-SB-4.0-5.0	SILVER	J	0.0456	0.230	PQL	mg/Kg	J (all detects)

Method: 7471B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-607-SA5D-SB-0.0-0.5	MERCURY	J	0.0121	0.0178	PQL	mg/Kg	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH119

Laboratory: LL

EDD Filename: PrepPH119_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-570-SA5D-SB-4.0-5.0	2-METHYLNAPHTHALENE	J	1.1	1.8	PQL	ug/Kg	J (all detects)
	ANTHRACENE	J	0.41	1.8	PQL	ug/Kg	
	BENZO(A)PYRENE	J	1.5	1.8	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	1.0	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	1.1	1.8	PQL	ug/Kg	
SL-607-SA5D-SB-0.0-0.5	NAPHTHALENE	J	0.82	1.8	PQL	ug/Kg	J (all detects)
SL-607-SA5D-SB-4.0-5.0	NAPHTHALENE	J	1.2	1.9	PQL	ug/Kg	J (all detects)

Enclosure II

EPA Level IV Data Validation Reports

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: October 1, 2013
LDC Report Date: December 18, 2013
Matrix: Soil
Parameters: Semivolatiles
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH119

Sample Identification

SL-570-SA5D-SB-0.0-0.5
SL-570-SA5D-SB-4.0-5.0
SL-564-SA5D-SB-0.0-0.5
SL-564-SA5D-SB-4.0-5.0
SL-607-SA5D-SB-0.0-0.5
SL-607-SA5D-SB-4.0-5.0

Introduction

This data review covers 6 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8270D using Selected Ion Monitoring (SIM) for Semivolatiles.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance was checked at 12 hour intervals. All ion abundance requirements were met.

III. Initial Calibration

Initial calibration was performed using required standard.

Percent relative standard deviations (%RSD) were less than or equal to 30.0% for all compounds.

Average relative response factors (RRF) for all compounds were within method and validation criteria.

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

Percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within the method criteria of less than or equal to 20.0% for calibration check compounds (CCCs) and 25.0% for all other compounds.

The percent differences (%D) of the second source calibration standard were less than or equal to 25.0% for all compounds.

All of the continuing calibration relative response factors (RRF) were within method and validation criteria.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No semivolatile contaminants were found in the method.

Sample EB1-100213 (from SDG PH120) was identified as an equipment blank. No semivolatile contaminants were found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
EB1-100213	10/2/13	Di-n-butylphthalate Diethylphthalate Bis(2-ethylhexyl)phthalate	0.18 ug/L 0.27 ug/L 0.059 ug/L	All samples in SDG PH119

Sample FB-041613 (from SDG PH032) was identified as a field blank. No semivolatile contaminants were found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB-041613	4/16/13	2-Methylnaphthalene Bis(2-ethylhexyl)phthalate Di-n-butylphthalate	0.012 ug/L 1.0 ug/L 0.11 ug/L	All samples in SDG PH119

Sample concentrations were compared to concentrations detected in the field blanks. The sample concentrations were either not detected or were significantly greater (>10X for common contaminants, >5X for other contaminants) than the concentrations found in the associated field blanks.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. Target Compound Identifications

All target compound identifications were within validation criteria.

XII. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH119	All compounds reported below the RL.	J (all detects)	A

XIII. Tentatively Identified Compounds (TICs)

Tentatively identified compounds were not reported by the laboratory.

XIV. System Performance

The system performance was acceptable.

XV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XVI. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Semivolatiles - Data Qualification Summary - SDG PH119**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-564-SA5D-SB-0.0-0.5 SL-564-SA5D-SB-4.0-5.0 SL-607-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-4.0-5.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Semivolatiles - Laboratory Blank Data Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Semivolatiles - Field Blank Data Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

LDC #: 30842C2b

VALIDATION COMPLETENESS WORKSHEET

Date: 12/3/13

SDG #: PH119

Level IV

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: BR

2nd Reviewer: SU

METHOD: GC/MS Polynuclear Aromatic Hydrocarbons (EPA SW 846 Method 8270C-SIM)
SVOCS

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 10/1/13
II.	GC/MS Instrument performance check	A	
III.	Initial calibration	A	RSD ≤ 30%
IV.	Continuing calibration/ICV	A	1W/CCV = 25%
V.	Blanks	A	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	N	Client spec.
VIII.	Laboratory control samples	A	LCS
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	A	
XI.	Target compound identification	A	
XII.	Compound quantitation/RL/LOQ/LODs	A	
XIII.	Tentatively identified compounds (TICs)	N	
XIV.	System performance	A	
XV.	Overall assessment of data	A	
XVI.	Field duplicates	N	
XVII.	Field blanks	SW	EB = EB1-100213 (PH120) FB = FB-0443 (PA032)

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: 861

1	SL-570-SA5D-SB-0.0-0.5	11	21	31	SBLKLA276
2	SL-570-SA5D-SB-4.0-5.0	12	22	32	
3	SL-564-SA5D-SB-0.0-0.5	13	23	33	
4	SL-564-SA5D-SB-4.0-5.0	14	24	34	
5	SL-607-SA5D-SB-0.0-0.5	15	25	35	
6	SL-607-SA5D-SB-4.0-5.0	16	26	36	
7		17	27	37	
8		18	28	38	
9		19	29	39	
10		20	30	40	

Method: Semivolatiles (EPA SW 846 Method 8270C)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. GC/MS Instrument performance check				
Were the DFTPP performance results reviewed and found to be within the specified criteria?	/			
Were all samples analyzed within the 12 hour clock criteria?	/			
III. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	/			
Were all percent relative standard deviations (%RSD) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	/			
Was a curve fit used for evaluation?		/		
Did the initial calibration meet the curve fit acceptance criteria of ≥ 0.990 ?			/	
Were all percent relative standard deviations (%RSD) $\leq 30\%$ and relative response factors (RRF) > 0.05 ?	/			
IV. Continuing calibration				
Was a continuing calibration standard analyzed at least once every 12 hours for each instrument?	/			
Were all percent differences (%D) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	/			
Were all percent differences (%D) $\leq 25\%$ and relative response factors (RRF) ≥ 0.05 ?	/			
V. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was a method blank analyzed for each matrix and concentration?	/			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.		/		
VI. Surrogate spikes				
Were all surrogate %R within QC limits?	/			
If 2 or more base neutral or acid surrogates were outside QC limits, was a reanalysis performed to confirm %R?			/	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?			/	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.			/	
Was a MS/MSD analyzed every 20 samples of each matrix?			/	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?			/	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			

Validation Area	Yes	No	NA	Findings/Comments
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
X. Internal standards				
Were internal standard area counts within -50% or +100% of the associated calibration standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were retention times within + 30 seconds from the associated calibration standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XI. Target compound identification				
Were relative retention times (RRT's) within + 0.06 RRT units of the standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did compound spectra meet specified EPA "Functional Guidelines" criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were chromatogram peaks verified and accounted for?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XII. Compound quantitation/RLs				
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were compound quantitation and RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XIII. Tentatively identified compounds (TICs)				
Were the major ions (> 10 percent relative intensity) in the reference spectrum evaluated in sample spectrum?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were relative intensities of the major ions within $\pm 20\%$ between the sample and the reference spectra?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Did the raw data indicate that the laboratory performed a library search for all required peaks in the chromatograms (samples and blanks)?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
XIV. System performance				
System performance was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XV. Overall assessment of data				
Overall assessment of data was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XVI. Field duplicates				
Field duplicate pairs were identified in this SDG.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field duplicates.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
XVII. Field blanks				
Field blanks were identified in this SDG.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field blanks.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	

VALIDATION FINDINGS WORKSHEET

METHOD: GC/MS SVOA

A. Phenol	T. 4-Chloroaniline	MM. 4-Chlorophenyl-phenyl ether	FFF. Di-n-octylphthalate	YYY. 2,3,5-Trimethylnaphthalene
B. Bis (2-chloroethyl) ether	U. Hexachlorobutadiene	NN. Fluorene	GGG. Benzo(b)fluoranthene	ZZZ. Perylene
C. 2-Chlorophenol	V. 4-Chloro-3-methylphenol	OO. 4-Nitroaniline	HHH. Benzo(k)fluoranthene	AAAA. Dibenzothiophene
D. 1,3-Dichlorobenzene	W. 2-Methylnaphthalene	PP. 4,6-Dinitro-2-methylphenol	III. Benzo(a)pyrene	BBBB. Benzo(a)fluoranthene
E. 1,4-Dichlorobenzene	X. Hexachlorocyclopentadiene	QQ. N-Nitrosodiphenylamine	JJJ. Indeno(1,2,3-cd)pyrene	CCCC. Benzo(b)fluorene
F. 1,2-Dichlorobenzene	Y. 2,4,6-Trichlorophenol	RR. 4-Bromophenyl-phenylether	KKK. Dibenz(a,h)anthracene	DDDD. cis/trans-Decalin
G. 2-Methylphenol	Z. 2,4,5-Trichlorophenol	SS. Hexachlorobenzene	LLL. Benzo(g,h,i)perylene	EEEE. Biphenyl
H. 2,2'-Oxybis(1-chloropropane)	AA. 2-Chloronaphthalene	TT. Pentachlorophenol	MMM. Bis(2-Chloroisopropyl)ether	FFFF. Retene
I. 4-Methylphenol	BB. 2-Nitroaniline	UU. Phenanthrene	NNN. Aniline	GGGG. C30-Hopane
J. N-Nitroso-di-n-propylamine	CC. Dimethylphthalate	VV. Anthracene	OOO. N-Nitrosodimethylamine	HHHH. 1-Methylphenanthrene
K. Hexachloroethane	DD. Acenaphthylene	WW. Carbazole	PPP. Benzoic Acid	IIII. 1,4-Dioxane
L. Nitrobenzene	EE. 2,6-Dinitrotoluene	XX. Di-n-butylphthalate	QQQ. Benzyl alcohol	JJJJ. Acetophenone
M. Isophorone	FF. 3-Nitroaniline	YY. Fluoranthene	RRR. Pyridine	KKKK. Atrazine
N. 2-Nitrophenol	GG. Acenaphthene	ZZ. Pyrene	SSS. Benzidine	LLLL. Benzaldehyde
O. 2,4-Dimethylphenol	HH. 2,4-Dinitrophenol	AAA. Butylbenzylphthalate	TTT. 1-Methylnaphthalene	MMMM. Caprolactam
P. Bis(2-chloroethoxy)methane	II. 4-Nitrophenol	BBB. 3,3'-Dichlorobenzidine	UUU. Benzo(b)thiophene	NNNN.
Q. 2,4-Dichlorophenol	JJ. Dibenzofuran	CCC. Benzo(a)anthracene	VVV. Benzonaphthothiophene	OOOO.
R. 1,2,4-Trichlorobenzene	KK. 2,4-Dinitrotoluene	DDD. Chrysene	WWW. Benzo(e)pyrene	PPPP.
S. Naphthalene	LL. Diethylphthalate	EEE. Bis(2-ethylhexyl)phthalate	XXX. 2,6-Dimethylnaphthalene	QQQQ.

VALIDATION FINDINGS WORKSHEET

Field Blanks

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

FB = FB-041613

Y N N/A Were field blanks identified in this SDG?

SDG # PH032

Y N N/A Were target compounds detected in the field blanks?

Blank units: ug/L Associated sample units: ug/L

Sampling date: 4/16/13

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All (ND + 75X)

Compound	Blank ID		Sample Identification						
	<u>FB</u>	<u>52</u>							
<u>W</u>	<u>0.012</u>	<u>0.06</u>							
<u>EEE</u>	<u>1.0</u>	<u>5.10</u>							
<u>XX</u>	<u>0.11</u>	<u>1.1</u>							

Blank units: ug/L Associated sample units: ug/L

EB = EBI-100213

Sampling date: 10/2/13

SDG # PH20

Field blank type: (circle one) Field Blank / Rinsate / Other EB Associated Samples: All (ND + 75X)

Compound	Blank ID		Sample Identification						
	<u>EB</u>	<u>52</u>							
<u>XX</u>	<u>0.18</u>	<u>1.8</u>							
<u>LL</u>	<u>0.27</u>	<u>2.7</u>							
<u>LEE</u>	<u>0.059</u>	<u>0.59</u>							

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Common contaminants such as the phthalates and TICs noted above that were detected in samples within ten times the associated field blank concentration were qualified as not detected, "U". Other contaminants within five times the field blank concentration were also qualified as not detected, "U".

LDC #: 30842C2b

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer BR
 2nd Reviewer SM

METHOD: GC/MS SVOA (EPA SW 846 Method 8270DSIM)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$$RRF = (A_x)(C_{is}) / (A_{is})(C_x)$$

average RRF = sum of the RRFs/number of standards

$$\%RSD = 100 * (S/X)$$

 A_x = Area of Compound C_x = Concentration of compound,

S= Standard deviation of the RRFs,

 A_{is} = Area of associated internal standard C_{is} = Concentration of internal standard

X = Mean of the RRFs

#	Standard ID	Calibration Date	Compound (IS)	Reported RRF (1 std)	Recalculated RRF (1 std)	Reported Average RRF (Initial)	Recalculated Average RRF (Initial)	Reported %RSD	Recalculated %RSD
1	HP10976	9/19/2013	N-Nitrosodimethylamine (IS1)	1.432	1.432	1.536	1.536	8	8
			Naphthalene (IS2)	1.030	1.030	1.042	1.042	2	2
			Fluorene (IS3)	1.335	1.335	1.312	1.312	2	2
			Anthracene (IS4)	1.125	1.125	1.113	1.1135	4	4
			Chrysene (IS5)	1.224	1.224	1.234	1.234	2	2
			Benzo(a)pyrene (IS6)	1.198	1.198	1.154	1.1545	3	3

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

METHOD: GC/MS SVOA (EPA SW 846 Method 8270DSIM)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

$$\% \text{ Difference} = 100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$$

$$\text{RRF} = (\text{Ax})(\text{Cis}) / (\text{Ais})(\text{Cx})$$

Where:

ave. RRF = initial calibration average RRF

RRF = continuing calibration RRF

Ax = Area of compound

Cx = Concentration of compound

Ais = Area of associated internal standard

Cis = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (IS)	Average RRF (Initial RRF)	Reported (CC RRF)	Recalculated (CC RRF)	Reported %D	Recalculated %D
1	ij0091	10/04/13	N-Nitrosodimethylamine (IS1)	1.536	1.553	1.553	1	1
			Naphthalene (IS2)	1.042	1.035	1.035	1	1
			Fluorene (IS3)	1.312	1.299	1.299	1	1
			Anthracene (IS4)	1.113	1.122	1.122	1	1
			Chrysene (IS5)	1.234	1.235	1.235	0	0
			Benzo(a)pyrene (IS6)	1.154	1.165	1.165	1	1

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

METHOD: GC/MS SVOA (EPA SW 846 Method 8270DSIM)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

$$\% \text{ Difference} = 100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$$

$$\text{RRF} = (\text{Ax})(\text{Cis}) / (\text{Ais})(\text{Cx})$$

Where:

ave. RRF = initial calibration average RRF

RRF = continuing calibration RRF

Ax = Area of compound

Cx = Concentration of compound

Ais = Area of associated internal standard

Cis = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (IS)	Average RRF (Initial RRF)	Reported (CC RRF)	Recalculated (CC RRF)	Reported %D	Recalculated %D
1	ij0121	10/08/13	N-Nitrosodimethylamine (IS1)	1.536	1.248	1.248	19	19
			Naphthalene (IS2)	1.042	1.061	1.061	2	2
			Fluorene (IS3)	1.312	1.291	1.291	2	2
			Anthracene (IS4)	1.113	1.108	1.108	0	0
			Chrysene (IS5)	1.234	1.151	1.151	7	7
			Benzo(a)pyrene (IS6)	1.154	1.139	1.139	1	1

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

METHOD: GC/MS Semivolatiles (EPA SW 846 Method 8270C)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
SS = Surrogate Spiked

Sample ID: 1

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5	Fluoranthene-d10 1.00	0.891	89	89	0
2-Fluorobiphenyl	Benzo(a)pyrene-d12 ↓	0.940	94	94	0
Terphenyl-d14	1-Methylnaphthalene-d10 ↓	1.017	102	102	0
Phenol-d5					
2-Fluorophenol					
2,4,6-Tribromophenol					
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5					
2-Fluorobiphenyl					
Terphenyl-d14					
Phenol-d5					
2-Fluorophenol					
2,4,6-Tribromophenol					
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5					
2-Fluorobiphenyl					
Terphenyl-d14					
Phenol-d5					
2-Fluorophenol					
2,4,6-Tribromophenol					
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

LDC #: 30842022

VALIDATION FINDINGS WORKSHEET

Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Page: 1 of 1
 Reviewer: BR
 2nd Reviewer: SH

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

% Recovery = $100 * (SC/SA)$

Where: SSC = Spike concentration
 SA = Spike added

RPD = $|LCS - LCSD| * 2 / (LCS + LCSD)$

LCS = Laboratory control sample percent recovery

LCSD = Laboratory control sample duplicate percent recovery

LCS/LCSD samples: 276LALCS

Compound	Spike Added (<u>µg/L</u>)		Spike Concentration (<u>µg/L</u>)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD	LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
					Reported	Recalc.	Reported	Recalc.	Reported	Recalculated
Phenol										
N-Nitroso-di-n-propylamine										
4-Chloro-3-methylphenol										
Acenaphthene	33.33	—	32.78	—	98	98	—	—	—	—
Pentachlorophenol										
Pyrene	33.33	—	33.67	—	101	101	—	—	—	—

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

Y N N/A
V N N/A

Were all reported results recalculated and verified for all level IV samples?
Were all recalculated results for detected target compounds agree within 10.0% of the reported results?

Concentration = (Ax)(Is)(Vj)(DF)(2.0) / (Ais)(RRF)(Vo)(Vj)(%S)

- Ax = Area of the characteristic ion (EICP) for the compound to be measured
Ais = Area of the characteristic ion (EICP) for the specific internal standard
Is = Amount of internal standard added in nanograms (ng)
Vo = Volume or weight of sample extract in milliliters (ml) or grams (g).
Vj = Volume of extract injected in microliters (ul)
Vt = Volume of the concentrated extract in microliters (ul)
Df = Dilution Factor.
%S = Percent solids, applicable to soil and solid matrices only.
2.0 = Factor of 2 to account for GPC cleanup

Example:

Sample I.D. 2, ODD:

ODD = 3 mg / kg

Conc. = (31534 / 364348) * 1 * 1000 * () * ()
= 2.60945152 mg/kg

Table with 5 columns: #, Sample ID, Compound, Reported Concentration, Calculated Concentration, Qualification. The table is mostly empty for data entry.

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: October 1, 2013
LDC Report Date: December 18, 2013
Matrix: Soil
Parameters: Chlorinated Pesticides
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH119

Sample Identification

SL-570-SA5D-SB-0.0-0.5
SL-570-SA5D-SB-4.0-5.0

Introduction

This data review covers 2 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8081B for Chlorinated Pesticides.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/ECD Instrument Performance Check

Instrument performance was acceptable unless noted otherwise under initial calibration and continuing calibration sections.

III. Initial Calibration

Initial calibration of single compounds was performed for the primary (quantitation) column and confirmation column as required by this method.

The percent relative standard deviations (%RSD) were less than or equal to 20.0% for all compounds.

Retention time windows were evaluated and considered technically acceptable.

IV. Continuing Calibration

Continuing calibration was performed at required frequencies.

The percent differences (%D) of calibration factors in continuing standard mixtures were within the 20.0% QC limits.

The percent differences (%D) of the second source calibration standard were less than or equal to 20.0% for all compounds.

Retention times (RT) of all compounds in the calibration standards were within QC limits.

The individual 4,4'-DDT and Endrin breakdowns (%BD) were less than or equal to 15.0%.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No chlorinated pesticide contaminants were found in the method blanks.

Sample EB1-100213 (from SDG PH120) was identified as an equipment blank. No chlorinated pesticide contaminants were found.

Sample FB-041613 (from SDG PH032) was identified as a field blank. No chlorinated pesticide contaminants were found.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Florisil Cartridge Check

Florisil cleanup was not required and therefore not performed in this SDG.

XI. GPC Calibration

GPC cleanup was not required and therefore not performed in this SDG.

XII. Target Compound Identification

All target compound identifications were within validation criteria.

XIII. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH119	All compounds reported below the RL.	J (all detects)	A

XIV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XV. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Chlorinated Pesticides - Data Qualification Summary - SDG PH119**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Chlorinated Pesticides - Laboratory Blank Data Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Chlorinated Pesticides - Field Blank Data Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

LDC #: 30842C3a

VALIDATION COMPLETENESS WORKSHEET

Date: 12/3/13

SDG #: PH119

Level IV

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: BR

2nd Reviewer: SM

METHOD: GC Chlorinated Pesticides (EPA SW846 Method 8081A)^B

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 10/1/13
II.	GC/ECD Instrument Performance Check	A	
III.	Initial calibration	A	RSD ≤ 20?
IV.	Continuing calibration/ICV	A	1(W) COV ≤ 20?
V.	Blanks	A	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	N	Client spec.
VIII.	Laboratory control samples	A	LCS
IX.	Regional quality assurance and quality control	N	
X.	Florisil cartridge check	N	
XI.	GPC Calibration	N	
XII.	Target compound identification	A	
XIII.	Compound quantitation/RL/LOQ/LODs	A	
XIV.	Overall assessment of data	A	
XV.	Field duplicates	N	
XVI.	Field blanks	ND	FB = FB-041613 (SDG # PH032) EB = EB1-100213 (SDG # PH120)

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

D = Duplicate
TB = Trip blank
EB = Equipment blank

Validated Samples: Soil

1	SL-570-SA5D-SB-0.0-0.5	11		21		31	PBLR 06276
2	SL-570-SA5D-SB-4.0-5.0	12		22		32	
3		13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

Method: Pesticides/PCBs (EPA SW 846 Method 8081/8082)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. GC/ECD Instrument performance check				
Was the instrument performance found to be acceptable?	/			
III. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	/			
Was a linear fit used for evaluation? If yes, were all percent relative standard deviations (%RSD) \leq 20%?	/			
Was a curve fit used for evaluation? If Yes, what was the acceptance criteria used?		/		
Did the initial calibration meet the curve fit acceptance criteria?			/	
Were the RT windows properly established?	/			
Were the required standard concentrations analyzed in the initial calibration?	/			
IV. Continuing calibration				
What type of continuing calibration calculation was performed? <u>X</u> %D or ___ %R	/			
Were Evaluation mix standards analyzed prior to the initial calibration and sample analysis?	/			
Were endrin and 4,4'-DDT breakdowns \leq 15%.0 for individual breakdown in the Evaluation mix standards?	/			
Was a continuing calibration analyzed daily?	/			
Were all percent differences (%D) \leq 20%.0 or percent recoveries 80-120%?	/			
Were all the retention times within the acceptance windows?	/			
V. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was a method blank analyzed for each matrix and concentration?	/			
Were extract cleanup blanks analyzed with every batch requiring clean-up?			/	
Was there contamination in the method blanks or clean-up blanks? If yes, please see the Blanks validation completeness worksheet.		/		
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?	/			
If the percent recovery (%R) of one or more surrogates was outside QC limits, was a reanalysis performed to confirm %R?			/	

Validation Area	Yes	No	NA	Findings/Comments
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?			/	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.		/		
Was a MS/MSD analyzed every 20 samples of each matrix?			/	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?			/	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per extraction batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	/			
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?		/		
Were the performance evaluation (PE) samples within the acceptance limits?			/	
X. Target compound identification				
Were the retention times of reported detects within the RT windows?	/			
XI. Compound quantitation/CRQLs				
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions, dry weight factors, and clean-up activities applicable to level IV validation?	/			
XII. System performance				
System performance was found to be acceptable.	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.		/		

VALIDATION FINDINGS WORKSHEET

METHOD: Pesticide/PCBs (EPASW 846 Method 8081/8082)

A. alpha-BHC	I. Dieldrin	Q. Endrin ketone	Y. Aroclor-1242	GG. Chlordane
B. beta-BHC	J. 4,4'-DDE	R. Endrin aldehyde	Z. Aroclor-1248	HH. Chlordane (Technical)
C. delta-BHC	K. Endrin	S. alpha-Chlordane	AA. Aroclor-1254	II. Aroclor 1262
D. gamma-BHC	L. Endosulfan II	T. gamma-Chlordane	BB. Aroclor-1260	JJ. Aroclor 1268
E. Heptachlor	M. 4,4'-DDD	U. Toxaphene	CC. 2,4'-DDD	KK. Oxychlordane
F. Aldrin	N. Endosulfan sulfate	V. Aroclor-1016	DD. 2,4'-DDE	LL. trans-Nonachlor
G. Heptachlor epoxide	O. 4,4'-DDT	W. Aroclor-1221	EE. 2,4'-DDT	MM. cis-Nonachlor
H. Endosulfan I	P. Methoxychlor	X. Aroclor-1232	FF. Hexachlorobenzene	NN.

Notes: _____

LDC#: 30842C3a

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: BR
 2nd Reviewer: SL

METHOD: GC Pesticides (EPA SW 846 Method 8081B)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$RRF = (A_x)(C_{is}) / (A_{is})(C_x)$ <p>average RRF = sum of the RRFs/number of standards</p> $\%RSD = 100 * (S/X)$	<p>Where</p> <p>A_x = Area of Compound</p> <p>C_x = Concentration of compound,</p> <p>S= Standard deviation of the RRFs,</p>	<p>A_{is} = Area of associated internal standard</p> <p>C_{is} = Concentration of internal standard</p> <p>X = Mean of the RRFs</p>
--	--	---

#	Standard ID	Calibration Date	Compound	Reported RRF (10 std)	Recalculated RRF (10 std)	Reported Average RRF (Initial)	Recalculated Average RRF (Initial)	Reported %RSD	Recalculated %RSD
1	ICAL	10/3/2013	Endosulfan I (RTX-CLP)	568000	567954	571000	571000	4	4
	H9191A		Methoxychlor (RTX-CLP)	240000	240156	236400	236400	5	5
			Endosulfan I (RTX-CLPII)	101000	100605	101000	100980	3	3
			Methoxychlor (RTX-CLPII)	59700	59742	59200	59160	2	2

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Calculation Verification

METHOD: GC Pesticides (EPA SW 846 Method 8081B)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

$\% \text{ Difference} = 100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$
 $\text{RRF} = (\text{Ax})(\text{Cis}) / (\text{Ais})(\text{Cx})$

Where:

ave. RRF = initial calibration average RRF
 RRF = continuing calibration RRF
 Ax = Area of compound,

Cx = Concentration of compound,
 Ais = Area of associated internal standard
 Cis = Concentration of internal standard

#	Standard ID	Calibration Date	Compound	Average CF/ Conc	Reported Conc/CF (CC)	Recalculated Conc/CF (CC)	Reported % D	Recalculated %D
1	MIXA3IX	10/7/2013 8:53	Endosulfan I (RTX-CLP)	10.00	10.05	10.05	0	0.5
			Methoxychlor (RTX-CLP)	100.0	97.69	97.71	2	2
			Endosulfan I (RTX-CLPII)	10.00	10.01	10.01	0	0
			Methoxychlor (RTX-CLPII)	100.0	97.59	97.55	2	2
2								

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

METHOD: GC Pesticides/PCBs (EPA SW 846 Method 8081/8082)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
SS = Surrogate Spiked

Sample ID: 1

Surrogate	Column	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
Tetrachloro-m-xylene						
Tetrachloro-m-xylene	RTX-CLP	10.1	7.863534	78	78	0
Decachlorobiphenyl	↓	10.2	9.065509	89	89	0
Decachlorobiphenyl						

Sample ID: _____

Surrogate	Column	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
Tetrachloro-m-xylene						
Tetrachloro-m-xylene						
Decachlorobiphenyl						
Decachlorobiphenyl						

Sample ID: _____

Surrogate	Column	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
Tetrachloro-m-xylene						
Tetrachloro-m-xylene						
Decachlorobiphenyl						
Decachlorobiphenyl						

Sample ID: _____

Surrogate	Column	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
Tetrachloro-m-xylene						
Tetrachloro-m-xylene						
Decachlorobiphenyl						
Decachlorobiphenyl						

Notes: _____

LDC #: 30842C3a

VALIDATION FINDINGS WORKSHEET

Laboratory Control Sample/Laboratory Control Sample Duplicate Results Verification

Page: 1 of 1
 Reviewer: BR
 2nd Reviewer: SH

METHOD: GC Pesticides/PCBs (EPA SW 846 Method 8081/8082)

The percent recoveries (%R) and Relative Percent difference (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

% Recovery = $100 * (SSC - SC) / SA$

Where: SSC = Spiked sample concentration
 SA = Spike added

SC = Concentration

RPD = $100 * |LCS - LCSD| / (LCS + LCSD)$

LCS = Laboratory control sample percent recovery

LCSD = Laboratory control sample duplicate percent recovery

LCS/LCSD samples: LCS 06274

Compound	Spike Added		Spiked Sample Concentration		LCS		LCSD		LCS/LCSD	
	<u>(ug/kg)</u>		<u>(ug/kg)</u>		Percent Recovery		Percent Recovery		RPD	
	LCS	LCSD	LCS	LCSD	Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
gamma-BHC	3.42	-	3.26	-	95	95	-	-	-	-
4,4'-DDT	7.15	-	7.34	-	103	103	-	-	-	-
Aroclor 1260										

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

METHOD: GC Pesticides/PCBs (EPA SW 846 Method 8081/8082)

Y N N/A
Y N N/A

Were all reported results recalculated and verified for all level IV samples?

Were all recalculated results for detected target compounds agree within 10.0% of the reported results?

Example:

Sample I.D. AI1 ND:

Conc. = _____

LCS 06276 RTX-CLP $\frac{5}{8}$

= Endosulfan I = 3.5 mg/kg

$$= \frac{(5996100)(10)}{(571043)(30)}$$

$$= 3.499780246 \text{ mg/kg}$$

#	Sample ID	Compound	Reported Concentration ()	Calculated Concentration ()	Qualification

Note: _____

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name: Santa Susana Field Laboratory

Collection Date: October 1, 2013

LDC Report Date: December 19, 2013

Matrix: Soil

Parameters: Polychlorinated Biphenyls

Validation Level: Level IV

Laboratory: Eurofins Lancaster Laboratories

Sample Delivery Group (SDG): PH119

Sample Identification

SL-570-SA5D-SB-0.0-0.5
SL-570-SA5D-SB-4.0-5.0
SL-564-SA5D-SB-0.0-0.5
SL-564-SA5D-SB-4.0-5.0
SL-607-SA5D-SB-0.0-0.5
SL-607-SA5D-SB-4.0-5.0

Introduction

This data review covers 6 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8082A for Polychlorinated Biphenyls.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/ECD Instrument Performance Check

Instrument performance was acceptable unless noted otherwise under initial calibration and continuing calibration sections.

III. Initial Calibration

Initial calibration of multicomponent compounds was performed for the primary (quantitation) column as required by the method.

The percent relative standard deviations (%RSD) were less than or equal to 20.0% for all compounds.

Retention time windows were evaluated and considered technically acceptable.

IV. Continuing Calibration

Continuing calibration was performed at required frequencies.

The percent differences (%D) of calibration factors in continuing standard mixtures were within the 20.0% QC limits.

The percent difference (%D) of the second source calibration standard were less than or equal to 20.0% for all compounds.

Retention times (RT) of all compounds in the calibration standards were within QC limits.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No polychlorinated biphenyl contaminants were found in the method blanks.

Sample EB-100213 (from SDG PH120) was identified as an equipment blank. No polychlorinated biphenyl contaminants were found.

Sample FB-041613 (from SDG PH032) was identified as a field blank. No polychlorinated biphenyl contaminants were found.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

LCS ID (Associated Samples)	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Affected Compound	Flag	A or P
LCS08278 (All samples in SDG PH119)	Aroclor-5442	-	90 (62-87)	-	Aroclor-5442 Aroclor-5432 Aroclor-5460	J (all detects) J (all detects) J (all detects)	P

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Florisil Cartridge Check

Florisil cleanup was not required and therefore not performed in this SDG.

XI. GPC Calibration

GPC cleanup was not required and therefore not performed in this SDG.

XII. Target Compound Identification

All target compound identifications were within validation criteria.

XIII. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH119	All compounds reported below the RL.	J (all detects)	A

XIV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XV. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Polychlorinated Biphenyls - Data Qualification Summary - SDG PH119**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-564-SA5D-SB-0.0-0.5 SL-564-SA5D-SB-4.0-5.0 SL-607-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-4.0-5.0	Aroclor-5442 Aroclor-5432 Aroclor-5460	J (all detects) J (all detects) J (all detects)	P	Laboratory control samples (%R) (L)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-564-SA5D-SB-0.0-0.5 SL-564-SA5D-SB-4.0-5.0 SL-607-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-4.0-5.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Polychlorinated Biphenyls - Laboratory Blank Data Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Polychlorinated Biphenyls - Field Blank Data Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

METHOD: GC Polychlorinated Biphenyls (EPA SW 846 Method 8082A)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 10/1/13
II.	GC/ECD Instrument Performance Check	N	
III.	Initial calibration	A	RSD ≤ 20%
IV.	Continuing calibration/ICV	A	1CV/ICV ≤ 20%
V.	Blanks	D	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	N	Client spec
VIII.	Laboratory control samples	SN	LCS/D
IX.	Regional quality assurance and quality control	N	
X.	Florisil cartridge check	N	
XI.	GPC Calibration	N	
XII.	Target compound identification	A	
XIII.	Compound quantitation/RL/LOQ/LODs	A	
XIV.	Overall assessment of data	A	
XV.	Field duplicates	N	
XVI.	Field blanks	ND	FB = FB-041613 (SD6# PA032)

EB = EB1-100213 (SD6# PH120)

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: SN

1	SL-570-SA5D-SB-0.0-0.5	11	21	31	PBLK 08278
2	SL-570-SA5D-SB-4.0-5.0	12	22	32	
3	SL-564-SA5D-SB-0.0-0.5	13	23	33	
4	SL-564-SA5D-SB-4.0-5.0	14	24	34	
5	SL-607-SA5D-SB-0.0-0.5	15	25	35	
6	SL-607-SA5D-SB-4.0-5.0	16	26	36	
7		17	27	37	
8		18	28	38	
9		19	29	39	
10		20	30	40	

Notes: _____

Method: Pesticides/PCBs (EPA SW 846 Method 8081/8082)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. GC/ECD Instrument performance check				
Was the instrument performance found to be acceptable?			/	
III. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	/			
Was a linear fit used for evaluation? If yes, were all percent relative standard deviations (%RSD) \leq 20%?	/			
Was a curve fit used for evaluation? If Yes, what was the acceptance criteria used?		/		
Did the initial calibration meet the curve fit acceptance criteria?			/	
Were the RT windows properly established?	/			
Were the required standard concentrations analyzed in the initial calibration?	/			
IV. Continuing calibration				
What type of continuing calibration calculation was performed? <u>X</u> %D or ___%R	/			
Were Evaluation mix standards analyzed prior to the initial calibration and sample analysis?			/	
Were endrin and 4,4'-DDT breakdowns \leq 15%.0 for individual breakdown in the Evaluation mix standards?			/	
Was a continuing calibration analyzed daily?	/			
Were all percent differences (%D) \leq 20%.0 or percent recoveries 80-120%?	/			
Were all the retention times within the acceptance windows?	/			
V. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was a method blank analyzed for each matrix and concentration?	/			
Were extract cleanup blanks analyzed with every batch requiring clean-up?			/	
Was there contamination in the method blanks or clean-up blanks? If yes, please see the Blanks validation completeness worksheet.			/	
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?	/			
If the percent recovery (%R) of one or more surrogates was outside QC limits, was a reanalysis performed to confirm %R?			/	

Validation Area	Yes	No	NA	Findings/Comments
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?			/	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.		/		
Was a MS/MSD analyzed every 20 samples of each matrix?			/	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?			/	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per extraction batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?		/		
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?		/		
Were the performance evaluation (PE) samples within the acceptance limits?			/	
X. Target compound identification				
Were the retention times of reported detects within the RT windows?	/			
XI. Compound quantitation/CRQLs				
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions, dry weight factors, and clean-up activities applicable to level IV validation?	/			
XII. System performance				
System performance was found to be acceptable.	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.		/		

VALIDATION FINDINGS WORKSHEET

METHOD: Pesticide/PCBs (EPASW 846 Method 8081/8082)

A. alpha-BHC	I. Dieldrin	Q. Endrin ketone	Y. Aroclor-1242	GG. Chlordane
B. beta-BHC	J. 4,4'-DDE	R. Endrin aldehyde	Z. Aroclor-1248	HH. Chlordane (Technical)
C. delta-BHC	K. Endrin	S. alpha-Chlordane	AA. Aroclor-1254	II. Aroclor 1262
D. gamma-BHC	L. Endosulfan II	T. gamma-Chlordane	BB. Aroclor-1260	JJ. Aroclor 1268
E. Heptachlor	M. 4,4'-DDD	U. Toxaphene	CC. 2,4'-DDD	KK. Oxychlordane
F. Aldrin	N. Endosulfan sulfate	V. Aroclor-1016	DD. 2,4'-DDE	LL. trans-Nonachlor
G. Heptachlor epoxide	O. 4,4'-DDT	W. Aroclor-1221	EE. 2,4'-DDT	MM. cis-Nonachlor
H. Endosulfan I	P. Methoxychlor	X. Aroclor-1232	FF. Hexachlorobenzene	NN.

Notes: _____

LDC #: 30842C26

VALIDATION FINDINGS WORKSHEET Laboratory Control Samples

Page: 1 of 1
Reviewer: BR
2nd Reviewer: SY

METHOD: GC Pesticides/PCBs (EPA SW 846 Method 8081/8082)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

N N/A Were a laboratory control samples (LCS) and laboratory control sample duplicate (LCSD) analyzed for each matrix in this SDG?

N N/A Were the LCS percent recoveries (%R) and relative percent differences (RPD) within the QC limits?

Level IV/D Only

N N/A Was a LCS analyzed every 20 samples for each matrix or whenever a sample extraction was performed?

#	Date	LCS/LCSD ID	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Associated Samples	Code: L Qualifications
		LCS08278	Arochlor 5442	()	90 (62-87)	()	All	J det/P (Qual RB-5442 PCB-5432 PCB-5460)
				()	()	()		
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LDC#: 30842C3b

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
Reviewer: BR
2nd Reviewer: DL

METHOD: GC PCBs (EPA SW 846 Method 8082A)

The calibration factors (CF), average CF, and relative standard deviation (%RSD) were recalculated for compounds identified below using the following calculations:

$$CF = A/C$$

average CF = sum of the CF/number of standards

$$\%RSD = 100 * (S/X)$$

Where:

A = Area of compound

C = Concentration of compound

S = Standard deviation of calibration factors

X = Mean of calibration factors

#	Standard ID	Calibration Date	Compound	Reported CF (500 std)	Recalculated CF (500 std)	Reported Average CF (Initial)	Recalculated Average CF (Initial)	Reported %RSD	Recalculated %RSD
1	ICAL 19843A	10/1/2013	PCB1260-1 (ZB-MultiR1)	2112	2112	2199	2199	5	5
			PCB1260-1 (ZB-MultiR2I)	2439	2439	2633	2633	10	10

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC#: 30842C3b

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Calculation Verification

Page: 1 of 1
Reviewer: BR
2nd Reviewer: ju

METHOD: GC PCBs (EPA SW 846 Method 8082)

The percent difference (%D) of the initial calibration average Calibration Factors (CF) and the continuing calibration percent difference (%D) values were recalculated for the compounds identified below using the following calculation:

$$\text{Percent difference (\%D)} = 100 * (N - C)/N$$

Where: N = Initial Calibration Factor or Nominal Amount
C = Calibration Factor from Continuing Calibration Standard or Calculated Amount

#	Standard ID	Calibration Date	Compound	Conc	Reported Conc (CCV)	Recalculated Conc (CCV)	Reported % D	Recalculated %D
1	AR163PU	10/7/2013	PCB1260 (ZB-MultiR1)	200	214.20	214.19	7	7
		21:31	PCB1260 (ZB-MultiR2)	200	205.96	205.97	3	3

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30842C36

VALIDATION FINDINGS WORKSHEET Surrogate Results Verification

Page: 1 of 1
Reviewer: BR
2nd reviewer: SM

METHOD: GC Pesticides/PCBs (EPA SW 846 Method 8081/8082)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
SS = Surrogate Spiked

Sample ID: 1

Surrogate	Column	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
Tetrachloro-m-xylene						
Tetrachloro-m-xylene	Z8 Multiresidue-1	10.1	10.782764	107	107	0
Decachlorobiphenyl	J	10.2	11.440549	113		
Decachlorobiphenyl						

Sample ID:

Surrogate	Column	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
Tetrachloro-m-xylene						
Tetrachloro-m-xylene						
Decachlorobiphenyl						
Decachlorobiphenyl						

Sample ID:

Surrogate	Column	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
Tetrachloro-m-xylene						
Tetrachloro-m-xylene						
Decachlorobiphenyl						
Decachlorobiphenyl						

Sample ID:

Surrogate	Column	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
Tetrachloro-m-xylene						
Tetrachloro-m-xylene						
Decachlorobiphenyl						
Decachlorobiphenyl						

Notes: _____

LDC #: 30842C36

VALIDATION FINDINGS WORKSHEET

Laboratory Control Sample/Laboratory Control Sample Duplicate Results Verification

Page: 1 of 1
 Reviewer: BR
 2nd Reviewer: JCA

METHOD: GC Pesticides/PCBs (EPA SW 846 Method 8081/8082)

The percent recoveries (%R) and Relative Percent difference (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

% Recovery = $100 * (SSC - SC) / SA$

Where: SSC = Spiked sample concentration
 SA = Spike added

SC = Concentration

RPD = $100 * |LCS - LCSD| / (LCS + LCSD)$

LCS = Laboratory control sample percent recovery

LCSD = Laboratory control sample duplicate percent recovery

LCS/LCSD samples: LCS 68278 / LCSD 68278

Compound	Spike Added (ug/kg)		Spiked Sample Concentration (ug/kg)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD	LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
					Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
gamma-BHC										
4,4'-DDT										
Aroclor 1260	167	-	193.64	-	116	116	-	-	-	-

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

METHOD: GC Pesticides/PCBs (EPA SW 846 Method 8081/8082)

- N N/A Were all reported results recalculated and verified for all level IV samples?
 N N/A Were all recalculated results for detected target compounds agree within 10.0% of the reported results?

ZB - Multi R1

$$PCB-1260-1 = \frac{1171316}{2199} = 532.66$$

$$1260-2 = 570.62$$

$$-3 = 592.14$$

$$-4 = 561.26$$

$$-5 = 629.12$$

$$-6 = 599.91$$

$$\text{Average} = 580.95$$

Example:

Sample I.D. ALL ND:

Conc. = _____

LC5 08278

$$BB = 193.66 \text{ ug/kg}$$

$$BB = \frac{580.95 \times 10}{30}$$

$$= 193.65 \text{ ug/kg}$$

#	Sample ID	Compound	Reported Concentration ()	Calculated Concentration ()	Qualification

Note: _____

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: October 1, 2013
LDC Report Date: December 4, 2013
Matrix: Soil
Parameters: Metals
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH119

Sample Identification

SL-570-SA5D-SB-0.0-0.5
SL-570-SA5D-SB-4.0-5.0
SL-564-SA5D-SB-0.0-0.5
SL-564-SA5D-SB-4.0-5.0
SL-607-SA5D-SB-0.0-0.5
SL-607-SA5D-SB-4.0-5.0

Introduction

This data review covers 6 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Methods 6010C, 6020A, and 7471B for Metals. The metals analyzed were Aluminum, Antimony, Arsenic, Barium, Beryllium, Boron, Cadmium, Calcium, Chromium, Cobalt, Copper, Iron, Lead, Lithium, Magnesium, Manganese, Mercury, Molybdenum, Nickel, Phosphorus, Potassium, Selenium, Silver, Sodium, Strontium, Thallium, Tin, Titanium, Vanadium, Zinc, and Zirconium.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review (January 2010).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. ICPMS Tune

The mass calibration was within 0.1 AMU and the percent relative standard deviation (%RSD) was less than or equal to 5%.

III. Calibration

The initial and continuing calibrations were performed at the required frequency.

The calibration standards criteria were met.

IV. Blanks

Method blanks were reviewed for each matrix as applicable. No metal contaminants were found in the initial, continuing and preparation blanks with the following exceptions:

Method Blank ID	Analyte	Maximum Concentration	Associated Samples
PB (prep blank)	Aluminum Calcium Tin Zinc	8.512 mg/Kg 15,220 mg/Kg 1.394 mg/Kg 1.376 mg/Kg	All samples in SDG PH119
ICB/CCB	Aluminum Copper Lead	47.7 ug/L 1.9 ug/L 2.2 ug/L	All samples in SDG PH119

Data qualification by the initial, continuing and preparation blanks (ICB/CCB/PBs) was based on the maximum contaminant concentration in the ICB/CCB/PBs in the analysis of each analyte. The sample concentrations were either not detected or were significantly greater (>5X blank contaminants) than the concentrations found in the associated method blanks with the following exceptions:

Sample	Analyte	Reported Concentration	Modified Final Concentration
SL-570-SA5D-SB-0.0-0.5	Tin	2.90 mg/Kg	2.90U mg/Kg
SL-570-SA5D-SB-4.0-5.0	Tin	0.977 mg/Kg	0.977U mg/Kg

Sample	Analyte	Reported Concentration	Modified Final Concentration
SL-564-SA5D-SB-0.0-0.5	Tin	2.75 mg/Kg	2.75U mg/Kg
SL-564-SA5D-SB-4.0-5.0	Tin	2.73 mg/Kg	2.73U mg/Kg
SL-607-SA5D-SB-0.0-0.5	Tin	3.04 mg/Kg	3.04U mg/Kg
SL-607-SA5D-SB-4.0-5.0	Tin	3.11 mg/Kg	3.11U mg/Kg

Sample EB1-100213 (from SDG PH120) was identified as an equipment blank. No metal contaminants were found with the following exceptions:

Blank ID	Sampling Date	Analyte	Concentration	Associated Samples
EB1-100213	10/2/13	Barium Boron Calcium Copper Molybdenum	0.00058 mg/L 0.0098 mg/L 0.0427 mg/L 0.0028 mg/L 0.0086 mg/L	All samples in SDG PH119

Sample FB-041613 (from SDG PH032) was identified as a field blank. No metal contaminants were found with the following exceptions:

Blank ID	Sampling Date	Analyte	Concentration	Associated Samples
FB-041613	4/11/13	Molybdenum Tin	0.0132 mg/L 0.0029 mg/L	All samples in SDG PH119

Sample concentrations were compared to concentrations detected in the field blanks. The sample concentrations were either not detected or were significantly greater (>5X blank contaminants) than the concentrations found in the associated field blanks with the following exceptions:

Sample	Analyte	Reported Concentration	Modified Final Concentration
SL-570-SA5D-SB-4.0-5.0	Tin	0.977 mg/Kg	0.977U mg/Kg

V. ICP Interference Check Sample (ICS) Analysis

The frequency of analysis was met.

The criteria for analysis were met.

VI. Matrix Spike Analysis

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

Spike ID (Associated Samples)	Analyte	MS (%R) (Limits)	MSD (%R) (Limits)	RPD (Limits)	Flag	A or P
SL-582-SA5D-SB-4.0-5.0MS/MSD (All samples in SDG PH119)	Antimony	41 (75-125)	43 (75-125)	-	J (all detects) UJ (all non-detects)	A
SL-582-SA5D-SB-4.0-5.0MS/MSD (All samples in SDG PH119)	Potassium	173 (75-125)	167 (75-125)	-	J (all detects)	A

VII. Duplicate Sample Analysis

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

IX. Internal Standards

All internal standard percent recoveries (%R) were within QC limits.

X. Furnace Atomic Absorption QC

Graphite furnace atomic absorption was not utilized in this SDG.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met.

XII. Sample Result Verification

All sample result verifications were acceptable.

All metals reported below the RL and above the MDL were qualified as follows:

Sample	Analyte	Flag	A or P
All samples in SDG PH119	All analytes reported below the RL and above the MDL.	J (all detects)	A

XIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Metals - Data Qualification Summary - SDG PH119**

SDG	Sample	Analyte	Flag	A or P	Reason (Code)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-564-SA5D-SB-0.0-0.5 SL-564-SA5D-SB-4.0-5.0 SL-607-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-4.0-5.0	Antimony	J (all detects) UJ (all non-detects)	A	Matrix spike/Matrix spike duplicate (%R) (Q)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-564-SA5D-SB-0.0-0.5 SL-564-SA5D-SB-4.0-5.0 SL-607-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-4.0-5.0	Potassium	J (all detects)	A	Matrix spike/Matrix spike duplicate (%R) (Q)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-564-SA5D-SB-0.0-0.5 SL-564-SA5D-SB-4.0-5.0 SL-607-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-4.0-5.0	All analytes reported below the RL and above the MDL.	J (all detects)	A	Sample result verification (Z)

**Santa Susana Field Laboratory
Metals - Laboratory Blank Data Qualification Summary - SDG PH119**

SDG	Sample	Analyte	Modified Final Concentration	A or P	Code
PH119	SL-570-SA5D-SB-0.0-0.5	Tin	2.90U mg/Kg	A	B
PH119	SL-570-SA5D-SB-4.0-5.0	Tin	0.977U mg/Kg	A	B
PH119	SL-564-SA5D-SB-0.0-0.5	Tin	2.75U mg/Kg	A	B
PH119	SL-564-SA5D-SB-4.0-5.0	Tin	2.73U mg/Kg	A	B
PH119	SL-607-SA5D-SB-0.0-0.5	Tin	3.04U mg/Kg	A	B
PH119	SL-607-SA5D-SB-4.0-5.0	Tin	3.11U mg/Kg	A	B

**Santa Susana Field Laboratory
Metals - Field Blank Data Qualification Summary - SDG PH119**

SDG	Sample	Analyte	Modified Final Concentration	A or P	Code
PH119	SL-570-SA5D-SB-4.0-5.0	Tin	0.977U mg/Kg	A	F

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification											
	FB-041613 (SDG: PH032)	Action Limit	2										
Mo	0.0132	6.60											
Sn	0.0029	1.45	0.977										

Sampling date: 10/2/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification											
	EB1-100213 (SDG: PH120)	Action Limit	No Qualifiers										
Ba	0.00058	0.29											
B	0.0098	4.9											
Ca	0.0427	21.35											
Cu	0.0028	1.4											
Mo	0.0086	4.3											

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

LDC #: 30842C4

VALIDATION FINDINGS WORKSHEET
Matrix Spike/Matrix Spike Duplicates

METHOD: Trace metals (EPA SW 846 Method 6010B/6020A/7000)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Y N N/A

Was a matrix spike analyzed for each matrix in this SDG?

Y N N/A

Were matrix spike percent recoveries (%R) within the control limits of 75-125? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.

Y N N/A

Were all duplicate sample relative percent differences (RPD) \leq 20% for water samples and \leq 35% for soil samples?

LEVEL IV ONLY:

Y N N/A

Were recalculated results acceptable? See Level IV Recalculation Worksheet for recalculations.

#	MS/MSD ID	Matrix	Analyte	MS %Recovery	MSD %Recovery	RPD (Limits)	Associated Samples	Qualifications
	SL582-SA5D-SB (QH118)	4.0-5.0mg/msd	Sb	41	43		All	J/UJA (Q)
			K	173	167		↓	Jdet/A ↓

Comments: _____

LDC #: 3084209

VALIDATION FINDINGS WORKSHEET
Initial and Continuing Calibration Calculation Verification

Page: 1 of 1
 Reviewer: GR
 2nd Reviewer: [Signature]

METHOD: Trace Metals (EPA SW 846 Method 6010/6020/7000)

An initial and continuing calibration verification percent recovery (%R) was recalculated for each type of analysis using the following formula:

$$\%R = \frac{\text{Found}}{\text{True}} \times 100$$
 Where, Found = concentration (in ug/L) of each analyte measured in the analysis of the ICV or CCV solution
 True = concentration (in ug/L) of each analyte in the ICV or CCV source

Standard ID	Type of Analysis	Element	Found (ug/L)	True (ug/L)	Recalculated	Reported	Acceptable (Y/N)
					%R	%R	
ICV	ICP (Initial calibration)	Li	609.56	600	101.6	101.6	Y ↓
ICV	ICP/MS (Initial calibration)	Sr	52.22	50	104.4	104.4	
ICV	CVAA (Initial calibration)	Hg	2.54	2.5		101.6	
CCV(15:32)	ICP (Continuing calibration)	V	522.22	500	104.4	104.4	
ICV(23:26)	ICP/MS (Continuing calibration)	Tl	25.51	25	102	102.0	
ICV(06:31)	CVAA (Continuing calibration)	Hg	1.03	1.0	103.0	103.0	
	GFAA (Initial calibration)						
	GFAA (Continuing calibration)						

Comments: Refer to Calibration Verification findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the calculated results.

LDC #: 3084209

VALIDATION FINDINGS WORKSHEET

Level IV Recalculation Worksheet

Page: 1 of 1
 Reviewer: 92
 2nd Reviewer: [Signature]

METHOD: Trace Metals (EPA SW 846 Method 6010/6020/7000)

Percent recoveries (%R) for an ICP interference check sample, a laboratory control sample and a matrix spike sample were recalculated using the following formula:

$$\%R = \frac{\text{Found}}{\text{True}} \times 100$$

Where, Found = Concentration of each analyte measured in the analysis of the sample. For the matrix spike calculation, Found = SSR (spiked sample result) - SR (sample result).
 True = Concentration of each analyte in the source.

A sample and duplicate relative percent difference (RPD) was recalculated using the following formula:

$$RPD = \frac{|S-D|}{(S+D)/2} \times 100$$

Where, S = Original sample concentration
 D = Duplicate sample concentration

An ICP serial dilution percent difference (%D) was recalculated using the following formula:

$$\%D = \frac{|I-SDR|}{I} \times 100$$

Where, I = Initial Sample Result (mg/L)
 SDR = Serial Dilution Result (mg/L) (Instrument Reading x 5)

Sample ID	Type of Analysis	Element	Found / S / I (units)	True / D / SDR (units)	Recalculated	Reported	Acceptable (Y/N)
					%R / RPD / %D	%R / RPD / %D	
ICSAS	ICP interference check	Mn	486.5	500	97.3	97.3	Y
LCS	Laboratory control sample	P	106325	100000	106	106	
SL-582-SA50-SB-4.0-5.0	Matrix spike	Mn	(SSR-SR) 182.0455	198.0198	92	92	
↓	Duplicate	Cr	30.1436	30.4535	1	1	
↓	ICP serial dilution	Ca	233954.73	239378.7	2	2	

Comments: Refer to appropriate worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30842C4

VALIDATION FINDINGS WORKSHEET

Sample Calculation Verification

Page: 1 of 1
Reviewer: OR
2nd reviewer: W

METHOD: Trace Metals (EPA SW 846 Method 6010/6020/7000)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- | | | | |
|---------------------------------------|----------------------------|------------------------------|--|
| <input checked="" type="checkbox"/> Y | <input type="checkbox"/> N | <input type="checkbox"/> N/A | Have results been reported and calculated correctly? |
| <input checked="" type="checkbox"/> Y | <input type="checkbox"/> N | <input type="checkbox"/> N/A | Are results within the calibrated range of the instruments and within the linear range of the ICP? |
| <input checked="" type="checkbox"/> Y | <input type="checkbox"/> N | <input type="checkbox"/> N/A | Are all detection limits below the CRDL? |

Detected analyte results for Ba were recalculated and verified using the following equation:

$$\text{Concentration} = \frac{(\text{RD})(\text{FV})(\text{Dil})}{(\text{In. Vol.})}$$

Recalculation:

- RD = Raw data concentration
- FV = Final volume (ml)
- In. Vol. = Initial volume (ml) or weight (G)
- Dil = Dilution factor

$$\frac{1.1839 \text{ mg/L} (50 \text{ mL}) (2)}{1.02 \text{ g} (0.917)} = 126.57 \text{ mg/kg}$$

#	Sample ID	Analyte	Reported Concentration (mg/kg)	Calculated Concentration (mg/kg)	Acceptable (Y/N)
	1	Ba	127	127	Y
	2	Mn	297	297	Y
	3	Hg	0.026	0.026	Y
	4	Se	0.20	0.20	Y
	5	Ti	1570	1570	Y
	6	Co	9.3	9.3	Y

Note: _____

LDC #: 30842C4

VALIDATION COMPLETENESS WORKSHEET

Date: 12/4/13

SDG #: PH119

Level IV

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: [Signature]

74713

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 10/1/13
II.	ICP/MS Tune	A	
III.	Calibration	A	
IV.	Blanks	SW	
V.	ICP Interference Check Sample (ICS) Analysis	A	
VI.	Matrix Spike Analysis	SW	MS/D (From PH118)
VII.	Duplicate Sample Analysis	A	Dup ↓
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	A	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	A	
XII.	Sample Result Verification	A	
XIII.	Overall Assessment of Data	A	
XIV.	Field Duplicates	N	
XV.	Field Blanks	SW	FB=FB-041613 EB=EB1-100213

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

(PH032)

(PH120)

Validated Samples:

Soil

1	SL-570-SA5D-SB-0.0-0.5	11		21		31	
2	SL-570-SA5D-SB-4.0-5.0	12		22		32	
3	SL-564-SA5D-SB-0.0-0.5	13		23		33	
4	SL-564-SA5D-SB-4.0-5.0	14		24		34	
5	SL-607-SA5D-SB-0.0-0.5	15		25		35	
6	SL-607-SA5D-SB-4.0-5.0	16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

Method: Metals (EPA SW 846 Method 6010B/7000/6020)

Validation Area	Yes	No	NA	Findings/Comments
<i>I. Technical holding times</i>				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
<i>II. ICP/MS Tune</i>				
Were all isotopes in the tuning solution mass resolution within 0.1 amu?	/			
Were %RSD of isotopes in the tuning solution $\leq 5\%$?	/			
<i>III. Calibration</i>				
Were all instruments calibrated daily, each set-up time?	/			
Were the proper number of standards used?	/			
Were all initial and continuing calibration verification %Rs within the 90-110% (80-120% for mercury) QC limits?	/			
Were all initial calibration correlation coefficients > 0.995 ?	/			
<i>IV. Blanks</i>				
Was a method blank associated with every sample in this SDG?	/			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	/			
<i>V. ICP Interference Check Sample</i>				
Were ICP interference check samples performed daily?	/			
Were the AB solution percent recoveries (%R) with the 80-120% QC limits?	/			
<i>VI. Matrix spike/Matrix spike duplicates</i>				
Were a matrix spike (MS) and duplicate (DUP) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD or MS/DUP. Soil / Water.	/			
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the 75-125 QC limits? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.		/		
Were the MS/MSD or duplicate relative percent differences (RPD) $\leq 20\%$ for waters and $\leq 35\%$ for soil samples? A control limit of $\pm RL$ ($\pm 2X RL$ for soil) was used for samples that were $\leq 5X$ the RL, including when only one of the duplicate sample values were $\leq 5X$ the RL.	/			
<i>VII. Laboratory control samples</i>				
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per extraction batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the 80-120% QC limits for water samples and laboratory established QC limits for soils?	/			

Validation Area	Yes	No	NA	Findings/Comments
VIII. Furnace Atomic Absorption QC				
If MSA was performed, was the correlation coefficients > 0.995?			/	
Do all applicable analyses have duplicate injections? (Level IV only)			/	
For sample concentrations > RL, are applicable duplicate injection RSD values < 20%? (Level IV only)			/	
Were analytical spike recoveries within the 85-115% QC limits?			/	
IX. ICP Serial Dilution				
Was an ICP serial dilution analyzed if analyte concentrations were > 50X the MDL (ICP)/>100X the MDL(ICP/MS)?	/			
Were all percent differences (%Ds) < 10%?	/			
Was there evidence of negative interference? If yes, professional judgement will be used to qualify the data.			/	
X. Internal Standards (EPA SW 846 Method 6020/EPA 200.8)				
Were all the percent recoveries (%R) within the 30-120% (6020)/60-125% (200.8) of the intensity of the internal standard in the associated initial calibration?	/			
If the %Rs were outside the criteria, was a reanalysis performed?			/	
XI. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?		/		
Were the performance evaluation (PE) samples within the acceptance limits?			/	
XII. Sample Result Verification				
Were RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target analytes were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target analytes were detected in the field blanks.	/			

**VALIDATION FINDINGS WORKSHEET
PB/ICB/CCB QUALIFIED SAMPLES**

METHOD: Trace metals (EPA SW 864 Method 6010B/6020/7000)

Soil preparation factor applied: 100x x 2xdil

Reason: B

Sample Concentration units, unless otherwise noted: mg/Kg

Associated Samples: All

Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/L)	Maximum ICB/CCB ^a (ug/L)	Action Level	Sample Identification											
					1	2	3	4	5	6						
Al	8.512		47.7	47.7												
Ca	15.220			76.1												
Cu			1.9	1.9												
Pb			2.2	2.2												
Sn	1.394			6.97	2.90	0.977	2.75	2.73	3.04	3.11						
Zn	1.376			6.88												

Samples with analyte concentrations within five times the associated ICB, CCB or PB concentration are listed above with the identifications from the Validation Completeness Worksheet. These sample results were qualified as not detected, "U".

Note : a - The listed analyte concentration is the highest ICB, CCB, or PB detected in the analysis of each element.

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: October 1, 2013
LDC Report Date: December 18, 2013
Matrix: Soil
Parameters: Herbicides
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories

Sample Delivery Group (SDG): PH119

Sample Identification

SL-570-SA5D-SB-0.0-0.5
SL-570-SA5D-SB-4.0-5.0
SL-570-SA5D-SB-0.0-0.5MS
SL-570-SA5D-SB-0.0-0.5MSD

Introduction

This data review covers 4 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8151A for Herbicides.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Initial Calibration

Initial calibration was performed as required by the method.

The percent relative standard deviations (%RSD) were less than or equal to 20.0% for all compounds.

Retention time windows were evaluated and considered technically acceptable.

III. Continuing Calibration

Continuing calibration was performed at the required frequencies.

The percent differences (%D) were less than or equal to 20.0% for all compounds with the following exceptions:

Date	Standard	Column	Compound	%D	Associated Samples	Flag	A or P
10/11/13	HERB3SY	MR-1	Dinoseb	26	PBLK19282	J (all detects) UJ (all non-detects)	A
10/11/13	HERB3SY	MR-2	Dinoseb	22	PBLK19282	J (all detects) UJ (all non-detects)	A
10/11/13	HERB3SZ	MR-1	Dinoseb	21	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-570-SA5D-SB-0.0-0.5MS SL-570-SA5D-SB-0.0-0.5MSD	J (all detects) UJ (all non-detects)	A
10/11/13	HERB3SZ	MR-2	Dinoseb	34	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-570-SA5D-SB-0.0-0.5MS SL-570-SA5D-SB-0.0-0.5MSD	J (all detects) UJ (all non-detects)	A

The percent differences (%D) of the second source calibration standard were less than or equal to 20.0% for all compounds.

Retention times (RT) of all compounds in the calibration standards were within QC limits.

IV. Blanks

Method blanks were reviewed for each matrix as applicable. No herbicide contaminants were found in the method blanks.

Sample EB1-100213 (from SDG PH120) was identified as an equipment blank. No herbicide contaminants were found.

Sample FB-041613 (from SDG PH032) was identified as a field blank. No herbicide contaminants were found.

V. Surrogate Recovery

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. Matrix Spike/Matrix Spike Duplicate

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Target Compound Identification

All target compound identifications were within validation criteria.

IX. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH119	All compounds reported below the RL.	J (all detects)	A

X. System Performance

The system performance was acceptable.

XI. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XII. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Herbicides - Data Qualification Summary - SDG PH119**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0	Dinoseb	J (all detects) UJ (all non-detects)	A	Continuing calibration (%D) (C)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Herbicides - Laboratory Blank Data Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Herbicides - Field Blank Data Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

LDC #: 30842C5

VALIDATION COMPLETENESS WORKSHEET

Date: 12/3/13

SDG #: PH119

Level IV

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: RA

2nd Reviewer: SA

METHOD: GC Herbicides (EPA SW 846 Method 8151A)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 10/1/13
II	Initial calibration	A	RSD ≤ 20%
III.	Calibration verification/ICV	SW	ICV/COV ≤ 20%
IV.	Blanks	A	
V	Surrogate recovery	A	
VI.	Matrix spike/Matrix spike duplicates	D	
VII.	Laboratory control samples	A	LCS
VIII.	Target compound identification	A	
IX.	Compound quantitation/RL/LOQ/LODs	A	
X.	System Performance	A	
XI.	Overall assessment of data	D	
XII.	Field duplicates	N	
XIII.	Field blanks	ND	FB = FB-041613 (SDG # PH132) EB = EB1-100213 (SDG # PH120)

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: Soil

1	SL-570-SA5D-SB-0.0-0.5	11		21		31	PBLK19282
2	SL-570-SA5D-SB-4.0-5.0	12		22		32	
3	#1MS	13		23		33	
4	#1MSD	14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes:

LDC #: 30842CS

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: BR
 2nd Reviewer: SM

Method: GC HPLC

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	/			
Was a linear fit used for evaluation? If yes, were all percent relative standard deviations (%RSD) \leq 20%?	/			
Was a curve fit used for evaluation? If Yes, what was the acceptance criteria used?			/	
Did the initial calibration meet the curve fit acceptance criteria?			/	
Were the RT windows properly established?	/			
IV. Continuing calibration				
What type of continuing calibration calculation was performed? <input checked="" type="checkbox"/> %D or <input type="checkbox"/> %R	/			
Was a continuing calibration analyzed daily?	/			
Were all percent differences (%D) \leq 20%.0 or percent recoveries 80-120%?	/			
Were all the retention times within the acceptance windows?			/	
V. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was a method blank analyzed for each matrix and concentration?	/			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.			/	
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?	/			
If the percent recovery (%R) of one or more surrogates was outside QC limits, was a reanalysis performed to confirm %R?			/	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?			/	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	/			
Was a MS/MSD analyzed every 20 samples of each matrix?	/			
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	/			
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per extraction batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	/			

LDC #: 3084205

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: BR
 2nd Reviewer: SA

Validation Area	Yes	No	NA	Findings/Comments
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?		/		
Were the performance evaluation (PE) samples within the acceptance limits?			/	
X. Target compound identification				
Were the retention times of reported detects within the RT windows?	/			
XI. Compound quantitation/CRQLs				
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XII. System performance				
System performance was found to be acceptable.	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.		/		

VALIDATION FINDINGS WORKSHEET

METHOD: GC HPLC

8310	8330	8151	8141	8141(Con't)	8021B
A. Acenaphthene	A. HMX	A. 2,4-D	A. Dichlorvos	V. Fensulfothion	V. Benzene
B. Acenaphthylene	B. RDX	B. 2,4-DB	B. Mevinphos	W. Bolstar	CC. Toluene
C. Anthracene	C. 1,3,5-Trinitrobenzene	C. 2,4,5-T	C. Demeton-O	X. EPN	EE. Ethylbenzene
D. Benzo(a)anthracene	D. 1,3-Dinitrobenzene	D. 2,4,5-TP	D. Demeton-S	Y. Azinphos-methyl	SSS. o-Xylene
E. Benzo(a)pyrene	E. Tetryl	E. Dinoseb	E. Ethoprop	Z. Coumaphos	RRR. m,p-Xylenes
F. Benzo(b)fluoranthene	F. Nitrobenzene	F. Dichlorprop	F. Naled	AA. Parathion	GG. Total xylenes
G. Benzo(g,h,i)perylene	G. 2,4,6-Trinitrotoluene	G. Dicamba	G. Sulfotepp	BB. Famphur	
H. Benzo(k)fluoranthene	H. 4-Amino-2,6-dinitrotoluene	H. Dalapon	H. Phorate	CC. Phosmet	
I. Chrysene	I. 2-Amino-4,6-dinitrotoluene	I. MCPP	I. Dimethoate	DD. Trifluralin	
J. Dibenz(a,h)anthracene	J. 2,4-Dinitrotoluene	J. MCPA	J. Diazinon	EE. Def	
K. Fluoranthene	K. 2,6-Dinitrotoluene	K. Pentachlorophenol	K. Disulfoton	FF. Prowl	
L. Fluorene	L. 2-Nitrotoluene	L. 2,4,5-TP (silvex)	L. Parathion-methyl	GG. Ethion	
M. Indeno(1,2,3-cd)pyrene	M. 3-Nitrotoluene	M. Silvex	M. Ronnel	HH. Tetrachlorvinphos	
N. Naphthalene	N. 4-Nitrotoluene		N. Malathion	II. Sulprofos	
O. Phenanthrene	O.		O. Chlorpyrifos		
P. Pyrene	P.		P. Fenthion		
Q.	Q		Q. Parathion-ethyl		
R.			R. Trichloronate		
S.			S. Merphos		
			T. Stirophos		
			U. Tokuthion		

Notes: _____

LDC #: 30842CS

VALIDATION FINDINGS WORKSHEET Continuing Calibration

METHOD: GC HPLC

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

N N/A Were continuing calibration standards analyzed at the required frequencies?

N N/A Did the continuing calibration standards meet the %D validation criteria of $\leq 20.0\%$?

Level IV Only

N N/A Were the retention times for all calibrated compounds within their respective acceptance windows?

#	Date	Standard ID	Detector/ Column	Compound	%D (Limit)	RT (limit)	Associated Samples	Col: C Qualifications
	10/11/13	CCV-HERB3SY	MR-1	E	for 26 26	()	PBLK19282	J/WJA
			MR-2	E	22	()		↓
						()		
						()		
	10/11/13	CCV-HERB3SZ	MR-1	E	21	()	1-4	J/WJA
			MR-2	E	34	()		↓
						()		
						()		
						()		
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						()		
						()		

LDC#: 30842C5

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: BR
 2nd Reviewer: SM

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$$RRF = (A_x)(C_{is}) / (A_{is})(C_x)$$

average RRF = sum of the RRFs/number of standards

$$\%RSD = 100 * (S/X)$$

A_x = Area of Compound A_{is} = Area of associated internal standard

C_x = Concentration of compound, C_{is} = Concentration of internal standard

S= Standard deviation of the RRF X = Mean of the RRFs

#	Standard ID	Calibration Date	Compound	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				RRF (RRF 176 std)	RRF (RRF 176 std)	Average RRF (Initial)	Average RRF (Initial)	%RSD	%RSD
1	ICAL	10/7/2013	2,4-D (MR-1)	1.38E-03	1.04E-03	1.34E-03	1.34E-03	5	5
	18273A		Dinoseb (MR-1)	2.74E-04	1.26E-03	3.16E-04	3.16E-04	9	9
			2,4-D (MR-2)	1.30E-03	1.21E-03	1.35E-03	1.35E-03	5	5
			Dinoseb (MR-2)	2.56E-04	1.07E-03	3.13E-04	3.13E-04	15	15

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC#: 30842C5

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Calculation Verification

Page: 1 of 1
 Reviewer: BR
 2nd Reviewer: 84

METHOD: GC Pesticides (EPA SW 846 Method 8081)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

% Difference = 100 * (ave. RRF - RRF)/ave. RRF
 RRF = (Ax)(Cis)/(Ais)(Cx)

Where: ave. RRF = initial calibration average RRF
 RRF = continuing calibration RRF
 Ax = Area of compound,
 Cx = Concentration of compound,
 Ais = Area of associated internal standard
 Cis = Concentration of internal standard

#	Standard ID	Calibration Date	Compound	Average RRF (Initial)	Reported	Recalculated	Reported	Recalculated
					RRF (CC)	RRF (CC)	% D	%D
1	HERB3SY	10/11/2013 14:32	2,4-D (MR-1)	176.00	179.10	179.10	2	2
			Dinoseb (MR-1)	100.70	127.33	127.41	26	26
			2,4-D (MR-2)	176.00	171.76	171.76	2	2
			Dinoseb (MR-2)	100.70	123.01	122.91	22	22
2	HERB3SZ	10/11/2013 17:31	2,4-D (MR-1)	176.00	180.10	180.10	2	2
			Dinoseb (MR-1)	100.70	121.85	121.93	21	21
			2,4-D (MR-2)	176.00	176.27	176.27	0	0
			Dinoseb (MR-2)	100.70	134.98	134.87	34	34

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results

LDC #: 308205

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

Page: 1 of 1
Reviewer: BR
2nd reviewer: 84

METHOD: GC HPLC

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
SS = Surrogate Spiked

Sample ID: 1

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
<u>2,4-Dichlorophenylacetic acid MR-1</u>		<u>669</u>	<u>58.048141</u>	<u>87</u>	<u>87</u>	<u>0</u>

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

LDC #: 30842CS

VALIDATION FINDINGS WORKSHEET

Matrix Spike/Matrix Spike Duplicates Results Verification

Page: 1 of 1

Reviewer: BR

2nd Reviewer: 84

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the matrix spike and matrix spike duplicate were recalculated for the compounds identified below using the following calculation:

$\% \text{Recovery} = 100 * (\text{SSC} - \text{SC}) / \text{SA}$

Where

SSC = Spiked sample concentration

MS = Matrix spike

SC = Sample concentration

MSD = Matrix spike duplicate

$\text{RPD} = ((\text{SSCMS} - \text{SSCMSD}) * 2) / (\text{SSCMS} + \text{SSCMSD}) * 100$

SA = Spike added

MS/MSD samples: 3/4

Compound	Spike Added (ug/kg)		Sample Conc. (ug/kg)	Spike Sample Concentration (ug/kg)		Matrix spike		Matrix Spike Duplicate		MS/MSD	
	MS	MSD		MS	MSD	Percent Recovery		Percent Recovery		RPD	
						Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)											
Diesel (8015)											
Benzene (8021B)											
Methane (RSK-175)											
2,4-D (8151)	83.7	83.7	0 ↓	82.92	84.39	99	99	101	101	82.3	2 1.8
Dinoseb (8151)	142	142	↓	48.32	62.0	34	34	44	44	25	25
Naphthalene (8310)											
Anthracene (8310)											
HMX (8330)											
2,4,6-Trinitrotoluene (8330)											

Comments: Refer to Matrix Spike/Matrix Spike Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3884205

VALIDATION FINDINGS WORKSHEET

Page: 1 of 1

Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Reviewer: BR

2nd Reviewer: Sy

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

%Recovery = 100 * (SSC - SC)/SA

Where SSC = Spiked sample concentration

SC = Sample concentration

RPD = (((SSCLCS - SSCLCSD} * 2) / (SSCLCS + SSCLCSD))*100

SA = Spike added

LCS = Laboratory Control Sample

LCSD = Laboratory Control Sample duplicate

LCS/LCSD samples: LCS 19282

Compound	Spike Added (<u>ug/kg</u>)		Spike Sample Concentration (<u>ug/kg</u>)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD	LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
					Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)										
Diesel (8015)										
Benzene (8021B)										
Methane (RSK-175)										
2,4-D (8151)	<u>83.7</u>	<u>-</u>	<u>87.17</u>	<u>-</u>	<u>104</u>	<u>104</u>	<u>-</u>	<u>-</u>	<u>-</u>	<u>-</u>
Dinoseb (8151)	<u>142</u>	<u>-</u>	<u>48.82</u>	<u>-</u>	<u>34</u>	<u>34</u>	<u>-</u>	<u>-</u>	<u>-</u>	<u>-</u>
Naphthalene (8310)										
Anthracene (8310)										
HMX (8330)										
2,4,6-Trinitrotoluene (8330)										

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3084205

VALIDATION FINDINGS WORKSHEET Sample Calculation Verification

Page: 1 of 1
Reviewer: BR
2nd Reviewer: [Signature]

METHOD: GC HPLC

N N/A Were all reported results recalculated and verified for all level IV samples?
 N N/A Were all recalculated results for detected target compounds within 10% of the reported results?

Concentration = $\frac{(A)(Fv)(Df)}{(RF)(Vs \text{ or } Ws)(\%S/100)}$

Example:

Sample ID: All Compound Name ND
LCS 19282 MR-1
2.4-D = 87.17
Concentration = $\frac{(163566)(1)(10)}{(46453)(0.001340864)(30)}$
= 87.17418172 mg/l

#	Sample ID	Compound	Reported Concentrations ()	Recalculated Results Concentrations ()	Qualifications

Comments: _____

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: October 1, 2013
LDC Report Date: December 4, 2013
Matrix: Soil
Parameters: Wet Chemistry
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH119

Sample Identification

SL-598-SA5D-SB-0.0-0.5
SL-598-SA5D-SB-4.0-5.0
SL-598-SA5D-SB-9.0-10.0
SL-598-SA5D-SB-14.0-15.0
SL-598-SA5D-SB-18.0-19.0
SL-598-SA5D-SB-0.0-0.5MS
SL-598-SA5D-SB-0.0-0.5DUP

Introduction

This data review covers 7 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA Method 300.0 for Fluoride.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review (January 2010).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Initial Calibration

All criteria for the initial calibration of each method were met.

III. Calibration Verification

Calibration verification frequency and analysis criteria were met for each method when applicable.

IV. Blanks

Method blanks were reviewed for each matrix as applicable. No contaminant concentrations were found in the initial, continuing and preparation blanks.

Sample EB1-100213 (from SDG PH120) was identified as an equipment blank. No contaminant concentrations were found.

Sample FB-041613 (from SDG PH032) was identified as a field blank. No contaminant concentrations were found.

V. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) analyses were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits with the following exceptions:

Spike ID (Associated Samples)	Analyte	%R (Limits)	Flag	A or P
SL-598-SA5D-SB-0.0-0.5MS (All samples in SDG PH119)	Fluoride	49 (80-120)	J (all detects) UJ (all non-detects)	A

VI. Duplicates

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

VII. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Sample Result Verification

All sample result verifications were acceptable.

All analytes reported below the RL and above the MDL were qualified as follows:

Sample	Analyte	Flag	A or P
All samples in SDG PH119	All analytes reported below the RL and above the MDL.	J (all detects)	A

IX. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

X. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Wet Chemistry - Data Qualification Summary - SDG PH119**

SDG	Sample	Analyte	Flag	A or P	Reason (Code)
PH119	SL-598-SA5D-SB-0.0-0.5 SL-598-SA5D-SB-4.0-5.0 SL-598-SA5D-SB-9.0-10.0 SL-598-SA5D-SB-14.0-15.0 SL-598-SA5D-SB-18.0-19.0	Fluoride	J (all detects) U (all non-detects)	A	Matrix spike (%R) (Q)
PH119	SL-598-SA5D-SB-0.0-0.5 SL-598-SA5D-SB-4.0-5.0 SL-598-SA5D-SB-9.0-10.0 SL-598-SA5D-SB-14.0-15.0 SL-598-SA5D-SB-18.0-19.0	All analytes reported below the RL and above the MDL.	J (all detects)	A	Sample result verification (Z)

**Santa Susana Field Laboratory
Wet Chemistry – Laboratory Blank Data Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Wet Chemistry - Field Blank Data Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

LDC #: 30842C6

VALIDATION COMPLETENESS WORKSHEET

Date: 12/4/13

SDG #: PH119

Level IV

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: AL

2nd Reviewer: [Signature]

METHOD: (Analyte) Fluoride (EPA Method 300.0)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 10/1/13
IIa.	Initial calibration	A	
IIb.	Calibration verification	A	
III.	Blanks	A	
IV.	Matrix Spike/Matrix Spike Duplicates	SW	ES = MS
V.	Duplicates	A/A	Dup
VI.	Laboratory control samples	A	LCS
VII.	Sample result verification	A	
VIII.	Overall assessment of data	A	
IX.	Field duplicates	N	
X.	Field blanks	ND	FB = FB-041613 EB = EB1-100213

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

(PH032)
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank
 (PH120)

Validated Samples: Soil

1	SL-598-SA5D-SB-0.0-0.5	11		21		31	
2	SL-598-SA5D-SB-4.0-5.0	12		22		32	
3	SL-598-SA5D-SB-9.0-10.0	13		23		33	
4	SL-598-SA5D-SB-14.0-15.0	14		24		34	
5	SL-598-SA5D-SB-18.0-19.0	15		25		35	
6	(X) MS	16		26		36	
7	↓ Dup	17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

Method: Inorganics (EPA Method See cover)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. Calibration				
Were all instruments calibrated daily, each set-up time?	/			
Were the proper number of standards used?	/			
Were all initial calibration correlation coefficients > 0.995?	/			
Were all initial and continuing calibration verification %Rs within the 90-110% QC limits?	/			
Were titrant checks performed as required? (Level IV only)			/	
Were balance checks performed as required? (Level IV only)			/	
III. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.		/		
IV. Matrix spike/Matrix spike duplicates and Duplicates				
Were a matrix spike (MS) and duplicate (DUP) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD or MS/DUP. Soil / Water.	/			
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the 75-125 QC limits? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.		/		
Were the MS/MSD or duplicate relative percent differences (RPD) ≤ 20% for waters and ≤ 35% for soil samples? A control limit of ≤ CRDL (≤ 2X CRDL for soil) was used for samples that were ≤ 5X the CRDL, including when only one of the duplicate sample values were ≤ 5X the CRDL.	/			
V. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per extraction batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the 80-120% (85-115% for Method 300.0) QC limits?	/			
VI. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?		/		
Were the performance evaluation (PE) samples within the acceptance limits?			/	

LDC #: 308426

VALIDATION FINDINGS CHECKLIST

Page: 2 of 3
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Validation Area	Yes	No	NA	Findings/Comments
VII. Sample Result Verification				
Were RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were detection limits < RL?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
VIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IX. Field duplicates				
Field duplicate pairs were identified in this SDG.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target analytes were detected in the field duplicates.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
X. Field blanks				
Field blanks were identified in this SDG.	<input checked="" type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target analytes were detected in the field blanks.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	

LDC #: 30812Cb

VALIDATION FINDINGS WORKSHEET
Matrix Spike Analysis

Page: (of)
Reviewer: g
2nd Reviewer: W

METHOD: Inorganics, Method See card

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- N N/A Was a matrix spike analyzed for each matrix in this SDG?
- N N/A Were matrix spike percent recoveries (%R) within the control limits of 75-125 (85-115% for Method 300.0)? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.

LEVEL IV ONLY:

- N N/A Were recalculated results acceptable? See Level IV Recalculation Worksheet for recalculations.

#	Date	Matrix Spike ID	Matrix	Analyte	%R	Associated Samples	Qualifications
		6	S	F	49 (80-120)	All	JLJ/A (R)

Comments: _____

LDC #: 308426

Validation Findings Worksheet
Initial and Continuing Calibration Calculation Verification

Page: 1 of 1
 Reviewer: aw
 2nd Reviewer: h

Method: Inorganics, Method See Cover

The correlation coefficient (r) for the calibration of F was recalculated. Calibration date: 9/18/13

An initial or continuing calibration verification percent recovery (%R) was recalculated for each type of analysis using the following formula:

$$\%R = \frac{\text{Found} \times 100}{\text{True}}$$

Where, Found = concentration of each analyte measured in the analysis of the ICV or CCV solution
 True = concentration of each analyte in the ICV or CCV source

Type of analysis	Analyte	Standard	Conc. (mg/L)	Area	Recalculated	Reported	Acceptable (Y/N)
					r or r ²	r or r ²	
Initial calibration	F	s1	0.0	0	0.999575	0.999603	Y
		s2	0.1	0.017			
		s3	0.4	0.071			
		s4	1	0.188			
		s5	2	0.377			
		s6	3	0.584			
Calibration verification		ICV	1.5	1.515993	101	101	
Calibration verification		CCV		1.517236	101	101	
Calibration verification	↓	↓	↓	1.505556	100	100	↓

Comments: Refer to Calibration Verification findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Level IV Recalculation Worksheet

METHOD: Inorganics, Method see cover

Percent recoveries (%R) for a laboratory control sample and a matrix spike sample were recalculated using the following formula:

$$\%R = \frac{\text{Found}}{\text{True}} \times 100$$
 Where, Found = concentration of each analyte measured in the analysis of the sample. For the matrix spike calculation, Found = SSR (spiked sample result) - SR (sample result).
True = concentration of each analyte in the source.

A sample and duplicate relative percent difference (RPD) was recalculated using the following formula:

$$RPD = \frac{|S-D|}{(S+D)/2} \times 100$$
 Where, S = Original sample concentration
D = Duplicate sample concentration

Sample ID	Type of Analysis	Element	Found / S (units)	True / D (units)	Recalculated	Reported	Acceptable (Y/N)
					%R / RPD	%R / RPD	
LCS	Laboratory control sample	F	5.16	S	103	103	Y
Aa 6	Matrix spike sample	↓	(SSR-SR) 2.44	4.98	49	49	↓
Aa 7	Duplicate sample	↓	2.01	1.96	4	4	↓

Comments: Refer to appropriate worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30842CC6

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

Page: 1 of 1
Reviewer: CR
2nd reviewer: [Signature]

METHOD: Inorganics, Method see cover

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- Y N N/A Have results been reported and calculated correctly?
- Y N N/A Are results within the calibrated range of the instruments?
- Y N N/A Are all detection limits below the CRQL?

Compound (analyte) results for F reported with a positive detect were recalculated and verified using the following equation:

Concentration =

Recalculation:

$$y = 0.195x - 0.0062$$

$$\frac{0.097 + 0.0062}{0.195} \times \frac{50 \text{ mL}}{0.89 (10.04 \text{ g})} = 2.96 \text{ mg/L}$$

#	Sample ID	Analyte	Reported Concentration (mg/L)	Calculated Concentration (mg/L)	Acceptable (Y/N)
0.934	1	F	2.2	2.2	Y
0.89	2		2.9	3.0	↓
0.872	3		2.5	2.5	
0.845	4		3.1	3.1	
0.852	5		3.8	3.9	

Note: _____

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: October 1, 2013
LDC Report Date: December 18, 2013
Matrix: Soil/Water
Parameters: Total Petroleum Hydrocarbons as Gasoline
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH119

Sample Identification

TB-100113
SL-570-SA5D-SB-4.0-5.0
SL-564-SA5D-SB-4.0-5.0
SL-607-SA5D-SB-4.0-5.0

Introduction

This data review covers 3 soil samples and one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8015B for Total Petroleum Hydrocarbons (TPH) as Gasoline.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Initial Calibration

Initial calibration of compounds was performed as required by the method.

The percent relative standard deviations (%RSD) of calibration factors for compounds were less than or equal to 20.0%.

III. Calibration Verification

Calibration verification was performed at required frequencies. The percent differences (%D) of amounts in continuing standard mixtures were within the 20.0% QC limits.

The percent differences (%D) of the second source calibration standard were less than or equal to 20.0% for all compounds.

IV. Blanks

Method blanks were reviewed for each matrix as applicable. No total petroleum hydrocarbons as gasoline contaminants were found in the method blanks.

Sample TB-100113 was identified as a trip blank. No total petroleum hydrocarbons as gasoline contaminants were found.

Sample EB1-100213 (from SDG PH120) was identified as an equipment blank. No total petroleum hydrocarbons as gasoline contaminants were found.

Sample FB-041613 (from SDG PH032) was identified as a field blank. No total petroleum hydrocarbons as gasoline contaminants were found.

V. Surrogate Recovery

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VII. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VIII. Target Compound Identification

All target compound identifications were within validation criteria.

IX. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH119	All compounds reported below the RL.	J (all detects)	A

X. System Performance

The system performance was acceptable.

XI. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XII. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Gasoline - Data Qualification Summary - SDG
 PH119**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH119	TB-100113 SL-570-SA5D-SB-4.0-5.0 SL-564-SA5D-SB-4.0-5.0 SL-607-SA5D-SB-4.0-5.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Gasoline - Laboratory Blank Data Qualification
 Summary - SDG PH119**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Gasoline - Field Blank Data Qualification
 Summary - SDG PH119**

No Sample Data Qualified in this SDG

LDC #: 30842C7

VALIDATION COMPLETENESS WORKSHEET

Date: 12/3/13

SDG #: PH119

Level IV

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: *SN*2nd Reviewer: *SN*

METHOD: GC TPH as Gasoline (EPA SW 846 Method 8015B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 10/1/13
II	Initial calibration	A	RSD $\leq 20\%$
III.	Calibration verification/ICV	A	ICV/CCV $\leq 20\%$
IV.	Blanks	A	
V	Surrogate recovery	A	
VI.	Matrix spike/Matrix spike duplicates	N	Client spec.
VII.	Laboratory control samples	A	LCS(7)
VIII.	Target compound identification	A	
IX.	Compound quantitation/RL/LOQ/LODs	A	
X.	System Performance	A	
XI.	Overall assessment of data	A	
XII.	Field duplicates	N	
XIII.	Field blanks	ND	TB=1 FB = FB-041613 (SDG # PH032) EB = EB1-100213 (SDG # PH120)

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

D = Duplicate
TB = Trip blank
EB = Equipment blank

Validated Samples: *Water + Soil*

1	TB-100113	<i>W</i>	11		21		31	<i>BLKYE</i>
2	SL-570-SA5D-SB-4.0-5.0		12		22		32	<i>BLKYU</i>
3	SL-564-SA5D-SB-4.0-5.0		13		23		33	
4	SL-607-SA5D-SB-4.0-5.0		14		24		34	
5			15		25		35	
6			16		26		36	
7			17		27		37	
8			18		28		38	
9			19		29		39	
10			20		30		40	

Notes: _____

Method: GC HPLC

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
II. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a linear fit used for evaluation? If yes, were all percent relative standard deviations (%RSD) \leq 20%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a curve fit used for evaluation? If Yes, what was the acceptance criteria used?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Did the initial calibration meet the curve fit acceptance criteria?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the RT windows properly established?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IV. Continuing calibration				
What type of continuing calibration calculation was performed? <input checked="" type="checkbox"/> %D or <input type="checkbox"/> %R	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a continuing calibration analyzed daily?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) \leq 20% or percent recoveries 80-120%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all the retention times within the acceptance windows?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
V. Blanks				
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank analyzed for each matrix and concentration?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
If the percent recovery (%R) of one or more surrogates was outside QC limits, was a reanalysis performed to confirm %R?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	

Validation Area	Yes	No	NA	Findings/Comments
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?		/		
Were the performance evaluation (PE) samples within the acceptance limits?			/	
X. Target compound identification				
Were the retention times of reported detects within the RT windows?	/			
XI. Compound quantitation/CRQLs				
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XII. System performance				
System performance was found to be acceptable.	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.		/		

LDC #: 30842C7

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page 1 of 1

Reviewer: BR

2nd Reviewer: Py

METHOD: GC X HPLC _____

The calibration factors (CF), average CF, and relative standard deviation (%RSD) were recalculated for compounds identified below using the following calculations:

$$CF = A/C$$

average CF = sum of the CF/number of standards

$$\%RSD = 100 * (S/X)$$

Where:

A = Area of compound

C = Concentration of compound

S = Standard deviation of calibration factors

X = Mean of calibration factors

#	Standard ID	Calibration Date	Compound	Reported CF (550 std)	Recalculated CF (550 std)	Reported Average CF (Initial)	Recalculated Average CF (Initial)	Reported %RSD	Recalculated %RSD
1	ICAL 11379F J&W DB-MTB	9/27/2013	GRO	53902	53902	55635	55635	4	4

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30842C7

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page 1 of 1

Reviewer: BR

2nd Reviewer: 82

METHOD: GC X HPLC _____

The calibration factors (CF), average CF, and relative standard deviation (%RSD) were recalculated for compounds identified below using the following calculations:

$$CF = A/C$$

$$\text{average CF} = \text{sum of the CF}/\text{number of standards}$$

$$\%RSD = 100 * (S/X)$$

Where:

A = Area of compound

C = Concentration of compound

S = Standard deviation of calibration factors

X = Mean of calibration factors

#	Standard ID	Calibration Date	Compound	Reported CF (550 std)	Recalculated CF (550 std)	Reported Average CF (Initial)	Recalculated Average CF (Initial)	Reported %RSD	Recalculated %RSD
1	ICAL 11002F J&W DB-VRX	9/16/2013	GRO	4534	4534	4650	4650	3	3

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30842C7

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Calculation Verification

Page: (of 1
Reviewer: BR
2nd Reviewer: 84

METHOD: GC _____ HPLC _____

The percent difference (%D) of the initial calibration average Calibration Factors (CF) and the continuing calibration percent difference (%D) values were recalculated for the compounds identified below using the following calculation:

$$\text{Percent difference (\%D)} = 100 * (N - C)/N$$

Where:

N = Initial Calibration Factor or Nominal Amount

C = Calibration Factor from Continuing Calibration Standard or Calculated Amount

#	Standard ID	Calibration Date	Compound	CCV Conc/CF	Reported Conc/CF	Recalculated Conc/CF	Reported % D	Recalculated %D
1	16281B.0002	10/8/2013	GRO	220.00	216.88	216.88	1	1
	11379F	16:16						
2	20277.0002	10/7/2013	GRO	550.00	600.40	600.40	9	9
	11002F	10:13						
3								

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3084207

VALIDATION FINDINGS WORKSHEET Surrogate Results Verification

Page: 1 of 1

Reviewer: BR

2nd reviewer: JK

METHOD: GC HPLC

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: $SF/SS * 100$

Where: SF = Surrogate Found
SS = Surrogate Spiked

Sample ID: 2

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
Trifluoromethane - F	J & W DB MFB	0.7455	0.691347	93	93	0

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

LDC #: 3084207

VALIDATION FINDINGS WORKSHEET

Page: 1 of 1

Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Reviewer: BR

2nd Reviewer: SC

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

%Recovery = 100 * (SSC - SC)/SA

Where SSC = Spiked sample concentration
SA = Spike added
LCS = Laboratory Control Sample

SC = Sample concentration

LCSD = Laboratory Control Sample duplicate

RPD = (((SSCLCS - SSCLCSD) * 2) / (SSCLCS + SSCLCSD)) * 100

LCS/LCSD samples: LCS FT / LCSD CK

Compound	Spike Added (<u>ug/L</u>)		Spike Sample Concentration (<u>ug/L</u>)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD	LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
					Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)	<u>1180</u>	<u>1100</u>	<u>1184.36</u>	<u>1208.3</u>	<u>108</u>	<u>108</u>	<u>110</u>	<u>110</u>	<u>2</u>	<u>1.5</u>
Diesel (8015)										
Benzene (8021B)										
Methane (RSK-175)										
2,4-D (8151)										
Dinoseb (8151)										
Naphthalene (8310)										
Anthracene (8310)										
HMX (8330)										
2,4,6-Trinitrotoluene (8330)										

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30842C7

VALIDATION FINDINGS WORKSHEET Sample Calculation Verification

Page: 1 of 1
Reviewer: BR
2nd Reviewer: SC

METHOD: GC HPLC

N N/A Were all reported results recalculated and verified for all level IV samples?
 N N/A Were all recalculated results for detected target compounds within 10% of the reported results?

Concentration = $\frac{(A)(Fv)(Df)}{(RF)(Vs \text{ or } Ws)(\%S/100)}$

Example:

Sample ID: A11 Compound Name: ND

LCSFT

GRD = 1189.36 ug/L
Concentration = _____

$$= \frac{(5530523)(1000)}{(4650)(1)(60)} = 1,189.359785 \text{ ug/L}$$

- A= Area or height of the compound to be measured
- Fv= Final Volume of extract
- Df= Dilution Factor
- RF= Average response factor of the compound
In the initial calibration
- Vs= Initial volume of the sample
- Ws= Initial weight of the sample
- %S= Percent Solid

#	Sample ID	Compound	Reported Concentrations ()	Recalculated Results Concentrations ()	Qualifications

Comments: _____

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: October 1, 2013
LDC Report Date: December 18, 2013
Matrix: Soil
Parameters: Total Petroleum Hydrocarbons as Extractables
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH119

Sample Identification

SL-570-SA5D-SB-0.0-0.5
SL-570-SA5D-SB-4.0-5.0
SL-564-SA5D-SB-0.0-0.5
SL-564-SA5D-SB-4.0-5.0
SL-607-SA5D-SB-0.0-0.5
SL-607-SA5D-SB-4.0-5.0
SL-570-SA5D-SB-0.0-0.5MS
SL-570-SA5D-SB-0.0-0.5MSD

Introduction

This data review covers 8 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8015B for Total Petroleum Hydrocarbons (TPH) as Extractables.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Initial Calibration

Initial calibration of compounds was performed as required by the method.

The percent relative standard deviations (%RSD) of calibration factors for compounds were less than or equal to 20.0%.

III. Calibration Verification

Calibration verification was performed at required frequencies. The percent differences (%D) of amounts in continuing standard mixtures were within the 20.0% QC limits.

The percent differences (%D) of the second source calibration standard were less than or equal to 20.0% for all compounds.

IV. Blanks

Method blanks were reviewed for each matrix as applicable. No total petroleum hydrocarbons as extractables contaminants were found in the method blanks.

Sample EB1-100213 (from SDG PH120) was identified as an equipment blank. No total petroleum hydrocarbons as extractable contaminants were found.

Sample FB-041613 (from SDG PH032) was identified as a field blank. No total petroleum hydrocarbons as extractable contaminants were found.

V. Surrogate Recovery

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VII. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Target Compound Identification

All target compound identifications were within validation criteria.

IX. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH119	All compounds reported below the RL.	J (all detects)	A

X. System Performance

The system performance was acceptable.

XI. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XII. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Extractables - Data Qualification Summary -
 SDG PH119**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-570-SA5D-SB-4.0-5.0 SL-564-SA5D-SB-0.0-0.5 SL-564-SA5D-SB-4.0-5.0 SL-607-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-4.0-5.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Extractables - Laboratory Blank Data
 Qualification Summary - SDG PH119**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Extractables - Field Blank Data Qualification
 Summary - SDG PH119**

No Sample Data Qualified in this SDG

LDC #: 30842C8

VALIDATION COMPLETENESS WORKSHEET

Date: 12/3/13

SDG #: PH119

Level IV

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: BR

2nd Reviewer: SM

METHOD: GC TPH as Extractables (EPA SW 846 Method 8015B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

Validation Area		Comments	
I.	Technical holding times	A	Sampling dates: 10/1/13
II.	Initial calibration	A	RSD ≤ 20%
III.	Calibration verification/ICV	D	1w/1cw ≤ 20%
IV.	Blanks	A	
V.	Surrogate recovery	A	
VI.	Matrix spike/Matrix spike duplicates	A	
VII.	Laboratory control samples	A	LCS
VIII.	Target compound identification	A	
IX.	Compound quantitation/RL/LOQ/LODs	A	
X.	System Performance	A	
XI.	Overall assessment of data	A	
XII.	Field duplicates	N	
XIII.	Field blanks	ND	FB = FB-041613 (SDG # PH032) EB = EB1-100213 (SDG # PH126)

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

D = Duplicate
TB = Trip blank
EB = Equipment blank

Validated Samples: Soil

1	SL-570-SA5D-SB-0.0-0.5	11	21	31	PBK38280
2	SL-570-SA5D-SB-4.0-5.0	12	22	32	
3	SL-564-SA5D-SB-0.0-0.5	13	23	33	
4	SL-564-SA5D-SB-4.0-5.0	14	24	34	
5	SL-607-SA5D-SB-0.0-0.5	15	25	35	
6	SL-607-SA5D-SB-4.0-5.0	16	26	36	
7	#1 MS	17	27	37	
8	#1 MSD	18	28	38	
9		19	29	39	
10		20	30	40	

Notes: _____

Method: GC HPLC

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
II. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a linear fit used for evaluation? If yes, were all percent relative standard deviations (%RSD) < 20%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a curve fit used for evaluation? If Yes, what was the acceptance criteria used?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Did the initial calibration meet the curve fit acceptance criteria?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the RT windows properly established?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IV. Continuing calibration				
What type of continuing calibration calculation was performed? <input checked="" type="checkbox"/> %D or <input type="checkbox"/> %R	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a continuing calibration analyzed daily?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) < 20% or percent recoveries 80-120%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all the retention times within the acceptance windows?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
V. Blanks				
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank analyzed for each matrix and concentration?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
If the percent recovery (%R) of one or more surrogates was outside QC limits, was a reanalysis performed to confirm %R?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	

LDC #: 30842c8

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: BR
 2nd Reviewer: SH

Validation Area	Yes	No	NA	Findings/Comments
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?		/		
Were the performance evaluation (PE) samples within the acceptance limits?			/	
X. Target compound identification				
Were the retention times of reported detects within the RT windows?	/			
XI. Compound quantitation/CRQLs				
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XII. System performance				
System performance was found to be acceptable.	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.		/		

LDC #: 30842C8

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page 1 of 1

Reviewer: BR

2nd Reviewer: 84

METHOD: GC X HPLC _____

The calibration factors (CF), average CF, and relative standard deviation (%RSD) were recalculated for compounds identified below using the following calculations:

$$CF = A/C$$

$$\text{average CF} = \text{sum of the CF}/\text{number of standards}$$

$$\%RSD = 100 * (S/X)$$

Where:

A = Area of compound

C = Concentration of compound

S = Standard deviation of calibration factors

X = Mean of calibration factors

#	Standard ID	Calibration Date	Compound	Reported CF (144 std)	Recalculated CF (144 std)	Reported Average CF (Initial)	Recalculated Average CF (Initial)	Reported %RSD	Recalculated %RSD
1	ICAL CP23-19879A ZB-5	8/19/2013	C8-C40	22576	22576	22983	22983	7	7

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30842C8

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Calculation Verification

Page: 1 of 1
Reviewer: BR
2nd Reviewer: 84

METHOD: GC HPLC

The percent difference (%D) of the initial calibration average Calibration Factors (CF) and the continuing calibration percent difference (%D) values were recalculated for the compounds identified below using the following calculation:

$$\text{Percent difference (\%D)} = 100 * (N - C)/N$$

Where:

N = Initial Calibration Factor or Nominal Amount

C = Calibration Factor from Continuing Calibration Standard or Calculated Amount

#	Standard ID	Calibration Date	Compound	CCV Conc/CF	Reported Conc/CF	Recalculated Conc/CF	Reported % D	Recalculated %D
1	J277.0093	10/8/2013	C8-C40	288.01	278.02	278.02	3	3
2								
3								

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3084208

VALIDATION FINDINGS WORKSHEET Surrogate Results Verification

Page: 1 of 1
Reviewer: BR
2nd reviewer: SM

METHOD: X GC HPLC

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
SS = Surrogate Spiked

Sample ID: 1

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
Chlorobenzene	ZB-5	2	1.680579	85	84	1
Orthotercphenyl	↓	2	1.476762	74	74	0

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

LDC #: 30842C8

VALIDATION FINDINGS WORKSHEET
Matrix Spike/Matrix Spike Duplicates Results Verification

METHOD: X GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the matrix spike and matrix spike duplicate were recalculated for the compounds identified below using the following calculation:

%Recovery = 100 * (SSC - SC)/SA

Where

SSC = Spiked sample concentration

MS = Matrix spike

SC = Sample concentration

MSD = Matrix spike duplicate

RPD = (((SSCMS - SSCMSD) * 2) / (SSCMS + SSCMSD)) * 100

SA = Spike added

MS/MSD samples: 7/8

Compound	Spike Added (mg/kg)		Sample Conc. (mg/kg)	Spike Sample Concentration (mg/kg)		Matrix spike		Matrix Spike Duplicate		MS/MSD	
	MS	MSD		MS	MSD	Percent Recovery		Percent Recovery		RPD	
						Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)											
Diesel (8015)											
Benzene (8021B)											
Methane (RSK-175)											
2,4-D (8151)											
Dinoseb (8151)											
Naphthalene (8310)											
Anthracene (8310)											
HMX (8330)											
2,4,6-Trinitrotoluene (8330)											
EFH (C30-C40)	15	15	0	8.8	8.33	59	59	54	54	5	55

Comments: Refer to Matrix Spike/Matrix Spike Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 38842C8

VALIDATION FINDINGS WORKSHEET

Page: 1 of 1

Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Reviewer: BR

2nd Reviewer: SM

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

%Recovery = 100 * (SSC - SC)/SA

Where SSC = Spiked sample concentration

SC = Sample concentration

RPD = (((SSCLCS - SSCLCSD) * 2) / (SSCLCS + SSCLCSD)) * 100

SA = Spike added

LCS = Laboratory Control Sample

LCSD = Laboratory Control Sample duplicate

LCS/LCSD samples: LCS38280

Compound	Spike Added (mg/kg)		Spike Sample Concentration (mg/kg)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD	LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
					Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)										
Diesel (8015)										
Benzene (8021B)										
Methane (RSK-175)										
2,4-D (8151)										
Dinoseb (8151)										
Naphthalene (8310)										
Anthracene (8310)										
HMX (8330)										
2,4,6-Trinitrotoluene (8330)										
ETD (C30-C40)	15	-	9-81	-	65	65	-	-	-	-

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3084208

VALIDATION FINDINGS WORKSHEET Sample Calculation Verification

Page: 1 of 1
Reviewer: BR
2nd Reviewer: SL

METHOD: GC HPLC

N N/A Were all reported results recalculated and verified for all level IV samples?
 N N/A Were all recalculated results for detected target compounds within 10% of the reported results?

Concentration = $\frac{(A)(Fv)(Df)}{(RF)(Vs \text{ or } Ws)(\%S/100)}$

Example:

Sample ID All Compound Name ND

$LC838280 = C30 - C40 = 9.8 \text{ mg/kg}$

Concentration = $\frac{(6762873)(1000)}{(22983)(30.1)(1000)}$
 $= 9.775928561 \text{ mg/kg}$

- A= Area or height of the compound to be measured
- Fv= Final Volume of extract
- Df= Dilution Factor
- RF= Average response factor of the compound
In the initial calibration
- Vs= Initial volume of the sample
- Ws= Initial weight of the sample
- %S= Percent Solid

#	Sample ID	Compound	Reported Concentrations ()	Recalculated Results Concentrations ()	Qualifications

Comments: _____

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: October 1, 2013
LDC Report Date: December 10, 2013
Matrix: Soil
Parameters: Dioxins/Dibenzofurans
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH119

Sample Identification

SL-570-SA5D-SB-0.0-0.5
SL-564-SA5D-SB-0.0-0.5
SL-607-SA5D-SB-0.0-0.5

Introduction

This data review covers 3 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA Method 1613B for Polychlorinated Dioxins/Dibenzofurans.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and the USEPA Contract Laboratory Program National Functional Guidelines for Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. HRGC/HRMS Instrument Performance Check

Instrument performance was checked at the required daily frequency.

The chromatographic resolution between 2,3,7,8-TCDD and the peaks representing any other unlabeled TCDD isomers was resolved with a valley of less than or equal to 25%.

PFK and static resolving power were within validation criteria.

III. Initial Calibration

A five point initial calibration was performed as required by the method.

Percent relative standard deviations (%RSD) were less than or equal to 20.0% for unlabeled compounds and less than or equal to 35.0% for labeled compounds.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

The minimum S/N ratio was greater than or equal to 10 for each unlabeled compound and labeled compound.

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

All of the continuing calibration percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within QC limits.

The percent differences (%D) of the second source calibration standard were within QC limits.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No polychlorinated dioxin/dibenzofuran contaminants were found in the method blanks with the following exceptions:

Method Blank ID	Extraction Date	Compound	Concentration	Associated Samples
BLK281002	10/8/13	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,6,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.0417 ng/Kg 0.0514 ng/Kg 0.0350 ng/Kg 0.0212 ng/Kg 0.0871 ng/Kg 0.306 ng/Kg 0.0743 ng/Kg 0.0383 ng/Kg 0.0397 ng/Kg 0.0378 ng/Kg 0.0495 ng/Kg 0.0565 ng/Kg 0.0654 ng/Kg 0.0445 ng/Kg 0.161 ng/Kg	All samples in SDG PH119

Sample concentrations were compared to concentrations detected in the method blanks. The sample concentrations were either not detected or were significantly greater (>5X blank contaminants) than the concentrations found in the associated method blanks with the following exceptions:

Sample	Compound	Reported Concentration	Modified Final Concentration
SL-570-SA5D-SB-0.0-0.5	1,2,3,7,8-PeCDD 1,2,3,6,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.0532 ng/Kg 0.0568 ng/Kg 0.0504 ng/Kg 0.235 ng/Kg 1.53 ng/Kg 0.0599 ng/Kg 0.0432 ng/Kg 0.0512 ng/Kg 0.0321 ng/Kg 0.0544 ng/Kg 0.0565 ng/Kg 0.0877 ng/Kg 0.0742 ng/Kg 0.167 ng/Kg	0.0532U ng/Kg 0.0568U ng/Kg 0.0504U ng/Kg 0.235U ng/Kg 1.53U ng/Kg 0.0599U ng/Kg 0.0432U ng/Kg 0.0512U ng/Kg 0.0321U ng/Kg 0.0544U ng/Kg 0.0565U ng/Kg 0.0877U ng/Kg 0.0742U ng/Kg 0.167U ng/Kg
SL-564-SA5D-SB-0.0-0.5	2,3,7,8-TCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.0495 ng/Kg 0.0569 ng/Kg 0.0954 ng/Kg 0.289 ng/Kg 0.0728 ng/Kg 0.0515 ng/Kg 0.0209 ng/Kg 0.0273 ng/Kg 0.0550 ng/Kg 0.0890 ng/Kg 0.0856 ng/Kg 0.0361 ng/Kg 0.188 ng/Kg	0.0495U ng/Kg 0.0569U ng/Kg 0.0954U ng/Kg 0.289U ng/Kg 0.0728U ng/Kg 0.0515U ng/Kg 0.0209U ng/Kg 0.0273U ng/Kg 0.0550U ng/Kg 0.0890U ng/Kg 0.0856U ng/Kg 0.0361U ng/Kg 0.188U ng/Kg

Sample	Compound	Reported Concentration	Modified Final Concentration
SL-607-SA5D-SB-0.0-0.5	1,2,3,7,8-PeCDD	0.0276 ng/Kg	0.0276U ng/Kg
	1,2,3,6,7,8-HxCDD	0.108 ng/Kg	0.108U ng/Kg
	1,2,3,4,6,7,8-HpCDD	0.239 ng/Kg	0.239U ng/Kg
	OCDD	0.701 ng/Kg	0.701U ng/Kg
	1,2,3,7,8-PeCDF	0.0885 ng/Kg	0.0885U ng/Kg
	2,3,4,7,8-PeCDF	0.0442 ng/Kg	0.0442U ng/Kg
	1,2,3,4,7,8-HxCDF	0.0311 ng/Kg	0.0311U ng/Kg
	1,2,3,6,7,8-HxCDF	0.0334 ng/Kg	0.0334U ng/Kg
	2,3,4,6,7,8-HxCDF	0.0348 ng/Kg	0.0348U ng/Kg
	1,2,3,4,6,7,8-HpCDF	0.0307 ng/Kg	0.0307U ng/Kg
	1,2,3,4,7,8,9-HpCDF	0.0229 ng/Kg	0.0229U ng/Kg
	OCDF	0.125 ng/Kg	0.125U ng/Kg

Sample EB1-100213 (from SDG PH120) was identified as an equipment blank. No polychlorinated dioxin/dibenzofuran contaminants were found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
EB1-100213	10/2/13	1,2,3,7,8-PeCDD	0.338 pg/L	All samples in SDG PH119
		1,2,3,6,7,8-HxCDD	0.227 pg/L	
		1,2,3,4,6,7,8-HpCDD	0.368 pg/L	
		OCDD	1.49 pg/L	
		2,3,7,8-TCDF	0.237 pg/L	
		1,2,3,7,8-PeCDF	0.348 pg/L	
		2,3,4,7,8-PeCDF	0.247 pg/L	
		1,2,3,4,7,8-HxCDF	0.201 pg/L	
		1,2,3,6,7,8-HxCDF	0.207 pg/L	
		1,2,3,7,8,9-HxCDF	0.334 pg/L	
		2,3,4,6,7,8-HxCDF	0.255 pg/L	
		1,2,3,4,6,7,8-HpCDF	0.288 pg/L	
		1,2,3,4,7,8,9-HpCDF	0.363 pg/L	
		OCDF	1.43 pg/L	

Sample FB-041613 (from SDG PH032) was identified as a field blank. No perchlorate was found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB-041613	4/16/13	1,2,3,7,8-PeCDD	0.434 pg/L	All samples in SDG PH119
		1,2,3,4,6,7,8-HpCDD	0.309 pg/L	
		OCDD	0.838 pg/L	
		1,2,3,7,8-PeCDF	0.324 pg/L	
		2,3,4,7,8-PeCDF	0.429 pg/L	
		1,2,3,4,7,8-HxCDF	0.257 pg/L	
		1,2,3,7,8,9-HxCDF	0.241 pg/L	
		2,3,4,6,7,8-HxCDF	0.284 pg/L	
		1,2,3,4,6,7,8-HpCDF	0.314 pg/L	
		OCDF	0.357 pg/L	

Sample concentrations were compared to concentrations detected in the field blanks. The sample concentrations were either not detected or were significantly greater (>5X for other contaminants) than the concentrations found in the associated field blanks.

VI. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VII. Ongoing Precision Recovery (OPR)

Ongoing precision recovery samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Regional Quality Assurance and Quality Control

Not applicable.

IX. Internal Standards

All internal standard recoveries were within QC limits.

X. Target Compound Identifications

All target compound identifications were within validation criteria.

XI. Compound Quantitation

All compound quantitations were within validation criteria with the following exceptions:

Sample	Compound	Finding	Criteria	Flag	A or P
SL-607-SA5D-SB-0.0-0.5	2,3,7,8-TCDF	2nd column confirmation was not performed for this compound.	This compound must be confirmed on the 2nd column per the method.	None	P

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH119	All compounds reported below the RL.	J (all detects)	A

XII. System Performance

The system performance was acceptable.

XIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Dioxins/Dibenzofurans - Data Qualification Summary - SDG PH119**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH119	SL-607-SA5D-SB-0.0-0.5	2,3,7,8-TCDF	None	P	Compound quantitation (column confirmation) (*XI)
PH119	SL-570-SA5D-SB-0.0-0.5 SL-564-SA5D-SB-0.0-0.5 SL-607-SA5D-SB-0.0-0.5	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Dioxins/Dibenzofurans - Laboratory Blank Data Qualification Summary - SDG PH119**

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
PH119	SL-570-SA5D-SB-0.0-0.5	1,2,3,7,8-PeCDD 1,2,3,6,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.0532U ng/Kg 0.0568U ng/Kg 0.0504U ng/Kg 0.235U ng/Kg 1.53U ng/Kg 0.0599U ng/Kg 0.0432U ng/Kg 0.0512U ng/Kg 0.0321U ng/Kg 0.0544U ng/Kg 0.0565U ng/Kg 0.0877U ng/Kg 0.0742U ng/Kg 0.167U ng/Kg	A	B
PH119	SL-564-SA5D-SB-0.0-0.5	2,3,7,8-TCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.0495U ng/Kg 0.0569U ng/Kg 0.0954U ng/Kg 0.289U ng/Kg 0.0728U ng/Kg 0.0515U ng/Kg 0.0209U ng/Kg 0.0273U ng/Kg 0.0550U ng/Kg 0.0890U ng/Kg 0.0856U ng/Kg 0.0361U ng/Kg 0.188U ng/Kg	A	B
PH119	SL-607-SA5D-SB-0.0-0.5	1,2,3,7,8-PeCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.0276U ng/Kg 0.108U ng/Kg 0.239U ng/Kg 0.701U ng/Kg 0.0885U ng/Kg 0.0442U ng/Kg 0.0311U ng/Kg 0.0334U ng/Kg 0.0348U ng/Kg 0.0307U ng/Kg 0.0229U ng/Kg 0.125U ng/Kg	A	B

Santa Susana Field Laboratory
Dioxins/Dibenzofurans - Field Blank Data Qualification Summary - SDG PH119

No Sample Data Qualified in this SDG

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 10/1/13
II.	HRGC/HRMS Instrument performance check	A	
III.	Initial calibration	A	≈ 20/35
IV.	Continuing Calibration	A	QC limits
V.	Blanks	SW	
VI.	Matrix spike/Matrix spike duplicates	N	client
VII.	Laboratory control samples	A	OPR
VIII.	Regional quality assurance and quality control	N	
IX.	Internal standards	A	
X.	Target compound identifications	A	
XI.	Compound quantitation/RL/LOQ/LODs-	SW	
XII.	System performance	A	
XIII.	Overall assessment of data	A	
XIV.	Field duplicates	N	
XV.	Field blanks	SW	EB=EB1-100213(PH120); FB=FB-041613(PH032)

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

D = Duplicate
TB = Trip blank
EB = Equipment blank

Validated Samples: *Soils*

1	SL-570-SA5D-SB-0.0-0.5	11		21		31	
2	SL-564-SA5D-SB-0.0-0.5	12		22		32	
3	SL-607-SA5D-SB-0.0-0.5	13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20	BLK 281002	30		40	

Notes: _____

Method: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. GC/MS Instrument performance check				
Was PFK exact mass 380.9760 verified?	/			
Were the retention time windows established for all homologues?	/			
Was the chromatographic resolution between 2,3,7,8-TCDD and peaks representing any other unlabeled TCDD isomers < 25% ?	/			
Is the static resolving power at least 10,000 (10% valley definition)?	/			
Was the mass resolution adequately check with PFK?	/			
Was the presence of 1,2,8,9-TCDD and 1,3,4,6,8-PeCDF verified?	/			
III. Initial calibration				
Was the initial calibration performed at 5 concentration levels?	/			
Were all percent relative standard deviations (%RSD) ≤ 20% for unlabeled compounds and < 35% for labeled compounds ?	/			
Did all calibration standards meet the Ion Abundance Ratio criteria?	/			
Was the signal to noise ratio for each target compound ≥ 2.5 and for each recovery and internal standard > 10?	/			
IV. Continuing calibration				
Was a routine calibration performed at the beginning and end of each 12 hour period?	/			
Were all the concentrations for the unlabeled compounds and labeled compounds within the QC limits (Method 1613B, Table 6)?	/			
Did all routine calibration standards meet the Ion Abundance Ratio criteria?	/			
V. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was a method blank performed for each matrix and concentration?			/	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet?	/			
VI. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.			/	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?			/	
VII. Laboratory control samples				
Was an LCS analyzed for this SDG?	/		/	
Was an LCS analyzed per extraction batch?			/	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	/			

VALIDATION FINDINGS CHECKLIST

Validation Area	Yes	No	NA	Findings/Comments
VIII. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?			/	
Were the performance evaluation (PE) samples within the acceptance limits?			/	
IX. Internal standards				
Were internal standard recoveries within the 25-150% criteria?	/			
Was the minimum S/N ratio of all internal standard peaks > 10?	/			
X. Target compound identification				
For 2,3,7,8 substituted congeners with associated labeled standards, were the retention times of the two quantitation peaks within -1 to 3 sec. of the RT of the labeled standard?	/			
For 2,3,7,8 substituted congeners without associated labeled standards, were the relative retention times of the two quantitation peaks within 0.005 time units of the RRT measured in the routine calibration?	/			
For non-2,3,7,8 substituted congeners, were the retention times of the two quantitation peaks within RT established in the performance check solution?			/	
Did compound spectra contain all characteristic ions listed in the table attached?	/			
Was the Ion Abundance Ratio for the two quantitation ions within criteria?		/		
Was the signal to noise ratio for each target compound and labeled standard > 2.5?	/			
Does the maximum intensity of each specified characteristic ion coincide within ± 2 seconds (includes labeled standards)?	/			
For PCDF identification, was any signal (S/N ≥ 2.5, at ± seconds RT) detected in the corresponding PCDF channel?		/		
Was an acceptable lock mass recorded and monitored?	/			
XI. Compound quantitation/CRQLs				
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	/			
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XII. System performance				
System performance was found to be acceptable.	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.			/	
Target compounds were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.	/			

VALIDATION FINDINGS WORKSHEET

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

A. 2,3,7,8-TCDD	F. 1,2,3,4,6,7,8-HpCDD	K. 1,2,3,4,7,8-HxCDF	P. 1,2,3,4,7,8,9-HpCDF	U. Total HpCDD
B. 1,2,3,7,8-PeCDD	G. OCDD	L. 1,2,3,6,7,8-HxCDF	Q. OCDF	V. Total TCDF
C. 1,2,3,4,7,8-HxCDD	H. 2,3,7,8-TCDF	M. 2,3,4,6,7,8-HxCDF	R. Total TCDD	W. Total PeCDF
D. 1,2,3,6,7,8-HxCDD	I. 1,2,3,7,8-PeCDF	N. 1,2,3,7,8,9-HxCDF	S. Total PeCDD	X. Total HxCDF
E. 1,2,3,7,8,9-HxCDD	J. 2,3,4,7,8-PeCDF	O. 1,2,3,4,6,7,8-HpCDF	T. Total HxCDD	Y. Total HpCDF

Notes: _____

VALIDATION FINDINGS WORKSHEET

Blanks

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

N N/A Were all samples associated with a method blank?

N N/A Was a method blank performed for each matrix and whenever a sample extraction was performed?

N N/A Was the method blank contaminated?

Blank extraction date: 10/08/13 Blank analysis date: 10/10/13 Associated samples: All *≥5x Qual U (B)*

Conc. units: ng/kg

Compound	Blank ID	Sample Identification							
		5x	1	2	3				
	BLK281002								
A	0.0417	0.209		0.0495*					
B	0.0514*	0.257	0.0532		0.0276*				
D	0.0350*	0.175	0.0568*		0.108*				
E	0.0212*	0.106	0.0504	0.0569*					
F	0.0871*	0.436	0.235*	0.0954*	0.239				
G	0.306	1.53	1.53	0.289*	0.701*				
I	0.0743*	0.372	0.0599*	0.0728	0.0885*				
J	0.0383*	0.192	0.0432*	0.0515*	0.0442				
K	0.0397*	0.199	0.0512*	0.0209*	0.0311*				
L	0.0378	0.189	0.0321*	0.0273*	0.0334*				
M	0.0495*	0.248	0.0544	0.0550*	0.0348*				
N	0.0565*	0.283	0.0565*	0.0890*					
O	0.0654*	0.327	0.0877*	0.0856*	0.0307				
P	0.0445*	0.223	0.0742*	0.0361*	0.0229*				
Q	0.161*	0.805	0.167*	0.188	0.125*				

*EMPC

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
All contaminants within five times the method blank concentration were qualified as not detected, "U".

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: HRGC/HRMS Dioxins (EPA Method 1613B)

Blank units: pg/L **Associated sample units:** ng/kg

Sampling date: 10/02/13

Field blank type: (circle one) Field Blank / Rinsate / Other: EB **Associated Samples:** All >5x

Compound	Blank ID	Sample Identification							
		5X							
	EB1-100213	5X							
B	0.338*	0.00169							
D	0.227*	0.00114							
F	0.368*	0.00184							
G	1.49	0.00745							
H	0.237*	0.00119							
I	0.348*	0.00174							
J	0.247*	0.00124							
K	0.201*	0.00101							
L	0.207*	0.00104							
N	0.334*	0.00167							
M	0.255*	0.00128							
O	0.288*	0.00144							
P	0.363*	0.001815							
Q	1.43*	0.00715							

* EMPC

EB1-100213 (PH120)

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 All contaminants within five times the blank concentration were qualified as not detected, "U".

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: HRGC/HRMS Dioxins (EPA Method 1613B)

Blank units: pg/L **Associated sample units:** ng/kg

Sampling date: 04/16/13

Field blank type: (circle one) Field Blank / Rinsate / Other: FB **Associated Samples:** All >5x

Compound	Blank ID	Sample Identification							
		5X							
	FB-041613	5X							
B	0.434*	0.00217							
F	0.309*	0.00155							
G	0.838*	0.00419							
I	0.324	0.00162							
J	0.429*	0.00215							
K	0.257	0.00129							
N	0.241*	0.00121							
M	0.284*	0.00142							
O	0.314*	0.00157							
Q	0.357*	0.00179							

* EMPC

FB-041613 (PH032)

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 All contaminants within five times the blank concentration were qualified as not detected, "U".

VALIDATION FINDINGS WORKSHEET
Compound Quantitation and Reported CRQLs

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1631B)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- N N/A Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound?
- N N/A Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary).

#	Date	Compound	Finding	Associated Samples	Qualifications
			no 2,3,7,8-TCDF confirmation	3	None / P

Comments: See sample calculation verification worksheet for recalculations

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$$RRF = (A_x)(C_{is}) / (A_{is})(C_x)$$

average RRF = sum of the RRFs/number of standards

$$\%RSD = 100 * (S/X)$$

A_x = Area of compound,

C_x = Concentration of compound,

S = Standard deviation of the RRFs,

A_{is} = Area of associated internal standard

C_{is} = Concentration of internal standard

X = Mean of the RRFs

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				Average RRF (initial)	Average RRF (initial)	RRF (CS3 std)	RRF (CS3 std)	%RSD	%RSD
1	ICAL	09/23/2013	2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	1.008	1.008	1.043	1.043	4.15	4.14
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	1.109	1.109	1.125	1.125	1.62	1.60
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	0.949	0.949	0.995	0.995	4.48	4.48
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	1.056	1.056	1.079	1.079	1.66	1.66
			OCDF (¹³ C-OCDF)	0.963	0.963	0.998	0.998	2.75	2.75
2			2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)						
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)						
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)						
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)						
			OCDF (¹³ C-OCDF)						
3			2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)						
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)						
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)						
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)						
			OCDF (¹³ C-OCDF)						

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET

Routine Calibration Results Verification

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

$$\% \text{ Difference} = 100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$$

$$\text{RRF} = (A_x)(C_{is}) / (A_{is})(C_x)$$

Where: ave. RRF = initial calibration average RRF

RRF = continuing calibration RRF

 A_x = Area of compound,

 A_{is} = Area of associated internal standard

 C_x = Concentration of compound,

 C_{is} = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Spiked Conc (ng/mL)	Reported	Recalculated	Reported	Recalculated
					Conc (ng/mL)	Conc (ng/mL)	%R	%R
1	CS3002	10-10-13	2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	10	9.82	9.82	98	98
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	10	10.27	10.27	103	103
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	50	52.49	52.49	105	105
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	50	51.80	51.78	104	104
			OCDF (¹³ C-OCDF)	100	103.35	103.38	103	103
2			2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	10				
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	10				
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	50				
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	50				
			OCDF (¹³ C-OCDF)	100				
3			2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	10				
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	10				
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	50				
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	50				
			OCDF (¹³ C-OCDF)	100				

Comments: Refer to Routine Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Laboratory Control Sample Results Verification

METHOD: GC/MS Dioxins/Dibenzofurans (EPA Method 1613B)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the laboratory control sample and laboratory control sample duplicate (if applicable) were recalculated for the compounds identified below using the following calculation:

% Recovery = 100 * SSC/SA

Where: SSC = Spiked sample concentration
 SA = Spike added

RPD = |LCS - LCSD| * 2 / (LCS + LCSD)

LCS = Laboratory control sample percent recovery

LCSD = Laboratory control sample duplicate percent recovery

LCS ID: OPR281002

Compound	Spike Added (ng/Kg)		Spiked Sample Concentration (ng/Kg)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD	LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
					Reported	Recalc	Reported	Recalc	Reported	Recalculated
2,3,7,8-TCDD	20	NA	19.1	NA	95	95				
1,2,3,7,8-PeCDD	100		97.9		98	98				
1,2,3,4,7,8-HxCDD	100		97.2		97	97				
1,2,3,4,7,8,9-HpCDF	100		95.8		94	96				
OCDF	200		192		96	96				

Comments: Refer to Laboratory Control Sample findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30842021

VALIDATION FINDINGS WORKSHEET

Sample Calculation Verification

Page 1 of 1
Reviewer: [Signature]
2nd reviewer: SW

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

- N N/A Were all reported results recalculated and verified for all level IV samples?
- N N/A Were all recalculated results for detected target compounds agree within 10.0% of the reported results?

$$\text{Concentration} = \frac{A_x(I_s)(DF)}{(A_{is})(RRF)(V_o)(\%S)}$$

- A_x = Area of the characteristic ion (EICP) for the compound to be measured
- A_{is} = Area of the characteristic ion (EICP) for the specific internal standard
- I_s = Amount of internal standard added in nanograms (ng)
- V_o = Volume or weight of sample extract in milliliters (ml) or grams (g).
- RRF = Relative Response Factor (average) from the initial calibration
- Df = Dilution Factor.
- %S = Percent solids, applicable to soil and solid matrices only.

Example:

Sample I.D. 1, Q:

$$\begin{aligned} \text{Conc.} &= \frac{(302+227) \times (4000) \times (1)}{(739798 + 662127) \times (0.963) \times (0.25) \times (0.917)} \\ &= 0.167 \text{ ng/Kg} \end{aligned}$$

#	Sample ID	Compound	Reported Concentration ()	Calculated Concentration ()	Qualification

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH120

Prepared for

CDM Smith
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
2701 Loker Ave West, Suite 220
Carlsbad, California 92010

December 23, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on October 2, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by Environmental Protection Agency (EPA) SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)

Pesticides by EPA SW 846 8081B

Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A

Metals by EPA SW 846 Method 6010C, 6020A, 7470A and 7471B

Herbicides by EPA SW 846 Method 8151A

Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M

TPH as Extractables by EPA SW 846 Method 8015M

Dioxins and Dibenzofurans by EPA Method 1613B

Wet Chemistry:

Hexavalent Chromium by EPA Method 7199

Fluoride by EPA Method 300.0

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), method blanks, trip blanks, equipment blanks, field blanks and field duplicate samples. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met with the exception of one sample for hexavalent chromium. The associated sample results were qualified as non-detected estimated (UJ). The details regarding the qualification of data are provided in Enclosure I.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of several blanks for pesticides, metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks with the following exceptions:

SDG/Method	Method Blank ID	Analyte	Maximum Concentration	Associated Samples
PH120/6010C	ICB/CCB	Molybdenum	3.3 ug/L	EB1-100213

Sample concentrations were compared to concentrations detected in the initial and continuing blanks. The sample concentrations were not detected or were significantly greater than the concentrations found in the associated blanks with the following exceptions:

SDG/Method	Sample	Compound	Reported Concentration	Modified Final Concentration
PH120/6010C	EB1-100213	Molybdenum	8.6 ug/L	8.6U ug/L

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pairs for fluoride. The associated sample results were qualified as detected estimated (J). The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of several LCS/LCSD pairs for SVOCs, herbicides and TPH as extractables. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution was not performed for this SDG.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH120	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pair was collected and analyzed for fluoride. RPDs were not within QC limits. The associated sample results were qualified as detected estimated (J). The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No volatile contaminants were found in the trip blank.

One equipment blank was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins, hexavalent chromium and fluoride. The equipment blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to equipment blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the equipment blanks were not qualified. The equipment blank outlier reports are presented in Enclosure I.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins, hexavalent chromium and fluoride. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1

Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
02-Oct-2013	TB1-100213	7222768	TB	5030B	8015M	III
02-Oct-2013	SL-594-SA5D-SB-0.0-0.5	7222765	N	METHOD	300.0	III
02-Oct-2013	SL-594-SA5D-SB-4.0-5.0	7222766	N	METHOD	300.0	III
02-Oct-2013	SL-594-SA5D-SB-7.0-8.0	7222767	N	METHOD	300.0	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5	7222763	N	3050B	6010C	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5	7222763	N	3050B	6020A	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5	7222763	N	3546	8015M	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5	7222763	N	3546	8081B	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5	7222763	N	3546	8082A	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5	7222763	N	3546	8270D SIM	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5	7222763	N	METHOD	1613B	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5	7222763	N	METHOD	7471B	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5MSD	P222763M242034A	MSD	3546	8081B	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5MSD	P222763M260436	MSD	3546	8270D SIM	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5MS	P222763R242019A	MS	3546	8081B	III
02-Oct-2013	SL-589-SA5D-SB-0.0-0.5MS	P222763R260403	MS	3546	8270D SIM	III
02-Oct-2013	SL-589-SA5D-SB-4.0-5.0	7222764	N	3050B	6010C	III
02-Oct-2013	SL-589-SA5D-SB-4.0-5.0	7222764	N	3050B	6020A	III
02-Oct-2013	SL-589-SA5D-SB-4.0-5.0	7222764	N	3546	8015M	III
02-Oct-2013	SL-589-SA5D-SB-4.0-5.0	7222764	N	3546	8081B	III
02-Oct-2013	SL-589-SA5D-SB-4.0-5.0	7222764	N	3546	8082A	III
02-Oct-2013	SL-589-SA5D-SB-4.0-5.0	7222764	N	3546	8270D SIM	III
02-Oct-2013	SL-589-SA5D-SB-4.0-5.0	7222764	N	5035A	8015M	III
02-Oct-2013	SL-589-SA5D-SB-4.0-5.0	7222764	N	METHOD	7471B	III
02-Oct-2013	SL-600-SA5D-SB-0.0-0.5	7222769	N	METHOD	300.0	III
02-Oct-2013	SL-600-SA5D-SB-0.0-0.5MS	7222770	MS	METHOD	300.0	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
02-Oct-2013	SL-600-SA5D-SB-0.0-0.5DUP	7222771	DUP	METHOD	300.0	III
02-Oct-2013	SL-900-SA5D-SB-0.0-0.5	7222772	FD	METHOD	300.0	III
02-Oct-2013	SL-600-SA5D-SB-4.0-5.0	7222773	N	METHOD	300.0	III
02-Oct-2013	SL-600-SA5D-SB-9.0-10.0	7222774	N	METHOD	300.0	III
02-Oct-2013	SL-600-SA5D-SB-14.0-15.0	7222775	N	METHOD	300.0	III
02-Oct-2013	EB1-100213	7222762	EB	3005A	6010C	III
02-Oct-2013	EB1-100213	7222762	EB	3510C	8015M	III
02-Oct-2013	EB1-100213	7222762	EB	3510C	8081B	III
02-Oct-2013	EB1-100213	7222762	EB	3510C	8082A	III
02-Oct-2013	EB1-100213	7222762	EB	3510C	8270D SIM	III
02-Oct-2013	EB1-100213	7222762	EB	5030B	8015M	III
02-Oct-2013	EB1-100213	7222762	EB	Gen Prep	300.0	III
02-Oct-2013	EB1-100213	7222762	EB	Gen Prep	7199	III
02-Oct-2013	EB1-100213	7222762	EB	M3010A	6020A	III
02-Oct-2013	EB1-100213	7222762	EB	METHOD	1613B	III
02-Oct-2013	EB1-100213	7222762	EB	METHOD	7470A	III
02-Oct-2013	EB1-100213	7222762	EB	METHOD	8151A	III
02-Oct-2013	SL-600-SA5D-SB-18.5-19.5	7222776	N	METHOD	300.0	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: GENCHEM
Method: 300.0 **Matrix:** SO

Sample ID: SL-594-SA5D-SB-0.0-0.5 Collected: 10/2/2013 9:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	0.76	J	0.41	MDL	1.0	PQL	mg/Kg	J	Z

Sample ID: SL-600-SA5D-SB-0.0-0.5 Collected: 10/2/2013 1:25:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	0.93	J	0.42	MDL	1.0	PQL	mg/Kg	J	Z, Q, FD

Sample ID: SL-600-SA5D-SB-14.0-15.0 Collected: 10/2/2013 1:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	7.1		0.46	MDL	1.1	PQL	mg/Kg	J	Q

Sample ID: SL-600-SA5D-SB-18.5-19.5 Collected: 10/2/2013 2:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	4.1		0.45	MDL	1.1	PQL	mg/Kg	J	Q

Sample ID: SL-600-SA5D-SB-9.0-10.0 Collected: 10/2/2013 1:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	9.3		0.46	MDL	1.2	PQL	mg/Kg	J	Q

Sample ID: SL-900-SA5D-SB-0.0-0.5 Collected: 10/2/2013 1:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	2.5		0.44	MDL	1.1	PQL	mg/Kg	J	FD

Method Category: METALS
Method: 6010C **Matrix:** AQ

Sample ID: EB1-100213 Collected: 10/2/2013 2:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BARIUM	0.00058	J	0.00033	MDL	0.0100	PQL	mg/L	J	Z
BORON	0.0098	J	0.0084	MDL	0.100	PQL	mg/L	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6010C	Matrix: AQ

Sample ID: EB1-100213 Collected: 10/2/2013 2:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	0.0427	J	0.0334	MDL	0.400	PQL	mg/L	J	Z
COPPER	0.0028	J	0.0027	MDL	0.0200	PQL	mg/L	J	Z
MOLYBDENUM	0.0086	J	0.0017	MDL	0.0200	PQL	mg/L	U	B

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-589-SA5D-SB-0.0-0.5 Collected: 10/2/2013 10:30:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CADMIUM	0.268	J	0.0810	MDL	1.07	PQL	mg/Kg	J	Z
TIN	3.05	J	0.235	MDL	10.7	PQL	mg/Kg	U	B

Sample ID: SL-589-SA5D-SB-4.0-5.0 Collected: 10/2/2013 10:40:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.932	J	0.0742	MDL	1.11	PQL	mg/Kg	J	Z
CADMIUM	0.372	J	0.0842	MDL	1.11	PQL	mg/Kg	J	Z
MOLYBDENUM	0.281	J	0.188	MDL	2.21	PQL	mg/Kg	U	F, F
TIN	2.86	J	0.244	MDL	11.1	PQL	mg/Kg	U	B

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-589-SA5D-SB-0.0-0.5 Collected: 10/2/2013 10:30:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.249	J	0.107	MDL	0.426	PQL	mg/Kg	J	Z

Sample ID: SL-589-SA5D-SB-0.0-0.5 Collected: 10/2/2013 10:30:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0665	J	0.0277	MDL	0.213	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

12/23/2013 8:59:16 AM

ADR version 1.7.0.207

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Data Qualifier Summary

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS									
Method:	6020A	Matrix:			SO					

Sample ID: SL-589-SA5D-SB-4.0-5.0 Collected: 10/2/2013 10:40:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.289	J	0.111	MDL	0.443	PQL	mg/Kg	J	Z

Sample ID: SL-589-SA5D-SB-4.0-5.0 Collected: 10/2/2013 10:40:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0485	J	0.0288	MDL	0.221	PQL	mg/Kg	J	Z

Method Category:	METALS									
Method:	7199	Matrix:			AQ					

Sample ID: EB1-100213 Collected: 10/2/2013 2:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	10.0	U	5.0	MDL	10.0	PQL	ug/L	UJ	H

Method Category:	SVOA									
Method:	1613B	Matrix:			AQ					

Sample ID: EB1-100213 Collected: 10/2/2013 2:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.368	JBQ	0.169	MDL	9.70	PQL	pg/L	U	B
1,2,3,4,6,7,8-HPCDF	0.288	JBQ	0.0778	MDL	9.70	PQL	pg/L	U	B
1,2,3,4,7,8,9-HPCDF	0.363	JBQ	0.0889	MDL	9.70	PQL	pg/L	U	B
1,2,3,4,7,8-HXCDF	0.201	JBQ	0.0844	MDL	9.70	PQL	pg/L	U	B
1,2,3,6,7,8-HXCDD	0.227	JBQ	0.183	MDL	9.70	PQL	pg/L	U	B
1,2,3,6,7,8-HXCDF	0.207	JBQ	0.0855	MDL	9.70	PQL	pg/L	U	B
1,2,3,7,8,9-HXCDF	0.334	JBQ	0.0871	MDL	9.70	PQL	pg/L	U	B
1,2,3,7,8-PECDD	0.338	JBQ	0.277	MDL	9.70	PQL	pg/L	U	B
1,2,3,7,8-PECDF	0.348	JBQ	0.135	MDL	9.70	PQL	pg/L	U	B
2,3,4,6,7,8-HXCDF	0.255	JBQ	0.0804	MDL	9.70	PQL	pg/L	U	B
2,3,4,7,8-PECDF	0.247	JBQ	0.119	MDL	9.70	PQL	pg/L	U	B
2,3,7,8-TCDF	0.237	JQ	0.228	MDL	1.94	PQL	pg/L	J	Z
OCDD	1.49	JB	0.343	MDL	19.4	PQL	pg/L	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

12/23/2013 8:59:16 AM

ADR version 1.7.0.207

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Data Qualifier Summary

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 1613B **Matrix:** AQ

Sample ID: EB1-100213 Collected: 10/2/2013 2:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
OCDF	1.43	JBQ	0.285	MDL	19.4	PQL	pg/L	U	B

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-589-SA5D-SB-0.0-0.5 Collected: 10/2/2013 10:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0854	JBQ	0.0596	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0915	JBQ	0.0258	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0707	JQ	0.0558	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.133	JQ	0.0600	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.0953	JQ	0.0619	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDF	0.139	JQ	0.0435	MDL	5.22	PQL	ng/Kg	J	Z
OCDD	0.858	JBQ	0.0576	MDL	10.4	PQL	ng/Kg	U	B
OCDF	0.219	JBQ	0.0764	MDL	10.4	PQL	ng/Kg	U	B

Method Category: SVOA
Method: 8015M **Matrix:** AQ

Sample ID: EB1-100213 Collected: 10/2/2013 2:00:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C8-C11)	0.097	U	0.048	MDL	0.097	PQL	mg/L	UJ	L

Method Category: SVOA
Method: 8270D SIM **Matrix:** AQ

Sample ID: EB1-100213 Collected: 10/2/2013 2:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
2-METHYLNAPHTHALENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
ACENAPHTHENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** AQ

Sample ID: EB1-100213 Collected: 10/2/2013 2:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ACENAPHTHYLENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
ANTHRACENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
BENZO(A)ANTHRACENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
BENZO(A)PYRENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
BENZO(B)FLUORANTHENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
BENZO(E)PYRENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
BENZO(G,H,I)PERYLENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	E
BENZO(K)FLUORANTHENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
BIS(2-ETHYLHEXYL)PHTHALATE	0.059	J	0.051	MDL	1.0	PQL	ug/L	J	Z, E
Butylbenzylphthalate	1.0	U	0.051	MDL	1.0	PQL	ug/L	UJ	E
CHRYSENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
Diethylphthalate	0.27	J	0.051	MDL	1.0	PQL	ug/L	J	Z, L, E
Dimethylphthalate	1.0	U	0.051	MDL	1.0	PQL	ug/L	UJ	E
Di-n-butylphthalate	0.18	J	0.051	MDL	1.0	PQL	ug/L	J	Z, L, E
FLUORANTHENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
FLUORENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E
NAPHTHALENE	0.051	U	0.030	MDL	0.051	PQL	ug/L	UJ	L, E
PHENANTHRENE	0.051	U	0.030	MDL	0.051	PQL	ug/L	UJ	L, E
PYRENE	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	L, E

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-589-SA5D-SB-4.0-5.0 Collected: 10/2/2013 10:40:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	1.0	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	1.0	J	0.37	MDL	1.8	PQL	ug/Kg	J	Z
FLUORANTHENE	1.4	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z
PHENANTHRENE	1.1	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z
PYRENE	1.3	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Calibration Blank Contamination
B	Method Blank Contamination
E	Laboratory Control Precision
F	Equipment Blank Contamination
F	Field Blank Contamination
FD	Field Duplicate Precision
H	Sampling to Analysis Estimation
L	Laboratory Control Precision
L	Laboratory Control Spike Lower Estimation
L	Laboratory Control Spike Upper Estimation
Q	Matrix Spike Lower Estimation
S	Surrogate/Tracer Recovery Lower Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I

EPA Level III ADR Outliers

(Including Manual Review Outliers)

Quality Control Outlier Reports

PH120

QC Outlier Report: HoldingTimes

Lab Reporting Batch ID: PH120
EDD Filename: PrepPH120_v1

Laboratory: LL
eQAPP Name: CDM_SSFL_131101_Lan

Method: 7199		Preparation Method: Gen Prep			
Matrix: AQ					
Sample ID	Type	Actual	Criteria	Units	Flag
EB1-100213 (RES)	Sampling To Analysis	25.25	24.00	HOURS	J(all detects) UJ(all non-detects)

Method Blank Outlier Report

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2810B371346	10/10/2013 1:46:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF OCDD OCDF	0.614 pg/L 0.745 pg/L 0.694 pg/L 0.571 pg/L 0.572 pg/L 0.559 pg/L 0.348 pg/L 0.600 pg/L 0.427 pg/L 0.502 pg/L 0.387 pg/L 0.364 pg/L 1.53 pg/L 2.35 pg/L	EB1-100213

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB1-100213(RES)	1,2,3,4,6,7,8-HPCDD	0.368 pg/L	0.368U pg/L
EB1-100213(RES)	1,2,3,4,6,7,8-HPCDF	0.288 pg/L	0.288U pg/L
EB1-100213(RES)	1,2,3,4,7,8,9-HPCDF	0.363 pg/L	0.363U pg/L
EB1-100213(RES)	1,2,3,4,7,8-HXCDF	0.201 pg/L	0.201U pg/L
EB1-100213(RES)	1,2,3,6,7,8-HXCDD	0.227 pg/L	0.227U pg/L
EB1-100213(RES)	1,2,3,6,7,8-HXCDF	0.207 pg/L	0.207U pg/L
EB1-100213(RES)	1,2,3,7,8,9-HXCDF	0.334 pg/L	0.334U pg/L
EB1-100213(RES)	1,2,3,7,8-PECDD	0.338 pg/L	0.338U pg/L
EB1-100213(RES)	1,2,3,7,8-PECDF	0.348 pg/L	0.348U pg/L
EB1-100213(RES)	2,3,4,6,7,8-HXCDF	0.255 pg/L	0.255U pg/L
EB1-100213(RES)	2,3,4,7,8-PECDF	0.247 pg/L	0.247U pg/L
EB1-100213(RES)	OCDD	1.49 pg/L	1.49U pg/L
EB1-100213(RES)	OCDF	1.43 pg/L	1.43U pg/L

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2830B371446	10/14/2013 2:46:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,6,7,8-HXCDF 1,2,3,7,8-PECDF OCDD OCDF	0.0753 ng/Kg 0.0423 ng/Kg 0.0948 ng/Kg 0.0800 ng/Kg 0.215 ng/Kg 0.137 ng/Kg	SL-589-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-589-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.0854 ng/Kg	0.0854U ng/Kg
SL-589-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0915 ng/Kg	0.0915U ng/Kg
SL-589-SA5D-SB-0.0-0.5(RES)	OCDD	0.858 ng/Kg	0.858U ng/Kg
SL-589-SA5D-SB-0.0-0.5(RES)	OCDF	0.219 ng/Kg	0.219U ng/Kg

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Method Blank Outlier Report

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P28135AB221117	10/12/2013 11:17:00 AM	PHOSPHORUS POTASSIUM ZINC	0.0547 mg/L 0.203 mg/L 0.0059 mg/L	EB1-100213

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P28137AB221240	10/14/2013 12:40:00 PM	ALUMINUM CALCIUM IRON MAGNESIUM TIN ZINC	9.86 mg/Kg 12.9 mg/Kg 3.84 mg/Kg 2.49 mg/Kg 1.41 mg/Kg 0.656 mg/Kg	SL-589-SA5D-SB-0.0-0.5 SL-589-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-589-SA5D-SB-0.0-0.5(REA)	TIN	3.05 mg/Kg	3.05U mg/Kg
SL-589-SA5D-SB-4.0-5.0(REA)	TIN	2.86 mg/Kg	2.86U mg/Kg

Method: 8081B
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P80803AB241052A	10/8/2013 10:52:00 AM	BETA-BHC	0.0046 ug/L	EB1-100213

Equipment Rinsate Blank Outlier Report

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C

Matrix: SO

Equipment Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
EB1-100213(RES)	10/2/2013 2:00:00 PM	BARIUM BORON CALCIUM COPPER MOLYBDENUM	0.00058 mg/L 0.0098 mg/L 0.0427 mg/L 0.0028 mg/L 0.0086 mg/L	SL-589-SA5D-SB-0.0-0.5 SL-589-SA5D-SB-4.0-5.0 SL-594-SA5D-SB-0.0-0.5 SL-594-SA5D-SB-4.0-5.0 SL-594-SA5D-SB-7.0-8.0 SL-600-SA5D-SB-0.0-0.5 SL-600-SA5D-SB-14.0-15.0 SL-600-SA5D-SB-18.5-19.5 SL-600-SA5D-SB-4.0-5.0 SL-600-SA5D-SB-9.0-10.0 SL-900-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-589-SA5D-SB-4.0-5.0(REA)	MOLYBDENUM	0.281 mg/Kg	0.281U mg/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041613(REA2/TOT)	4/16/2013 3:15:00 PM	MOLYBDENUM TIN	0.0132 mg/L 0.0029 mg/L	SL-589-SA5D-SB-0.0-0.5 SL-589-SA5D-SB-4.0-5.0 SL-594-SA5D-SB-0.0-0.5 SL-594-SA5D-SB-4.0-5.0 SL-594-SA5D-SB-7.0-8.0 SL-600-SA5D-SB-0.0-0.5 SL-600-SA5D-SB-14.0-15.0 SL-600-SA5D-SB-18.5-19.5 SL-600-SA5D-SB-4.0-5.0 SL-600-SA5D-SB-9.0-10.0 SL-900-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-589-SA5D-SB-4.0-5.0(REA)	MOLYBDENUM	0.281 mg/Kg	0.281U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 300.0

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-600-SA5D-SB-0.0-0.5MS (SL-600-SA5D-SB-0.0-0.5 SL-600-SA5D-SB-14.0-15.0 SL-600-SA5D-SB-18.5-19.5 SL-600-SA5D-SB-9.0-10.0)	FLUORIDE	36	-	80.00-120.00	-	FLUORIDE	J (all detects) UJ (all non-detects)

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8015M
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P32808AQ321759A P32808AY321820A (EB1-100213)	EFH (C8-C11)	64	65	70.00-130.00	-	EFH (C8-C11)	J (all detects) UJ (all non-detects)

Method: 8151A
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P32827AQ241903A P32827AY241930A (EB1-100213)	DINOSEB	238	238	16.00-163.00	-	DINOSEB	J(all detects)

Method: 8270D SIM
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P2WILCSQ262241 P2WILCSY262310 (EB1-100213)	1-METHYLNAPHTHALENE	60	-	80.00-126.00	55 (30.00)	1-METHYLNAPHTHALENE	J(all detects) UJ(all non-detects)
	2-METHYLNAPHTHALENE	58	-	81.00-124.00	56 (30.00)	2-METHYLNAPHTHALENE	
	ACENAPHTHENE	57	-	77.00-118.00	53 (30.00)	ACENAPHTHENE	
	ACENAPHTHYLENE	57	-	80.00-123.00	59 (30.00)	ACENAPHTHYLENE	
	ANTHRACENE	60	-	78.00-123.00	53 (30.00)	ANTHRACENE	
	BENZO(A)ANTHRACENE	62	-	73.00-127.00	50 (30.00)	BENZO(A)ANTHRACENE	
	BENZO(A)PYRENE	65	-	72.00-120.00	47 (30.00)	BENZO(A)PYRENE	
	BENZO(B)FLUORANTHENE	66	-	79.00-136.00	47 (30.00)	BENZO(B)FLUORANTHENE	
	BENZO(E)PYRENE	60	-	70.00-130.00	49 (30.00)	BENZO(E)PYRENE	
	BENZO(G,H,I)PERYLENE	-	-	64.00-130.00	33 (30.00)	BENZO(G,H,I)PERYLENE	
	BENZO(K)FLUORANTHENE	71	-	73.00-131.00	45 (30.00)	BENZO(K)FLUORANTHENE	
	BIS(2-ETHYLHEXYL)PHTHALAT	-	-	70.00-143.00	58 (30.00)	BIS(2-ETHYLHEXYL)PHTHALA	
	Butylbenzylphthalate	-	-	40.00-138.00	50 (30.00)	Butylbenzylphthalate	
	CHRYSENE	63	-	76.00-125.00	45 (30.00)	CHRYSENE	
	Diethylphthalate	63	-	64.00-128.00	49 (30.00)	Diethylphthalate	
	Dimethylphthalate	-	-	23.00-139.00	50 (30.00)	Dimethylphthalate	
	Di-n-butylphthalate	61	-	64.00-141.00	50 (30.00)	Di-n-butylphthalate	
	FLUORANTHENE	61	-	79.00-124.00	51 (30.00)	FLUORANTHENE	
	FLUORENE	55	-	74.00-115.00	55 (30.00)	FLUORENE	
	NAPHTHALENE	57	-	75.00-120.00	56 (30.00)	NAPHTHALENE	
	PHENANTHRENE	58	-	75.00-120.00	52 (30.00)	PHENANTHRENE	
	PYRENE	61	-	71.00-130.00	48 (30.00)	PYRENE	

Field Duplicate RPD Report

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 160.3M

Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-600-SA5D-SB-0.0-0.5	SL-900-SA5D-SB-0.0-0.5			
MOISTURE	5.0	8.6	53		No Qualifiers Applied

Method: 300.0

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-600-SA5D-SB-0.0-0.5	SL-900-SA5D-SB-0.0-0.5			
FLUORIDE	0.93	2.5	92	50.00	J(all detects) UJ(all non-detects)

Method: 9045M

Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-600-SA5D-SB-0.0-0.5	SL-900-SA5D-SB-0.0-0.5			
PH	6.83	7.26	6	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-100213	1,2,3,4,6,7,8-HPCDD	JBQ	0.368	9.70	PQL	pg/L	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.288	9.70	PQL	pg/L	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.363	9.70	PQL	pg/L	
	1,2,3,4,7,8-HXCDF	JBQ	0.201	9.70	PQL	pg/L	
	1,2,3,6,7,8-HXCDD	JBQ	0.227	9.70	PQL	pg/L	
	1,2,3,6,7,8-HXCDF	JBQ	0.207	9.70	PQL	pg/L	
	1,2,3,7,8,9-HXCDF	JBQ	0.334	9.70	PQL	pg/L	
	1,2,3,7,8-PECDD	JBQ	0.338	9.70	PQL	pg/L	
	1,2,3,7,8-PECDF	JBQ	0.348	9.70	PQL	pg/L	
	2,3,4,6,7,8-HXCDF	JBQ	0.255	9.70	PQL	pg/L	
	2,3,4,7,8-PECDF	JBQ	0.247	9.70	PQL	pg/L	
	2,3,7,8-TCDF	JQ	0.237	1.94	PQL	pg/L	
	OCDD	JB	1.49	19.4	PQL	pg/L	
	OCDF	JBQ	1.43	19.4	PQL	pg/L	

Method: 6010C

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-100213	BARIUM	J	0.00058	0.0100	PQL	mg/L	J (all detects)
	BORON	J	0.0098	0.100	PQL	mg/L	
	CALCIUM	J	0.0427	0.400	PQL	mg/L	
	COPPER	J	0.0028	0.0200	PQL	mg/L	
	MOLYBDENUM	J	0.0086	0.0200	PQL	mg/L	

Method: 8270D SIM

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-100213	BIS(2-ETHYLHEXYL)PHTHALATE	J	0.059	1.0	PQL	ug/L	J (all detects)
	Diethylphthalate	J	0.27	1.0	PQL	ug/L	
	Di-n-butylphthalate	J	0.18	1.0	PQL	ug/L	

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-589-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.0854	5.22	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0915	5.22	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0707	5.22	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JQ	0.133	5.22	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JQ	0.0953	5.22	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.139	5.22	PQL	ng/Kg	
	OCDD	JBQ	0.858	10.4	PQL	ng/Kg	
	OCDF	JBQ	0.219	10.4	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH120

Laboratory: LL

EDD Filename: PrepPH120_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 300.0

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-594-SA5D-SB-0.0-0.5	FLUORIDE	J	0.76	1.0	PQL	mg/Kg	J (all detects)
SL-600-SA5D-SB-0.0-0.5	FLUORIDE	J	0.93	1.0	PQL	mg/Kg	J (all detects)

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-589-SA5D-SB-0.0-0.5	CADMIUM	J	0.268	1.07	PQL	mg/Kg	J (all detects)
	TIN	J	3.05	10.7	PQL	mg/Kg	
SL-589-SA5D-SB-4.0-5.0	BERYLLIUM	J	0.932	1.11	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.372	1.11	PQL	mg/Kg	
	MOLYBDENUM	J	0.281	2.21	PQL	mg/Kg	
	TIN	J	2.86	11.1	PQL	mg/Kg	

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-589-SA5D-SB-0.0-0.5	SELENIUM	J	0.249	0.426	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0665	0.213	PQL	mg/Kg	
SL-589-SA5D-SB-4.0-5.0	SELENIUM	J	0.289	0.443	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0485	0.221	PQL	mg/Kg	

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-589-SA5D-SB-4.0-5.0	BENZO(B)FLUORANTHENE	J	1.0	1.8	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	1.0	1.8	PQL	ug/Kg	
	FLUORANTHENE	J	1.4	1.8	PQL	ug/Kg	
	PHENANTHRENE	J	1.1	1.8	PQL	ug/Kg	
	PYRENE	J	1.3	1.8	PQL	ug/Kg	

LDC #: 30842D4

VALIDATION COMPLETENESS WORKSHEET

Date: 12/4/13

SDG #: PH120

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: CL

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	-	Sampling dates: 10/2/13
II.	ICP/MS Tune	-	
III.	Calibration	-	
IV.	Blanks	SW	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	N	CS
VII.	Duplicate Sample Analysis	N	CS
VIII.	Laboratory Control Samples (LCS)	N	LCS/9
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	N	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	N	
XV.	Field Blanks	SW	FB = FB-011613 EB = EB-1002B a

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples:

soil/water

1	EB-100213 w	11		21		31	
2	SL-589-SA5D-SB-0.0-0.5 S	12		22		32	
3	SL-589-SA5D-SB-4.0-5.0 J	13		23		33	
4		14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 30842D4

**VALIDATION FINDINGS WORKSHEET
PB/ICB/CCB QUALIFIED SAMPLES**

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: Trace metals (EPA SW 864 Method 6010B/6020/7000)

Soil preparation factor applied: 100x Reason: B

Sample Concentration units, unless otherwise noted: ug/L Associated Samples: All Water

					Sample Identification										
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/L)	Maximum ICB/CCB ^a (ug/l)	Action Level	1										
Mo			3.3	16.5	8.6										

Samples with analyte concentrations within five times the associated ICB, CCB or PB concentration are listed above with the identifications from the Validation Completeness Worksheet. These sample results were qualified as not detected, "U".

Note : a - The listed analyte concentration is the highest ICB, CCB, or PB detected in the analysis of each element.

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: Associated Samples: All Soil

Analyte	Blank ID	Sample Identification											
	FB-041613 (SDG: PH032)	Action Limit	2										
Mo	0.0132	6.60	0.28										
Sn	0.0029	1.45											

Sampling date: 10/2/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: Associated Samples: All Soil

Analyte	Blank ID	Sample Identification											
	EB1-100213 (SDG: PH120)	Action Limit	2										
Ba	0.00058	0.29											
B	0.0098	4.9											
Ca	0.0427	21.35											
Cu	0.0028	1.4											
Mo	0.0086	4.3	See FB										

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH122

Prepared for

CDM Smith
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
2701 Loker Ave West, Suite 220
Carlsbad, California 92010

December 23, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on October 3, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by Environmental Protection Agency (EPA) SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A, and 7471B
Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M
TPH as Extractables by EPA SW 846 Method 8015M
Dioxins and Dibenzofurans by EPA Method 1613B

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), method blanks, trip blanks, equipment blanks, and field blanks samples. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VIII. Laboratory Duplicates Sample

The laboratory has indicated that there were no laboratory duplicate (DUP) analyses specified for the samples in this SDG, and therefore laboratory duplicate analyses were not performed for this SDG.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution was not performed for this SDG.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH122	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

No field duplicates were identified in this SDG.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No volatile contaminants were found in the trip blank.

One equipment blank (from SDG PH120) was collected and analyzed for SVOCs, PCBs, metals, TPH as gasoline, TPH as extractables and dioxins. The equipment blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to equipment blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the equipment blanks were not qualified. The equipment blank outlier reports are presented in Enclosure I.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, PCBs, metals, TPH as gasoline, TPH as extractables and dioxins. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
03-Oct-2013	TB-100313	7224842	TB	5030B	8015M	III
03-Oct-2013	SL-502-SA5D-SB-0.0-0.5	7224840	N	3050B	6010C	III
03-Oct-2013	SL-502-SA5D-SB-0.0-0.5	7224840	N	3050B	6020A	III
03-Oct-2013	SL-502-SA5D-SB-0.0-0.5	7224840	N	3546	8015M	III
03-Oct-2013	SL-502-SA5D-SB-0.0-0.5	7224840	N	3546	8082A	III
03-Oct-2013	SL-502-SA5D-SB-0.0-0.5	7224840	N	3546	8270D SIM	III
03-Oct-2013	SL-502-SA5D-SB-0.0-0.5	7224840	N	METHOD	1613B	III
03-Oct-2013	SL-502-SA5D-SB-0.0-0.5	7224840	N	METHOD	7471B	III
03-Oct-2013	SL-502-SA5D-SB-0.0-0.5MSD	P224840M242250A	MSD	3546	8082A	III
03-Oct-2013	SL-502-SA5D-SB-0.0-0.5MS	P224840R242231A	MS	3546	8082A	III
03-Oct-2013	SL-502-SA5D-SB-6.5-7.5	7224841	N	3050B	6010C	III
03-Oct-2013	SL-502-SA5D-SB-6.5-7.5	7224841	N	3050B	6020A	III
03-Oct-2013	SL-502-SA5D-SB-6.5-7.5	7224841	N	3546	8015M	III
03-Oct-2013	SL-502-SA5D-SB-6.5-7.5	7224841	N	3546	8082A	III
03-Oct-2013	SL-502-SA5D-SB-6.5-7.5	7224841	N	3546	8270D SIM	III
03-Oct-2013	SL-502-SA5D-SB-6.5-7.5	7224841	N	5035A	8015M	III
03-Oct-2013	SL-502-SA5D-SB-6.5-7.5	7224841	N	METHOD	7471B	III
03-Oct-2013	SL-508-SA5D-SB-0.0-0.5	7224843	N	3050B	6010C	III
03-Oct-2013	SL-508-SA5D-SB-0.0-0.5	7224843	N	3050B	6020A	III
03-Oct-2013	SL-508-SA5D-SB-0.0-0.5	7224843	N	3546	8015M	III
03-Oct-2013	SL-508-SA5D-SB-0.0-0.5	7224843	N	3546	8082A	III
03-Oct-2013	SL-508-SA5D-SB-0.0-0.5	7224843	N	3546	8270D SIM	III
03-Oct-2013	SL-508-SA5D-SB-0.0-0.5	7224843	N	METHOD	1613B	III
03-Oct-2013	SL-508-SA5D-SB-0.0-0.5	7224843	N	METHOD	7471B	III
03-Oct-2013	SL-508-SA5D-SB-4.0-5.0	7224844	N	3050B	6010C	III
03-Oct-2013	SL-508-SA5D-SB-4.0-5.0	7224844	N	3050B	6020A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
03-Oct-2013	SL-508-SA5D-SB-4.0-5.0	7224844	N	3546	8015M	III
03-Oct-2013	SL-508-SA5D-SB-4.0-5.0	7224844	N	3546	8082A	III
03-Oct-2013	SL-508-SA5D-SB-4.0-5.0	7224844	N	3546	8270D SIM	III
03-Oct-2013	SL-508-SA5D-SB-4.0-5.0	7224844	N	5035A	8015M	III
03-Oct-2013	SL-508-SA5D-SB-4.0-5.0	7224844	N	METHOD	7471B	III
03-Oct-2013	SL-508-SA5D-SB-9.0-10.0	7224845	N	3050B	6010C	III
03-Oct-2013	SL-508-SA5D-SB-9.0-10.0	7224845	N	3050B	6020A	III
03-Oct-2013	SL-508-SA5D-SB-9.0-10.0	7224845	N	3546	8015M	III
03-Oct-2013	SL-508-SA5D-SB-9.0-10.0	7224845	N	3546	8082A	III
03-Oct-2013	SL-508-SA5D-SB-9.0-10.0	7224845	N	3546	8270D SIM	III
03-Oct-2013	SL-508-SA5D-SB-9.0-10.0	7224845	N	5035A	8015M	III
03-Oct-2013	SL-508-SA5D-SB-9.0-10.0	7224845	N	METHOD	7471B	III
03-Oct-2013	SL-508-SA5D-SB-14.0-15.0	7224846	N	3050B	6010C	III
03-Oct-2013	SL-508-SA5D-SB-14.0-15.0	7224846	N	3050B	6020A	III
03-Oct-2013	SL-508-SA5D-SB-14.0-15.0	7224846	N	3546	8015M	III
03-Oct-2013	SL-508-SA5D-SB-14.0-15.0	7224846	N	3546	8082A	III
03-Oct-2013	SL-508-SA5D-SB-14.0-15.0	7224846	N	3546	8270D SIM	III
03-Oct-2013	SL-508-SA5D-SB-14.0-15.0	7224846	N	5035A	8015M	III
03-Oct-2013	SL-508-SA5D-SB-14.0-15.0	7224846	N	METHOD	7471B	III
03-Oct-2013	SL-508-SA5D-SB-19.0-20.0	7224847	N	3050B	6010C	III
03-Oct-2013	SL-508-SA5D-SB-19.0-20.0	7224847	N	3050B	6020A	III
03-Oct-2013	SL-508-SA5D-SB-19.0-20.0	7224847	N	3546	8015M	III
03-Oct-2013	SL-508-SA5D-SB-19.0-20.0	7224847	N	3546	8082A	III
03-Oct-2013	SL-508-SA5D-SB-19.0-20.0	7224847	N	3546	8270D SIM	III
03-Oct-2013	SL-508-SA5D-SB-19.0-20.0	7224847	N	5035A	8015M	III
03-Oct-2013	SL-508-SA5D-SB-19.0-20.0	7224847	N	METHOD	7471B	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
03-Oct-2013	SL-508-SA5D-SB-23.0-24.0	7224848	N	3050B	6010C	III
03-Oct-2013	SL-508-SA5D-SB-23.0-24.0	7224848	N	3050B	6020A	III
03-Oct-2013	SL-508-SA5D-SB-23.0-24.0	7224848	N	3546	8015M	III
03-Oct-2013	SL-508-SA5D-SB-23.0-24.0	7224848	N	3546	8082A	III
03-Oct-2013	SL-508-SA5D-SB-23.0-24.0	7224848	N	3546	8270D SIM	III
03-Oct-2013	SL-508-SA5D-SB-23.0-24.0	7224848	N	5035A	8015M	III
03-Oct-2013	SL-508-SA5D-SB-23.0-24.0	7224848	N	METHOD	7471B	III
03-Oct-2013	SL-507-SA5D-SB-0.0-0.5	7224849	N	3050B	6010C	III
03-Oct-2013	SL-507-SA5D-SB-0.0-0.5	7224849	N	3050B	6020A	III
03-Oct-2013	SL-507-SA5D-SB-0.0-0.5	7224849	N	3546	8015M	III
03-Oct-2013	SL-507-SA5D-SB-0.0-0.5	7224849	N	3546	8082A	III
03-Oct-2013	SL-507-SA5D-SB-0.0-0.5	7224849	N	3546	8270D SIM	III
03-Oct-2013	SL-507-SA5D-SB-0.0-0.5	7224849	N	METHOD	1613B	III
03-Oct-2013	SL-507-SA5D-SB-0.0-0.5	7224849	N	METHOD	7471B	III
03-Oct-2013	SL-507-SA5D-SB-4.0-5.0	7224850	N	3050B	6010C	III
03-Oct-2013	SL-507-SA5D-SB-4.0-5.0	7224850	N	3050B	6020A	III
03-Oct-2013	SL-507-SA5D-SB-4.0-5.0	7224850	N	3546	8015M	III
03-Oct-2013	SL-507-SA5D-SB-4.0-5.0	7224850	N	3546	8082A	III
03-Oct-2013	SL-507-SA5D-SB-4.0-5.0	7224850	N	3546	8270D SIM	III
03-Oct-2013	SL-507-SA5D-SB-4.0-5.0	7224850	N	5035A	8015M	III
03-Oct-2013	SL-507-SA5D-SB-4.0-5.0	7224850	N	METHOD	7471B	III
03-Oct-2013	SL-507-SA5D-SB-10.5-11.5	7224851	N	3050B	6010C	III
03-Oct-2013	SL-507-SA5D-SB-10.5-11.5	7224851	N	3050B	6020A	III
03-Oct-2013	SL-507-SA5D-SB-10.5-11.5	7224851	N	3546	8015M	III
03-Oct-2013	SL-507-SA5D-SB-10.5-11.5	7224851	N	3546	8082A	III
03-Oct-2013	SL-507-SA5D-SB-10.5-11.5	7224851	N	3546	8270D SIM	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
03-Oct-2013	SL-507-SA5D-SB-10.5-11.5	7224851	N	5035A	8015M	III
03-Oct-2013	SL-507-SA5D-SB-10.5-11.5	7224851	N	METHOD	7471B	III
03-Oct-2013	SL-608-SA5D-SB-0.0-0.5	7224852	N	3050B	6010C	III
03-Oct-2013	SL-608-SA5D-SB-0.0-0.5	7224852	N	3050B	6020A	III
03-Oct-2013	SL-608-SA5D-SB-0.0-0.5	7224852	N	3546	8015M	III
03-Oct-2013	SL-608-SA5D-SB-0.0-0.5	7224852	N	3546	8082A	III
03-Oct-2013	SL-608-SA5D-SB-0.0-0.5	7224852	N	3546	8270D SIM	III
03-Oct-2013	SL-608-SA5D-SB-0.0-0.5	7224852	N	METHOD	1613B	III
03-Oct-2013	SL-608-SA5D-SB-0.0-0.5	7224852	N	METHOD	7471B	III
03-Oct-2013	SL-608-SA5D-SB-4.0-5.0	7224853	N	3050B	6010C	III
03-Oct-2013	SL-608-SA5D-SB-4.0-5.0	7224853	N	3050B	6020A	III
03-Oct-2013	SL-608-SA5D-SB-4.0-5.0	7224853	N	3546	8015M	III
03-Oct-2013	SL-608-SA5D-SB-4.0-5.0	7224853	N	3546	8082A	III
03-Oct-2013	SL-608-SA5D-SB-4.0-5.0	7224853	N	3546	8270D SIM	III
03-Oct-2013	SL-608-SA5D-SB-4.0-5.0	7224853	N	5035A	8015M	III
03-Oct-2013	SL-608-SA5D-SB-4.0-5.0	7224853	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS

Method: 6010C

Matrix: SO

Sample ID: SL-502-SA5D-SB-0.0-0.5

Collected: 10/3/2013 8:40:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.849	J	0.0676	MDL	1.01	PQL	mg/Kg	J	Z
CADMIUM	0.518	J	0.0767	MDL	1.01	PQL	mg/Kg	J	Z
TIN	2.44	J	0.222	MDL	10.1	PQL	mg/Kg	U	B
MOLYBDENUM	4.55		0.172	MDL	2.02	PQL	mg/Kg	U	F

Sample ID: SL-502-SA5D-SB-6.5-7.5

Collected: 10/3/2013 8:50:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BORON	9.05	J	0.918	MDL	10.9	PQL	mg/Kg	J	Z
CADMIUM	0.343	J	0.0831	MDL	1.09	PQL	mg/Kg	J	Z
SODIUM	100	J	18.3	MDL	109	PQL	mg/Kg	J	Z
TIN	3.14	J	0.240	MDL	10.9	PQL	mg/Kg	U	B

Sample ID: SL-507-SA5D-SB-0.0-0.5

Collected: 10/3/2013 12:30:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	3.51	J	0.718	MDL	4.11	PQL	mg/Kg	J	Z
BERYLLIUM	0.785	J	0.0688	MDL	1.03	PQL	mg/Kg	J	Z
CADMIUM	0.690	J	0.0780	MDL	1.03	PQL	mg/Kg	J	Z
MOLYBDENUM	0.348	J	0.174	MDL	2.05	PQL	mg/Kg	U	F, F
SODIUM	84.3	J	17.1	MDL	103	PQL	mg/Kg	J	Z
TIN	2.71	J	0.226	MDL	10.3	PQL	mg/Kg	U	B

Sample ID: SL-507-SA5D-SB-10.5-11.5

Collected: 10/3/2013 12:50:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.546	J	0.0700	MDL	1.05	PQL	mg/Kg	J	Z
BORON	7.72	J	0.878	MDL	10.5	PQL	mg/Kg	J	Z
CADMIUM	0.224	J	0.0794	MDL	1.05	PQL	mg/Kg	J	Z
TIN	2.36	J	0.230	MDL	10.5	PQL	mg/Kg	U	B
Zirconium	3.82	J	0.878	MDL	5.23	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-507-SA5D-SB-4.0-5.0 Collected: 10/3/2013 12:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	4.07	J	0.773	MDL	4.42	PQL	mg/Kg	J	Z
BERYLLIUM	1.01	J	0.0740	MDL	1.10	PQL	mg/Kg	J	Z
CADMIUM	0.423	J	0.0839	MDL	1.10	PQL	mg/Kg	J	Z
TIN	2.97	J	0.243	MDL	11.0	PQL	mg/Kg	U	B

Sample ID: SL-508-SA5D-SB-0.0-0.5 Collected: 10/3/2013 9:25:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	3.72	J	0.722	MDL	4.13	PQL	mg/Kg	J	Z
BERYLLIUM	0.959	J	0.0691	MDL	1.03	PQL	mg/Kg	J	Z
CADMIUM	0.474	J	0.0784	MDL	1.03	PQL	mg/Kg	J	Z
MOLYBDENUM	0.209	J	0.175	MDL	2.06	PQL	mg/Kg	U	F, F
SODIUM	82.0	J	17.2	MDL	103	PQL	mg/Kg	J	Z
TIN	2.66	J	0.227	MDL	10.3	PQL	mg/Kg	U	B

Sample ID: SL-508-SA5D-SB-14.0-15.0 Collected: 10/3/2013 9:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.764	J	0.0735	MDL	1.10	PQL	mg/Kg	J	Z
BORON	8.07	J	0.921	MDL	11.0	PQL	mg/Kg	J	Z
CADMIUM	0.214	J	0.0833	MDL	1.10	PQL	mg/Kg	J	Z
TIN	2.77	J	0.241	MDL	11.0	PQL	mg/Kg	U	B

Sample ID: SL-508-SA5D-SB-19.0-20.0 Collected: 10/3/2013 10:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	4.22	J	0.755	MDL	4.31	PQL	mg/Kg	J	Z
BERYLLIUM	0.722	J	0.0723	MDL	1.08	PQL	mg/Kg	J	Z
BORON	8.19	J	0.906	MDL	10.8	PQL	mg/Kg	J	Z
CADMIUM	0.302	J	0.0820	MDL	1.08	PQL	mg/Kg	J	Z
TIN	2.73	J	0.237	MDL	10.8	PQL	mg/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS

Method: 6010C

Matrix: SO

Sample ID: SL-508-SA5D-SB-23.0-24.0

Collected: 10/3/2013 10:15:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	3.45	J	0.737	MDL	4.21	PQL	mg/Kg	J	Z
BERYLLIUM	0.574	J	0.0706	MDL	1.05	PQL	mg/Kg	J	Z
BORON	6.19	J	0.885	MDL	10.5	PQL	mg/Kg	J	Z
CADMIUM	0.272	J	0.0800	MDL	1.05	PQL	mg/Kg	J	Z
TIN	2.47	J	0.232	MDL	10.5	PQL	mg/Kg	U	B
Zirconium	3.76	J	0.885	MDL	5.27	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-4.0-5.0

Collected: 10/3/2013 9:35:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	3.46	J	0.786	MDL	4.49	PQL	mg/Kg	J	Z
BERYLLIUM	0.957	J	0.0752	MDL	1.12	PQL	mg/Kg	J	Z
CADMIUM	0.395	J	0.0853	MDL	1.12	PQL	mg/Kg	J	Z
TIN	2.69	J	0.247	MDL	11.2	PQL	mg/Kg	U	B

Sample ID: SL-508-SA5D-SB-9.0-10.0

Collected: 10/3/2013 9:45:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	4.26	J	0.781	MDL	4.46	PQL	mg/Kg	J	Z
BERYLLIUM	0.843	J	0.0747	MDL	1.12	PQL	mg/Kg	J	Z
BORON	10.7	J	0.937	MDL	11.2	PQL	mg/Kg	J	Z
CADMIUM	0.510	J	0.0848	MDL	1.12	PQL	mg/Kg	J	Z
TIN	2.60	J	0.245	MDL	11.2	PQL	mg/Kg	U	B

Sample ID: SL-608-SA5D-SB-0.0-0.5

Collected: 10/3/2013 1:40:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	3.44	J	0.737	MDL	4.21	PQL	mg/Kg	J	Z
BERYLLIUM	1.01	J	0.0705	MDL	1.05	PQL	mg/Kg	J	Z
CADMIUM	0.450	J	0.0800	MDL	1.05	PQL	mg/Kg	J	Z
MOLYBDENUM	0.184	J	0.179	MDL	2.10	PQL	mg/Kg	U	F, F
TIN	2.61	J	0.231	MDL	10.5	PQL	mg/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-608-SA5D-SB-4.0-5.0 Collected: 10/3/2013 1:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	4.10	J	0.760	MDL	4.34	PQL	mg/Kg	J	Z
BERYLLIUM	0.938	J	0.0728	MDL	1.09	PQL	mg/Kg	J	Z
CADMIUM	0.466	J	0.0825	MDL	1.09	PQL	mg/Kg	J	Z
TIN	2.75	J	0.239	MDL	10.9	PQL	mg/Kg	U	B

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-502-SA5D-SB-0.0-0.5 Collected: 10/3/2013 8:40:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0591	J	0.0262	MDL	0.202	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA5D-SB-6.5-7.5 Collected: 10/3/2013 8:50:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.188	J	0.109	MDL	0.437	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA5D-SB-6.5-7.5 Collected: 10/3/2013 8:50:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0566	J	0.0284	MDL	0.219	PQL	mg/Kg	J	Z

Sample ID: SL-507-SA5D-SB-0.0-0.5 Collected: 10/3/2013 12:30:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.260	J	0.103	MDL	0.411	PQL	mg/Kg	J	Z

Sample ID: SL-507-SA5D-SB-0.0-0.5 Collected: 10/3/2013 12:30:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.110	J	0.0267	MDL	0.205	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-507-SA5D-SB-4.0-5.0 Collected: 10/3/2013 12:40:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.127	J	0.110	MDL	0.442	PQL	mg/Kg	J	Z

Sample ID: SL-507-SA5D-SB-4.0-5.0 Collected: 10/3/2013 12:40:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0418	J	0.0287	MDL	0.221	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-0.0-0.5 Collected: 10/3/2013 9:25:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.314	J	0.103	MDL	0.413	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-0.0-0.5 Collected: 10/3/2013 9:25:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0578	J	0.0268	MDL	0.206	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-14.0-15.0 Collected: 10/3/2013 9:55:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0300	J	0.0285	MDL	0.219	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-19.0-20.0 Collected: 10/3/2013 10:05:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0399	J	0.0280	MDL	0.216	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-23.0-24.0 Collected: 10/3/2013 10:15:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0406	J	0.0274	MDL	0.211	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-4.0-5.0 Collected: 10/3/2013 9:35:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.189	J	0.112	MDL	0.449	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-508-SA5D-SB-4.0-5.0 Collected: 10/3/2013 9:35:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0664	J	0.0292	MDL	0.224	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-9.0-10.0 Collected: 10/3/2013 9:45:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0426	J	0.0290	MDL	0.223	PQL	mg/Kg	J	Z

Sample ID: SL-608-SA5D-SB-0.0-0.5 Collected: 10/3/2013 1:40:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.240	J	0.105	MDL	0.421	PQL	mg/Kg	J	Z

Sample ID: SL-608-SA5D-SB-0.0-0.5 Collected: 10/3/2013 1:40:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0572	J	0.0274	MDL	0.210	PQL	mg/Kg	J	Z

Sample ID: SL-608-SA5D-SB-4.0-5.0 Collected: 10/3/2013 1:50:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.148	J	0.109	MDL	0.434	PQL	mg/Kg	J	Z

Sample ID: SL-608-SA5D-SB-4.0-5.0 Collected: 10/3/2013 1:50:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0465	J	0.0282	MDL	0.217	PQL	mg/Kg	J	Z

Method Category: METALS
Method: 7471B **Matrix:** SO

Sample ID: SL-507-SA5D-SB-10.5-11.5 Collected: 10/3/2013 12:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0127	J	0.0104	MDL	0.0174	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 7471B **Matrix:** SO

Sample ID: SL-507-SA5D-SB-4.0-5.0 Collected: 10/3/2013 12:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0160	J	0.0111	MDL	0.0184	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-14.0-15.0 Collected: 10/3/2013 9:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0133	J	0.0111	MDL	0.0186	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-19.0-20.0 Collected: 10/3/2013 10:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0119	J	0.0110	MDL	0.0184	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-23.0-24.0 Collected: 10/3/2013 10:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0111	J	0.0101	MDL	0.0169	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-4.0-5.0 Collected: 10/3/2013 9:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0148	J	0.0107	MDL	0.0179	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA5D-SB-9.0-10.0 Collected: 10/3/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0125	J	0.0107	MDL	0.0178	PQL	mg/Kg	J	Z

Sample ID: SL-608-SA5D-SB-0.0-0.5 Collected: 10/3/2013 1:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0102	J	0.0099	MDL	0.0165	PQL	mg/Kg	J	Z

Sample ID: SL-608-SA5D-SB-4.0-5.0 Collected: 10/3/2013 1:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0123	J	0.0110	MDL	0.0184	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: METALS
Method: 7471B **Matrix:** SO

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-502-SA5D-SB-0.0-0.5 Collected: 10/3/2013 8:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	2.81	JB	0.0260	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.261	J	0.0269	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.361	J	0.0422	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.332	J	0.0293	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.970	J	0.0444	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.367	JB	0.0285	MDL	5.07	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.807	J	0.0417	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.524	J	0.0302	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.442	JQ	0.0331	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.846	JB	0.0262	MDL	5.07	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.356	J	0.0294	MDL	5.07	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.575	J	0.0236	MDL	5.07	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.150	J	0.0279	MDL	1.01	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.310	JQ	0.0598	MDL	1.01	PQL	ng/Kg	J	Z
OCDF	6.33	JB	0.0247	MDL	10.1	PQL	ng/Kg	J	Z

Sample ID: SL-507-SA5D-SB-0.0-0.5 Collected: 10/3/2013 12:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	4.31	JB	0.0723	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.196	J	0.0722	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.557	JQ	0.143	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.377	J	0.0988	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	1.60	J	0.150	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.398	JB	0.0940	MDL	5.20	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	1.41	J	0.133	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.920	J	0.0816	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.361	JQ	0.113	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	1.27	JBQ	0.100	MDL	5.20	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.380	J	0.0918	MDL	5.20	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.589	J	0.163	MDL	1.04	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA

Method: 1613B

Matrix: SO

Sample ID: SL-508-SA5D-SB-0.0-0.5

Collected: 10/3/2013 9:25:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	2.04	JBQ	0.122	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.426	JB	0.0408	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.105	J	0.0555	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.0866	JQ	0.0756	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.146	JQ	0.0712	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.463	JQ	0.0773	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.305	JBQ	0.0644	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.733	J	0.0739	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	1.20	J	0.0694	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.272	JQ	0.132	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	3.31	JB	0.0922	MDL	5.29	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0997	JQ	0.0631	MDL	5.29	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.136	JQ	0.0845	MDL	5.29	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.234	JQ	0.184	MDL	1.06	PQL	ng/Kg	J	Z
OCDF	0.947	JBQ	0.109	MDL	10.6	PQL	ng/Kg	J	Z

Sample ID: SL-608-SA5D-SB-0.0-0.5

Collected: 10/3/2013 1:40:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.228	JB	0.108	MDL	5.09	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0750	JBQ	0.0566	MDL	5.09	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.126	JQ	0.0886	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.175	JQ	0.105	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.285	J	0.103	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.590	J	0.108	MDL	5.09	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0897	JQ	0.0885	MDL	5.09	PQL	ng/Kg	J	Z
OCDD	1.66	JBQ	0.114	MDL	10.2	PQL	ng/Kg	J	Z
OCDF	0.257	JBQ	0.143	MDL	10.2	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8015M **Matrix:** SO

Sample ID: SL-502-SA5D-SB-0.0-0.5 Collected: 10/3/2013 8:40:00 Analysis Type: RES Dilution: 20

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	48	J	42	MDL	100	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 8082A **Matrix:** SO

Sample ID: SL-502-SA5D-SB-0.0-0.5 Collected: 10/3/2013 8:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	7.4	J	4.6	MDL	18	PQL	ug/Kg	J	Z

Sample ID: SL-507-SA5D-SB-0.0-0.5 Collected: 10/3/2013 12:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	10	J	4.6	MDL	18	PQL	ug/Kg	J	Z
AROCLOR 1260	7.2	J	4.1	MDL	18	PQL	ug/Kg	J	Z
Aroclor 5460	13	J	11	MDL	35	PQL	ug/Kg	J	Z

Sample ID: SL-508-SA5D-SB-19.0-20.0 Collected: 10/3/2013 10:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	8.9	J	4.8	MDL	19	PQL	ug/Kg	J	Z

Sample ID: SL-508-SA5D-SB-23.0-24.0 Collected: 10/3/2013 10:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	7.8	J	4.7	MDL	18	PQL	ug/Kg	J	Z

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-502-SA5D-SB-0.0-0.5 Collected: 10/3/2013 8:40:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(E)PYRENE	46	J	35	MDL	180	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-502-SA5D-SB-6.5-7.5 Collected: 10/3/2013 8:50:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	8.8	J	7.4	MDL	19	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	9.0	J	7.4	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	12	J	3.7	MDL	19	PQL	ug/Kg	J	Z

Sample ID: SL-507-SA5D-SB-0.0-0.5 Collected: 10/3/2013 12:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BIS(2-ETHYLHEXYL)PHTHALATE	68	J	63	MDL	190	PQL	ug/Kg	J	Z

Sample ID: SL-508-SA5D-SB-0.0-0.5 Collected: 10/3/2013 9:25:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.81	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(B)FLUORANTHENE	1.3	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
FLUORANTHENE	1.7	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	1.3	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
PHENANTHRENE	1.2	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
PYRENE	1.1	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-508-SA5D-SB-14.0-15.0 Collected: 10/3/2013 9:55:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTHRACENE	0.81	J	0.37	MDL	1.9	PQL	ug/Kg	J	Z
BENZO(E)PYRENE	6.9	J	3.7	MDL	19	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Method Blank Contamination
E	Laboratory Duplicate Precision
F	Equipment Blank Contamination
F	Field Blank Contamination
Q	Laboratory Duplicate Precision
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH122

Method Blank Outlier Report

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2830B371446	10/14/2013 2:46:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,6,7,8-HXCDF 1,2,3,7,8-PECDF OCDD OCDF	0.0753 ng/Kg 0.0423 ng/Kg 0.0948 ng/Kg 0.0800 ng/Kg 0.215 ng/Kg 0.137 ng/Kg	SL-502-SA5D-SB-0.0-0.5 SL-507-SA5D-SB-0.0-0.5 SL-508-SA5D-SB-0.0-0.5 SL-608-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-502-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.367 ng/Kg	0.367U ng/Kg
SL-507-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.398 ng/Kg	0.398U ng/Kg
SL-508-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.305 ng/Kg	0.305U ng/Kg
SL-608-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.228 ng/Kg	0.228U ng/Kg
SL-608-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0750 ng/Kg	0.0750U ng/Kg
SL-608-SA5D-SB-0.0-0.5(RES)	OCDF	0.257 ng/Kg	0.257U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P28137AB221240	10/14/2013 12:40:00 PM	ALUMINUM CALCIUM IRON MAGNESIUM TIN ZINC	9.86 mg/Kg 12.9 mg/Kg 3.84 mg/Kg 2.49 mg/Kg 1.41 mg/Kg 0.656 mg/Kg	SL-502-SA5D-SB-0.0-0.5 SL-502-SA5D-SB-6.5-7.5 SL-507-SA5D-SB-0.0-0.5 SL-507-SA5D-SB-10.5-11.5 SL-507-SA5D-SB-4.0-5.0 SL-508-SA5D-SB-0.0-0.5 SL-508-SA5D-SB-14.0-15.0 SL-508-SA5D-SB-19.0-20.0 SL-508-SA5D-SB-23.0-24.0 SL-508-SA5D-SB-4.0-5.0 SL-508-SA5D-SB-9.0-10.0 SL-608-SA5D-SB-0.0-0.5 SL-608-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-502-SA5D-SB-0.0-0.5(RES)	TIN	2.44 mg/Kg	2.44U mg/Kg
SL-502-SA5D-SB-6.5-7.5(RES)	TIN	3.14 mg/Kg	3.14U mg/Kg
SL-507-SA5D-SB-0.0-0.5(RES)	TIN	2.71 mg/Kg	2.71U mg/Kg
SL-507-SA5D-SB-10.5-11.5(RES)	TIN	2.36 mg/Kg	2.36U mg/Kg
SL-507-SA5D-SB-4.0-5.0(RES)	TIN	2.97 mg/Kg	2.97U mg/Kg
SL-508-SA5D-SB-0.0-0.5(RES)	TIN	2.66 mg/Kg	2.66U mg/Kg
SL-508-SA5D-SB-14.0-15.0(RES)	TIN	2.77 mg/Kg	2.77U mg/Kg
SL-508-SA5D-SB-19.0-20.0(RES)	TIN	2.73 mg/Kg	2.73U mg/Kg
SL-508-SA5D-SB-23.0-24.0(RES)	TIN	2.47 mg/Kg	2.47U mg/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

12/18/2013 2:33:48 PM

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Method Blank Outlier Report

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-508-SA5D-SB-4.0-5.0(RES)	TIN	2.69 mg/Kg	2.69U mg/Kg
SL-508-SA5D-SB-9.0-10.0(RES)	TIN	2.60 mg/Kg	2.60U mg/Kg
SL-608-SA5D-SB-0.0-0.5(RES)	TIN	2.61 mg/Kg	2.61U mg/Kg
SL-608-SA5D-SB-4.0-5.0(RES)	TIN	2.75 mg/Kg	2.75U mg/Kg

Equipment Rinsate Blank Outlier Report

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

Equipment Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
EB1-100213(RES)	10/2/2013 2:00:00 PM	BARIUM BORON CALCIUM COPPER MOLYBDENUM	0.00058 mg/L 0.0098 mg/L 0.0427 mg/L 0.0028 mg/L 0.0086 mg/L	SL-502-SA5D-SB-0.0-0.5 SL-502-SA5D-SB-6.5-7.5 SL-507-SA5D-SB-0.0-0.5 SL-507-SA5D-SB-10.5-11.5 SL-507-SA5D-SB-4.0-5.0 SL-508-SA5D-SB-0.0-0.5 SL-508-SA5D-SB-14.0-15.0 SL-508-SA5D-SB-19.0-20.0 SL-508-SA5D-SB-23.0-24.0 SL-508-SA5D-SB-4.0-5.0 SL-508-SA5D-SB-9.0-10.0 SL-608-SA5D-SB-0.0-0.5 SL-608-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-507-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	0.348 mg/Kg	0.348U mg/Kg
SL-508-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	0.209 mg/Kg	0.209U mg/Kg
SL-608-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	0.184 mg/Kg	0.184U mg/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041613(REA2/TOT)	4/16/2013 3:15:00 PM	MOLYBDENUM TIN	0.0132 mg/L 0.0029 mg/L	SL-502-SA5D-SB-0.0-0.5 SL-502-SA5D-SB-6.5-7.5 SL-507-SA5D-SB-0.0-0.5 SL-507-SA5D-SB-10.5-11.5 SL-507-SA5D-SB-4.0-5.0 SL-508-SA5D-SB-0.0-0.5 SL-508-SA5D-SB-14.0-15.0 SL-508-SA5D-SB-19.0-20.0 SL-508-SA5D-SB-23.0-24.0 SL-508-SA5D-SB-4.0-5.0 SL-508-SA5D-SB-9.0-10.0 SL-608-SA5D-SB-0.0-0.5 SL-608-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-502-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	4.55 mg/Kg	4.55U mg/Kg
SL-507-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	0.348 mg/Kg	0.348U mg/Kg
SL-508-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	0.209 mg/Kg	0.209U mg/Kg
SL-608-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	0.184 mg/Kg	0.184U mg/Kg

Reporting Limit Outliers

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-502-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	2.81	5.07	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	J	0.261	5.07	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	J	0.361	5.07	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	J	0.332	5.07	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	0.970	5.07	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.367	5.07	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	J	0.807	5.07	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.524	5.07	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.442	5.07	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.846	5.07	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	J	0.356	5.07	PQL	ng/Kg	
	2,3,4,7,8-PECDF	J	0.575	5.07	PQL	ng/Kg	
	2,3,7,8-TCDD	J	0.150	1.01	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.310	1.01	PQL	ng/Kg	
	OCDF	JB	6.33	10.1	PQL	ng/Kg	
SL-507-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	4.31	5.20	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	J	0.196	5.20	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.557	5.20	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	J	0.377	5.20	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	1.60	5.20	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.398	5.20	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	J	1.41	5.20	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.920	5.20	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.361	5.20	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	1.27	5.20	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	J	0.380	5.20	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.589	1.04	PQL	ng/Kg	
SL-508-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	2.04	5.29	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.426	5.29	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	J	0.105	5.29	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0866	5.29	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JQ	0.146	5.29	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.463	5.29	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.305	5.29	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	J	0.733	5.29	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	1.20	5.29	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.272	5.29	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	3.31	5.29	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JQ	0.0997	5.29	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.136	5.29	PQL	ng/Kg	
2,3,7,8-TCDF	JQ	0.234	1.06	PQL	ng/Kg		
OCDF	JBQ	0.947	10.6	PQL	ng/Kg		
SL-608-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	0.228	5.09	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0750	5.09	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JQ	0.126	5.09	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.175	5.09	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	J	0.285	5.09	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.590	5.09	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JQ	0.0897	5.09	PQL	ng/Kg	
	OCD	JBQ	1.66	10.2	PQL	ng/Kg	
	OCDF	JBQ	0.257	10.2	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-502-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.849	1.01	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.518	1.01	PQL	mg/Kg	
	TIN	J	2.44	10.1	PQL	mg/Kg	
SL-502-SA5D-SB-6.5-7.5	BORON	J	9.05	10.9	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.343	1.09	PQL	mg/Kg	
	SODIUM	J	100	109	PQL	mg/Kg	
	TIN	J	3.14	10.9	PQL	mg/Kg	
SL-507-SA5D-SB-0.0-0.5	ARSENIC	J	3.51	4.11	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.785	1.03	PQL	mg/Kg	
	CADMIUM	J	0.690	1.03	PQL	mg/Kg	
	MOLYBDENUM	J	0.348	2.05	PQL	mg/Kg	
	SODIUM	J	84.3	103	PQL	mg/Kg	
SL-507-SA5D-SB-10.5-11.5	TIN	J	2.71	10.3	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.546	1.05	PQL	mg/Kg	
	BORON	J	7.72	10.5	PQL	mg/Kg	
	CADMIUM	J	0.224	1.05	PQL	mg/Kg	
	TIN	J	2.36	10.5	PQL	mg/Kg	
SL-507-SA5D-SB-4.0-5.0	Zirconium	J	3.82	5.23	PQL	mg/Kg	J (all detects)
	ARSENIC	J	4.07	4.42	PQL	mg/Kg	
	BERYLLIUM	J	1.01	1.10	PQL	mg/Kg	
	CADMIUM	J	0.423	1.10	PQL	mg/Kg	
SL-508-SA5D-SB-0.0-0.5	TIN	J	2.97	11.0	PQL	mg/Kg	J (all detects)
	ARSENIC	J	3.72	4.13	PQL	mg/Kg	
	BERYLLIUM	J	0.959	1.03	PQL	mg/Kg	
	CADMIUM	J	0.474	1.03	PQL	mg/Kg	
	MOLYBDENUM	J	0.209	2.06	PQL	mg/Kg	
SL-508-SA5D-SB-14.0-15.0	SODIUM	J	82.0	103	PQL	mg/Kg	J (all detects)
	TIN	J	2.66	10.3	PQL	mg/Kg	
	BERYLLIUM	J	0.764	1.10	PQL	mg/Kg	
	BORON	J	8.07	11.0	PQL	mg/Kg	
	CADMIUM	J	0.214	1.10	PQL	mg/Kg	
SL-508-SA5D-SB-19.0-20.0	TIN	J	2.77	11.0	PQL	mg/Kg	J (all detects)
	ARSENIC	J	4.22	4.31	PQL	mg/Kg	
	BERYLLIUM	J	0.722	1.08	PQL	mg/Kg	
	BORON	J	8.19	10.8	PQL	mg/Kg	
	CADMIUM	J	0.302	1.08	PQL	mg/Kg	
SL-508-SA5D-SB-23.0-24.0	TIN	J	2.73	10.8	PQL	mg/Kg	J (all detects)
	ARSENIC	J	3.45	4.21	PQL	mg/Kg	
	BERYLLIUM	J	0.574	1.05	PQL	mg/Kg	
	BORON	J	6.19	10.5	PQL	mg/Kg	
	CADMIUM	J	0.272	1.05	PQL	mg/Kg	
	TIN	J	2.47	10.5	PQL	mg/Kg	
SL-508-SA5D-SB-4.0-5.0	Zirconium	J	3.76	5.27	PQL	mg/Kg	J (all detects)
	ARSENIC	J	3.46	4.49	PQL	mg/Kg	
	BERYLLIUM	J	0.957	1.12	PQL	mg/Kg	
	CADMIUM	J	0.395	1.12	PQL	mg/Kg	
SL-508-SA5D-SB-9.0-10.0	TIN	J	2.69	11.2	PQL	mg/Kg	J (all detects)
	ARSENIC	J	4.26	4.46	PQL	mg/Kg	
	BERYLLIUM	J	0.843	1.12	PQL	mg/Kg	
	BORON	J	10.7	11.2	PQL	mg/Kg	
	CADMIUM	J	0.510	1.12	PQL	mg/Kg	
	TIN	J	2.60	11.2	PQL	mg/Kg	

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Reporting Limit Outliers

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-608-SA5D-SB-0.0-0.5	ARSENIC	J	3.44	4.21	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	1.01	1.05	PQL	mg/Kg	
	CADMIUM	J	0.450	1.05	PQL	mg/Kg	
	MOLYBDENUM	J	0.184	2.10	PQL	mg/Kg	
	TIN	J	2.61	10.5	PQL	mg/Kg	
SL-608-SA5D-SB-4.0-5.0	ARSENIC	J	4.10	4.34	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.938	1.09	PQL	mg/Kg	
	CADMIUM	J	0.466	1.09	PQL	mg/Kg	
	TIN	J	2.75	10.9	PQL	mg/Kg	

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-502-SA5D-SB-0.0-0.5	SILVER	J	0.0591	0.202	PQL	mg/Kg	J (all detects)
SL-502-SA5D-SB-6.5-7.5	SELENIUM	J	0.188	0.437	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0566	0.219	PQL	mg/Kg	
SL-507-SA5D-SB-0.0-0.5	SELENIUM	J	0.260	0.411	PQL	mg/Kg	J (all detects)
	SILVER	J	0.110	0.205	PQL	mg/Kg	
SL-507-SA5D-SB-4.0-5.0	SELENIUM	J	0.127	0.442	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0418	0.221	PQL	mg/Kg	
SL-508-SA5D-SB-0.0-0.5	SELENIUM	J	0.314	0.413	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0578	0.206	PQL	mg/Kg	
SL-508-SA5D-SB-14.0-15.0	SILVER	J	0.0300	0.219	PQL	mg/Kg	J (all detects)
SL-508-SA5D-SB-19.0-20.0	SILVER	J	0.0399	0.216	PQL	mg/Kg	J (all detects)
SL-508-SA5D-SB-23.0-24.0	SILVER	J	0.0406	0.211	PQL	mg/Kg	J (all detects)
SL-508-SA5D-SB-4.0-5.0	SELENIUM	J	0.189	0.449	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0664	0.224	PQL	mg/Kg	
SL-508-SA5D-SB-9.0-10.0	SILVER	J	0.0426	0.223	PQL	mg/Kg	J (all detects)
SL-608-SA5D-SB-0.0-0.5	SELENIUM	J	0.240	0.421	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0572	0.210	PQL	mg/Kg	
SL-608-SA5D-SB-4.0-5.0	SELENIUM	J	0.148	0.434	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0465	0.217	PQL	mg/Kg	

Method: 7471B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-507-SA5D-SB-10.5-11.5	MERCURY	J	0.0127	0.0174	PQL	mg/Kg	J (all detects)
SL-507-SA5D-SB-4.0-5.0	MERCURY	J	0.0160	0.0184	PQL	mg/Kg	J (all detects)
SL-508-SA5D-SB-14.0-15.0	MERCURY	J	0.0133	0.0186	PQL	mg/Kg	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 7471B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-508-SA5D-SB-19.0-20.0	MERCURY	J	0.0119	0.0184	PQL	mg/Kg	J (all detects)
SL-508-SA5D-SB-23.0-24.0	MERCURY	J	0.0111	0.0169	PQL	mg/Kg	J (all detects)
SL-508-SA5D-SB-4.0-5.0	MERCURY	J	0.0148	0.0179	PQL	mg/Kg	J (all detects)
SL-508-SA5D-SB-9.0-10.0	MERCURY	J	0.0125	0.0178	PQL	mg/Kg	J (all detects)
SL-608-SA5D-SB-0.0-0.5	MERCURY	J	0.0102	0.0165	PQL	mg/Kg	J (all detects)
SL-608-SA5D-SB-4.0-5.0	MERCURY	J	0.0123	0.0184	PQL	mg/Kg	J (all detects)

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-502-SA5D-SB-0.0-0.5	EFH (C15-C20)	J	48	100	PQL	mg/Kg	J (all detects)

Method: 8082A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-502-SA5D-SB-0.0-0.5	AROCLOR 1254	J	7.4	18	PQL	ug/Kg	J (all detects)
SL-507-SA5D-SB-0.0-0.5	AROCLOR 1254	J	10	18	PQL	ug/Kg	J (all detects)
	AROCLOR 1260	J	7.2	18	PQL	ug/Kg	
	Aroclor 5460	J	13	35	PQL	ug/Kg	
SL-508-SA5D-SB-19.0-20.0	AROCLOR 1254	J	8.9	19	PQL	ug/Kg	J (all detects)
SL-508-SA5D-SB-23.0-24.0	AROCLOR 1254	J	7.8	18	PQL	ug/Kg	J (all detects)

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-502-SA5D-SB-0.0-0.5	BENZO(E)PYRENE	J	46	180	PQL	ug/Kg	J (all detects)
SL-502-SA5D-SB-6.5-7.5	BENZO(B)FLUORANTHENE	J	8.8	19	PQL	ug/Kg	J (all detects)
	BENZO(G,H,I)PERYLENE	J	9.0	19	PQL	ug/Kg	
	CHRYSENE	J	12	19	PQL	ug/Kg	
SL-507-SA5D-SB-0.0-0.5	BIS(2-ETHYLHEXYL)PHTHALATE	J	68	190	PQL	ug/Kg	J (all detects)
SL-508-SA5D-SB-0.0-0.5	2-METHYLNAPHTHALENE	J	0.81	1.8	PQL	ug/Kg	J (all detects)
	BENZO(B)FLUORANTHENE	J	1.3	1.8	PQL	ug/Kg	
	FLUORANTHENE	J	1.7	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	1.3	1.8	PQL	ug/Kg	
	PHENANTHRENE	J	1.2	1.8	PQL	ug/Kg	
	PYRENE	J	1.1	1.8	PQL	ug/Kg	

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Reporting Limit Outliers

Lab Reporting Batch ID: PH122

Laboratory: LL

EDD Filename: PrepPH122_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8270D SIM

Matrix: SO

<i>SampleID</i>	<i>Analyte</i>	<i>Lab Qual</i>	<i>Result</i>	<i>Reporting Limit</i>	<i>RL Type</i>	<i>Units</i>	<i>Flag</i>
SL-508-SA5D-SB-14.0-15.0	ANTHRACENE	J	0.81	1.9	PQL	ug/Kg	J (all detects)
	BENZO(E)PYRENE	J	6.9	19	PQL	ug/Kg	

LDC #: 30842E4

VALIDATION COMPLETENESS WORKSHEET

Date: 12/4/13

SDG #: PH120 Z

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: oz

2nd Reviewer:

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	—	Sampling dates: 10/3/13
II.	ICP/MS Tune	—	
III.	Calibration	—	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	—	
VI.	Matrix Spike Analysis	N	CS
VII.	Duplicate Sample Analysis	N	CS
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	N	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	N	
XV.	Field Blanks	SW	FB = FB-041613 EB = EB-100213

Note: A = Acceptable ND = No compounds detected D = Duplicate (PH120)
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: Soil

1	SL-502-SA5D-SB-0.0-0.5	11	SL-507-SA5D-SB-10.5-11.5	21		31	
2	SL-502-SA5D-SB-6.5-7.5	12	SL-608-SA5D-SB-0.0-0.5	22		32	
3	SL-508-SA5D-SB-0.0-0.5	13	SL-608-SA5D-SB-4.0-5.0	23		33	
4	SL-508-SA5D-SB-4.0-5.0	14		24		34	
5	SL-508-SA5D-SB-9.0-10.0	15		25		35	
6	SL-508-SA5D-SB-14.0-15.0	16		26		36	
7	SL-508-SA5D-SB-19.0-20.0	17		27		37	
8	SL-508-SA5D-SB-23.0-24.0	18		28		38	
9	SL-507-SA5D-SB-0.0-0.5	19		29		39	
10	SL-507-SA5D-SB-4.0-5.0	20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F
Sampling date: 4/11/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification													
	FB-041613 (SDG: PH032)	Action Limit	1	3	9	12									
Mo	0.0132	6.60	4.5	0.21	0.35	0.18									
Sn	0.0029	1.45													

Sampling date: 10/2/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification													
	EB1-100213 (SDG: PH120)	Action Limit	3	9	12										
Ba	0.00058	0.29													
B	0.0098	4.9													
Ca	0.0427	21.35													
Cu	0.0028	1.4													
Mo	0.0086	4.3	See FB	See FB	See FB										

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH123

Prepared for

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Prepared by

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December 23, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on October 7, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by Environmental Protection Agency (EPA) SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)

Pesticides by EPA SW 846 Method 8081B

Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A

Metals by EPA SW 846 Method 6010C, 6020A and 7471B

Herbicides by EPA SW 846 Method 8151A

Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M

TPH as Extractables by EPA SW 846 Method 8015M

Dioxins and Dibenzofurans by EPA Method 1613B

Wet Chemistry:

Fluoride by EPA Method 300.0

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), method blanks, trip blanks, equipment blanks and field blanks. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for fluoride. The associated sample results were qualified as detected estimated (J). The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution was not performed for this SDG.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH123	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

No field duplicates were identified in this SDG.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No volatile contaminants were found in the trip blank.

One equipment blank (from SDG PH124) was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins and fluoride. The equipment blank had detections for SVOCs, metals and dioxins. The associated sample results were not detected or were significantly greater than the concentrations found in the equipment blanks, therefore no data were qualified.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins and fluoride. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1

Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
07-Oct-2013	TB-100713	7228020	TB	5030B	8015M	III
07-Oct-2013	SL-575-SA5D-SB-0.0-0.5	7228023	N	3546	8270D SIM	III
07-Oct-2013	SL-575-SA5D-SB-0.0-0.5	7228023	N	3550B	8151A	III
07-Oct-2013	SL-575-SA5D-SB-3.0-4.0	7228024	N	3546	8270D SIM	III
07-Oct-2013	SL-575-SA5D-SB-3.0-4.0	7228024	N	3550B	8151A	III
07-Oct-2013	SL-567-SA5D-SB-0.0-0.5	7228021	N	3050B	6010C	III
07-Oct-2013	SL-567-SA5D-SB-0.0-0.5	7228021	N	3050B	6020A	III
07-Oct-2013	SL-567-SA5D-SB-0.0-0.5	7228021	N	3546	8015M	III
07-Oct-2013	SL-567-SA5D-SB-0.0-0.5	7228021	N	3546	8081B	III
07-Oct-2013	SL-567-SA5D-SB-0.0-0.5	7228021	N	3546	8082A	III
07-Oct-2013	SL-567-SA5D-SB-0.0-0.5	7228021	N	3546	8270D SIM	III
07-Oct-2013	SL-567-SA5D-SB-0.0-0.5	7228021	N	3550B	8151A	III
07-Oct-2013	SL-567-SA5D-SB-0.0-0.5	7228021	N	METHOD	1613B	III
07-Oct-2013	SL-567-SA5D-SB-0.0-0.5	7228021	N	METHOD	7471B	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5	7228022	N	3050B	6010C	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5	7228022	N	3050B	6020A	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5	7228022	N	3546	8015M	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5	7228022	N	3546	8081B	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5	7228022	N	3546	8082A	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5	7228022	N	3546	8270D SIM	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5	7228022	N	3550B	8151A	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5	7228022	N	5035A	8015M	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5	7228022	N	METHOD	7471B	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5MSD	P228022M260559	MSD	3546	8270D SIM	III
07-Oct-2013	SL-567-SA5D-SB-3.5-4.5MS	P228022R260527	MS	3546	8270D SIM	III
07-Oct-2013	SL-593-SA5D-SB-0.0-0.5	7228025	N	METHOD	300.0	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
07-Oct-2013	SL-593-SA5D-SB-0.0-0.5DUP	P228025D272355A	DUP	METHOD	300.0	III
07-Oct-2013	SL-593-SA5D-SB-0.0-0.5MS	P228025R270011A	MS	METHOD	300.0	III
07-Oct-2013	SL-593-SA5D-SB-3.5-4.5	7228026	N	METHOD	300.0	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH123

Laboratory: LL

EDD Filename: PrepPH123_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	GENCHEM	
Method:	300.0	Matrix: SO

Sample ID: SL-593-SA5D-SB-0.0-0.5 Collected: 10/7/2013 12:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	0.74		0.41	MDL	0.51	PQL	mg/Kg	J	Q

Sample ID: SL-593-SA5D-SB-3.5-4.5 Collected: 10/7/2013 1:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORIDE	3.4		0.43	MDL	0.54	PQL	mg/Kg	J	Q

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-567-SA5D-SB-0.0-0.5 Collected: 10/7/2013 9:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	3.70	J	0.723	MDL	4.13	PQL	mg/Kg	J	Z
BERYLLIUM	0.665	J	0.0692	MDL	1.03	PQL	mg/Kg	J	Z
CADMIUM	0.701	J	0.0785	MDL	1.03	PQL	mg/Kg	J	Z
MOLYBDENUM	0.600	J	0.176	MDL	2.07	PQL	mg/Kg	U	F
TIN	2.34	J	0.227	MDL	10.3	PQL	mg/Kg	U	B

Sample ID: SL-567-SA5D-SB-3.5-4.5 Collected: 10/7/2013 10:20:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.682	J	0.0712	MDL	1.06	PQL	mg/Kg	J	Z
CADMIUM	0.507	J	0.0807	MDL	1.06	PQL	mg/Kg	J	Z
TIN	2.39	J	0.234	MDL	10.6	PQL	mg/Kg	U	B

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-567-SA5D-SB-0.0-0.5 Collected: 10/7/2013 9:50:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.401	J	0.103	MDL	0.413	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH123

Laboratory: LL

EDD Filename: PrepPH123_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-567-SA5D-SB-0.0-0.5	Collected: 10/7/2013 9:50:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0788	J	0.0269	MDL	0.207	PQL	mg/Kg	J	Z

Sample ID: SL-567-SA5D-SB-3.5-4.5	Collected: 10/7/2013 10:20:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0421	J	0.0276	MDL	0.212	PQL	mg/Kg	J	Z

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-567-SA5D-SB-0.0-0.5	Collected: 10/7/2013 9:50:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	1.15	JB	0.0717	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.118	J	0.0910	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.151	JQ	0.0966	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDF	0.126	JQ	0.0998	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.248	J	0.0968	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.275	JBQ	0.0919	MDL	5.07	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0938	JQ	0.0908	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDF	0.357	J	0.102	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.260	JQ	0.154	MDL	5.07	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	1.54	JB	0.126	MDL	5.07	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HxCDF	0.262	J	0.0937	MDL	5.07	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.237	JQ	0.117	MDL	5.07	PQL	ng/Kg	J	Z
OCDF	2.28	JB	0.104	MDL	10.1	PQL	ng/Kg	J	Z

Method Category:	SVOA	
Method:	8015M	Matrix: SO

Sample ID: SL-567-SA5D-SB-0.0-0.5	Collected: 10/7/2013 9:50:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	2.9	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH123

Laboratory: LL

EDD Filename: PrepPH123_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	SVOA								
Method:	8081B	Matrix:	SO						

Sample ID: SL-567-SA5D-SB-0.0-0.5 Collected: 10/7/2013 9:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
4,4'-DDE	0.86	J	0.34	MDL	1.8	PQL	ug/Kg	J	Z
4,4'-DDT	1.1	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-567-SA5D-SB-3.5-4.5 Collected: 10/7/2013 10:20:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
4,4'-DDE	0.83	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z

Method Category:	SVOA								
Method:	8270D SIM	Matrix:	SO						

Sample ID: SL-567-SA5D-SB-3.5-4.5 Collected: 10/7/2013 10:20:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.88	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
Butylbenzylphthalate	9.6	J	6.5	MDL	20	PQL	ug/Kg	J	Z
CHRYSENE	1.0	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	1.0	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-575-SA5D-SB-0.0-0.5 Collected: 10/7/2013 8:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.81	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(B)FLUORANTHENE	1.0	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
CHRYSENE	1.4	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
DIBENZO(A,H)ANTHRACENE	1.0	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
FLUORANTHENE	0.90	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
NAPHTHALENE	1.1	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
PHENANTHRENE	0.71	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	0.71	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-575-SA5D-SB-3.0-4.0 Collected: 10/7/2013 9:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	0.49	J	0.35	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH123

Laboratory: LL

EDD Filename: PrepPH123_v1

eQAPP Name: CDM_SSFL_131101_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Method Blank Contamination
F	Field Blank Contamination
Q	Matrix Spike Lower Rejection
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH123

Method Blank Outlier Report

Lab Reporting Batch ID: PH123

Laboratory: LL

EDD Filename: PrepPH123_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2830B371446	10/14/2013 2:46:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,6,7,8-HXCDF 1,2,3,7,8-PECDF OCDD OCDF	0.0753 ng/Kg 0.0423 ng/Kg 0.0948 ng/Kg 0.0800 ng/Kg 0.215 ng/Kg 0.137 ng/Kg	SL-567-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-567-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.275 ng/Kg	0.275U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P28137AB221240	10/14/2013 12:40:00 PM	ALUMINUM CALCIUM IRON MAGNESIUM TIN ZINC	9.86 mg/Kg 12.9 mg/Kg 3.84 mg/Kg 2.49 mg/Kg 1.41 mg/Kg 0.656 mg/Kg	SL-567-SA5D-SB-0.0-0.5 SL-567-SA5D-SB-3.5-4.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-567-SA5D-SB-0.0-0.5(RES)	TIN	2.34 mg/Kg	2.34U mg/Kg
SL-567-SA5D-SB-3.5-4.5(RES)	TIN	2.39 mg/Kg	2.39U mg/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH123

Laboratory: LL

EDD Filename: PrepPH123_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C

Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041613(REA2/TOT)	4/16/2013 3:15:00 PM	MOLYBDENUM TIN	0.0132 mg/L 0.0029 mg/L	SL-567-SA5D-SB-0.0-0.5 SL-567-SA5D-SB-3.5-4.5 SL-575-SA5D-SB-0.0-0.5 SL-575-SA5D-SB-3.0-4.0 SL-593-SA5D-SB-0.0-0.5 SL-593-SA5D-SB-3.5-4.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-567-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	0.600 mg/Kg	0.600U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH123

Laboratory: LL

EDD Filename: PrepPH123_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 300.0

Matrix: SO

<i>QC Sample ID (Associated Samples)</i>	<i>Compound</i>	<i>MS %R</i>	<i>MSD %R</i>	<i>%R Limits</i>	<i>RPD (Limits)</i>	<i>Affected Compounds</i>	<i>Flag</i>
SL-593-SA5D-SB-0.0-0.5MS (SL-593-SA5D-SB-0.0-0.5 SL-593-SA5D-SB-3.5-4.5)	FLUORIDE	14	-	80.00-120.00	-	FLUORIDE	J (all detects) R (all non-detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH123

Laboratory: LL

EDD Filename: PrepPH123_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-567-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	1.15	5.07	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	J	0.118	5.07	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.151	5.07	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JQ	0.126	5.07	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	0.248	5.07	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.275	5.07	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JQ	0.0938	5.07	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.357	5.07	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.260	5.07	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	1.54	5.07	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	J	0.262	5.07	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.237	5.07	PQL	ng/Kg	
	OCDF	JB	2.28	10.1	PQL	ng/Kg	

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-567-SA5D-SB-0.0-0.5	ARSENIC	J	3.70	4.13	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.665	1.03	PQL	mg/Kg	
	CADMIUM	J	0.701	1.03	PQL	mg/Kg	
	MOLYBDENUM	J	0.600	2.07	PQL	mg/Kg	
	TIN	J	2.34	10.3	PQL	mg/Kg	
SL-567-SA5D-SB-3.5-4.5	BERYLLIUM	J	0.682	1.06	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.507	1.06	PQL	mg/Kg	
	TIN	J	2.39	10.6	PQL	mg/Kg	

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-567-SA5D-SB-0.0-0.5	SELENIUM	J	0.401	0.413	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0788	0.207	PQL	mg/Kg	
SL-567-SA5D-SB-3.5-4.5	SILVER	J	0.0421	0.212	PQL	mg/Kg	J (all detects)

Method: 8015M
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-567-SA5D-SB-0.0-0.5	EFH (C15-C20)	J	2.9	5.2	PQL	mg/Kg	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH123

Laboratory: LL

EDD Filename: PrepPH123_v1

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8081B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-567-SA5D-SB-0.0-0.5	4,4'-DDE	J	0.86	1.8	PQL	ug/Kg	J (all detects)
	4,4'-DDT	J	1.1	1.8	PQL	ug/Kg	
SL-567-SA5D-SB-3.5-4.5	4,4'-DDE	J	0.83	1.8	PQL	ug/Kg	J (all detects)

Method: 8270D SIM
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-567-SA5D-SB-3.5-4.5	BENZO(B)FLUORANTHENE	J	0.88	1.8	PQL	ug/Kg	J (all detects)
	Butylbenzylphthalate	J	9.6	20	PQL	ug/Kg	
	CHRYSENE	J	1.0	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	1.0	1.8	PQL	ug/Kg	
SL-575-SA5D-SB-0.0-0.5	2-METHYLNAPHTHALENE	J	0.81	1.7	PQL	ug/Kg	J (all detects)
	BENZO(B)FLUORANTHENE	J	1.0	1.7	PQL	ug/Kg	
	CHRYSENE	J	1.4	1.7	PQL	ug/Kg	
	DIBENZO(A,H)ANTHRACENE	J	1.0	1.7	PQL	ug/Kg	
	FLUORANTHENE	J	0.90	1.7	PQL	ug/Kg	
	NAPHTHALENE	J	1.1	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	0.71	1.7	PQL	ug/Kg	
PYRENE	J	0.71	1.7	PQL	ug/Kg		
SL-575-SA5D-SB-3.0-4.0	CHRYSENE	J	0.49	1.7	PQL	ug/Kg	J (all detects)

LDC #: 30842F4

VALIDATION COMPLETENESS WORKSHEET

Date: 12/4/13

SDG #: PH123

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: ea

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	-	Sampling dates: 10/7/13
II.	ICP/MS Tune	-	
III.	Calibration	-	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	N	CS
VII.	Duplicate Sample Analysis	N	CS
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	N	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	N	
XV.	Field Blanks	SW	FB = FB-04/613 EB = EB-100813

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

(PH123) (PH124)

Validated Samples:

Soil

1	SL-567-SA5D-SB-0.0-0.5	11	21	31
2	SL-567-SA5D-SB-3.5-4.5	12	22	32
3	SL-575-SA5D-SB-0.0-0.5	13	23	33
4	SL-575-SA5D-SB-3.0-4.0	14	24	34
5		15	25	35
6		16	26	36
7		17	27	37
8		18	28	38
9		19	29	39
10		20	30	40

Notes: _____

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg **Reason:** F
Sampling date: 4/11/13 **Soil factor applied:** 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ **Associated Samples:** All

Analyte	Blank ID	Sample Identification											
	FB-041613 (SDG: PH032)	Action Limit	1										
Mo	0.0132	6.60	0.60										
Sn	0.0029	1.45											

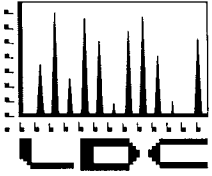
Sampling date: 10/8/13 **Soil factor applied:** 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ **Associated Samples:** All

Analyte	Blank ID	Sample Identification											
	EB-100813 (SDG: PH124)	Action Limit	No Qualifiers										
B	0.0085	4.25											
Ca	0.0487	24.35											

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

SAMPLE DELIVERY GROUP

PH124



LABORATORY DATA CONSULTANTS, INC.
2701 Loker Ave. West, Suite 220, Carlsbad, CA 92010 Bus: 760-827-1100 Fax: 760-827-1099

CDM
555 17th Street, Suite 1100
Denver, CO 80202
ATTN: Mrs. Cherie Zakowski

December 18, 2013

SUBJECT: Santa Susana Field Laboratory, Subarea 5D Data Validation

Dear Mrs. Zakowski,

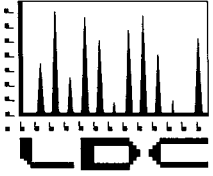
Enclosed is the final validation report for the fractions listed below. This SDG was received on November 19, 2013. Attachment 1 is a summary of the samples that were reviewed for each analysis.

LDC Project # 30854:

<u>SDG #</u>	<u>Fraction</u>
PH124	Semivolatiles, Chlorinated Pesticides, Polychlorinated Biphenyls, Metals, Herbicides, Wet Chemistry, Total Petroleum Hydrocarbons as Gasoline, Total Petroleum Hydrocarbons as Extractables, Dioxins/Dibenzofurans

The data validation was performed under Level III guidelines. The analyses were validated using the following documents, as applicable to each method:

- Quality Assurance Project Plan for Santa Susana Field Laboratory, RCRA Facility Investigation, Surficial Media Operable Unit, March 2009, Revision 4
- USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review, June 2008
- USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review, January 2010
- Polychlorinated Dioxins/Dibenzofurans Data Review, September 2005
- EPA SW 846, Third Edition, Test Methods for Evaluating Solid Waste, update 1, July 1992; update IIA, August 1993; update II, September 1994; update IIB, January 1995; update III, December 1996; update IIIA, April 1998; IIIB, November 2004; Update IV, February 2007



Please feel free to contact us if you have any questions.

Sincerely,

Shauna McKellar
Project Manager/Chemist

90/10 ADR/IV **LDC #30854 (CDM Federal Programs-Chantilly VA / Santa Susana Field Laboratory, Subarea 5D)**

LDC	SDG#	DATE REC'D	(4) DATE DUE	SVOA (8270D -SIM)		Pest. (8081B)		PCBs (8082A)		Metals & Hg (SW846)		Herbs. (8151A)		TPH-G (8015M)		TPH-E (8015M)		Dioxins (1613B)		Cr(VI) (7199)		F (300.0)																							
				W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S						
A	PH124	11/19/13	12/19/13	1	3	1	0	1	3	1	3	1	0	2	2	1	3	1	1	1	0	1	6																						
Total	T/SM			1	3	1	0	1	3	1	3	1	0	2	2	1	3	1	1	1	0	1	6	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	32	

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH124

Prepared for

CDM Smith
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
2701 Loker Ave West, Suite 220
Carlsbad, California 92010

December 18, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on October 8, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by Environmental Protection Agency (EPA) SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)

Pesticides by EPA SW 846 8081B

Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A

Metals by EPA SW 846 Method 6010C, 6020A, 7470A and 7471B

Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M

TPH as Extractables by EPA SW 846 Method 8015M

Dioxins and Dibenzofurans by EPA Method 1613B

Wet Chemistry:

Hexavalent Chromium by EPA Method 7199

Fluoride by EPA Method 300.0

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks and field blanks. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of several blanks for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks with the following exceptions:

SDG/ Method	Method Blank ID	Analyte	Maximum Concentration	Associated Samples
PH124/ 6010C	ICB/CCB	Molybdenum	2.3 ug/L	SL-561-SA5D-SB-0.0-0.5 SL-561-SA5D-SB-4.0-5.0 SL-561-SA5D-SB-9.0-10.0

Sample concentrations were compared to concentrations detected in the initial and continuing blanks. The sample concentrations were not detected or were significantly greater than the concentrations found in the associated blanks with the following exceptions:

SDG/Method	Sample	Compound	Reported Concentration	Modified Final Concentration
PH124/ 6010C	SL-561-SA5D-SB-0.0-0.5	Molybdenum	1.2 mg/Kg	1.2U mg/Kg
PH124/ 6010C	SL-561-SA5D-SB-4.0-5.0	Molybdenum	0.27 mg/Kg	0.27U mg/Kg
PH124/ 6010C	SL-561-SA5D-SB-9.0-10.0	Molybdenum	0.35 mg/Kg	0.35U mg/Kg

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits with the exception of one sample for PCBs. No data were qualified due to high %Rs since the associated results were non-detected.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of two LCS/LCSD pairs for herbicides and TPH as extractables. The associated sample results were qualified as non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met with the following exceptions:

Diluted Sample	Analyte	%D (Limits)	Associated Samples	Flag	A or P
SL-561-SA5D-SB-0.0-0.5	Strontium	16 (≤10)	All soil samples in SDG PH124	J (all detects) UJ (all non-detects)	A

The associated sample results were qualified as detected estimated (J).

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified

requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH124	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

No field duplicates were identified in this SDG.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No volatile contaminants were found in the trip blank.

One equipment blank was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins, hexavalent chromium and fluoride. The equipment blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to equipment blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the equipment blanks were not qualified. The equipment blank outlier reports are presented in Enclosure I.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins, hexavalent chromium and fluoride. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
08-Oct-2013	TB-100813	7229586	TB	5030B	8015M	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5	7229587	N	3050B	6010C	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5	7229587	N	3050B	6020A	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5	7229587	N	3546	8015M	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5	7229587	N	3546	8082A	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5	7229587	N	3546	8270D SIM	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5	7229587	N	METHOD	1613B	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5	7229587	N	METHOD	300.0	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5	7229587	N	METHOD	7471B	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5DUP	P229587D220920	DUP	METHOD	7471B	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5DUP	P229587D221415A	DUP	3050B	6020A	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5DUP	P229587D221415B	DUP	3050B	6020A	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5DUP	P229587D221942	DUP	3050B	6010C	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5MSD	P229587M220507	MSD	3050B	6010C	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5MSD	P229587M220924	MSD	METHOD	7471B	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5MSD	P229587M221420A	MSD	3050B	6020A	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5MSD	P229587M221420B	MSD	3050B	6020A	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5MSD	P229587M240822A	MSD	3546	8082A	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5MS	P229587R220922	MS	METHOD	7471B	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5MS	P229587R221417A	MS	3050B	6020A	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5MS	P229587R221417B	MS	3050B	6020A	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5MS	P229587R221946	MS	3050B	6010C	III
08-Oct-2013	SL-561-SA5D-SB-0.0-0.5MS	P229587R240804A	MS	3546	8082A	III
08-Oct-2013	SL-561-SA5D-SB-4.0-5.0	7229588	N	3050B	6010C	III
08-Oct-2013	SL-561-SA5D-SB-4.0-5.0	7229588	N	3050B	6020A	III
08-Oct-2013	SL-561-SA5D-SB-4.0-5.0	7229588	N	3546	8015M	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
08-Oct-2013	SL-561-SA5D-SB-4.0-5.0	7229588	N	3546	8082A	III
08-Oct-2013	SL-561-SA5D-SB-4.0-5.0	7229588	N	3546	8270D SIM	III
08-Oct-2013	SL-561-SA5D-SB-4.0-5.0	7229588	N	5035A	8015M	III
08-Oct-2013	SL-561-SA5D-SB-4.0-5.0	7229588	N	METHOD	300.0	III
08-Oct-2013	SL-561-SA5D-SB-4.0-5.0	7229588	N	METHOD	7471B	III
08-Oct-2013	SL-561-SA5D-SB-9.0-10.0	7229589	N	3050B	6010C	III
08-Oct-2013	SL-561-SA5D-SB-9.0-10.0	7229589	N	3050B	6020A	III
08-Oct-2013	SL-561-SA5D-SB-9.0-10.0	7229589	N	3546	8015M	III
08-Oct-2013	SL-561-SA5D-SB-9.0-10.0	7229589	N	3546	8082A	III
08-Oct-2013	SL-561-SA5D-SB-9.0-10.0	7229589	N	3546	8270D SIM	III
08-Oct-2013	SL-561-SA5D-SB-9.0-10.0	7229589	N	5035A	8015M	III
08-Oct-2013	SL-561-SA5D-SB-9.0-10.0	7229589	N	METHOD	300.0	III
08-Oct-2013	SL-561-SA5D-SB-9.0-10.0	7229589	N	METHOD	7471B	III
08-Oct-2013	SL-602-SA5D-SB-0.0-0.5	7229590	N	METHOD	300.0	III
08-Oct-2013	SL-602-SA5D-SB-4.0-5.0	7229591	N	METHOD	300.0	III
08-Oct-2013	SL-602-SA5D-SB-9.0-10.0	7229592	N	METHOD	300.0	III
08-Oct-2013	EB-100813	7229585	EB	3005A	6010C	III
08-Oct-2013	EB-100813	7229585	EB	3510C	8015M	III
08-Oct-2013	EB-100813	7229585	EB	3510C	8081B	III
08-Oct-2013	EB-100813	7229585	EB	3510C	8082A	III
08-Oct-2013	EB-100813	7229585	EB	3510C	8270D SIM	III
08-Oct-2013	EB-100813	7229585	EB	5030B	8015M	III
08-Oct-2013	EB-100813	7229585	EB	Gen Prep	300.0	III
08-Oct-2013	EB-100813	7229585	EB	Gen Prep	7199	III
08-Oct-2013	EB-100813	7229585	EB	M3010A	6020A	III
08-Oct-2013	EB-100813	7229585	EB	METHOD	1613B	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
08-Oct-2013	EB-100813	7229585	EB	METHOD	7470A	III
08-Oct-2013	EB-100813	7229585	EB	METHOD	8151A	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6010C	Matrix: AQ

Sample ID: EB-100813	Collected: 10/8/2013 2:00:00	Analysis Type: RES	Dilution: 1
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Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	0.0487	J	0.0334	MDL	0.400	PQL	mg/L	U	B
BORON	0.0085	J	0.0084	MDL	0.100	PQL	mg/L	U	B

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-561-SA5D-SB-0.0-0.5	Collected: 10/8/2013 8:40:00	Analysis Type: REA	Dilution: 1
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Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.12	U	0.763	MDL	4.12	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.993	J	0.0691	MDL	1.03	PQL	mg/Kg	J	Z
BORON	5.60	J	0.866	MDL	10.3	PQL	mg/Kg	J	Z
CHROMIUM	39.0		0.165	MDL	3.09	PQL	mg/Kg	J	Q, E
MOLYBDENUM	1.21	J	0.175	MDL	2.06	PQL	mg/Kg	U	B, F
NICKEL	21.4		0.134	MDL	2.06	PQL	mg/Kg	J	E, Q
SODIUM	83.7	J	17.2	MDL	103	PQL	mg/Kg	J	Z
TIN	3.12	J	0.227	MDL	10.3	PQL	mg/Kg	U	B
Zirconium	3.09	J	0.866	MDL	5.15	PQL	mg/Kg	J	Z

Sample ID: SL-561-SA5D-SB-0.0-0.5	Collected: 10/8/2013 8:40:00	Analysis Type: REA3	Dilution: 1
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Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
VANADIUM	69.3		0.134	MDL	1.03	PQL	mg/Kg	J	Q

Sample ID: SL-561-SA5D-SB-4.0-5.0	Collected: 10/8/2013 9:00:00	Analysis Type: REA	Dilution: 1
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Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.30	U	0.795	MDL	4.30	PQL	mg/Kg	UJ	Q
BERYLLIUM	1.02	J	0.0720	MDL	1.07	PQL	mg/Kg	J	Z
BORON	2.19	J	0.903	MDL	10.7	PQL	mg/Kg	U	F
CHROMIUM	38.1		0.172	MDL	3.22	PQL	mg/Kg	J	Q, E
MOLYBDENUM	0.270	J	0.183	MDL	2.15	PQL	mg/Kg	U	B, F
NICKEL	22.1		0.140	MDL	2.15	PQL	mg/Kg	J	E, Q
SODIUM	86.2	J	18.0	MDL	107	PQL	mg/Kg	J	Z
TIN	3.09	J	0.236	MDL	10.7	PQL	mg/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS		
Method:	6010C	Matrix:	SO

Sample ID: SL-561-SA5D-SB-4.0-5.0		Collected: 10/8/2013 9:00:00		Analysis Type: REA			Dilution: 1		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
Zirconium	2.58	J	0.903	MDL	5.37	PQL	mg/Kg	J	Z

Sample ID: SL-561-SA5D-SB-4.0-5.0		Collected: 10/8/2013 9:00:00		Analysis Type: REA3			Dilution: 1		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
VANADIUM	70.7		0.140	MDL	1.07	PQL	mg/Kg	J	Q

Sample ID: SL-561-SA5D-SB-9.0-10.0		Collected: 10/8/2013 10:10:00		Analysis Type: REA			Dilution: 1		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.26	U	0.788	MDL	4.26	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.910	J	0.0713	MDL	1.06	PQL	mg/Kg	J	Z
BORON	3.77	J	0.894	MDL	10.6	PQL	mg/Kg	U	F
CADMIUM	0.0979	J	0.0809	MDL	1.06	PQL	mg/Kg	J	Z
CHROMIUM	38.9		0.170	MDL	3.19	PQL	mg/Kg	J	Q, E
MOLYBDENUM	0.353	J	0.181	MDL	2.13	PQL	mg/Kg	U	B, F
NICKEL	22.2		0.138	MDL	2.13	PQL	mg/Kg	J	E, Q
TIN	2.94	J	0.234	MDL	10.6	PQL	mg/Kg	U	B
Zirconium	3.30	J	0.894	MDL	5.32	PQL	mg/Kg	J	Z

Sample ID: SL-561-SA5D-SB-9.0-10.0		Collected: 10/8/2013 10:10:00		Analysis Type: REA3			Dilution: 1		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
VANADIUM	67.5		0.138	MDL	1.06	PQL	mg/Kg	J	Q

Method Category:	METALS		
Method:	6020A	Matrix:	SO

Sample ID: SL-561-SA5D-SB-0.0-0.5		Collected: 10/8/2013 8:40:00		Analysis Type: REA			Dilution: 2		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.401	J	0.103	MDL	0.412	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-561-SA5D-SB-0.0-0.5 Collected: 10/8/2013 8:40:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0551	J	0.0268	MDL	0.206	PQL	mg/Kg	J	Z
STRONTIUM	35.1		0.0701	MDL	0.412	PQL	mg/Kg	J	A

Sample ID: SL-561-SA5D-SB-4.0-5.0 Collected: 10/8/2013 9:00:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.297	J	0.107	MDL	0.430	PQL	mg/Kg	J	Z

Sample ID: SL-561-SA5D-SB-4.0-5.0 Collected: 10/8/2013 9:00:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0563	J	0.0279	MDL	0.215	PQL	mg/Kg	J	Z
STRONTIUM	45.7		0.0731	MDL	0.430	PQL	mg/Kg	J	A

Sample ID: SL-561-SA5D-SB-9.0-10.0 Collected: 10/8/2013 10:10:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.257	J	0.106	MDL	0.426	PQL	mg/Kg	J	Z

Sample ID: SL-561-SA5D-SB-9.0-10.0 Collected: 10/8/2013 10:10:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0428	J	0.0277	MDL	0.213	PQL	mg/Kg	J	Z
STRONTIUM	52.3		0.0724	MDL	0.426	PQL	mg/Kg	J	A

Method Category:	SVOA	
Method:	1613B	Matrix: AQ

Sample ID: EB-100813 Collected: 10/8/2013 2:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.806	JBQ	0.170	MDL	9.67	PQL	pg/L	U	B
1,2,3,4,6,7,8-HPCDF	0.770	JB	0.0795	MDL	9.67	PQL	pg/L	U	B
1,2,3,4,7,8,9-HPCDF	0.514	JB	0.0880	MDL	9.67	PQL	pg/L	U	B
1,2,3,4,7,8-HxCDD	0.378	JBQ	0.161	MDL	9.67	PQL	pg/L	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA

Method: 1613B

Matrix: AQ

Sample ID: EB-100813

Collected: 10/8/2013 2:00:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HXCDF	0.564	JBQ	0.124	MDL	9.67	PQL	pg/L	U	B
1,2,3,6,7,8-HXCDD	0.472	JBQ	0.173	MDL	9.67	PQL	pg/L	U	B
1,2,3,6,7,8-HXCDF	0.596	JBQ	0.121	MDL	9.67	PQL	pg/L	U	B
1,2,3,7,8,9-HXCDD	0.382	JBQ	0.156	MDL	9.67	PQL	pg/L	U	B
1,2,3,7,8,9-HXCDF	0.575	JBQ	0.128	MDL	9.67	PQL	pg/L	U	B
1,2,3,7,8-PECDF	0.774	JBQ	0.138	MDL	9.67	PQL	pg/L	U	B
2,3,4,6,7,8-HXCDF	0.603	JBQ	0.120	MDL	9.67	PQL	pg/L	U	B
2,3,4,7,8-PECDF	0.455	JB	0.124	MDL	9.67	PQL	pg/L	U	B
OCDD	1.70	JB	0.201	MDL	19.3	PQL	pg/L	U	B
OCDF	1.54	JB	0.232	MDL	19.3	PQL	pg/L	U	B

Method Category: SVOA

Method: 1613B

Matrix: SO

Sample ID: SL-561-SA5D-SB-0.0-0.5

Collected: 10/8/2013 8:40:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	3.81	JB	0.141	MDL	5.16	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.629	JB	0.0505	MDL	5.16	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.149	JBQ	0.0769	MDL	5.16	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.116	JQ	0.0799	MDL	5.16	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.116	JBQ	0.0762	MDL	5.16	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.163	JQ	0.0840	MDL	5.16	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0913	JQ	0.0688	MDL	5.16	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.219	JQ	0.0817	MDL	5.16	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.179	JB	0.0649	MDL	5.16	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.230	JQ	0.189	MDL	1.03	PQL	ng/Kg	J	Z
OCDF	1.28	JBQ	0.0939	MDL	10.3	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method Category: SVOA
Method: 8015M **Matrix:** AQ

Sample ID: EB-100813 **Collected:** 10/8/2013 2:00:00 **Analysis Type:** RES **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C8-C11)	0.096	U	0.048	MDL	0.096	PQL	mg/L	UJ	L

Method Category: SVOA
Method: 8015M **Matrix:** SO

Sample ID: SL-561-SA5D-SB-4.0-5.0 **Collected:** 10/8/2013 9:00:00 **Analysis Type:** REA **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	5.1	J	2.2	MDL	5.5	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 8270D SIM **Matrix:** AQ

Sample ID: EB-100813 **Collected:** 10/8/2013 2:00:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BIS(2-ETHYLHEXYL)PHTHALATE	0.13	J	0.052	MDL	1.0	PQL	ug/L	U	B
Diethylphthalate	0.29	J	0.052	MDL	1.0	PQL	ug/L	J	Z
Di-n-butylphthalate	0.17	J	0.052	MDL	1.0	PQL	ug/L	J	Z

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-561-SA5D-SB-0.0-0.5 **Collected:** 10/8/2013 8:40:00 **Analysis Type:** RES **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	1.2	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-561-SA5D-SB-0.0-0.5 **Collected:** 10/8/2013 8:40:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ACENAPHTHYLENE	0.42	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(A)ANTHRACENE	1.4	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	1.4	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-561-SA5D-SB-0.0-0.5 Collected: 10/8/2013 8:40:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(G,H,I)PERYLENE	1.4	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	9.4	J	6.3	MDL	19	PQL	ug/Kg	J	Z
INDENO(1,2,3-CD)PYRENE	1.2	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-561-SA5D-SB-4.0-5.0 Collected: 10/8/2013 9:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.75	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	0.54	J	0.37	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	0.93	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-561-SA5D-SB-9.0-10.0 Collected: 10/8/2013 10:10:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BIS(2-ETHYLHEXYL)PHTHALATE	11	J	6.5	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	0.45	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	1.0	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH124
EDD Filename: PH124_v1.

Laboratory: LL
eQAPP Name: CDM_SSFL_131101_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
*#	Professional Judgment
A	ICP Serial Dilution
B	Calibration Blank Contamination
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Equipment Blank Contamination
F	Field Blank Contamination
L	Laboratory Control Spike Lower Estimation
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH124

Method Blank Outlier Report

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2870B372231	10/16/2013 10:31:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF 2,3,7,8-TCDF OCDD OCDF	0.857 pg/L 0.818 pg/L 0.760 pg/L 0.696 pg/L 0.643 pg/L 0.399 pg/L 0.462 pg/L 0.472 pg/L 0.653 pg/L 0.426 pg/L 0.841 pg/L 0.519 pg/L 0.625 pg/L 0.412 pg/L 1.75 pg/L 1.79 pg/L	EB-100813

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB-100813(RES)	1,2,3,4,6,7,8-HPCDD	0.806 pg/L	0.806U pg/L
EB-100813(RES)	1,2,3,4,6,7,8-HPCDF	0.770 pg/L	0.770U pg/L
EB-100813(RES)	1,2,3,4,7,8,9-HPCDF	0.514 pg/L	0.514U pg/L
EB-100813(RES)	1,2,3,4,7,8-HxCDD	0.378 pg/L	0.378U pg/L
EB-100813(RES)	1,2,3,4,7,8-HXCDF	0.564 pg/L	0.564U pg/L
EB-100813(RES)	1,2,3,6,7,8-HXCDD	0.472 pg/L	0.472U pg/L
EB-100813(RES)	1,2,3,6,7,8-HXCDF	0.596 pg/L	0.596U pg/L
EB-100813(RES)	1,2,3,7,8,9-HXCDD	0.382 pg/L	0.382U pg/L
EB-100813(RES)	1,2,3,7,8,9-HXCDF	0.575 pg/L	0.575U pg/L
EB-100813(RES)	1,2,3,7,8-PECDF	0.774 pg/L	0.774U pg/L
EB-100813(RES)	2,3,4,6,7,8-HXCDF	0.603 pg/L	0.603U pg/L
EB-100813(RES)	2,3,4,7,8-PECDF	0.455 pg/L	0.455U pg/L
EB-100813(RES)	OCDD	1.70 pg/L	1.70U pg/L
EB-100813(RES)	OCDF	1.54 pg/L	1.54U pg/L

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2870B371343	10/17/2013 1:43:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HXCDF 2,3,4,6,7,8-HXCDF OCDD OCDF	0.0612 ng/Kg 0.0498 ng/Kg 0.101 ng/Kg 0.0571 ng/Kg 0.0492 ng/Kg 0.214 ng/Kg 0.191 ng/Kg	SL-561-SA5D-SB-0.0-0.5

Method Blank Outlier Report

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-561-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.149 ng/Kg	0.149U ng/Kg
SL-561-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.116 ng/Kg	0.116U ng/Kg
SL-561-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.179 ng/Kg	0.179U ng/Kg

Method: 6010C
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P29035AB222022	10/18/2013 8:22:00 PM	ALUMINUM BORON CALCIUM MAGNESIUM ZINC	0.0833 mg/L 0.0099 mg/L 0.175 mg/L 0.0196 mg/L 0.0086 mg/L	EB-100813

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB-100813(RES)	BORON	0.0085 mg/L	0.0085U mg/L
EB-100813(RES)	CALCIUM	0.0487 mg/L	0.0487U mg/L

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P29037AB221926	10/19/2013 7:26:00 PM	CALCIUM POTASSIUM TIN	3.41 mg/Kg 10.0 mg/Kg 1.49 mg/Kg	SL-561-SA5D-SB-0.0-0.5 SL-561-SA5D-SB-4.0-5.0 SL-561-SA5D-SB-9.0-10.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-561-SA5D-SB-0.0-0.5(REA)	TIN	3.12 mg/Kg	3.12U mg/Kg
SL-561-SA5D-SB-4.0-5.0(REA)	TIN	3.09 mg/Kg	3.09U mg/Kg
SL-561-SA5D-SB-9.0-10.0(REA)	TIN	2.94 mg/Kg	2.94U mg/Kg

Method: 8270D SIM
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
PLKWB28B261827	10/14/2013 6:27:00 PM	BIS(2-ETHYLHEXYL)PHTHALATE	0.083 ug/L	EB-100813

Method Blank Outlier Report

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8270D SIM
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB-100813(RES)	BIS(2-ETHYLHEXYL)PHTHALATE	0.13 ug/L	1.0U ug/L

Equipment Rinsate Blank Outlier Report

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

Equipment Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
EB-100813(RES)	10/8/2013 2:00:00 PM	BORON CALCIUM	0.0085 mg/L 0.0487 mg/L	SL-561-SA5D-SB-0.0-0-0.5 SL-561-SA5D-SB-4.0-5.0 SL-561-SA5D-SB-9.0-10.0 SL-602-SA5D-SB-0.0-0-0.5 SL-602-SA5D-SB-4.0-5.0 SL-602-SA5D-SB-9.0-10.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-561-SA5D-SB-4.0-5.0(REA)	BORON	2.19 mg/Kg	2.19U mg/Kg
SL-561-SA5D-SB-9.0-10.0(REA)	BORON	3.77 mg/Kg	3.77U mg/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041613(REA2/TOT)	4/16/2013 3:15:00 PM	MOLYBDENUM TIN	0.0132 mg/L 0.0029 mg/L	SL-561-SA5D-SB-0.0-0.5 SL-561-SA5D-SB-4.0-5.0 SL-561-SA5D-SB-9.0-10.0 SL-602-SA5D-SB-0.0-0.5 SL-602-SA5D-SB-4.0-5.0 SL-602-SA5D-SB-9.0-10.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-561-SA5D-SB-0.0-0.5(REA)	MOLYBDENUM	1.21 mg/Kg	1.21U mg/Kg
SL-561-SA5D-SB-4.0-5.0(REA)	MOLYBDENUM	0.270 mg/Kg	0.270U mg/Kg
SL-561-SA5D-SB-9.0-10.0(REA)	MOLYBDENUM	0.353 mg/Kg	0.353U mg/Kg

Surrogate Outlier Report

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8082A

Matrix: SO

<i>Sample ID (Analysis Type)</i>	<i>Surrogate</i>	<i>Sample % Recovery</i>	<i>% Recovery Limits</i>	<i>Affected Compounds</i>	<i>Flag</i>
SL-561-SA5D-SB-4.0-5.0	DECACHLOROBIPHENYL	123	45.00-120.00	All Target Analytes	J (all detects)

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-561-SA5D-SB-0.0-0.5MS (TOT)	ALUMINUM	4351	5460	75.00-125.00	-	ALUMINUM	J (all detects) Al, Ca, Fe, Mg, Mn, K, Ti, No Qual, >4x
SL-561-SA5D-SB-0.0-0.5MSD (TOT)	CALCIUM	-	149	75.00-125.00	-	CALCIUM	
(SL-561-SA5D-SB-0.0-0.5)	IRON	1046	2202	75.00-125.00	-	IRON	
SL-561-SA5D-SB-4.0-5.0	MAGNESIUM	404	534	75.00-125.00	-	MAGNESIUM	
SL-561-SA5D-SB-9.0-10.0)	MANGANESE	-	157	75.00-125.00	-	MANGANESE	
	POTASSIUM	147	186	75.00-125.00	-	POTASSIUM	
	TITANIUM	488	698	75.00-125.00	-	TITANIUM	
	VANADIUM	-	136	75.00-125.00	-	VANADIUM	
SL-561-SA5D-SB-0.0-0.5MS (TOT)	ANTIMONY	57	60	75.00-125.00	-	ANTIMONY	J(all detects) UJ(all non-detects) P, No Qual, >4x
SL-561-SA5D-SB-0.0-0.5MSD (TOT)	CHROMIUM	673	134	75.00-125.00	91 (20.00)	CHROMIUM	
(SL-561-SA5D-SB-0.0-0.5)	NICKEL	188	-	75.00-125.00	49 (20.00)	NICKEL	
SL-561-SA5D-SB-4.0-5.0	PHOSPHORUS	40	57	75.00-125.00	-	PHOSPHORUS	
SL-561-SA5D-SB-9.0-10.0)							

Method: 6020A

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-561-SA5D-SB-0.0-0.5MS (TOT)	STRONTIUM	62	72	75.00-125.00	-	STRONTIUM	No Qual, >4x
SL-561-SA5D-SB-0.0-0.5MSD (TOT)							
(SL-561-SA5D-SB-0.0-0.5)							
SL-561-SA5D-SB-4.0-5.0							
SL-561-SA5D-SB-9.0-10.0)							

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-561-SA5D-SB-0.0-0.5DUP (TOT) (SL-561-SA5D-SB-0.0-0.5 SL-561-SA5D-SB-4.0-5.0 SL-561-SA5D-SB-9.0-10.0)	CADMIUM MOLYBDENUM	200 77	20.00 20.00	No Qual, OK by Difference

Method: 7471B
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-561-SA5D-SB-0.0-0.5DUP (TOT) (SL-561-SA5D-SB-0.0-0.5 SL-561-SA5D-SB-4.0-5.0 SL-561-SA5D-SB-9.0-10.0)	MERCURY	35	20.00	No Qual, OK by Difference

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8151A

Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P32827AQ241903A P32827AY241930A (EB-100813)	DINOSEB	238	238	16.00-163.00	-	DINOSEB	J (all detects)

Method: 8015M

Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P32870AQ320635A (EB-100813)	EFH (C8-C11)	69	-	70.00-130.00	-	EFH (C8-C11)	J(all detects) UJ(all non-detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 1613B

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB-100813	1,2,3,4,6,7,8-HPCDD	JBQ	0.806	9.67	PQL	pg/L	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.770	9.67	PQL	pg/L	
	1,2,3,4,7,8,9-HPCDF	JB	0.514	9.67	PQL	pg/L	
	1,2,3,4,7,8-HxCDD	JBQ	0.378	9.67	PQL	pg/L	
	1,2,3,4,7,8-HXCDF	JBQ	0.564	9.67	PQL	pg/L	
	1,2,3,6,7,8-HXCDD	JBQ	0.472	9.67	PQL	pg/L	
	1,2,3,6,7,8-HXCDF	JBQ	0.596	9.67	PQL	pg/L	
	1,2,3,7,8,9-HXCDD	JBQ	0.382	9.67	PQL	pg/L	
	1,2,3,7,8,9-HXCDF	JBQ	0.575	9.67	PQL	pg/L	
	1,2,3,7,8-PECDF	JBQ	0.774	9.67	PQL	pg/L	
	2,3,4,6,7,8-HXCDF	JBQ	0.603	9.67	PQL	pg/L	
	2,3,4,7,8-PECDF	JB	0.455	9.67	PQL	pg/L	
	OCDD	JB	1.70	19.3	PQL	pg/L	
	OCDF	JB	1.54	19.3	PQL	pg/L	

Method: 6010C

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB-100813	BORON	J	0.0085	0.100	PQL	mg/L	J (all detects)
	CALCIUM	J	0.0487	0.400	PQL	mg/L	

Method: 8270D SIM

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB-100813	BIS(2-ETHYLHEXYL)PHTHALATE	J	0.13	1.0	PQL	ug/L	J (all detects)
	Diethylphthalate	J	0.29	1.0	PQL	ug/L	
	Di-n-butylphthalate	J	0.17	1.0	PQL	ug/L	

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	3.81	5.16	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.629	5.16	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.149	5.16	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.116	5.16	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.116	5.16	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.163	5.16	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JQ	0.0913	5.16	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JQ	0.219	5.16	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.179	5.16	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.230	1.03	PQL	ng/Kg	
	OCDF	JBQ	1.28	10.3	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.993	1.03	PQL	mg/Kg	J (all detects)
	BORON	J	5.60	10.3	PQL	mg/Kg	
	MOLYBDENUM	J	1.21	2.06	PQL	mg/Kg	
	SODIUM	J	83.7	103	PQL	mg/Kg	
	TIN	J	3.12	10.3	PQL	mg/Kg	
	Zirconium	J	3.09	5.15	PQL	mg/Kg	
SL-561-SA5D-SB-4.0-5.0	BERYLLIUM	J	1.02	1.07	PQL	mg/Kg	J (all detects)
	BORON	J	2.19	10.7	PQL	mg/Kg	
	MOLYBDENUM	J	0.270	2.15	PQL	mg/Kg	
	SODIUM	J	86.2	107	PQL	mg/Kg	
	TIN	J	3.09	10.7	PQL	mg/Kg	
	Zirconium	J	2.58	5.37	PQL	mg/Kg	
SL-561-SA5D-SB-9.0-10.0	BERYLLIUM	J	0.910	1.06	PQL	mg/Kg	J (all detects)
	BORON	J	3.77	10.6	PQL	mg/Kg	
	CADIUM	J	0.0979	1.06	PQL	mg/Kg	
	MOLYBDENUM	J	0.353	2.13	PQL	mg/Kg	
	TIN	J	2.94	10.6	PQL	mg/Kg	
	Zirconium	J	3.30	5.32	PQL	mg/Kg	

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA5D-SB-0.0-0.5	SELENIUM	J	0.401	0.412	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0551	0.206	PQL	mg/Kg	
SL-561-SA5D-SB-4.0-5.0	SELENIUM	J	0.297	0.430	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0563	0.215	PQL	mg/Kg	
SL-561-SA5D-SB-9.0-10.0	SELENIUM	J	0.257	0.426	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0428	0.213	PQL	mg/Kg	

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA5D-SB-4.0-5.0	EFH (C21-C30)	J	5.1	5.5	PQL	mg/Kg	J (all detects)

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA5D-SB-0.0-0.5	1-METHYLNAPHTHALENE	J	1.2	1.8	PQL	ug/Kg	J (all detects)
	ACENAPHTHYLENE	J	0.42	1.8	PQL	ug/Kg	
	BENZO(A)ANTHRACENE	J	1.4	1.8	PQL	ug/Kg	
	BENZO(A)PYRENE	J	1.4	1.8	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	1.4	1.8	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	9.4	19	PQL	ug/Kg	
	INDENO(1,2,3-CD)PYRENE	J	1.2	1.8	PQL	ug/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH124

Laboratory: LL

EDD Filename: PH124_v1.

eQAPP Name: CDM_SSFL_131101_Lan

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA5D-SB-4.0-5.0	2-METHYLNAPHTHALENE	J	0.75	1.8	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	0.54	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	0.93	1.8	PQL	ug/Kg	
SL-561-SA5D-SB-9.0-10.0	BIS(2-ETHYLHEXYL)PHTHALATE	J	11	19	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	0.45	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	1.0	1.8	PQL	ug/Kg	

LDC #: 30854A4

VALIDATION COMPLETENESS WORKSHEET

Date: 12/4/13

SDG #: PH124

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: *al*

2nd Reviewer: *✓*

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	-	Sampling dates: 10/8/13
II.	ICP/MS Tune	-	
III.	Calibration	-	
IV.	Blanks	SW	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	SW	SW ^a MS/P
VII.	Duplicate Sample Analysis	SW	DUP
VIII.	Laboratory Control Samples (LCS)	N	LCS/D
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	SW	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	N	
XV.	Field Blanks	SW	FB = FB-041613 EB = 1

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet
 ND = No compounds detected
 R = Rinsate
 FB = Field blank
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples:

1	EB-100813	11		21		31	
2	SL-561-SA5D-SB-0.0-0.5	12		22		32	
3	SL-561-SA5D-SB-4.0-5.0	13		23		33	
4	SL-561-SA5D-SB-9.0-10.0	14		24		34	
5	(MS)	15		25		35	
6	MSD	16		26		36	
7	DUP	17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

LDC #: 30854A4

VALIDATION FINDINGS WORKSHEET
PB/ICB/CCB QUALIFIED SAMPLES

Page: 1 of 1
Reviewer:
2nd Reviewer:

METHOD: Trace metals (EPA SW 864 Method 6010B/6020/7000)

Soil preparation factor applied: 100x Reason: B

Sample Concentration units, unless otherwise noted: mg/Kg Associated Samples: All Soil

					Sample Identification												
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/l)	Maximum ICB/CCB ^a (ug/l)	Action Level	2	3	4										
Mo			2.3	2.3	1.21	0.270	0.353										

Samples with analyte concentrations within five times the associated ICB, CCB or PB concentration are listed above with the identifications from the Validation Completeness Worksheet. These sample results were qualified as not detected, "U".

Note : a - The listed analyte concentration is the highest ICB, CCB, or PB detected in the analysis of each element.

VALIDATION FINDINGS WORKSHEET Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All Soil

Analyte	Blank ID	Sample Identification											
	FB-041613 (SDG: PH032)	Action Limit	2	3	4								
Mo	0.0132	6.60	1.21	0.270	0.353								
Sn	0.0029	1.45											

Sampling date: 10/8/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All Soil

Analyte	Blank ID	Sample Identification											
	EB-100813 (SDG: PH124)	Action Limit	3	4									
B	0.0085	4.25	2.19	3.87									
Ca	0.0487	24.35											

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".



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QUALITY ASSURANCE SUMMARY
FORM 5A (MS/MSD)
MATRIX SPIKE/MATRIX SPIKE DUPLICATE
SDG No.: PH124
Matrix: SOIL Level: LOW
(low/med):

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Background Lab Sample ID: 7229587BKG Matrix Spike Lab Sample ID: 7229587MS Matrix Spike Duplicate Lab Sample ID: 7229587MSD
Batch Id(s): P29037A, P29038B

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Control Limit		M
		Result	C	Result	C	Result	C				%R	Q	%R	Q		%R	RPD	
Aluminum		25591.2608		34121.9500		36296.3706		196.0784	196.0784	MG/KG	4351	Q	5460	Q	6		20	P
Antimony		0.7255	U	27.7324		29.2971		49.0196	49.0196	MG/KG	57	N	60	N	5	75 - 125	20	P
Arsenic		4.4794		19.1069		20.0843		14.7059	14.7059	MG/KG	99		106		5	75 - 125	20	P
Barium		110.1373		300.2284		307.0637		196.0784	196.0784	MG/KG	97		100		2	75 - 125	20	P
Beryllium		0.9441	B	5.9873		6.1382		4.9020	4.9020	MG/KG	103		106		2	75 - 125	20	P
Boron		5.3225	B	184.1667		189.5020		196.0784	196.0784	MG/KG	91		94		3	75 - 125	20	P
Cadmium		0.0745	U	4.6941		4.7314		4.9020	4.9020	MG/KG	96		97		1	75 - 125	20	P
Calcium		7956.3961		8279.1020		8539.4108		392.1569	392.1569	MG/KG	82		149		3		20	P
Chromium		37.0588		168.9833		63.3735		19.6078	19.6078	MG/KG	673	N	134	N	91	* 75 - 125	20	P
Cobalt		9.7833		57.0735		56.7245		49.0196	49.0196	MG/KG	96		96		1	75 - 125	20	P
Copper		26.9588		54.2196		56.2029		24.5098	24.5098	MG/KG	111		119		4	75 - 125	20	P
Iron		34653.5196		35679.2598		36812.0098		98.0392	98.0392	MG/KG	1046		2202		3		20	P
Lead		14.1608		27.5941		28.1637		14.7059	14.7059	MG/KG	91		95		2	75 - 125	20	P
Lithium		24.1980		125.2588		127.6196		98.0392	98.0392	MG/KG	103		105		2	75 - 125	20	P
Magnesium		7593.8775		8385.0941		8641.4755		196.0784	196.0784	MG/KG	404		534		3		20	P
Manganese		416.2735		468.7333		493.2176		49.0196	49.0196	MG/KG	107		157		5		20	P
Mercury		0.0210		0.1884		0.1910		0.1599	0.1638	MG/KG	105		104		1	65 - 135	20	CV
Molybdenum		1.1471	B	182.8078		186.4892		196.0784	196.0784	MG/KG	93		95		2	75 - 125	20	P
Nickel		20.3608		112.4941		68.2010		49.0196	49.0196	MG/KG	188	N	98		49	* 75 - 125	20	P
Phosphorus		717.0196		756.7137		772.6059		98.0392	98.0392	MG/KG	40		57		2		20	P
Potassium		6613.1824		8055.7598		8440.7647		980.3922	980.3922	MG/KG	147		186		5		20	P
Selenium	78	0.3810	B	2.3084		2.3759		1.9608	1.9608	MG/KG	98		102		3	75 - 125	20	MS
Silver	107	0.0524	B	10.6225		10.7278		9.8039	9.8039	MG/KG	108		109		1	75 - 125	20	MS
Sodium		79.5971	B	1034.1971		1048.2392		980.3922	980.3922	MG/KG	97		99		1	75 - 125	20	P
Strontium	88	33.3455		38.2090		38.9669		7.8431	7.8431	MG/KG	62		72		2		20	MS
Thallium	203	0.3982		0.7616		0.8090		0.3922	0.3922	MG/KG	93		105		6	75 - 125	20	MS

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry	CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL, B= Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS
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QUALITY ASSURANCE SUMMARY
 FORM 5A (MS/MSD)
 MATRIX SPIKE/MATRIX SPIKE DUPLICATE
 SDG No.: PH124
 Matrix: SOIL Level (low/med): LOW

Background Lab Sample ID: 7229587BKG Matrix Spike Lab Sample ID: 7229587MS Matrix Spike Duplicate Lab Sample ID: 7229587MSD
 Batch Id(s): P29037A, P29038B

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Q	Control Limit			
		Result	C	Result	C	Result	C				%R	Q	%R	Q			%R	RPD	M	
Tin		2.9696	B	357.6088		360.3108		392.1569	392.1569	MG/KG	90		91		1		75 - 125	20	P	
Titanium		1178.6010		1657.3804		1862.8157		98.0392	98.0392	MG/KG	488		698		12				20	P
Vanadium		65.9098		126.9431		132.6049		49.0196	49.0196	MG/KG	125		136	N	4		75 - 125	20	P	
Zinc		83.7510		132.7069		135.5176		49.0196	49.0196	MG/KG	100		106		2		75 - 125	20	P	
Zirconium		2.9431	B	87.0853		86.4912		98.0392	98.0392	MG/KG	86		85		1		75 - 125	20	P	

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL, B= Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS
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QUALITY ASSURANCE SUMMARY

FORM 6

DUPLICATES

SDG No.: PH124

Matrix: SOIL Level (low/med): LOW

Background Lab Sample ID: 7229587BKG

Duplicate Lab Sample ID: 7229587DUP

Batch ID(s): P29037A, P29038B

Concentration Units: MG/KG

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			25591.2608		26309.8275		3		P
Antimony			-1.8167	B	-2.0225	B	-11		P
Arsenic		3.9	4.4794		5.1392		14		P
Barium			110.1373		110.6559		0		P
Beryllium			0.9441	B	0.9510	B	1		P
Boron			5.3225	B	5.3118	B	0		P
Cadmium			0.0745	B	0.1088	B	200		P
Calcium			7956.3961		7913.3667		1		P
Chromium			37.0588		37.8667		2		P
Cobalt			9.7833		10.0745		3		P
Copper			26.9588		27.1029		1		P
Iron			34653.5196		34449.1667		1		P
Lead		2.9	14.1608		13.3049		6		P
Lithium			24.1980		24.7441		2		P
Magnesium			7593.8775		7662.7618		1		P
Manganese			416.2735		421.9382		1		P
Mercury		0.0	0.0240	B	0.0147	B	35		CV
Molybdenum			1.1471	B	0.5108	B	77		P
Nickel			20.3608		20.8559		2		P
Phosphorus			717.0196		662.8794		8		P
Potassium			6613.1824		6597.1265		0		P
Selenium	78		0.3810	B	0.3894	B	2		MS
Silver	107		0.0524	B	0.0555	B	6		MS
Sodium			79.5971	B	78.8814	B	1		P
Strontium	88		33.3455		32.3031		3		MS
Thallium	203	0.2	0.3982		0.3659		8		MS
Tin			2.9696	B	2.9755	B	0		P
Titanium			1178.6010		1227.5245		4		P
Vanadium			65.9098		67.0647		2		P
Zinc			83.7510		83.2637		1		P
Zirconium			2.9431	B	3.5784	B	19		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).

The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

ok by difference

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer</p> <p>MS = ICP Mass Spectrometry</p> <p>CV = Cold Vapor</p> <p>AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U= Below MDL</p> <p>B= Below LOQ</p> <p>FLAGS:</p> <p>Duplicate Out of Spec</p>
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Background Lab Sample ID: 7229587BKG
 Batch ID(s): P29037A
 Concentration Units: UG/L

Serial Dilution Lab Sample ID: 7229587L
 SL-561-SASD-SB-00-01

Analyte	Mass	Initial Sample Result (I)	C	Serial Dilution Result (S)	C	% Diff.	Q	M
Aluminum		261030.8600		267886.3000		3		P
Antimony		7.4000	U	37.0000	U			P
Arsenic		45.6900		62.7000	B	37		P
Barium		1123.4000		1184.1500		5		P
Beryllium		9.6300	B	9.2500	B	4		P
Boron		54.2900	B	168.8000	B	211		P
Cadmium		0.7600	U	5.5500	B	100		P
Calcium		81155.2400		84618.3000		4		P
Chromium		378.0000		384.4500		2		P
Cobalt		99.7900		108.8000		9		P
Copper		274.9800		270.2000		2		P
Iron		70693.1800		72837.8500		3		P
Lead		144.4400		168.2000		16		P
Lithium		246.8200		254.2500		3		P
Magnesium		77457.5500		79468.4500		3		P
Manganese		4245.9900		4487.0500		6		P
Molybdenum		11.7000	B	29.3000	B	150		P
Nickel		207.6800		227.4500		10		P
Phosphorus		7313.6000		7355.7000		1		P
Potassium		67454.4600		70764.0500		5		P
Selenium	78	1.9430	B	2.5000	U	100		MS
Silver	107	0.2670	B	0.6500	U	100		MS
Sodium		811.8900	B	835.0000	U	100		P
Strontium	88	170.0620		142.7650		16		MS
Thallium	203	2.0310		2.0000	B	2		MS
Tin		30.2900	B	32.0000	B	6		P
Titanium		12021.7300		11814.7500		2		P
Vanadium		672.2800		685.7500		2		P
Zinc		854.2600		883.4000		3		P
Zirconium		30.0200	B	42.0000	U	100		P

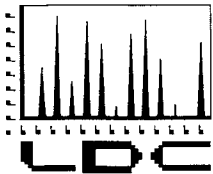
NOTE: An E in column Q indicates the presence of a chemical or physical interference in the matrix when the % difference is greater than 10%. This applies only when (I) is greater than or equal to 50x MDL for ICP, 100x MDL for ICP-MS (6020), 50x MDL for ICP-MS (200.8), or 25x MDL for GFAA.

(Handwritten signature)

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: E = Matrix Effects exist as proven by Serial Dilution or Spiked Dilution
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SAMPLE DELIVERY GROUP

PH127



CDM
555 17th Street, Suite 1100
Denver, CO 80202
ATTN: Mrs. Cherie Zakowski

January 24, 2014

SUBJECT: Santa Susana Field Laboratory, Subarea 5D Data Validation

Dear Mrs. Zakowski,

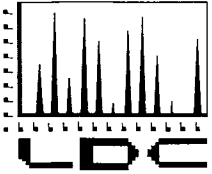
Enclosed is the final validation report for the fractions listed below. This SDG was received on December 31, 2013. Attachment 1 is a summary of the samples that were reviewed for each analysis.

LDC Project # 31098:

<u>SDG #</u>	<u>Fraction</u>
PH127	Semivolatiles, Polynuclear Aromatic Hydrocarbons, Chlorinated Pesticides, Polychlorinated Biphenyls, Metals, Herbicides, Wet Chemistry, Total Petroleum Hydrocarbons as Gasoline, Total Petroleum Hydrocarbons as Extractables, Dioxins/Dibenzofurans

The data validation was performed under Level III guidelines. The analyses were validated using the following documents, as applicable to each method:

- Quality Assurance Project Plan for Santa Susana Field Laboratory, RCRA Facility Investigation, Surficial Media Operable Unit, March 2009, Revision 4
- USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review, June 2008
- USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review, January 2010
- Polychlorinated Dioxins/Dibenzofurans Data Review, September 2005
- EPA SW 846, Third Edition, Test Methods for Evaluating Solid Waste, update 1, July 1992; update IIA, August 1993; update II, September 1994; update IIB, January 1995; update III, December 1996; update IIIA, April 1998; IIIB, November 2004; Update IV, February 2007



Please feel free to contact us if you have any questions.

Sincerely,

A handwritten signature in black ink, appearing to read 'Shauna McKellar', written in a cursive style.

Shauna McKellar
Project Manager/Chemist

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH127

Prepared for

CDM Smith
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
2701 Loker Ave West, Suite 220
Carlsbad, California 92010

January 24, 2014

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on November 12, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan (QAPP) for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by Environmental Protection Agency (EPA) SW 846 Method 8270D
SVOCs by EPA SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Pesticides by EPA SW 846 Method 8081B
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Herbicides by EPA SW 846 Method 8151A
Metals by EPA SW 846 Method 6010C, 6020A, 7470A and 7471B
Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M
TPH as Extractables by EPA SW 846 Method 8015M
Dioxins and Dibenzofurans by EPA Method 1613B

Wet Chemistry:

Hexavalent Chromium by EPA Method 7199

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks and field duplicate samples. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of several blanks for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks with the following exceptions:

SDG/ Method	Method Blank ID	Analyte	Maximum Concentration	Associated Samples
PH127/ 6010C	ICB/CCB	Magnesium	34.0 ug/L	EB1-111213

Sample concentrations were compared to concentrations detected in the initial and continuing blanks. The sample concentrations were not detected or were significantly greater than the concentrations found in the associated blanks with the following exceptions:

SDG/Method	Sample	Compound	Reported Concentration	Modified Final Concentration
PH127/ 6010C	EB1-111213	Magnesium	.0168 mg/L	0.0168 mg/L

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits with the exception of one sample for TPH as extractables. No data were qualified due to high %Rs since the associated results were non-detected.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for SVOCs and metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of several LCS/LCSD pairs for herbicides and TPH as extractables. The associated sample results were qualified as non-detected estimated (UJ). The details regarding the qualification of data are provided in Enclosure I.

The lab indicated that SVOC analytes 3,5-dimethylphenol and 2,6-dichlorophenol were not spiked in the LCS/LCSD samples in the preparation batch containing sample EB1-111213. Since any re-analysis would have occurred outside of the method holding time, these analytes were reported for the sample from the original run. The associated sample results were qualified as non-detected estimated (UJ).

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH127	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pair was collected and analyzed for SVOCs, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables and hexavalent chromium. All RPDs were within QC limits with the exception of one metal and TPH as gasoline. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No contaminants were found in the trip blank.

One equipment blank was collected and analyzed for SVOCs, pesticides, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins and hexavalent chromium. The equipment blank had detections for SVOCs, dioxins, and metals. The sample results were not detected or were significantly greater than the concentrations found in the equipment blank, therefore no data were qualified.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, pesticides, PCBs, herbicides, metals, TPH as gasoline, TPH as extractables, dioxins, and hexavalent chromium. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1

Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
12-Nov-2013	TB1-111213	7275675	TB	5030B	8015M	III
12-Nov-2013	SL-560-SA5D-SB-0.0-0.5	7275662	N	3050B	6010C	III
12-Nov-2013	SL-560-SA5D-SB-0.0-0.5	7275662	N	3050B	6020A	III
12-Nov-2013	SL-560-SA5D-SB-0.0-0.5	7275662	N	3060A	7199	III
12-Nov-2013	SL-560-SA5D-SB-0.0-0.5	7275662	N	3546	8015M	III
12-Nov-2013	SL-560-SA5D-SB-0.0-0.5	7275662	N	3546	8082A	III
12-Nov-2013	SL-560-SA5D-SB-0.0-0.5	7275662	N	3546	8270D	III
12-Nov-2013	SL-560-SA5D-SB-0.0-0.5	7275662	N	3546	8270D SIM	III
12-Nov-2013	SL-560-SA5D-SB-0.0-0.5	7275662	N	3550B	8151A	III
12-Nov-2013	SL-560-SA5D-SB-0.0-0.5	7275662	N	METHOD	1613B	III
12-Nov-2013	SL-560-SA5D-SB-0.0-0.5	7275662	N	METHOD	7471B	III
12-Nov-2013	SL-860-SA5D-SB-4.0-5.0	7275671	FD	3050B	6010C	III
12-Nov-2013	SL-860-SA5D-SB-4.0-5.0	7275671	FD	3050B	6020A	III
12-Nov-2013	SL-860-SA5D-SB-4.0-5.0	7275671	FD	3060A	7199	III
12-Nov-2013	SL-860-SA5D-SB-4.0-5.0	7275671	FD	3546	8015M	III
12-Nov-2013	SL-860-SA5D-SB-4.0-5.0	7275671	FD	3546	8082A	III
12-Nov-2013	SL-860-SA5D-SB-4.0-5.0	7275671	FD	3546	8270D	III
12-Nov-2013	SL-860-SA5D-SB-4.0-5.0	7275671	FD	3546	8270D SIM	III
12-Nov-2013	SL-860-SA5D-SB-4.0-5.0	7275671	FD	3550B	8151A	III
12-Nov-2013	SL-860-SA5D-SB-4.0-5.0	7275671	FD	5035A	8015M	III
12-Nov-2013	SL-860-SA5D-SB-4.0-5.0	7275671	FD	METHOD	7471B	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0	7275663	N	3050B	6010C	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0	7275663	N	3050B	6020A	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0	7275663	N	3060A	7199	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0	7275663	N	3546	8015M	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0	7275663	N	3546	8082A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0	7275663	N	3546	8270D	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0	7275663	N	3546	8270D SIM	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0	7275663	N	3550B	8151A	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0	7275663	N	5035A	8015M	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0	7275663	N	METHOD	7471B	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275664	MS	3050B	6010C	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275664	MS	3050B	6020A	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275664	MS	3546	8015M	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275664	MS	3546	8082A	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275664	MS	3546	8270D	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275664	MS	3546	8270D SIM	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275664	MS	3550B	8151A	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275664	MS	5035A	8015M	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275664	MS	METHOD	7471B	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MSD	7275665	MSD	3050B	6010C	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MSD	7275665	MSD	3050B	6020A	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MSD	7275665	MSD	3546	8015M	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MSD	7275665	MSD	3546	8082A	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MSD	7275665	MSD	3546	8270D	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MSD	7275665	MSD	3546	8270D SIM	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MSD	7275665	MSD	3550B	8151A	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MSD	7275665	MSD	5035A	8015M	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MSD	7275665	MSD	METHOD	7471B	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275666	MS	3060A	7199	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0MS	7275667	MS	3060A	7199	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0DUP	7275669	DUP	3050B	6010C	III

III = EPA Level 3 Data Review
IV = EPA Level 4 Data Validation

N = Normal Sample
FD = Field Duplicate

TB = Trip Blank
FB = Field Blank

MS = Matrix Spike
MSD = Matrix Spike Duplicate

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0DUP	7275669	DUP	3050B	6020A	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0DUP	7275669	DUP	3060A	7199	III
12-Nov-2013	SL-560-SA5D-SB-4.0-5.0DUP	7275669	DUP	METHOD	7471B	III
12-Nov-2013	SL-560-SA5D-SB-7.0-8.0	7275670	N	3050B	6010C	III
12-Nov-2013	SL-560-SA5D-SB-7.0-8.0	7275670	N	3050B	6020A	III
12-Nov-2013	SL-560-SA5D-SB-7.0-8.0	7275670	N	3060A	7199	III
12-Nov-2013	SL-560-SA5D-SB-7.0-8.0	7275670	N	3546	8015M	III
12-Nov-2013	SL-560-SA5D-SB-7.0-8.0	7275670	N	3546	8082A	III
12-Nov-2013	SL-560-SA5D-SB-7.0-8.0	7275670	N	3546	8270D	III
12-Nov-2013	SL-560-SA5D-SB-7.0-8.0	7275670	N	3546	8270D SIM	III
12-Nov-2013	SL-560-SA5D-SB-7.0-8.0	7275670	N	3550B	8151A	III
12-Nov-2013	SL-560-SA5D-SB-7.0-8.0	7275670	N	5035A	8015M	III
12-Nov-2013	SL-560-SA5D-SB-7.0-8.0	7275670	N	METHOD	7471B	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5	7275672	N	3050B	6010C	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5	7275672	N	3050B	6020A	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5	7275672	N	3546	8015M	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5	7275672	N	3546	8081B	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5	7275672	N	3546	8082A	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5	7275672	N	3546	8270D	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5	7275672	N	3546	8270D SIM	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5	7275672	N	METHOD	1613B	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5	7275672	N	METHOD	7471B	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5MSD	P275672M241619A	MSD	3546	8081B	III
12-Nov-2013	SL-549-SA5D-SB-0.0-0.5MS	P275672R241604A	MS	3546	8081B	III
12-Nov-2013	SL-549-SA5D-SB-4.0-5.0	7275673	N	3050B	6010C	III
12-Nov-2013	SL-549-SA5D-SB-4.0-5.0	7275673	N	3050B	6020A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
12-Nov-2013	SL-549-SA5D-SB-4.0-5.0	7275673	N	3546	8015M	III
12-Nov-2013	SL-549-SA5D-SB-4.0-5.0	7275673	N	3546	8081B	III
12-Nov-2013	SL-549-SA5D-SB-4.0-5.0	7275673	N	3546	8082A	III
12-Nov-2013	SL-549-SA5D-SB-4.0-5.0	7275673	N	3546	8270D	III
12-Nov-2013	SL-549-SA5D-SB-4.0-5.0	7275673	N	3546	8270D SIM	III
12-Nov-2013	SL-549-SA5D-SB-4.0-5.0	7275673	N	5035A	8015M	III
12-Nov-2013	SL-549-SA5D-SB-4.0-5.0	7275673	N	METHOD	1613B	III
12-Nov-2013	SL-549-SA5D-SB-4.0-5.0	7275673	N	METHOD	7471B	III
12-Nov-2013	SL-549-SA5D-SB-11.0-12.0	7275674	N	3050B	6010C	III
12-Nov-2013	SL-549-SA5D-SB-11.0-12.0	7275674	N	3050B	6020A	III
12-Nov-2013	SL-549-SA5D-SB-11.0-12.0	7275674	N	3546	8015M	III
12-Nov-2013	SL-549-SA5D-SB-11.0-12.0	7275674	N	3546	8081B	III
12-Nov-2013	SL-549-SA5D-SB-11.0-12.0	7275674	N	3546	8082A	III
12-Nov-2013	SL-549-SA5D-SB-11.0-12.0	7275674	N	3546	8270D	III
12-Nov-2013	SL-549-SA5D-SB-11.0-12.0	7275674	N	3546	8270D SIM	III
12-Nov-2013	SL-549-SA5D-SB-11.0-12.0	7275674	N	5035A	8015M	III
12-Nov-2013	SL-549-SA5D-SB-11.0-12.0	7275674	N	METHOD	1613B	III
12-Nov-2013	SL-549-SA5D-SB-11.0-12.0	7275674	N	METHOD	7471B	III
12-Nov-2013	EB1-111213	7275661	EB	3005A	6010C	III
12-Nov-2013	EB1-111213	7275661	EB	3510C	8015M	III
12-Nov-2013	EB1-111213	7275661	EB	3510C	8081B	III
12-Nov-2013	EB1-111213	7275661	EB	3510C	8082A	III
12-Nov-2013	EB1-111213	7275661	EB	3510C	8270D	III
12-Nov-2013	EB1-111213	7275661	EB	3510C	8270D SIM	III
12-Nov-2013	EB1-111213	7275661	EB	5030B	8015M	III
12-Nov-2013	EB1-111213	7275661	EB	Gen Prep	7199	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
12-Nov-2013	EB1-111213	7275661	EB	M3010A	6020A	III
12-Nov-2013	EB1-111213	7275661	EB	METHOD	1613B	III
12-Nov-2013	EB1-111213	7275661	EB	METHOD	7470A	III
12-Nov-2013	EB1-111213	7275661	EB	METHOD	8151A	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS
Method: 6010C **Matrix:** AQ

Sample ID: EB1-111213 Collected: 11/12/2013 3:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BARIUM	0.00034	J	0.00033	MDL	0.0100	PQL	mg/L	J	Z
CALCIUM	0.0349	J	0.0334	MDL	0.400	PQL	mg/L	J	Z
MAGNESIUM	0.0168	J	0.0167	MDL	0.200	PQL	mg/L	U	B

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-549-SA5D-SB-0.0-0.5 Collected: 11/12/2013 10:25:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	19000		7.58	MDL	42.1	PQL	mg/Kg	J	E
ANTIMONY	4.21	U	0.778	MDL	4.21	PQL	mg/Kg	UJ	Q, E
ARSENIC	11.8		0.736	MDL	4.21	PQL	mg/Kg	J	Q, E
BERYLLIUM	0.839	J	0.0705	MDL	1.05	PQL	mg/Kg	J	Z
BORON	3.76	J	0.884	MDL	10.5	PQL	mg/Kg	J	Z
CALCIUM	2800		3.51	MDL	21.0	PQL	mg/Kg	J	E
CHROMIUM	27.5		0.168	MDL	3.16	PQL	mg/Kg	J	Q, E
COPPER	12.5		0.305	MDL	2.10	PQL	mg/Kg	J	Q, E
IRON	25500		3.81	MDL	42.1	PQL	mg/Kg	J	E
LEAD	8.15		0.526	MDL	3.16	PQL	mg/Kg	J	Q, E
MAGNESIUM	5190		1.76	MDL	10.5	PQL	mg/Kg	J	E
MANGANESE	311		0.0873	MDL	1.05	PQL	mg/Kg	J	Q
MOLYBDENUM	0.429	J	0.179	MDL	2.10	PQL	mg/Kg	U	F
PHOSPHORUS	318		3.04	MDL	10.5	PQL	mg/Kg	J	Q, E
POTASSIUM	4140		8.77	MDL	105	PQL	mg/Kg	J	Q
SODIUM	88.5	J	17.6	MDL	105	PQL	mg/Kg	J	Z
TIN	2.91	J	0.231	MDL	10.5	PQL	mg/Kg	U	B
VANADIUM	47.4		0.137	MDL	1.05	PQL	mg/Kg	J	Q
ZINC	61.4		0.210	MDL	4.21	PQL	mg/Kg	J	Q

Sample ID: SL-549-SA5D-SB-11.0-12.0 Collected: 11/12/2013 11:05:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	14200		7.72	MDL	42.8	PQL	mg/Kg	J	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.28	U	0.792	MDL	4.28	PQL	mg/Kg	UJ	Q, E
ARSENIC	13.2		0.750	MDL	4.28	PQL	mg/Kg	J	Q, E
BERYLLIUM	0.793	J	0.0717	MDL	1.07	PQL	mg/Kg	J	Z
BORON	2.24	J	0.899	MDL	10.7	PQL	mg/Kg	J	Z
CALCIUM	2100		3.58	MDL	21.4	PQL	mg/Kg	J	E
CHROMIUM	20.6		0.171	MDL	3.21	PQL	mg/Kg	J	Q, E
COPPER	6.19		0.311	MDL	2.14	PQL	mg/Kg	J	Q, E
IRON	23300		3.88	MDL	42.8	PQL	mg/Kg	J	E
LEAD	7.15		0.535	MDL	3.21	PQL	mg/Kg	J	Q, E
MAGNESIUM	3810		1.79	MDL	10.7	PQL	mg/Kg	J	E
MANGANESE	127		0.0889	MDL	1.07	PQL	mg/Kg	J	Q
MOLYBDENUM	0.262	J	0.182	MDL	2.14	PQL	mg/Kg	U	F
PHOSPHORUS	101		3.09	MDL	10.7	PQL	mg/Kg	J	Q, E
POTASSIUM	1490		8.93	MDL	107	PQL	mg/Kg	J	Q
TIN	2.76	J	0.236	MDL	10.7	PQL	mg/Kg	U	B
VANADIUM	42.6		0.139	MDL	1.07	PQL	mg/Kg	J	Q
ZINC	38.4		0.214	MDL	4.28	PQL	mg/Kg	J	Q
Zirconium	4.10	J	0.899	MDL	5.35	PQL	mg/Kg	J	Z

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	22000		7.88	MDL	43.7	PQL	mg/Kg	J	E
ANTIMONY	4.37	U	0.809	MDL	4.37	PQL	mg/Kg	UJ	Q, E
ARSENIC	13.3		0.765	MDL	4.37	PQL	mg/Kg	J	Q, E
BERYLLIUM	0.972	J	0.0733	MDL	1.09	PQL	mg/Kg	J	Z
BORON	3.24	J	0.918	MDL	10.9	PQL	mg/Kg	J	Z
CALCIUM	2520		3.65	MDL	21.9	PQL	mg/Kg	J	E
CHROMIUM	27.9		0.175	MDL	3.28	PQL	mg/Kg	J	Q, E
COPPER	11.0		0.317	MDL	2.19	PQL	mg/Kg	J	Q, E
IRON	26800		3.96	MDL	43.7	PQL	mg/Kg	J	E
LEAD	7.86		0.547	MDL	3.28	PQL	mg/Kg	J	Q, E
MAGNESIUM	5390		1.83	MDL	10.9	PQL	mg/Kg	J	E
MANGANESE	336		0.0907	MDL	1.09	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-549-SA5D-SB-4.0-5.0 Collected: 11/12/2013 10:40:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MOLYBDENUM	0.314	J	0.186	MDL	2.19	PQL	mg/Kg	U	F
PHOSPHORUS	229		3.16	MDL	10.9	PQL	mg/Kg	J	Q, E
POTASSIUM	3450		9.12	MDL	109	PQL	mg/Kg	J	Q
TIN	3.16	J	0.241	MDL	10.9	PQL	mg/Kg	U	B
VANADIUM	50.1		0.142	MDL	1.09	PQL	mg/Kg	J	Q
ZINC	58.8		0.219	MDL	4.37	PQL	mg/Kg	J	Q

Sample ID: SL-560-SA5D-SB-0.0-0.5 Collected: 11/12/2013 8:35:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	35900		7.82	MDL	43.4	PQL	mg/Kg	J	E
ANTIMONY	4.34	U	0.803	MDL	4.34	PQL	mg/Kg	UJ	Q, E
ARSENIC	9.70		0.759	MDL	4.34	PQL	mg/Kg	J	Q, E
BORON	8.20	J	0.911	MDL	10.8	PQL	mg/Kg	J	Z
CALCIUM	6910		3.62	MDL	21.7	PQL	mg/Kg	J	E
CHROMIUM	42.4		0.174	MDL	3.25	PQL	mg/Kg	J	Q, E
COPPER	14.9		0.315	MDL	2.17	PQL	mg/Kg	J	Q, E
LEAD	12.2		0.542	MDL	3.25	PQL	mg/Kg	J	Q, E
MAGNESIUM	8150		1.81	MDL	10.8	PQL	mg/Kg	J	E
MANGANESE	487		0.0900	MDL	1.08	PQL	mg/Kg	J	Q
MOLYBDENUM	0.227	J	0.184	MDL	2.17	PQL	mg/Kg	U	F
PHOSPHORUS	308		3.14	MDL	10.8	PQL	mg/Kg	J	Q, E
POTASSIUM	4120		9.05	MDL	108	PQL	mg/Kg	J	Q
TIN	3.29	J	0.239	MDL	10.8	PQL	mg/Kg	U	B
VANADIUM	80.0		0.141	MDL	1.08	PQL	mg/Kg	J	Q
ZINC	81.7		0.217	MDL	4.34	PQL	mg/Kg	J	Q

Sample ID: SL-560-SA5D-SB-0.0-0.5 Collected: 11/12/2013 8:35:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	39300		7.85	MDL	86.8	PQL	mg/Kg	J	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS

Method: 6010C

Matrix: SO

Sample ID: SL-560-SA5D-SB-4.0-5.0

Collected: 11/12/2013 9:20:00

Analysis Type: REA

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	14400		7.22	MDL	40.1	PQL	mg/Kg	J	E
ANTIMONY	4.01	U	0.741	MDL	4.01	PQL	mg/Kg	UJ	Q, E
ARSENIC	5.64		0.701	MDL	4.01	PQL	mg/Kg	J	Q, E
BERYLLIUM	0.668	J	0.0671	MDL	1.00	PQL	mg/Kg	J	Z
BORON	1.90	J	0.842	MDL	10.0	PQL	mg/Kg	J	Z
CALCIUM	2270		3.35	MDL	20.0	PQL	mg/Kg	J	E
CHROMIUM	16.4		0.160	MDL	3.01	PQL	mg/Kg	J	Q, E
COPPER	5.94		0.291	MDL	2.00	PQL	mg/Kg	J	Q, E
IRON	21200		3.63	MDL	40.1	PQL	mg/Kg	J	E
LEAD	6.33		0.501	MDL	3.01	PQL	mg/Kg	J	Q, E
MAGNESIUM	3520		1.67	MDL	10.0	PQL	mg/Kg	J	E
MANGANESE	148		0.0832	MDL	1.00	PQL	mg/Kg	J	Q
MOLYBDENUM	0.261	J	0.170	MDL	2.00	PQL	mg/Kg	UJ	FD, F
PHOSPHORUS	134		2.90	MDL	10.0	PQL	mg/Kg	J	Q, E
POTASSIUM	2100		8.36	MDL	100	PQL	mg/Kg	J	Q
TIN	2.97	J	0.220	MDL	10.0	PQL	mg/Kg	U	B
VANADIUM	33.1		0.130	MDL	1.00	PQL	mg/Kg	J	Q
ZINC	45.6		0.200	MDL	4.01	PQL	mg/Kg	J	Q
Zirconium	4.11	J	0.842	MDL	5.01	PQL	mg/Kg	J	Z

Sample ID: SL-560-SA5D-SB-7.0-8.0

Collected: 11/12/2013 9:35:00

Analysis Type: REA

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	10700		7.17	MDL	39.8	PQL	mg/Kg	J	E
ANTIMONY	3.98	U	0.736	MDL	3.98	PQL	mg/Kg	UJ	Q, E
ARSENIC	5.76		0.696	MDL	3.98	PQL	mg/Kg	J	Q, E
BERYLLIUM	0.601	J	0.0666	MDL	0.994	PQL	mg/Kg	J	Z
BORON	1.76	J	0.835	MDL	9.94	PQL	mg/Kg	J	Z
CALCIUM	2420		3.32	MDL	19.9	PQL	mg/Kg	J	E
CHROMIUM	19.4		0.159	MDL	2.98	PQL	mg/Kg	J	Q, E
COPPER	8.67		0.288	MDL	1.99	PQL	mg/Kg	J	Q, E
IRON	15700		3.60	MDL	39.8	PQL	mg/Kg	J	E
LEAD	6.10		0.497	MDL	2.98	PQL	mg/Kg	J	Q, E
MAGNESIUM	2770		1.66	MDL	9.94	PQL	mg/Kg	J	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-560-SA5D-SB-7.0-8.0 Collected: 11/12/2013 9:35:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MANGANESE	103		0.0825	MDL	0.994	PQL	mg/Kg	J	Q
MOLYBDENUM	0.213	J	0.169	MDL	1.99	PQL	mg/Kg	U	F
PHOSPHORUS	169		2.87	MDL	9.94	PQL	mg/Kg	J	Q, E
POTASSIUM	1400		8.29	MDL	99.4	PQL	mg/Kg	J	Q
TIN	2.36	J	0.219	MDL	9.94	PQL	mg/Kg	U	B
VANADIUM	28.1		0.129	MDL	0.994	PQL	mg/Kg	J	Q
ZINC	35.2		0.199	MDL	3.98	PQL	mg/Kg	J	Q
Zirconium	3.30	J	0.835	MDL	4.97	PQL	mg/Kg	J	Z

Sample ID: SL-860-SA5D-SB-4.0-5.0 Collected: 11/12/2013 9:05:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	13300		7.22	MDL	40.1	PQL	mg/Kg	J	E
ANTIMONY	4.01	U	0.741	MDL	4.01	PQL	mg/Kg	UJ	Q, E
ARSENIC	5.98		0.701	MDL	4.01	PQL	mg/Kg	J	Q, E
BERYLLIUM	0.618	J	0.0671	MDL	1.00	PQL	mg/Kg	J	Z
BORON	1.94	J	0.841	MDL	10.0	PQL	mg/Kg	J	Z
CALCIUM	2770		3.35	MDL	20.0	PQL	mg/Kg	J	E
CHROMIUM	18.2		0.160	MDL	3.00	PQL	mg/Kg	J	Q, E
COPPER	6.97		0.290	MDL	2.00	PQL	mg/Kg	J	Q, E
IRON	21400		3.63	MDL	40.1	PQL	mg/Kg	J	E
LEAD	6.82		0.501	MDL	3.00	PQL	mg/Kg	J	Q, E
MAGNESIUM	3760		1.67	MDL	10.0	PQL	mg/Kg	J	E
MANGANESE	160		0.0831	MDL	1.00	PQL	mg/Kg	J	Q
MOLYBDENUM	2.00	U	0.170	MDL	2.00	PQL	mg/Kg	UJ	FD
PHOSPHORUS	198		2.89	MDL	10.0	PQL	mg/Kg	J	Q, E
POTASSIUM	2510		8.35	MDL	100	PQL	mg/Kg	J	Q
TIN	2.85	J	0.220	MDL	10.0	PQL	mg/Kg	U	B
VANADIUM	32.7		0.130	MDL	1.00	PQL	mg/Kg	J	Q
ZINC	49.9		0.200	MDL	4.01	PQL	mg/Kg	J	Q
Zirconium	3.96	J	0.841	MDL	5.01	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS

Method: 6020A

Matrix: SO

Sample ID: SL-549-SA5D-SB-0.0-0.5

Collected: 11/12/2013 10:25:00

Analysis Type: REA2

Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0281	J	0.0274	MDL	0.210	PQL	mg/Kg	J	Z
STRONTIUM	28.4		0.0715	MDL	0.421	PQL	mg/Kg	J	Q, E
THALLIUM	0.372		0.0316	MDL	0.210	PQL	mg/Kg	J	Q

Sample ID: SL-549-SA5D-SB-0.0-0.5

Collected: 11/12/2013 10:25:00

Analysis Type: REA3

Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.234	J	0.105	MDL	0.421	PQL	mg/Kg	J	Z

Sample ID: SL-549-SA5D-SB-11.0-12.0

Collected: 11/12/2013 11:05:00

Analysis Type: REA2

Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0425	J	0.0278	MDL	0.214	PQL	mg/Kg	J	Z
STRONTIUM	19.9		0.0728	MDL	0.428	PQL	mg/Kg	J	Q, E
THALLIUM	0.251		0.0321	MDL	0.214	PQL	mg/Kg	J	Q

Sample ID: SL-549-SA5D-SB-4.0-5.0

Collected: 11/12/2013 10:40:00

Analysis Type: REA2

Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0498	J	0.0284	MDL	0.219	PQL	mg/Kg	J	Z
STRONTIUM	27.3		0.0743	MDL	0.437	PQL	mg/Kg	J	Q, E
THALLIUM	0.368		0.0328	MDL	0.219	PQL	mg/Kg	J	Q

Sample ID: SL-549-SA5D-SB-4.0-5.0

Collected: 11/12/2013 10:40:00

Analysis Type: REA3

Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.123	J	0.109	MDL	0.437	PQL	mg/Kg	J	Z

Sample ID: SL-560-SA5D-SB-0.0-0.5

Collected: 11/12/2013 8:35:00

Analysis Type: REA2

Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0568	J	0.0282	MDL	0.217	PQL	mg/Kg	J	Z
STRONTIUM	36.4		0.0738	MDL	0.434	PQL	mg/Kg	J	Q, E
THALLIUM	0.421		0.0325	MDL	0.217	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS
Method: 6020A Matrix: SO

Sample ID: SL-560-SA5D-SB-0.0-0.5 Collected: 11/12/2013 8:35:00 Analysis Type: REA3 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.231	J	0.108	MDL	0.434	PQL	mg/Kg	J	Z

Sample ID: SL-560-SA5D-SB-4.0-5.0 Collected: 11/12/2013 9:20:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	13.2		0.0681	MDL	0.401	PQL	mg/Kg	J	Q, E
THALLIUM	0.224		0.0301	MDL	0.200	PQL	mg/Kg	J	Q

Sample ID: SL-560-SA5D-SB-7.0-8.0 Collected: 11/12/2013 9:35:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	11.4		0.0676	MDL	0.398	PQL	mg/Kg	J	Q, E
THALLIUM	0.245		0.0298	MDL	0.199	PQL	mg/Kg	J	Q

Sample ID: SL-860-SA5D-SB-4.0-5.0 Collected: 11/12/2013 9:05:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	14.4		0.0681	MDL	0.401	PQL	mg/Kg	J	Q, E
THALLIUM	0.291		0.0300	MDL	0.200	PQL	mg/Kg	J	Q

Method Category: METALS
Method: 7199 Matrix: SO

Sample ID: SL-560-SA5D-SB-4.0-5.0 Collected: 11/12/2013 9:20:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.33	J	0.15	MDL	0.42	PQL	mg/Kg	J	Z

Sample ID: SL-560-SA5D-SB-7.0-8.0 Collected: 11/12/2013 9:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.32	J	0.14	MDL	0.41	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS
Method: 7199 **Matrix:** SO

Sample ID: SL-860-SA5D-SB-4.0-5.0 Collected: 11/12/2013 9:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.34	J	0.15	MDL	0.43	PQL	mg/Kg	J	Z

Method Category: METALS
Method: 7471B **Matrix:** SO

Sample ID: SL-549-SA5D-SB-11.0-12.0 Collected: 11/12/2013 11:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0124	J	0.0106	MDL	0.0176	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 1613B **Matrix:** AQ

Sample ID: EB1-111213 Collected: 11/12/2013 3:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.00	JBQ	0.342	MDL	9.95	PQL	pg/L	U	B
1,2,3,4,6,7,8-HPCDF	0.885	JBQ	0.113	MDL	9.95	PQL	pg/L	U	B
1,2,3,4,7,8,9-HPCDF	0.846	JBQ	0.133	MDL	9.95	PQL	pg/L	U	B
1,2,3,4,7,8-HXCDF	0.641	JBQ	0.229	MDL	9.95	PQL	pg/L	U	B
1,2,3,6,7,8-HXCDD	0.661	JBQ	0.292	MDL	9.95	PQL	pg/L	U	B
1,2,3,6,7,8-HXCDF	0.622	JBQ	0.219	MDL	9.95	PQL	pg/L	U	B
1,2,3,7,8,9-HXCDF	0.756	JBQ	0.244	MDL	9.95	PQL	pg/L	U	B
1,2,3,7,8-PECDD	1.28	JB	0.390	MDL	9.95	PQL	pg/L	U	B
1,2,3,7,8-PECDF	0.895	JBQ	0.230	MDL	9.95	PQL	pg/L	U	B
2,3,4,6,7,8-HXCDF	0.664	JBQ	0.217	MDL	9.95	PQL	pg/L	U	B
2,3,4,7,8-PECDF	0.345	JBQ	0.211	MDL	9.95	PQL	pg/L	U	B
2,3,7,8-TCDD	0.717	JBQ	0.522	MDL	1.99	PQL	pg/L	U	B
OCDD	0.851	JBQ	0.366	MDL	19.9	PQL	pg/L	U	B
OCDF	1.81	JBQ	0.461	MDL	19.9	PQL	pg/L	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-549-SA5D-SB-0.0-0.5 Collected: 11/12/2013 10:25:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	2.13	JB	0.0441	MDL	5.28	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.564	JBQ	0.0153	MDL	5.28	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.120	JBQ	0.0410	MDL	5.28	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0851	JB	0.0471	MDL	5.28	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0668	JBQ	0.0389	MDL	5.28	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.741	JBQ	0.0495	MDL	5.28	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.120	JB	0.0308	MDL	5.28	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	1.15	JB	0.0477	MDL	5.28	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	1.60	JB	0.0484	MDL	5.28	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.148	JBQ	0.0594	MDL	5.28	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.609	JB	0.0322	MDL	5.28	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.114	JBQ	0.0336	MDL	5.28	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.108	JBQ	0.0344	MDL	5.28	PQL	ng/Kg	U	B
OCDF	1.37	JBQ	0.0897	MDL	10.6	PQL	ng/Kg	J	Z

Sample ID: SL-549-SA5D-SB-11.0-12.0 Collected: 11/12/2013 11:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0901	JBQ	0.0223	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0689	JBQ	0.00802	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0647	JBQ	0.0202	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0704	JB	0.0236	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.143	JBQ	0.0176	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.0868	JB	0.0257	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.117	JBQ	0.0136	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0673	JBQ	0.0231	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.115	JBQ	0.0182	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.194	JBQ	0.0355	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.282	JB	0.0184	MDL	5.53	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0818	JB	0.0164	MDL	5.53	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.251	JBQ	0.0202	MDL	5.53	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.117	JBQ	0.0379	MDL	1.11	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.0622	JQ	0.0351	MDL	1.11	PQL	ng/Kg	J	Z
OCDD	0.326	JB	0.0314	MDL	11.1	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-549-SA5D-SB-11.0-12.0 Collected: 11/12/2013 11:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
OCDF	0.267	JBQ	0.0440	MDL	11.1	PQL	ng/Kg	U	B

Sample ID: SL-549-SA5D-SB-4.0-5.0 Collected: 11/12/2013 10:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.521	JB	0.0383	MDL	5.61	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.158	JBQ	0.0109	MDL	5.61	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0547	JBQ	0.0324	MDL	5.61	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0601	JB	0.0274	MDL	5.61	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.107	JBQ	0.0247	MDL	5.61	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.136	JBQ	0.0287	MDL	5.61	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDF	0.100	JB	0.0196	MDL	5.61	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0891	JBQ	0.0280	MDL	5.61	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDF	0.200	JBQ	0.0305	MDL	5.61	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.204	JBQ	0.0416	MDL	5.61	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.223	JBQ	0.0242	MDL	5.61	PQL	ng/Kg	U	B
2,3,4,6,7,8-HxCDF	0.0671	JB	0.0229	MDL	5.61	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.153	JBQ	0.0277	MDL	5.61	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0685	JBQ	0.0422	MDL	1.12	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.0495	J	0.0471	MDL	1.12	PQL	ng/Kg	J	Z
OCDD	4.58	JB	0.0396	MDL	11.2	PQL	ng/Kg	J	Z
OCDF	0.660	JB	0.0734	MDL	11.2	PQL	ng/Kg	U	B

Sample ID: SL-560-SA5D-SB-0.0-0.5 Collected: 11/12/2013 8:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.230	JBQ	0.0342	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0577	JBQ	0.0125	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.136	JBQ	0.0364	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.0548	JBQ	0.0265	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.129	JBQ	0.0344	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDF	0.0729	JB	0.0209	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.145	JB	0.0330	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDF	0.298	JBQ	0.0316	MDL	5.34	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0938	JBQ	0.0493	MDL	5.34	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA
Method: 1613B **Matrix: SO**

Sample ID: SL-560-SA5D-SB-0.0-0.5 Collected: 11/12/2013 8:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,7,8-PECDF	0.129	JBQ	0.0287	MDL	5.34	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.102	JBQ	0.0241	MDL	5.34	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0461	JBQ	0.0322	MDL	5.34	PQL	ng/Kg	U	B
OCDD	0.895	JB	0.0447	MDL	10.7	PQL	ng/Kg	U	B
OCDF	0.413	JB	0.0826	MDL	10.7	PQL	ng/Kg	U	B

Method Category: SVOA
Method: 8015M **Matrix: AQ**

Sample ID: EB1-111213 Collected: 11/12/2013 3:00:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C30-C40)	0.49	U	0.097	MDL	0.49	PQL	mg/L	UJ	L
EFH (C8-C11)	0.097	U	0.049	MDL	0.097	PQL	mg/L	UJ	L

Method Category: SVOA
Method: 8015M **Matrix: SO**

Sample ID: SL-560-SA5D-SB-4.0-5.0 Collected: 11/12/2013 9:20:00 Analysis Type: RES Dilution: 26.04

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
GASOLINE RANGE ORGANICS (C5-C12)	0.4	J	0.2	MDL	1.1	PQL	mg/Kg	J	Z, FD

Sample ID: SL-560-SA5D-SB-7.0-8.0 Collected: 11/12/2013 9:35:00 Analysis Type: RES Dilution: 23.72

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
GASOLINE RANGE ORGANICS (C5-C12)	0.3	J	0.2	MDL	1	PQL	mg/Kg	J	Z

Sample ID: SL-860-SA5D-SB-4.0-5.0 Collected: 11/12/2013 9:05:00 Analysis Type: RES Dilution: 24.7

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
GASOLINE RANGE ORGANICS (C5-C12)	1.0	U	0.2	MDL	1.0	PQL	mg/Kg	UJ	FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA
Method: 8081B **Matrix:** SO

Sample ID: SL-549-SA5D-SB-0.0-0.5 Collected: 11/12/2013 10:25:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
4,4'-DDE	0.49	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z
4,4'-DDT	1.3	J	0.38	MDL	1.8	PQL	ug/Kg	J	Z

Method Category: SVOA
Method: 8151A **Matrix:** AQ

Sample ID: EB1-111213 Collected: 11/12/2013 3:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,4,5-T	0.050	U	0.015	MDL	0.050	PQL	ug/L	UJ	E
2,4,5-TP (Silvex)	0.050	U	0.0099	MDL	0.050	PQL	ug/L	UJ	E
2,4-D	0.50	U	0.16	MDL	0.50	PQL	ug/L	UJ	E
2,4-DB	0.99	U	0.30	MDL	0.99	PQL	ug/L	UJ	E
DALAPON	1.2	U	0.25	MDL	1.2	PQL	ug/L	UJ	E
DICAMBA	0.30	U	0.079	MDL	0.30	PQL	ug/L	UJ	E
DICHLOROPROP	0.50	U	0.16	MDL	0.50	PQL	ug/L	UJ	E
MCPA	200	U	50	MDL	200	PQL	ug/L	UJ	E
MCPP	200	U	50	MDL	200	PQL	ug/L	UJ	E

Method Category: SVOA
Method: 8270D **Matrix:** AQ

Sample ID: EB1-111213 Collected: 11/12/2013 3:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
3,5-Dimethylphenol	10	U	3	MDL	10	PQL	ug/L	UJ	L

Sample ID: EB1-111213 Collected: 11/12/2013 3:00:00 Analysis Type: RES-ACID Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,6-DICHLOROPHENOL	1	U	0.5	MDL	1	PQL	ug/L	UJ	L

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** AQ

Sample ID: EB1-111213 Collected: 11/12/2013 3:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
Diethylphthalate	0.37	J	0.051	MDL	1.0	PQL	ug/L	U	B
Di-n-butylphthalate	0.19	J	0.051	MDL	1.0	PQL	ug/L	U	B

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-549-SA5D-SB-0.0-0.5 Collected: 11/12/2013 10:25:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	1.3	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	1.1	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Calibration Blank Contamination
B	Method Blank Contamination
E	Laboratory Control Precision
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Field Blank Contamination
FD	Field Duplicate Precision
L	Laboratory Control Precision
L	Laboratory Control Spike Lower Estimation
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Lower Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH127

Method Blank Outlier Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 1613B
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK3230B370934	11/22/2013 9:34:00 AM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDD 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDD 1,2,3,7,8,9-HxCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HxCDF 2,3,4,7,8-PECDF 2,3,7,8-TCDD OCDD OCDF	0.616 pg/L 0.958 pg/L 0.986 pg/L 0.655 pg/L 0.912 pg/L 0.743 pg/L 1.05 pg/L 0.571 pg/L 1.12 pg/L 0.830 pg/L 1.16 pg/L 1.11 pg/L 0.618 pg/L 0.726 pg/L 2.62 pg/L 2.54 pg/L	EB1-111213

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB1-111213(RES)	1,2,3,4,6,7,8-HPCDD	1.00 pg/L	1.00U pg/L
EB1-111213(RES)	1,2,3,4,6,7,8-HPCDF	0.885 pg/L	0.885U pg/L
EB1-111213(RES)	1,2,3,4,7,8,9-HPCDF	0.846 pg/L	0.846U pg/L
EB1-111213(RES)	1,2,3,4,7,8-HxCDF	0.641 pg/L	0.641U pg/L
EB1-111213(RES)	1,2,3,6,7,8-HxCDD	0.661 pg/L	0.661U pg/L
EB1-111213(RES)	1,2,3,6,7,8-HxCDF	0.622 pg/L	0.622U pg/L
EB1-111213(RES)	1,2,3,7,8,9-HxCDF	0.756 pg/L	0.756U pg/L
EB1-111213(RES)	1,2,3,7,8-PECDD	1.28 pg/L	1.28U pg/L
EB1-111213(RES)	1,2,3,7,8-PECDF	0.895 pg/L	0.895U pg/L
EB1-111213(RES)	2,3,4,6,7,8-HxCDF	0.664 pg/L	0.664U pg/L
EB1-111213(RES)	2,3,4,7,8-PECDF	0.345 pg/L	0.345U pg/L
EB1-111213(RES)	2,3,7,8-TCDD	0.717 pg/L	0.717U pg/L
EB1-111213(RES)	OCDD	0.851 pg/L	0.851U pg/L
EB1-111213(RES)	OCDF	1.81 pg/L	1.81U pg/L

Method Blank Outlier Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK3240B371252	11/24/2013 12:52:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF 2,3,7,8-TCDD OCDD OCDF	0.0721 ng/Kg 0.0746 ng/Kg 0.0711 ng/Kg 0.0412 ng/Kg 0.0890 ng/Kg 0.0407 ng/Kg 0.0519 ng/Kg 0.0742 ng/Kg 0.0575 ng/Kg 0.0804 ng/Kg 0.121 ng/Kg 0.0893 ng/Kg 0.0714 ng/Kg 0.0670 ng/Kg 0.213 ng/Kg 0.233 ng/Kg	SL-549-SA5D-SB-0.0-0.5 SL-549-SA5D-SB-11.0-12.0 SL-549-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-549-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.120 ng/Kg	0.120U ng/Kg
SL-549-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0851 ng/Kg	0.0851U ng/Kg
SL-549-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.0668 ng/Kg	0.0668U ng/Kg
SL-549-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.120 ng/Kg	0.120U ng/Kg
SL-549-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.148 ng/Kg	0.148U ng/Kg
SL-549-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.114 ng/Kg	0.114U ng/Kg
SL-549-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.108 ng/Kg	0.108U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,4,6,7,8-HPCDD	0.0901 ng/Kg	0.0901U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0689 ng/Kg	0.0689U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0647 ng/Kg	0.0647U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,4,7,8-HxCDD	0.0704 ng/Kg	0.0704U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,4,7,8-HxCDF	0.143 ng/Kg	0.143U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,6,7,8-HXCDD	0.0868 ng/Kg	0.0868U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,6,7,8-HXCDF	0.117 ng/Kg	0.117U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,7,8,9-HXCDD	0.0673 ng/Kg	0.0673U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,7,8,9-HXCDF	0.115 ng/Kg	0.115U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,7,8-PECDD	0.194 ng/Kg	0.194U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	1,2,3,7,8-PECDF	0.282 ng/Kg	0.282U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	2,3,4,6,7,8-HXCDF	0.0818 ng/Kg	0.0818U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	2,3,4,7,8-PECDF	0.251 ng/Kg	0.251U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	2,3,7,8-TCDD	0.117 ng/Kg	0.117U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	OCDD	0.326 ng/Kg	0.326U ng/Kg
SL-549-SA5D-SB-11.0-12.0(RES)	OCDF	0.267 ng/Kg	0.267U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.158 ng/Kg	0.158U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0547 ng/Kg	0.0547U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDD	0.0601 ng/Kg	0.0601U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDF	0.107 ng/Kg	0.107U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDD	0.136 ng/Kg	0.136U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDF	0.100 ng/Kg	0.100U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-549-SA5D-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDD	0.0891 ng/Kg	0.0891U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDF	0.200 ng/Kg	0.200U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	1,2,3,7,8-PECDD	0.204 ng/Kg	0.204U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.223 ng/Kg	0.223U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0671 ng/Kg	0.0671U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	2,3,4,7,8-PECDF	0.153 ng/Kg	0.153U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	2,3,7,8-TCDD	0.0685 ng/Kg	0.0685U ng/Kg
SL-549-SA5D-SB-4.0-5.0(RES)	OCDF	0.660 ng/Kg	0.660U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.230 ng/Kg	0.230U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0577 ng/Kg	0.0577U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.136 ng/Kg	0.136U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0548 ng/Kg	0.0548U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.129 ng/Kg	0.129U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0729 ng/Kg	0.0729U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.145 ng/Kg	0.145U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.0938 ng/Kg	0.0938U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.129 ng/Kg	0.129U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.102 ng/Kg	0.102U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.0461 ng/Kg	0.0461U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	OCDD	0.895 ng/Kg	0.895U ng/Kg
SL-560-SA5D-SB-0.0-0.5(RES)	OCDF	0.413 ng/Kg	0.413U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P32537CB222019	11/22/2013 8:19:00 PM	ALUMINUM CALCIUM IRON MAGNESIUM TIN	19.2 mg/Kg 22.5 mg/Kg 8.11 mg/Kg 19.6 mg/Kg 1.56 mg/Kg	SL-549-SA5D-SB-0.0-0.5 SL-549-SA5D-SB-11.0-12.0 SL-549-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-0.0-0.5 SL-560-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-7.0-8.0 SL-860-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-549-SA5D-SB-0.0-0.5(REA)	TIN	2.91 mg/Kg	2.91U mg/Kg
SL-549-SA5D-SB-11.0-12.0(REA)	TIN	2.76 mg/Kg	2.76U mg/Kg
SL-549-SA5D-SB-4.0-5.0(REA)	TIN	3.16 mg/Kg	3.16U mg/Kg
SL-560-SA5D-SB-0.0-0.5(REA)	TIN	3.29 mg/Kg	3.29U mg/Kg
SL-560-SA5D-SB-4.0-5.0(REA)	TIN	2.97 mg/Kg	2.97U mg/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-560-SA5D-SB-7.0-8.0(REA)	TIN	2.36 mg/Kg	2.36U mg/Kg
SL-860-SA5D-SB-4.0-5.0(REA)	TIN	2.85 mg/Kg	2.85U mg/Kg

Method: 8270D SIM
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
PLKWA32B261437	11/23/2013 2:37:00 PM	BIS(2-ETHYLHEXYL)PHTHALATE Diethylphthalate Di-n-butylphthalate	0.096 ug/L 0.056 ug/L 0.057 ug/L	EB1-111213

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB1-111213(RES)	Diethylphthalate	0.37 ug/L	1.0U ug/L
EB1-111213(RES)	Di-n-butylphthalate	0.19 ug/L	1.0U ug/L

Field Blank Outlier Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041613(REA2/TOT)	4/16/2013 3:15:00 PM	MOLYBDENUM TIN	0.0132 mg/L 0.0029 mg/L	SL-549-SA5D-SB-0.0-0.5 SL-549-SA5D-SB-11.0-12.0 SL-549-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-0.0-0.5 SL-560-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-7.0-8.0 SL-860-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-549-SA5D-SB-0.0-0.5(REA)	MOLYBDENUM	0.429 mg/Kg	0.429U mg/Kg
SL-549-SA5D-SB-11.0-12.0(REA)	MOLYBDENUM	0.262 mg/Kg	0.262U mg/Kg
SL-549-SA5D-SB-4.0-5.0(REA)	MOLYBDENUM	0.314 mg/Kg	0.314U mg/Kg
SL-560-SA5D-SB-0.0-0.5(REA)	MOLYBDENUM	0.227 mg/Kg	0.227U mg/Kg
SL-560-SA5D-SB-4.0-5.0(REA)	MOLYBDENUM	0.261 mg/Kg	0.261U mg/Kg
SL-560-SA5D-SB-7.0-8.0(REA)	MOLYBDENUM	0.213 mg/Kg	0.213U mg/Kg

Surrogate Outlier Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 8015M

Matrix: SO

<i>Sample ID (Analysis Type)</i>	<i>Surrogate</i>	<i>Sample % Recovery</i>	<i>% Recovery Limits</i>	<i>Affected Compounds</i>	<i>Flag</i>
SL-549-SA5D-SB- 11.0-12.0 (REA)	CHLOROBENZENE	153	37.00-125.00	All Target Analytes	J (all detects)

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-560-SA5D-SB-4.0-5.0MS (TOT) SL-560-SA5D-SB-4.0-5.0MSD (TOT) (SL-549-SA5D-SB-0.0-0.5 SL-549-SA5D-SB-11.0-12.0 SL-549-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-0.0-0.5 SL-560-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-7.0-8.0 SL-860-SA5D-SB-4.0-5.0)	MANGANESE POTASSIUM TITANIUM VANADIUM ZINC	- 161 260 - -	192 191 444 132 147	75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00	- - - - -	MANGANESE POTASSIUM TITANIUM VANADIUM ZINC	J (all detects) Ti, No Qual, >4x
SL-560-SA5D-SB-4.0-5.0MS (TOT) SL-560-SA5D-SB-4.0-5.0MSD (TOT) (SL-549-SA5D-SB-0.0-0.5 SL-549-SA5D-SB-11.0-12.0 SL-549-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-0.0-0.5 SL-560-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-7.0-8.0 SL-860-SA5D-SB-4.0-5.0)	ALUMINUM IRON	-117 -1719	2431 9674	75.00-125.00 75.00-125.00	30 (20.00) 45 (20.00)	ALUMINUM IRON	J(all detects) UJ(all non-detects) No Qual, %R >4x
SL-560-SA5D-SB-4.0-5.0MS (TOT) SL-560-SA5D-SB-4.0-5.0MSD (TOT) (SL-549-SA5D-SB-0.0-0.5 SL-549-SA5D-SB-11.0-12.0 SL-549-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-0.0-0.5 SL-560-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-7.0-8.0 SL-860-SA5D-SB-4.0-5.0)	ANTIMONY ARSENIC CALCIUM CHROMIUM COPPER LEAD MAGNESIUM PHOSPHORUS	62 - 131 - - - 196 260	48 129 792 144 127 127 1134 137	75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00	30 (20.00) 22 (20.00) 63 (20.00) 26 (20.00) 23 (20.00) 24 (20.00) 38 (20.00) 39 (20.00)	ANTIMONY ARSENIC CALCIUM CHROMIUM COPPER LEAD MAGNESIUM PHOSPHORUS	J(all detects) UJ(all non-detects) Ca, Mg, No Qual, %R >4x

Method: 6020A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-560-SA5D-SB-4.0-5.0MSD (TOT) (SL-549-SA5D-SB-0.0-0.5 SL-549-SA5D-SB-11.0-12.0 SL-549-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-0.0-0.5 SL-560-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-7.0-8.0 SL-860-SA5D-SB-4.0-5.0)	THALLIUM	-	141	75.00-125.00	-	THALLIUM	J(all detects)
SL-560-SA5D-SB-4.0-5.0MSD (TOT) (SL-549-SA5D-SB-0.0-0.5 SL-549-SA5D-SB-11.0-12.0 SL-549-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-0.0-0.5 SL-560-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-7.0-8.0 SL-860-SA5D-SB-4.0-5.0)	STRONTIUM	-	236	75.00-125.00	34 (20.00)	STRONTIUM	J(all detects) UJ(all non-detects)

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 8270D

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-560-SA5D-SB-4.0-5.0MSD (SL-560-SA5D-SB-4.0-5.0)	3,3'-DICHLOROBENZIDINE	-	-	10.00-143.00	43 (30.00)	3,3'-DICHLOROBENZIDINE	J(all detects)

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6010C

Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-560-SA5D-SB-4.0-5.0DUP (TOT) (SL-549-SA5D-SB-0.0-0.5 SL-549-SA5D-SB-11.0-12.0 SL-549-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-0.0-0.5 SL-560-SA5D-SB-4.0-5.0 SL-560-SA5D-SB-7.0-8.0 SL-860-SA5D-SB-4.0-5.0)	MOLYBDENUM	200	20.00	No Qual, OK by Difference

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 8151A

Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P33203AQ241337A P33203AY241404A (EB1-111213)	DINOSEB	168	223	16.00-163.00	-	DINOSEB	J (all detects)
P33203AY241404A (EB1-111213)	2,4,5-T 2,4,5-TP (Silvex) 2,4-D 2,4-DB DALAPON DICAMBA DICHLOROPROP MCPA MCPD	- - - - - - - - -	- - - - - - - - -	55.00-169.00 58.00-155.00 68.00-155.00 50.00-163.00 39.00-115.00 55.00-163.00 89.00-162.00 68.00-154.00 46.00-173.00	34 (30.00) 32 (30.00) 35 (30.00) 35 (30.00) 48 (30.00) 33 (30.00) 31 (30.00) 39 (30.00) 57 (30.00)	2,4,5-T 2,4,5-TP (Silvex) 2,4-D 2,4-DB DALAPON DICAMBA DICHLOROPROP MCPA MCPD	J(all detects) UJ(all non-detects)

Method: 8015M

Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P33228AQ321533A P33228AY321554A (EB1-111213)	EFH (C30-C40) EFH (C8-C11)	45 55	56 65	70.00-130.00 70.00-130.00	- -	EFH (C30-C40) EFH (C8-C11)	J(all detects) UJ(all non-detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 160.3M

Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-560-SA5D-SB-4.0-5.0	SL-860-SA5D-SB-4.0-5.0			
MOISTURE	3.1	4.0	25		No Qualifiers Applied

Method: 6010C

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-560-SA5D-SB-4.0-5.0 (TOT)	SL-860-SA5D-SB-4.0-5.0 (TOT)			
ALUMINUM	14400	13300	8	50.00	No Qualifiers Applied
ARSENIC	5.64	5.98	6	50.00	
BARIUM	61.7	61.3	1	50.00	
BERYLLIUM	0.668	0.618	8	50.00	
BORON	1.90	1.94	2	50.00	
CALCIUM	2270	2770	20	50.00	
CHROMIUM	16.4	18.2	10	50.00	
COBALT	3.18	3.53	10	50.00	
COPPER	5.94	6.97	16	50.00	
IRON	21200	21400	1	50.00	
LEAD	6.33	6.82	7	50.00	
LITHIUM	12.7	12.5	2	50.00	
MAGNESIUM	3520	3760	7	50.00	
MANGANESE	148	160	8	50.00	
NICKEL	8.12	9.08	11	50.00	
PHOSPHORUS	134	198	39	50.00	
POTASSIUM	2100	2510	18	50.00	
SODIUM	132	136	3	50.00	
TIN	2.97	2.85	4	50.00	
TITANIUM	989	1070	8	50.00	
VANADIUM	33.1	32.7	1	50.00	
ZINC	45.6	49.9	9	50.00	
Zirconium	4.11	3.96	4	50.00	
MOLYBDENUM	0.261	2.00 U	200	50.00	J(all detects) UJ(all non-detects)

Method: 6020A

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-560-SA5D-SB-4.0-5.0 (TOT)	SL-860-SA5D-SB-4.0-5.0 (TOT)			
STRONTIUM	13.2	14.4	9	50.00	No Qualifiers Applied
THALLIUM	0.224	0.291	26	50.00	

Method: 7199

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-560-SA5D-SB-4.0-5.0 (TOT)	SL-860-SA5D-SB-4.0-5.0 (TOT)			
HEXAVALENT CHROMIUM	0.33	0.34	3	50.00	No Qualifiers Applied

Field Duplicate RPD Report

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 8015M

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-560-SA5D-SB-4.0-5.0	SL-860-SA5D-SB-4.0-5.0			
GASOLINE RANGE ORGANICS (C5-C12)	0.4	1.0 U	200	50.00	J(all detects) UJ(all non-detects)

Method: 9045M

Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-560-SA5D-SB-4.0-5.0	SL-860-SA5D-SB-4.0-5.0			
PH	7.40	8.50	14	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 1613B

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-111213	1,2,3,4,6,7,8-HPCDD	JBQ	1.00	9.95	PQL	pg/L	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.885	9.95	PQL	pg/L	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.846	9.95	PQL	pg/L	
	1,2,3,4,7,8-HXCDF	JBQ	0.641	9.95	PQL	pg/L	
	1,2,3,6,7,8-HXCDD	JBQ	0.661	9.95	PQL	pg/L	
	1,2,3,6,7,8-HXCDF	JBQ	0.622	9.95	PQL	pg/L	
	1,2,3,7,8,9-HXCDF	JBQ	0.756	9.95	PQL	pg/L	
	1,2,3,7,8-PECDD	JB	1.28	9.95	PQL	pg/L	
	1,2,3,7,8-PECDF	JBQ	0.895	9.95	PQL	pg/L	
	2,3,4,6,7,8-HXCDF	JBQ	0.664	9.95	PQL	pg/L	
	2,3,4,7,8-PECDF	JBQ	0.345	9.95	PQL	pg/L	
	2,3,7,8-TCDD	JBQ	0.717	1.99	PQL	pg/L	
	OCDD	JBQ	0.851	19.9	PQL	pg/L	
	OCDF	JBQ	1.81	19.9	PQL	pg/L	

Method: 6010C

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-111213	BARIUM	J	0.00034	0.0100	PQL	mg/L	J (all detects)
	CALCIUM	J	0.0349	0.400	PQL	mg/L	
	MAGNESIUM	J	0.0168	0.200	PQL	mg/L	

Method: 8270D SIM

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-111213	Diethylphthalate	J	0.37	1.0	PQL	ug/L	J (all detects)
	Di-n-butylphthalate	J	0.19	1.0	PQL	ug/L	

Reporting Limit Outliers

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-549-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	2.13	5.28	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.564	5.28	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.120	5.28	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0851	5.28	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0668	5.28	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.741	5.28	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.120	5.28	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	1.15	5.28	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	1.60	5.28	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.148	5.28	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.609	5.28	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.114	5.28	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.108	5.28	PQL	ng/Kg	
	OCDF	JBQ	1.37	10.6	PQL	ng/Kg	
SL-549-SA5D-SB-11.0-12.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.0901	5.53	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0689	5.53	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0647	5.53	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0704	5.53	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.143	5.53	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.0868	5.53	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.117	5.53	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.0673	5.53	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.115	5.53	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.194	5.53	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.282	5.53	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.0818	5.53	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.251	5.53	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.117	1.11	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0622	1.11	PQL	ng/Kg	
	OCDD	JB	0.326	11.1	PQL	ng/Kg	
OCDF	JBQ	0.267	11.1	PQL	ng/Kg		
SL-549-SA5D-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	0.521	5.61	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.158	5.61	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0547	5.61	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0601	5.61	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.107	5.61	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.136	5.61	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.100	5.61	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.0891	5.61	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.200	5.61	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.204	5.61	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.223	5.61	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.0671	5.61	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.153	5.61	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.0685	1.12	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.0495	1.12	PQL	ng/Kg	
	OCDD	JB	4.58	11.2	PQL	ng/Kg	
	OCDF	JB	0.660	11.2	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-560-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.230	5.34	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0577	5.34	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.136	5.34	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0548	5.34	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.129	5.34	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.0729	5.34	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.145	5.34	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.298	5.34	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.0938	5.34	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.129	5.34	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.102	5.34	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0461	5.34	PQL	ng/Kg	
	OCDD	JB	0.895	10.7	PQL	ng/Kg	
	OCDF	JB	0.413	10.7	PQL	ng/Kg	

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-549-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.839	1.05	PQL	mg/Kg	J (all detects)
	BORON	J	3.76	10.5	PQL	mg/Kg	
	MOLYBDENUM	J	0.429	2.10	PQL	mg/Kg	
	SODIUM	J	88.5	105	PQL	mg/Kg	
	TIN	J	2.91	10.5	PQL	mg/Kg	
SL-549-SA5D-SB-11.0-12.0	BERYLLIUM	J	0.793	1.07	PQL	mg/Kg	J (all detects)
	BORON	J	2.24	10.7	PQL	mg/Kg	
	MOLYBDENUM	J	0.262	2.14	PQL	mg/Kg	
	TIN	J	2.76	10.7	PQL	mg/Kg	
	Zirconium	J	4.10	5.35	PQL	mg/Kg	
SL-549-SA5D-SB-4.0-5.0	BERYLLIUM	J	0.972	1.09	PQL	mg/Kg	J (all detects)
	BORON	J	3.24	10.9	PQL	mg/Kg	
	MOLYBDENUM	J	0.314	2.19	PQL	mg/Kg	
	TIN	J	3.16	10.9	PQL	mg/Kg	
SL-560-SA5D-SB-0.0-0.5	BORON	J	8.20	10.8	PQL	mg/Kg	J (all detects)
	MOLYBDENUM	J	0.227	2.17	PQL	mg/Kg	
	TIN	J	3.29	10.8	PQL	mg/Kg	
SL-560-SA5D-SB-4.0-5.0	BERYLLIUM	J	0.668	1.00	PQL	mg/Kg	J (all detects)
	BORON	J	1.90	10.0	PQL	mg/Kg	
	MOLYBDENUM	J	0.261	2.00	PQL	mg/Kg	
	TIN	J	2.97	10.0	PQL	mg/Kg	
	Zirconium	J	4.11	5.01	PQL	mg/Kg	
SL-560-SA5D-SB-7.0-8.0	BERYLLIUM	J	0.601	0.994	PQL	mg/Kg	J (all detects)
	BORON	J	1.76	9.94	PQL	mg/Kg	
	MOLYBDENUM	J	0.213	1.99	PQL	mg/Kg	
	TIN	J	2.36	9.94	PQL	mg/Kg	
	Zirconium	J	3.30	4.97	PQL	mg/Kg	
SL-860-SA5D-SB-4.0-5.0	BERYLLIUM	J	0.618	1.00	PQL	mg/Kg	J (all detects)
	BORON	J	1.94	10.0	PQL	mg/Kg	
	TIN	J	2.85	10.0	PQL	mg/Kg	
	Zirconium	J	3.96	5.01	PQL	mg/Kg	

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Reporting Limit Outliers

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-549-SA5D-SB-0.0-0.5	SELENIUM	J	0.234	0.421	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0281	0.210	PQL	mg/Kg	
SL-549-SA5D-SB-11.0-12.0	SILVER	J	0.0425	0.214	PQL	mg/Kg	J (all detects)
SL-549-SA5D-SB-4.0-5.0	SELENIUM	J	0.123	0.437	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0498	0.219	PQL	mg/Kg	
SL-560-SA5D-SB-0.0-0.5	SELENIUM	J	0.231	0.434	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0568	0.217	PQL	mg/Kg	

Method: 7199

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-560-SA5D-SB-4.0-5.0	HEXAVALENT CHROMIUM	J	0.33	0.42	PQL	mg/Kg	J (all detects)
SL-560-SA5D-SB-7.0-8.0	HEXAVALENT CHROMIUM	J	0.32	0.41	PQL	mg/Kg	J (all detects)
SL-860-SA5D-SB-4.0-5.0	HEXAVALENT CHROMIUM	J	0.34	0.43	PQL	mg/Kg	J (all detects)

Method: 7471B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-549-SA5D-SB-11.0-12.0	MERCURY	J	0.0124	0.0176	PQL	mg/Kg	J (all detects)

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-560-SA5D-SB-4.0-5.0	GASOLINE RANGE ORGANICS (C5-C12)	J	0.4	1.1	PQL	mg/Kg	J (all detects)
SL-560-SA5D-SB-7.0-8.0	GASOLINE RANGE ORGANICS (C5-C12)	J	0.3	1	PQL	mg/Kg	J (all detects)

Method: 8081B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-549-SA5D-SB-0.0-0.5	4,4'-DDE	J	0.49	1.8	PQL	ug/Kg	J (all detects)
	4,4'-DDT	J	1.3	1.8	PQL	ug/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH127

Laboratory: LL

EDD Filename: PH127_v1.

eQAPP Name: CDM_SSFL_140113_Lan

Method: 8270D SIM

Matrix: SO

<i>SampleID</i>	<i>Analyte</i>	<i>Lab Qual</i>	<i>Result</i>	<i>Reporting Limit</i>	<i>RL Type</i>	<i>Units</i>	<i>Flag</i>
SL-549-SA5D-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	1.3	1.8	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	1.1	1.8	PQL	ug/Kg	

LDC #: 31098A4

VALIDATION COMPLETENESS WORKSHEET

Date: 1/17/13

SDG #: PH127

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: [Signature]

2nd Reviewer: _____

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 11/12/13
II.	ICP/MS Tune	+	
III.	Calibration	-	
IV.	Blanks	SW	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	SW	MS/D
VII.	Duplicate Sample Analysis	SW	DUP
VIII.	Laboratory Control Samples (LCS)	A	
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	A	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	-	(3,5)
XV.	Field Blanks	SW	EB=1 FB=FB-011613

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

(PH032)

Validated Samples: soil/water

1	EB1-111213	11	SL-560-SA5D-SB-4.0-5.0DUP	21	31
2	SL-560-SA5D-SB-0.0-0.5	12		22	32
3	SL-560-SA5D-SB-4.0-5.0	13		23	33
4	SL-560-SA5D-SB-7.0-8.0	14		24	34
5	SL-860-SA5D-SB-4.0-5.0	15		25	35
6	SL-549-SA5D-SB-0.0-0.5	16		26	36
7	SL-549-SA5D-SB-4.0-5.0	17		27	37
8	SL-549-SA5D-SB-11.0-12.0	18		28	38
9	SL-560-SA5D-SB-4.0-5.0MS	19		29	39
10	SL-560-SA5D-SB-4.0-5.0MSD	20		30	40

Notes: _____

VALIDATION FINDINGS WORKSHEET
PB/ICB/CCB QUALIFIED SAMPLES

METHOD: Trace metals (EPA SW 864 Method 6010B/6020/7000)

Soil preparation factor applied: _____

Reason: B

Sample Concentration units, unless otherwise noted: mg/L

Associated Samples: All Water

					Sample Identification											
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/L)	Maximum ICB/CCB ^a (ug/L)	Action Level	1											
Mg			34.0	0.17	0.0168											

Samples with analyte concentrations within five times the associated ICB, CCB or PB concentration are listed above with the identifications from the Validation Completeness Worksheet. These sample results were qualified as not detected, "U".

Note : a - The listed analyte concentration is the highest ICB, CCB, or PB detected in the analysis of each element.

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/16/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All Soil

Analyte	Blank ID	Sample Identification												
	FB-041613 (SDG: PH032)	Action Limit	2	3	4	6	7	8						
Mo	0.0132	6.60	0.227	0.262	0.213	0.429	0.314	0.262						
Sn	0.0029	1.45												

Sampling date: 9/25/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All Soil

Analyte	Blank ID	Sample Identification												
	EB1-111213 (SDG: PH127)	Action Limit	No Qualifiers											
Ba	0.00034	0.17												
Ca	0.0349	17.45												
Mg	0.0168	8.4												

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

74X

Al, Fe, Mg out
by RPD (J/S)

QUALITY ASSURANCE SUMMARY

FORM 5A (MS/MSD)

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

SDG No.: PH127

Matrix: SOIL

Level

(low/med):

LOW

Background Lab Sample ID: 7275663BKG Matrix Spike Lab Sample ID: 7275664MS Matrix Spike Duplicate Lab Sample ID: 7275665MSD

Batch Id(s): P32537C, P32538A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit		M		
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q		%R	RPD
Aluminum		13981.8359		13748.3850		18656.2942		200.0000	192.3077	MG/KG	-117		2431		30		20	P	
Antimony		0.7184	U	31.0450		22.9019		50.0000	48.0769	MG/KG	62	N	48	N	30	*	75 - 125	20	P
Arsenic		5.4699		19.2540		24.0308		15.0000	14.4231	MG/KG	92		129	N	22	*	75 - 125	20	P
Barium		59.7456		264.7610		279.5750		200.0000	192.3077	MG/KG	103		114		5		75 - 125	20	P
Beryllium		0.6476	B	5.6920		5.8875		5.0000	4.8077	MG/KG	101		109		3		75 - 125	20	P
Boron		1.8379	B	198.4720		191.4298		200.0000	192.3077	MG/KG	98		99		4		75 - 125	20	P
Cadmium		0.0738	U	4.8280		4.4731		5.0000	4.8077	MG/KG	97		93		8		75 - 125	20	P
Calcium		2195.6961		2719.3290		5242.4529		400.0000	384.6154	MG/KG	131		792		63			20	P
Chromium		15.8466		33.5030		43.4865		20.0000	19.2308	MG/KG	88		144	N	26	*	75 - 125	20	P
Cobalt		3.0845		52.0790		50.7115		50.0000	48.0769	MG/KG	98		99		3		75 - 125	20	P
Copper		5.7553		28.9250		36.2760		25.0000	24.0385	MG/KG	93		127	N	23	*	75 - 125	20	P
Iron		20556.9563		18838.1370		29858.8365		100.0000	96.1538	MG/KG	-1719		9674		45			20	P
Lead		6.1320		19.2830		24.4962		15.0000	14.4231	MG/KG	88		127	N	24	*	75 - 125	20	P
Lithium		12.2864		113.4940		114.0663		100.0000	96.1538	MG/KG	101		106		1		75 - 125	20	P
Magnesium		3407.0524		3798.5880		5587.1740		200.0000	192.3077	MG/KG	196		1134		88			20	P
Manganese		143.0126		205.0920		235.2606		50.0000	48.0769	MG/KG	124		192	N	14		75 - 125	20	P
Mercury		0.0098	U	0.1748		0.1682		0.1629	0.1593	MG/KG	107		106		4		65 - 135	20	CV
Molybdenum		0.2524	B	190.6210		179.9683		200.0000	192.3077	MG/KG	95		93		6		75 - 125	20	P
Nickel		7.8689		56.3590		59.2019		50.0000	48.0769	MG/KG	97		107		5		75 - 125	20	P
Phosphorus		129.6689		389.1840		261.2423		100.0000	96.1538	MG/KG	260	N	137	N	39	*	75 - 125	20	P
Potassium		2036.5107		3650.8770		3876.6019		1000.0000	961.5385	MG/KG	161	N	191	N	6		75 - 125	20	P
Selenium	78	0.0971	U	2.2712		2.2165		2.0000	1.9231	MG/KG	114		115		2		75 - 125	20	MS
Silver	107	0.0252	U	11.1060		9.9135		10.0000	9.6154	MG/KG	111		103		11		75 - 125	20	MS
Sodium		128.1476		1089.7000		1126.9308		1000.0000	961.5385	MG/KG	96		104		3		75 - 125	20	P
Strontium	88	12.7883		22.0000		30.9615		8.0000	7.6923	MG/KG	115		236	N	34	*	75 - 125	20	MS
Thallium	203	0.2173		0.6452		0.7604		0.4000	0.3846	MG/KG	107		141	N	16		75 - 125	20	MS

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor</p> <p>MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U= Below MDL, B= Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>
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Lancaster
Laboratories

74x

QUALITY ASSURANCE SUMMARY
FORM 5A (MS/MSD)
MATRIX SPIKE/MATRIX SPIKE DUPLICATE
SDG No.: PH127
Matrix: SOIL Level (low/med): LOW

Background Lab Sample ID: 7275663BKG Matrix Spike Lab Sample ID: 7275664MS Matrix Spike Duplicate Lab Sample ID: 7275665MSD
Batch Id(s): P32537C, P32538A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit				
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q	%R	RPD	M
Thallium	203	0.2173		0.6452		0.7604		0.4000	0.3846	MG/KG	107		141	N	16	75 - 125	20	MS	
Tin		2.8748	B	364.0900		338.7173		400.0000	384.6154	MG/KG	90		87		7	75 - 125	20	P	
Titanium		957.8806		1217.9040		1384.8683		100.0000	96.1538	MG/KG	260		444		13			20	P
Vanadium		32.0680		79.8930		95.6635		50.0000	48.0769	MG/KG	96		132	N	18	75 - 125	20	P	
Zinc		44.2058		98.9000		114.6510		50.0000	48.0769	MG/KG	109		147	N	15	75 - 125	20	P	
Zirconium		3.9835	B	101.6370		98.7712		100.0000	96.1538	MG/KG	98		99		3	75 - 125	20	P	

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U = Below MDL, B = Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>
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Background Lab Sample ID: 7275663BKG
 Batch ID(s): P32537C, P32538A
 Concentration Units: MG/KG

Duplicate Lab Sample ID: 7275669DUP

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			13981.8359		13961.2431		0		P
Antimony			0.7184	U	0.7255	U			P
Arsenic		3.9	5.4699		4.9804		9		P
Barium			59.7456		59.5814		0		P
Beryllium			0.6476	B	0.6206	B	4		P
Boron			1.8379	B	1.7294	B	6		P
Cadmium			0.0738	U	0.0745	U			P
Calcium			2195.6961		2162.1451		2		P
Chromium			15.8466		15.8108		0		P
Cobalt		1.0	3.0845		3.1833		3		P
Copper		1.9	5.7553		5.6108		3		P
Iron			20556.9563		20154.6010		2		P
Lead		2.9	6.1320		5.7853		6		P
Lithium		3.9	12.2864		12.4235		1		P
Magnesium			3407.0524		3402.8127		0		P
Manganese			143.0126		149.2167		4		P
Mercury			0.0098	U	0.0097	U			CV
Molybdenum			0.2524	B	0.1667	U	200		P
Nickel		1.9	7.8689		7.9275		1		P
Phosphorus			129.6689		130.5373		1		P
Potassium			2036.5107		2144.0049		5		P
Selenium	78		0.0971	U	0.0980	U			MS
Silver	107		0.0252	U	0.0255	U			MS
Sodium		97.1	128.1476		130.3686		2		P
Strontium	88		12.7883		13.8392		8		MS
Thallium	203	0.2	0.2173		0.2357		8		MS
Tin			2.8748	B	2.9147	B	1		P
Titanium			957.8806		989.9784		3		P
Vanadium			32.0680		31.4922		2		P
Zinc			44.2058		43.3049		2		P
Zirconium			3.9835	B	4.4422	B	11		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).
 The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

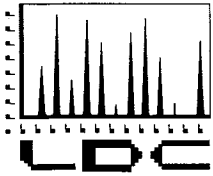
ok by difference

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: * = Duplicate Out of Spec
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SAMPLE DELIVERY GROUP

PH130



LABORATORY DATA CONSULTANTS, INC.

2701 Loker Ave. West, Suite 220, Carlsbad, CA 92010 Bus: 760-827-1100 Fax: 760-827-1099

CDM
555 17th Street, Suite 1100
Denver, CO 80202
ATTN: Mrs. Cherie Zakowski

January 24, 2014

SUBJECT: Santa Susana Field Laboratory, Subarea 5D Data Validation

Dear Mrs. Zakowski,

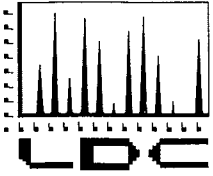
Enclosed is the final validation report for the fractions listed below. This SDG was received on January 3, 2014. Attachment 1 is a summary of the samples that were reviewed for each analysis.

LDC Project # 31106:

<u>SDG #</u>	<u>Fraction</u>
PH130	Polynuclear Aromatic Hydrocarbons, Polychlorinated Biphenyls, Metals, Herbicides, Wet Chemistry, Total Petroleum Hydrocarbons as Gasoline, Total Petroleum Hydrocarbons as Extractables, Dioxins/Dibenzofurans

The data validation was performed under Level III guidelines. The analyses were validated using the following documents, as applicable to each method:

- Quality Assurance Project Plan for Santa Susana Field Laboratory, RCRA Facility Investigation, Surficial Media Operable Unit, March 2009, Revision 4
- USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review, June 2008
- USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review, January 2010
- Polychlorinated Dioxins/Dibenzofurans Data Review, September 2005
- EPA SW 846, Third Edition, Test Methods for Evaluating Solid Waste, update 1, July 1992; update IIA, August 1993; update II, September 1994; update IIB, January 1995; update III, December 1996; update IIIA, April 1998; IIIB, November 2004; Update IV, February 2007



Please feel free to contact us if you have any questions.

Sincerely,

Shauna McKellar
Project Manager/Chemist

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 5D

SDG: PH130

Prepared for

CDM Smith
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
2701 Loker Ave West, Suite 220
Carlsbad, California 92010

January 24, 2014

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on November 15, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan (QAPP) for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOCs) by Environmental Protection Agency (EPA) SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)

Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A

Herbicides by EPA SW 846 Method 8151A

Metals by EPA SW 846 Method 6010C, 6020A and 7471B

Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M

TPH as Extractables by EPA SW 846 Method 8015M

Dioxins and Dibenzofurans by EPA Method 1613B

Wet Chemistry:

Hexavalent Chromium by EPA Method 7199

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks and field duplicate samples. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met with the exception of four samples for TPH as gasoline. The associated sample results were qualified as non-detected estimated (UJ). The details regarding the qualification of data are provided in Enclosure I.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for metals and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks with the following exceptions:

SDG/ Method	Method Blank ID	Analyte	Maximum Concentration	Associated Samples
PH130/ 6020A	ICB/CCB	Silver	0.25 ug/L	All samples in SDG PH130
PH130/ 6020A	ICB/CCB	Thallium	0.35 ug/L	All samples in SDG PH130

Sample concentrations were compared to concentrations detected in the initial and continuing blanks. The sample concentrations were not detected or were significantly greater than the concentrations found in the associated blanks with the following exceptions:

SDG/ Method	Sample	Compound	Reported Concentration	Modified Final Concentration
PH130/ 6020A	SL-516-SA5D-SB-0.0-0.5	Silver Thallium	0.0478 mg/kg 0.322 mg/kg	0.0478U mg/kg 0.322U mg/kg
PH130/ 6020A	SL-516-SA5D-SB-4.0-5.0	Silver Thallium	0.0648 mg/kg 0.301 mg/kg	0.0648U mg/kg 0.301U mg/kg

SDG/ Method	Sample	Compound	Reported Concentration	Modified Final Concentration
PH130/ 6020A	SL-816-SA5D-SB-4.0-5.0	Silver Thallium	0.0614 mg/kg 0.289 mg/kg	0.0614U mg/kg 0.289U mg/kg
PH130/ 6020A	SL-516-SA5D-SB-9.0-10.0	Silver Thallium	0.0472 mg/kg 0.292 mg/kg	0.0472U mg/kg 0.292U mg/kg
PH130/ 6020A	SL-516-SA5D-SB-13.0-14.0	Thallium	0.287 mg/kg	0.287U mg/kg
PH130/ 6020A	SL-609-SA5D-SB-0.0-0.5	Silver Thallium	0.0283 mg/kg 0.266 mg/kg	0.0283U mg/kg 0.266U mg/kg

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits with the exception of one sample for PCBs. No data were qualified due to high %Rs since the associated results were non-detected.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits with the exception of one DUP for metals. The associated sample results were qualified as detected estimated (J). The details regarding the qualification of data are provided in Enclosure I.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of several LCS/LCSD pairs for SVOCs and herbicides. The dinoseb result in samples SL-516-SA5D-SB-0.0-0.5, SL-516-SA5D-SB-13.0-14.0, SL-516-SA5D-SB-4.0-5.0, SL-516-SA5D-SB-9.0-10.0 and SL-816-SA5D-SB-4.0-5.0 were qualified as rejected (R) due to LCS/LCSD %Rs grossly outside of limits (i.e., $\leq 10\%$). The details regarding the qualification of data are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH130	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pair was collected and analyzed for SVOCs, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins and hexavalent chromium. All RPDs were within QC limits with the exception of several metals, dioxins and hexavalent chromium. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No contaminants were found in the trip blank.

One equipment blank (from SDG PH127) was collected and analyzed for SVOCs, PCBs, metals, herbicides, TPH as gasoline, TPH as extractables, dioxins and hexavalent chromium. The equipment blank had detections for SVOCs, metals and dioxins. The sample results were not detected or were significantly greater than the concentrations found in the equipment blank, therefore no data were qualified.

One field blank (from SDG PH032) was collected and analyzed for SVOCs, PCBs, herbicides, metals, TPH as gasoline, TPH as extractables, dioxins, and hexavalent chromium. The field blank had detections for SVOCs, metals and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

Five herbicide results were rejected due to LCS/LCSD %Rs grossly outside of QC limits. These results are not useable for all purposes.

The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. The remainder of the data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1

Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
15-Nov-2013	TB-111513	7280196	TB	5030B	8015M	III
15-Nov-2013	SL-516-SA5D-SB-0.0-0.5	7280197	N	3050B	6010C	III
15-Nov-2013	SL-516-SA5D-SB-0.0-0.5	7280197	N	3050B	6020A	III
15-Nov-2013	SL-516-SA5D-SB-0.0-0.5	7280197	N	3060A	7199	III
15-Nov-2013	SL-516-SA5D-SB-0.0-0.5	7280197	N	3546	8015M	III
15-Nov-2013	SL-516-SA5D-SB-0.0-0.5	7280197	N	3546	8082A	III
15-Nov-2013	SL-516-SA5D-SB-0.0-0.5	7280197	N	3546	8270D SIM	III
15-Nov-2013	SL-516-SA5D-SB-0.0-0.5	7280197	N	3550B	8151A	III
15-Nov-2013	SL-516-SA5D-SB-0.0-0.5	7280197	N	METHOD	1613B	III
15-Nov-2013	SL-516-SA5D-SB-0.0-0.5	7280197	N	METHOD	7471B	III
15-Nov-2013	SL-816-SA5D-SB-4.0-5.0	7280205	FD	3050B	6010C	III
15-Nov-2013	SL-816-SA5D-SB-4.0-5.0	7280205	FD	3050B	6020A	III
15-Nov-2013	SL-816-SA5D-SB-4.0-5.0	7280205	FD	3060A	7199	III
15-Nov-2013	SL-816-SA5D-SB-4.0-5.0	7280205	FD	3546	8015M	III
15-Nov-2013	SL-816-SA5D-SB-4.0-5.0	7280205	FD	3546	8082A	III
15-Nov-2013	SL-816-SA5D-SB-4.0-5.0	7280205	FD	3546	8270D SIM	III
15-Nov-2013	SL-816-SA5D-SB-4.0-5.0	7280205	FD	3550B	8151A	III
15-Nov-2013	SL-816-SA5D-SB-4.0-5.0	7280205	FD	5035A	8015M	III
15-Nov-2013	SL-816-SA5D-SB-4.0-5.0	7280205	FD	METHOD	1613B	III
15-Nov-2013	SL-816-SA5D-SB-4.0-5.0	7280205	FD	METHOD	7471B	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0	7280198	N	3050B	6010C	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0	7280198	N	3050B	6020A	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0	7280198	N	3060A	7199	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0	7280198	N	3546	8015M	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0	7280198	N	3546	8082A	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0	7280198	N	3546	8270D SIM	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0	7280198	N	3550B	8151A	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0	7280198	N	5035A	8015M	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0	7280198	N	METHOD	1613B	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0	7280198	N	METHOD	7471B	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280199	MS	3050B	6010C	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280199	MS	3050B	6020A	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280199	MS	3546	8015M	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280199	MS	3546	8082A	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280199	MS	3546	8270D SIM	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280199	MS	3550B	8151A	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280199	MS	5035A	8015M	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280199	MS	METHOD	1613B	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280199	MS	METHOD	7471B	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MSD	7280200	MSD	3050B	6010C	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MSD	7280200	MSD	3050B	6020A	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MSD	7280200	MSD	3546	8015M	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MSD	7280200	MSD	3546	8082A	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MSD	7280200	MSD	3546	8270D SIM	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MSD	7280200	MSD	3550B	8151A	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MSD	7280200	MSD	5035A	8015M	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MSD	7280200	MSD	METHOD	1613B	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MSD	7280200	MSD	METHOD	7471B	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280201	MS	3060A	7199	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0MS	7280202	MS	3060A	7199	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0DUP	7280204	DUP	3050B	6010C	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0DUP	7280204	DUP	3050B	6020A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0DUP	7280204	DUP	3060A	7199	III
15-Nov-2013	SL-516-SA5D-SB-4.0-5.0DUP	7280204	DUP	METHOD	7471B	III
15-Nov-2013	SL-516-SA5D-SB-9.0-10.0	7280206	N	3050B	6010C	III
15-Nov-2013	SL-516-SA5D-SB-9.0-10.0	7280206	N	3050B	6020A	III
15-Nov-2013	SL-516-SA5D-SB-9.0-10.0	7280206	N	3060A	7199	III
15-Nov-2013	SL-516-SA5D-SB-9.0-10.0	7280206	N	3546	8015M	III
15-Nov-2013	SL-516-SA5D-SB-9.0-10.0	7280206	N	3546	8082A	III
15-Nov-2013	SL-516-SA5D-SB-9.0-10.0	7280206	N	3546	8270D SIM	III
15-Nov-2013	SL-516-SA5D-SB-9.0-10.0	7280206	N	3550B	8151A	III
15-Nov-2013	SL-516-SA5D-SB-9.0-10.0	7280206	N	5035A	8015M	III
15-Nov-2013	SL-516-SA5D-SB-9.0-10.0	7280206	N	METHOD	1613B	III
15-Nov-2013	SL-516-SA5D-SB-9.0-10.0	7280206	N	METHOD	7471B	III
15-Nov-2013	SL-516-SA5D-SB-13.0-14.0	7280207	N	3050B	6010C	III
15-Nov-2013	SL-516-SA5D-SB-13.0-14.0	7280207	N	3050B	6020A	III
15-Nov-2013	SL-516-SA5D-SB-13.0-14.0	7280207	N	3060A	7199	III
15-Nov-2013	SL-516-SA5D-SB-13.0-14.0	7280207	N	3546	8015M	III
15-Nov-2013	SL-516-SA5D-SB-13.0-14.0	7280207	N	3546	8082A	III
15-Nov-2013	SL-516-SA5D-SB-13.0-14.0	7280207	N	3546	8270D SIM	III
15-Nov-2013	SL-516-SA5D-SB-13.0-14.0	7280207	N	3550B	8151A	III
15-Nov-2013	SL-516-SA5D-SB-13.0-14.0	7280207	N	5035A	8015M	III
15-Nov-2013	SL-516-SA5D-SB-13.0-14.0	7280207	N	METHOD	1613B	III
15-Nov-2013	SL-516-SA5D-SB-13.0-14.0	7280207	N	METHOD	7471B	III
15-Nov-2013	SL-609-SA5D-SB-0.0-0.5	7280208	N	3050B	6010C	III
15-Nov-2013	SL-609-SA5D-SB-0.0-0.5	7280208	N	3050B	6020A	III
15-Nov-2013	SL-609-SA5D-SB-0.0-0.5	7280208	N	3546	8015M	III
15-Nov-2013	SL-609-SA5D-SB-0.0-0.5	7280208	N	3546	8082A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
15-Nov-2013	SL-609-SA5D-SB-0.0-0.5	7280208	N	3546	8270D SIM	III
15-Nov-2013	SL-609-SA5D-SB-0.0-0.5	7280208	N	METHOD	1613B	III
15-Nov-2013	SL-609-SA5D-SB-0.0-0.5	7280208	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS

Method: 6010C

Matrix: SO

Sample ID: SL-516-SA5D-SB-0.0-0.5

Collected: 11/15/2013 8:35:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.19	U	0.776	MDL	4.19	PQL	mg/Kg	UJ	Q
BARIUM	118		0.0346	MDL	1.05	PQL	mg/Kg	J	Q
BERYLLIUM	0.899	J	0.0702	MDL	1.05	PQL	mg/Kg	J	Z
BORON	3.75	J	0.881	MDL	10.5	PQL	mg/Kg	J	Z, E
CALCIUM	3220		3.50	MDL	21.0	PQL	mg/Kg	J	E
MANGANESE	366		0.0870	MDL	1.05	PQL	mg/Kg	J	E
MOLYBDENUM	0.366	J	0.178	MDL	2.10	PQL	mg/Kg	U	F
TIN	2.94	J	0.231	MDL	10.5	PQL	mg/Kg	U	B
Zirconium	4.90	J	0.881	MDL	5.24	PQL	mg/Kg	J	Z
ALUMINUM	22800		7.56	MDL	41.9	PQL	mg/Kg	J	E

Sample ID: SL-516-SA5D-SB-13.0-14.0

Collected: 11/15/2013 9:08:00

Analysis Type: REA

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.00	U	0.739	MDL	4.00	PQL	mg/Kg	UJ	Q
MOLYBDENUM	0.509	J	0.170	MDL	2.00	PQL	mg/Kg	U	F

Sample ID: SL-516-SA5D-SB-13.0-14.0

Collected: 11/15/2013 9:08:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	15700		7.20	MDL	40.0	PQL	mg/Kg	J	E
BARIUM	77.0		0.0330	MDL	0.999	PQL	mg/Kg	J	Q
BERYLLIUM	0.650	J	0.0669	MDL	0.999	PQL	mg/Kg	J	Z
BORON	1.78	J	0.839	MDL	9.99	PQL	mg/Kg	J	Z, E
CALCIUM	4020		3.34	MDL	20.0	PQL	mg/Kg	J	E
MANGANESE	273		0.0829	MDL	0.999	PQL	mg/Kg	J	E
Zirconium	3.74	J	0.839	MDL	4.99	PQL	mg/Kg	J	Z

Sample ID: SL-516-SA5D-SB-4.0-5.0

Collected: 11/15/2013 8:45:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	26200		7.89	MDL	43.8	PQL	mg/Kg	J	E
ANTIMONY	4.38	U	0.810	MDL	4.38	PQL	mg/Kg	UJ	Q
BARIUM	136		0.0361	MDL	1.09	PQL	mg/Kg	J	Q
BERYLLIUM	0.984	J	0.0733	MDL	1.09	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-516-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BORON	2.58	J	0.919	MDL	10.9	PQL	mg/Kg	J	Z, E
CALCIUM	4330		3.65	MDL	21.9	PQL	mg/Kg	J	E
MANGANESE	425		0.0908	MDL	1.09	PQL	mg/Kg	J	E, FD
MOLYBDENUM	0.543	J	0.186	MDL	2.19	PQL	mg/Kg	UJ	FD, F
TIN	3.31	J	0.241	MDL	10.9	PQL	mg/Kg	U	B
Zirconium	4.99	J	0.919	MDL	5.47	PQL	mg/Kg	J	Z

Sample ID: SL-516-SA5D-SB-9.0-10.0 Collected: 11/15/2013 8:56:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	23900		7.78	MDL	43.1	PQL	mg/Kg	J	E
ANTIMONY	4.31	U	0.798	MDL	4.31	PQL	mg/Kg	UJ	Q
BARIUM	97.3		0.0356	MDL	1.08	PQL	mg/Kg	J	Q
BERYLLIUM	1.01	J	0.0723	MDL	1.08	PQL	mg/Kg	J	Z
BORON	1.67	J	0.906	MDL	10.8	PQL	mg/Kg	J	Z, E
CALCIUM	2990		3.60	MDL	21.6	PQL	mg/Kg	J	E
MANGANESE	274		0.0895	MDL	1.08	PQL	mg/Kg	J	E
MOLYBDENUM	0.366	J	0.183	MDL	2.16	PQL	mg/Kg	U	F
TIN	3.07	J	0.237	MDL	10.8	PQL	mg/Kg	U	B
Zirconium	4.73	J	0.906	MDL	5.39	PQL	mg/Kg	J	Z

Sample ID: SL-609-SA5D-SB-0.0-0.5 Collected: 11/15/2013 12:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	14700		7.51	MDL	41.7	PQL	mg/Kg	J	E
ANTIMONY	4.17	U	0.771	MDL	4.17	PQL	mg/Kg	UJ	Q
BARIUM	96.8		0.0344	MDL	1.04	PQL	mg/Kg	J	Q
BERYLLIUM	0.600	J	0.0698	MDL	1.04	PQL	mg/Kg	J	Z
BORON	1.27	J	0.875	MDL	10.4	PQL	mg/Kg	J	Z, E
CALCIUM	3700		3.48	MDL	20.8	PQL	mg/Kg	J	E
MANGANESE	333		0.0865	MDL	1.04	PQL	mg/Kg	J	E
TIN	2.57	J	0.229	MDL	10.4	PQL	mg/Kg	U	B
Zirconium	4.77	J	0.875	MDL	5.21	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS
Method: 6010C Matrix: SO

Sample ID: SL-816-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ALUMINUM	24400		7.80	MDL	43.2	PQL	mg/Kg	J	E
ANTIMONY	4.32	U	0.800	MDL	4.32	PQL	mg/Kg	UJ	Q
BARIUM	106		0.0357	MDL	1.08	PQL	mg/Kg	J	Q
BERYLLIUM	1.01	J	0.0724	MDL	1.08	PQL	mg/Kg	J	Z
BORON	1.71	J	0.908	MDL	10.8	PQL	mg/Kg	J	Z, E
CALCIUM	3670		3.61	MDL	21.6	PQL	mg/Kg	J	E
MANGANESE	243		0.0897	MDL	1.08	PQL	mg/Kg	J	E, FD
MOLYBDENUM	0.294	J	0.184	MDL	2.16	PQL	mg/Kg	UJ	FD, F
TIN	3.07	J	0.238	MDL	10.8	PQL	mg/Kg	U	B
Zirconium	5.10	J	0.908	MDL	5.41	PQL	mg/Kg	J	Z

Method Category: METALS
Method: 6020A Matrix: SO

Sample ID: SL-516-SA5D-SB-0.0-0.5 Collected: 11/15/2013 8:35:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.253	J	0.105	MDL	0.419	PQL	mg/Kg	J	Z

Sample ID: SL-516-SA5D-SB-0.0-0.5 Collected: 11/15/2013 8:35:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0478	J	0.0273	MDL	0.210	PQL	mg/Kg	U	B
STRONTIUM	25.7		0.0713	MDL	0.419	PQL	mg/Kg	J	Q, E
THALLIUM	0.322		0.0315	MDL	0.210	PQL	mg/Kg	U	B

Sample ID: SL-516-SA5D-SB-13.0-14.0 Collected: 11/15/2013 9:08:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.160	J	0.0999	MDL	0.400	PQL	mg/Kg	J	Z

Sample ID: SL-516-SA5D-SB-13.0-14.0 Collected: 11/15/2013 9:08:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	19.3		0.0679	MDL	0.400	PQL	mg/Kg	J	Q, E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-516-SA5D-SB-13.0-14.0 Collected: 11/15/2013 9:08:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
THALLIUM	0.287		0.0300	MDL	0.200	PQL	mg/Kg	U	B

Sample ID: SL-516-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:45:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.167	J	0.109	MDL	0.438	PQL	mg/Kg	J	Z

Sample ID: SL-516-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:45:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0648	J	0.0284	MDL	0.219	PQL	mg/Kg	U	B
STRONTIUM	27.8		0.0744	MDL	0.438	PQL	mg/Kg	J	Q, E
THALLIUM	0.301		0.0328	MDL	0.219	PQL	mg/Kg	U	B

Sample ID: SL-516-SA5D-SB-9.0-10.0 Collected: 11/15/2013 8:56:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.146	J	0.108	MDL	0.431	PQL	mg/Kg	J	Z

Sample ID: SL-516-SA5D-SB-9.0-10.0 Collected: 11/15/2013 8:56:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0472	J	0.0280	MDL	0.216	PQL	mg/Kg	U	B
STRONTIUM	23.0		0.0733	MDL	0.431	PQL	mg/Kg	J	Q, E
THALLIUM	0.292		0.0324	MDL	0.216	PQL	mg/Kg	U	B

Sample ID: SL-609-SA5D-SB-0.0-0.5 Collected: 11/15/2013 12:50:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0283	J	0.0271	MDL	0.208	PQL	mg/Kg	U	B
STRONTIUM	20.2		0.0708	MDL	0.417	PQL	mg/Kg	J	Q, E
THALLIUM	0.266		0.0313	MDL	0.208	PQL	mg/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-816-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:40:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.147	J	0.108	MDL	0.432	PQL	mg/Kg	J	Z

Sample ID: SL-816-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:40:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0614	J	0.0281	MDL	0.216	PQL	mg/Kg	U	B
STRONTIUM	25.9		0.0735	MDL	0.432	PQL	mg/Kg	J	Q, E
THALLIUM	0.289		0.0324	MDL	0.216	PQL	mg/Kg	U	B

Method Category: METALS
Method: 7199 **Matrix:** SO

Sample ID: SL-516-SA5D-SB-13.0-14.0 Collected: 11/15/2013 9:08:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.26	J	0.14	MDL	0.41	PQL	mg/Kg	J	Z

Sample ID: SL-516-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.44	U	0.15	MDL	0.44	PQL	mg/Kg	UJ	FD

Sample ID: SL-816-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.16	J	0.16	MDL	0.45	PQL	mg/Kg	J	Z, FD

Method Category: METALS
Method: 7471B **Matrix:** SO

Sample ID: SL-516-SA5D-SB-13.0-14.0 Collected: 11/15/2013 9:08:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0103	J	0.0101	MDL	0.0168	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA

Method: 1613B

Matrix: SO

Sample ID: SL-516-SA5D-SB-0.0-0.5

Collected: 11/15/2013 8:35:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.330	JB	0.0482	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0969	JB	0.0183	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.105	JB	0.0359	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0427	JB	0.0386	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0747	JB	0.0348	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.271	JBQ	0.0415	MDL	5.31	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0549	JQ	0.0305	MDL	5.31	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.421	JB	0.0369	MDL	5.31	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.677	J	0.0397	MDL	5.31	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.102	JQ	0.0539	MDL	5.31	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.168	JBQ	0.0317	MDL	5.31	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.118	JBQ	0.0312	MDL	5.31	PQL	ng/Kg	U	B
OCDD	3.39	JB	0.0518	MDL	10.6	PQL	ng/Kg	U	B
OCDF	0.365	JB	0.0727	MDL	10.6	PQL	ng/Kg	U	B

Sample ID: SL-516-SA5D-SB-13.0-14.0

Collected: 11/15/2013 9:08:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.137	JBQ	0.0277	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0921	JBQ	0.0121	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0501	JB	0.0183	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0505	JBQ	0.0226	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0758	JB	0.0197	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0367	JBQ	0.0244	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0515	J	0.0178	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.0443	JBQ	0.0226	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0384	J	0.0198	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0548	JQ	0.0358	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.0757	JBQ	0.0234	MDL	5.10	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0484	JB	0.0186	MDL	5.10	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0686	JBQ	0.0222	MDL	5.10	PQL	ng/Kg	U	B
OCDD	1.40	JB	0.0320	MDL	10.2	PQL	ng/Kg	U	B
OCDF	0.156	JBQ	0.0363	MDL	10.2	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-516-SA5D-SB-4.0-5.0	Collected: 11/15/2013 8:45:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.313	JB	0.0470	MDL	5.44	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.112	JBQ	0.0203	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,4,7,8,9-HPCDF	0.0665	JBQ	0.0339	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,4,7,8-HxCDD	0.0899	JBQ	0.0422	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,4,7,8-HXCDF	0.122	JBQ	0.0376	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,6,7,8-HXCDD	0.0667	JB	0.0441	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,6,7,8-HXCDF	0.0446	JQ	0.0344	MDL	5.44	PQL	ng/Kg	J	Z, FD
1,2,3,7,8,9-HXCDD	0.121	JBQ	0.0435	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,7,8,9-HXCDF	0.146	JQ	0.0466	MDL	5.44	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0967	JQ	0.0596	MDL	5.44	PQL	ng/Kg	J	Z, FD
1,2,3,7,8-PECDF	0.136	JB	0.0358	MDL	5.44	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0666	JBQ	0.0376	MDL	5.44	PQL	ng/Kg	UJ	B, FD
2,3,4,7,8-PECDF	0.0691	JB	0.0346	MDL	5.44	PQL	ng/Kg	UJ	B, FD
2,3,7,8-TCDD	1.09	U	0.0787	MDL	1.09	PQL	ng/Kg	UJ	FD
2,3,7,8-TCDF	1.09	U	0.0659	MDL	1.09	PQL	ng/Kg	UJ	FD
OCDD	1.85	JB	0.0549	MDL	10.9	PQL	ng/Kg	UJ	B, FD
OCDF	0.281	JB	0.0769	MDL	10.9	PQL	ng/Kg	U	B

Sample ID: SL-516-SA5D-SB-9.0-10.0	Collected: 11/15/2013 8:56:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.846	JB	0.0439	MDL	5.35	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.216	JB	0.0182	MDL	5.35	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.107	JBQ	0.0282	MDL	5.35	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0581	JBQ	0.0317	MDL	5.35	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.149	JBQ	0.0347	MDL	5.35	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.173	JBQ	0.0328	MDL	5.35	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.122	JQ	0.0295	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.216	JB	0.0307	MDL	5.35	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.205	JQ	0.0360	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0896	JQ	0.0446	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.303	JB	0.0329	MDL	5.35	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.133	JB	0.0310	MDL	5.35	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.122	JBQ	0.0329	MDL	5.35	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA

Method: 1613B

Matrix: SO

Sample ID: SL-516-SA5D-SB-9.0-10.0

Collected: 11/15/2013 8:56:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,3,7,8-TCDF	0.0992	JB	0.0672	MDL	1.07	PQL	ng/Kg	U	B
OCDD	9.55	JB	0.0407	MDL	10.7	PQL	ng/Kg	J	Z
OCDF	0.601	JBQ	0.0463	MDL	10.7	PQL	ng/Kg	U	B

Sample ID: SL-609-SA5D-SB-0.0-0.5

Collected: 11/15/2013 12:50:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.580	JBQ	0.0278	MDL	5.17	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.544	JB	0.0136	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.439	JB	0.0278	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.452	JB	0.0364	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDF	0.393	JB	0.0345	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.397	JB	0.0390	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.481	JQ	0.0275	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.524	JB	0.0388	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDF	0.492	J	0.0402	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.414	J	0.0422	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.501	JB	0.0298	MDL	5.17	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HxCDF	0.418	JB	0.0305	MDL	5.17	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.336	JBQ	0.0310	MDL	5.17	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0640	JBQ	0.0555	MDL	1.03	PQL	ng/Kg	U	B
OCDD	1.49	JB	0.0322	MDL	10.3	PQL	ng/Kg	U	B
OCDF	1.05	JBQ	0.0488	MDL	10.3	PQL	ng/Kg	U	B

Sample ID: SL-816-SA5D-SB-4.0-5.0

Collected: 11/15/2013 8:40:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.280	JB	0.0335	MDL	5.44	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.236	JB	0.0182	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,4,7,8,9-HPCDF	0.205	JB	0.0269	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,4,7,8-HxCDD	0.195	JBQ	0.0367	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,4,7,8-HxCDF	0.213	JBQ	0.0337	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,6,7,8-HxCDD	0.228	JBQ	0.0414	MDL	5.44	PQL	ng/Kg	UJ	B, FD
1,2,3,6,7,8-HxCDF	0.195	J	0.0311	MDL	5.44	PQL	ng/Kg	J	Z, FD
1,2,3,7,8,9-HxCDD	0.224	JBQ	0.0374	MDL	5.44	PQL	ng/Kg	UJ	B, FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-816-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,7,8,9-HXCDF	0.222	JQ	0.0342	MDL	5.44	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.172	J	0.0516	MDL	5.44	PQL	ng/Kg	J	Z, FD
1,2,3,7,8-PECDF	0.214	JB	0.0313	MDL	5.44	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.224	JB	0.0315	MDL	5.44	PQL	ng/Kg	UJ	B, FD
2,3,4,7,8-PECDF	0.223	JBQ	0.0342	MDL	5.44	PQL	ng/Kg	UJ	B, FD
2,3,7,8-TCDD	0.111	JBQ	0.0693	MDL	1.09	PQL	ng/Kg	UJ	B, FD
2,3,7,8-TCDF	0.0787	JB	0.0683	MDL	1.09	PQL	ng/Kg	UJ	B, FD
OCDD	1.03	JB	0.0398	MDL	10.9	PQL	ng/Kg	UJ	B, FD
OCDF	0.462	JB	0.0545	MDL	10.9	PQL	ng/Kg	U	B

Method Category: SVOA
Method: 8015M **Matrix:** SO

Sample ID: SL-516-SA5D-SB-0.0-0.5 Collected: 11/15/2013 8:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	3.9	J	2.1	MDL	5.3	PQL	mg/Kg	J	Z
EFH (C30-C40)	6.2	J	4.3	MDL	11	PQL	mg/Kg	J	Z

Sample ID: SL-516-SA5D-SB-13.0-14.0 Collected: 11/15/2013 9:08:00 Analysis Type: REA Dilution: 24.75

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
GASOLINE RANGE ORGANICS (C5-C12)	1.0	U	0.2	MDL	1.0	PQL	mg/Kg	UJ	H

Sample ID: SL-516-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:45:00 Analysis Type: REA Dilution: 26.04

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
GASOLINE RANGE ORGANICS (C5-C12)	1.2	U	0.2	MDL	1.2	PQL	mg/Kg	UJ	H

Sample ID: SL-516-SA5D-SB-9.0-10.0 Collected: 11/15/2013 8:56:00 Analysis Type: REA Dilution: 23.23

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
GASOLINE RANGE ORGANICS (C5-C12)	1.0	U	0.2	MDL	1.0	PQL	mg/Kg	UJ	H

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA
Method: 8015M **Matrix:** SO

Sample ID: SL-609-SA5D-SB-0.0-0.5 Collected: 11/15/2013 12:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	2.6	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z
EFH (C30-C40)	5.5	J	4.2	MDL	10	PQL	mg/Kg	J	Z

Sample ID: SL-816-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:40:00 Analysis Type: REA Dilution: 26.21

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
GASOLINE RANGE ORGANICS (C5-C12)	1.2	U	0.2	MDL	1.2	PQL	mg/Kg	UJ	H

Method Category: SVOA
Method: 8151A **Matrix:** SO

Sample ID: SL-516-SA5D-SB-0.0-0.5 Collected: 11/15/2013 8:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	25	U	9.6	MDL	25	PQL	ug/Kg	R	L

Sample ID: SL-516-SA5D-SB-13.0-14.0 Collected: 11/15/2013 9:08:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	25	U	9.2	MDL	25	PQL	ug/Kg	R	L

Sample ID: SL-516-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	26	U	9.9	MDL	26	PQL	ug/Kg	R	L

Sample ID: SL-516-SA5D-SB-9.0-10.0 Collected: 11/15/2013 8:56:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	26	U	9.9	MDL	26	PQL	ug/Kg	R	L

Sample ID: SL-816-SA5D-SB-4.0-5.0 Collected: 11/15/2013 8:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DINOSEB	26	U	9.9	MDL	26	PQL	ug/Kg	R	L

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-516-SA5D-SB-13.0-14.0 Collected: 11/15/2013 9:08:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.91	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(B)FLUORANTHENE	0.71	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	1.3	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
CHRYSENE	0.96	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
FLUORANTHENE	1.1	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
NAPHTHALENE	1.4	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
PHENANTHRENE	1.1	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	0.87	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PrepPH130

eQAPP Name: CDM_SSFL_140113_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Calibration Blank Contamination
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Field Blank Contamination
FD	Field Duplicate Precision
H	Extraction to Analysis Estimation
L	Laboratory Control Spike Lower Rejection
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH130

QC Outlier Report: HoldingTimes

Lab Reporting Batch ID: PH130
EDD Filename: PH130

Laboratory: LL
eQAPP Name: CDM_SSFL_140113_Lan

Method: 8015M

Preparation Method: 5035A

Matrix: SO

Sample ID	Type	Actual	Criteria	Units	Flag
SL-516-SA5D-SB-13.0-14.0 (REA)	Extraction To Analysis	10.00	7.00	DAYS	J (all detects)
SL-516-SA5D-SB-4.0-5.0 (REA)		11.00	7.00	DAYS	UJ (all non-detects)
SL-516-SA5D-SB-4.0-5.0MS (REA)		11.00	7.00	DAYS	
SL-516-SA5D-SB-4.0-5.0MSD (REA)		11.00	7.00	DAYS	
SL-516-SA5D-SB-9.0-10.0 (REA)		10.00	7.00	DAYS	
SL-816-SA5D-SB-4.0-5.0 (REA)		10.00	7.00	DAYS	

Method Blank Outlier Report

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK3290B372249	11/27/2013 10:49:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDD 1,2,3,7,8,9-HXCDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF 2,3,7,8-TCDD 2,3,7,8-TCDF OCDD OCDF	0.211 ng/Kg 0.0819 ng/Kg 0.0771 ng/Kg 0.0502 ng/Kg 0.0660 ng/Kg 0.0706 ng/Kg 0.0838 ng/Kg 0.0746 ng/Kg 0.0762 ng/Kg 0.0903 ng/Kg 0.166 ng/Kg 0.101 ng/Kg 1.58 ng/Kg 0.234 ng/Kg	SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-516-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.330 ng/Kg	0.330U ng/Kg
SL-516-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0969 ng/Kg	0.0969U ng/Kg
SL-516-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.105 ng/Kg	0.105U ng/Kg
SL-516-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0427 ng/Kg	0.0427U ng/Kg
SL-516-SA5D-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0747 ng/Kg	0.0747U ng/Kg
SL-516-SA5D-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.271 ng/Kg	0.271U ng/Kg
SL-516-SA5D-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.168 ng/Kg	0.168U ng/Kg
SL-516-SA5D-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.118 ng/Kg	0.118U ng/Kg
SL-516-SA5D-SB-0.0-0.5(RES)	OCDD	3.39 ng/Kg	3.39U ng/Kg
SL-516-SA5D-SB-0.0-0.5(RES)	OCDF	0.365 ng/Kg	0.365U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	1,2,3,4,6,7,8-HPCDD	0.137 ng/Kg	0.137U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0921 ng/Kg	0.0921U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0501 ng/Kg	0.0501U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	1,2,3,4,7,8-HxCDD	0.0505 ng/Kg	0.0505U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	1,2,3,4,7,8-HXCDF	0.0758 ng/Kg	0.0758U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	1,2,3,6,7,8-HXCDD	0.0367 ng/Kg	0.0367U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	1,2,3,7,8,9-HXCDD	0.0443 ng/Kg	0.0443U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	1,2,3,7,8-PECDF	0.0757 ng/Kg	0.0757U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	2,3,4,6,7,8-HXCDF	0.0484 ng/Kg	0.0484U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	2,3,4,7,8-PECDF	0.0686 ng/Kg	0.0686U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	OCDD	1.40 ng/Kg	1.40U ng/Kg
SL-516-SA5D-SB-13.0-14.0(RES)	OCDF	0.156 ng/Kg	0.156U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDD	0.313 ng/Kg	0.313U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.112 ng/Kg	0.112U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0665 ng/Kg	0.0665U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDD	0.0899 ng/Kg	0.0899U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	1,2,3,4,7,8-HXCDF	0.122 ng/Kg	0.122U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDD	0.0667 ng/Kg	0.0667U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDD	0.121 ng/Kg	0.121U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.136 ng/Kg	0.136U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 1613B

Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-516-SA5D-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0666 ng/Kg	0.0666U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	2,3,4,7,8-PECDF	0.0691 ng/Kg	0.0691U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	OCDD	1.85 ng/Kg	1.85U ng/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	OCDF	0.281 ng/Kg	0.281U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	1,2,3,4,6,7,8-HPCDD	0.846 ng/Kg	0.846U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	1,2,3,4,6,7,8-HPCDF	0.216 ng/Kg	0.216U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	1,2,3,4,7,8,9-HPCDF	0.107 ng/Kg	0.107U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	1,2,3,4,7,8-HxCDD	0.0581 ng/Kg	0.0581U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	1,2,3,4,7,8-HXCDF	0.149 ng/Kg	0.149U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	1,2,3,6,7,8-HxCDD	0.173 ng/Kg	0.173U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	1,2,3,7,8,9-HxCDD	0.216 ng/Kg	0.216U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	1,2,3,7,8-PECDF	0.303 ng/Kg	0.303U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	2,3,4,6,7,8-HXCDF	0.133 ng/Kg	0.133U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	2,3,4,7,8-PECDF	0.122 ng/Kg	0.122U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	2,3,7,8-TCDF	0.0992 ng/Kg	0.0992U ng/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	OCDF	0.601 ng/Kg	0.601U ng/Kg
SL-609-SA5D-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.580 ng/Kg	0.580U ng/Kg
SL-609-SA5D-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.336 ng/Kg	0.336U ng/Kg
SL-609-SA5D-SB-0.0-0.5(RES)	2,3,7,8-TCDD	0.0640 ng/Kg	0.0640U ng/Kg
SL-609-SA5D-SB-0.0-0.5(RES)	OCDD	1.49 ng/Kg	1.49U ng/Kg
SL-609-SA5D-SB-0.0-0.5(RES)	OCDF	1.05 ng/Kg	1.05U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDD	0.280 ng/Kg	0.280U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.236 ng/Kg	0.236U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.205 ng/Kg	0.205U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDD	0.195 ng/Kg	0.195U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	1,2,3,4,7,8-HXCDF	0.213 ng/Kg	0.213U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	1,2,3,6,7,8-HxCDD	0.228 ng/Kg	0.228U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDD	0.224 ng/Kg	0.224U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.214 ng/Kg	0.214U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.224 ng/Kg	0.224U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	2,3,4,7,8-PECDF	0.223 ng/Kg	0.223U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	2,3,7,8-TCDD	0.111 ng/Kg	0.111U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	2,3,7,8-TCDF	0.0787 ng/Kg	0.0787U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	OCDD	1.03 ng/Kg	1.03U ng/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	OCDF	0.462 ng/Kg	0.462U ng/Kg

Method Blank Outlier Report

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P32937AB220213	11/27/2013 2:13:00 AM	TIN	1.55 mg/Kg	SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-516-SA5D-SB-0.0-0.5(RES)	TIN	2.94 mg/Kg	2.94U mg/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	TIN	3.31 mg/Kg	3.31U mg/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	TIN	3.07 mg/Kg	3.07U mg/Kg
SL-609-SA5D-SB-0.0-0.5(RES)	TIN	2.57 mg/Kg	2.57U mg/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	TIN	3.07 mg/Kg	3.07U mg/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041613(REA2/TOT)	4/16/2013 3:15:00 PM	MOLYBDENUM TIN	0.0132 mg/L 0.0029 mg/L	SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-516-SA5D-SB-0.0-0.5(RES)	MOLYBDENUM	0.366 mg/Kg	0.366U mg/Kg
SL-516-SA5D-SB-13.0-14.0(REA)	MOLYBDENUM	0.509 mg/Kg	0.509U mg/Kg
SL-516-SA5D-SB-4.0-5.0(RES)	MOLYBDENUM	0.543 mg/Kg	0.543U mg/Kg
SL-516-SA5D-SB-9.0-10.0(RES)	MOLYBDENUM	0.366 mg/Kg	0.366U mg/Kg
SL-816-SA5D-SB-4.0-5.0(RES)	MOLYBDENUM	0.294 mg/Kg	0.294U mg/Kg

Surrogate Outlier Report

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 8082A

Matrix: SO

<i>Sample ID (Analysis Type)</i>	<i>Surrogate</i>	<i>Sample % Recovery</i>	<i>% Recovery Limits</i>	<i>Affected Compounds</i>	<i>Flag</i>
SL-816-SA5D-SB-4.0-5.0	DECACHLOROBIPHENYL	122	45.00-120.00	All Target Analytes	J (all detects)

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-516-SA5D-SB-4.0-5.0MS (TOT) SL-516-SA5D-SB-4.0-5.0MSD (TOT) (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0)	TITANIUM	212	172	75.00-125.00	-	TITANIUM	No Qual, >4x
SL-516-SA5D-SB-4.0-5.0MS (TOT) SL-516-SA5D-SB-4.0-5.0MSD (TOT) (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0)	ALUMINUM CALCIUM IRON MAGNESIUM MANGANESE	-1771 -304 -2043 5 -217	2164 -29 1200 193 -97	75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00	32 (20.00) 33 (20.00) - - -	ALUMINUM CALCIUM IRON MAGNESIUM MANGANESE	J(all detects) UJ(all non-detects) Al, Ca, No Qual %R >4x Fe, Mg, Mn, No Qual, >4x
SL-516-SA5D-SB-4.0-5.0MS (TOT) SL-516-SA5D-SB-4.0-5.0MSD (TOT) (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0)	ANTIMONY BARIUM	32 63	30 -	75.00-125.00 75.00-125.00	- -	ANTIMONY BARIUM	J(all detects) UJ(all non-detects)

Method: 6020A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-516-SA5D-SB-4.0-5.0MS (TOT) SL-516-SA5D-SB-4.0-5.0MSD (TOT) (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0)	STRONTIUM	-13	-	75.00-125.00	30 (20.00)	STRONTIUM	J(all detects) R(all non-detects) Post Spike = 351%

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6010C

Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-516-SA5D-SB-4.0-5.0DUP (TOT) (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0)	BORON MANGANESE	23 63	20.00 20.00	J (all detects) UJ (all non-detects)

Method: 6020A

Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-516-SA5D-SB-4.0-5.0DUP (TOT) (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0)	SILVER	35	20.00	No Qual, OK by Difference

Method: 7199

Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-516-SA5D-SB-4.0-5.0DUP (TOT) (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-816-SA5D-SB-4.0-5.0)	HEXAVALENT CHROMIUM	200	20.00	No Qual, OK by Difference

Method: 7471B

Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-516-SA5D-SB-4.0-5.0DUP (TOT) (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0)	MERCURY	51	20.00	No Qual, OK by Difference

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 8151A

Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P33270AQ241905A (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-816-SA5D-SB-4.0-5.0)	2,4,5-T 2,4-D	136 129	- -	58.00-135.00 59.00-122.00	- -	2,4,5-T 2,4-D	J (all detects)
P33270AQ241905A (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-816-SA5D-SB-4.0-5.0)	DINOSEB	7	-	10.00-36.00	-	DINOSEB	J(all detects) R(all non-detects)

Method: 8270D SIM

Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P5LDLCSQ261212 (SL-516-SA5D-SB-0.0-0.5 SL-516-SA5D-SB-13.0-14.0 SL-516-SA5D-SB-4.0-5.0 SL-516-SA5D-SB-9.0-10.0 SL-609-SA5D-SB-0.0-0.5 SL-816-SA5D-SB-4.0-5.0)	BIS(2-ETHYLHEXYL)PHTHALAT	127	-	79.00-121.00	-	BIS(2-ETHYLHEXYL)PHTHALA	J(all detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 160.3M

Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-516-SA5D-SB-4.0-5.0	SL-816-SA5D-SB-4.0-5.0			
MOISTURE	9.5	10.2	7		No Qualifiers Applied

Method: 1613B

Matrix: SO

Analyte	Concentration (ng/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-516-SA5D-SB-4.0-5.0	SL-816-SA5D-SB-4.0-5.0			
1,2,3,4,6,7,8-HPCDD	0.313	0.280	11	50.00	No Qualifiers Applied
1,2,3,7,8,9-HXCDF	0.146	0.222	41	50.00	
1,2,3,7,8-PECDF	0.136	0.214	45	50.00	
OCDF	0.281	0.462	49	50.00	
1,2,3,4,6,7,8-HPCDF	0.112	0.236	71	50.00	J(all detects) UJ(all non-detects)
1,2,3,4,7,8,9-HPCDF	0.0665	0.205	102	50.00	
1,2,3,4,7,8-HxCDD	0.0899	0.195	74	50.00	
1,2,3,4,7,8-HXCDF	0.122	0.213	54	50.00	
1,2,3,6,7,8-HxCDD	0.0667	0.228	109	50.00	
1,2,3,6,7,8-HXCDF	0.0446	0.195	126	50.00	
1,2,3,7,8,9-HxCDD	0.121	0.224	60	50.00	
1,2,3,7,8-PECDD	0.0967	0.172	56	50.00	
2,3,4,6,7,8-HXCDF	0.0666	0.224	108	50.00	
2,3,4,7,8-PECDF	0.0691	0.223	105	50.00	
2,3,7,8-TCDD	1.09 U	0.111	200	50.00	
2,3,7,8-TCDF	1.09 U	0.0787	200	50.00	
OCDD	1.85	1.03	57	50.00	

Method: 6010C

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-516-SA5D-SB-4.0-5.0 (TOT)	SL-816-SA5D-SB-4.0-5.0 (TOT)			
ALUMINUM	26200	24400	7	50.00	No Qualifiers Applied
ARSENIC	7.49	7.34	2	50.00	
BARIUM	136	106	25	50.00	
BERYLLIUM	0.984	1.01	3	50.00	
BORON	2.58	1.71	41	50.00	
CALCIUM	4330	3670	16	50.00	
CHROMIUM	25.2	25.6	2	50.00	
COBALT	7.02	6.98	1	50.00	
COPPER	12.1	11.4	6	50.00	
IRON	25400	26500	4	50.00	
LEAD	7.84	8.66	10	50.00	
LITHIUM	20.5	19.0	8	50.00	
MAGNESIUM	4960	5110	3	50.00	
NICKEL	14.3	12.9	10	50.00	
PHOSPHORUS	132	124	6	50.00	
POTASSIUM	2220	2270	2	50.00	
SODIUM	481	539	11	50.00	
TIN	3.31	3.07	8	50.00	
TITANIUM	1140	1050	8	50.00	
VANADIUM	49.7	48.6	2	50.00	
ZINC	48.1	51.9	8	50.00	
Zirconium	4.99	5.10	2	50.00	
MANGANESE	425	243	54	50.00	J(all detects)
MOLYBDENUM	0.543	0.294	59	50.00	

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Field Duplicate RPD Report

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6020A

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-516-SA5D-SB-4.0-5.0 (TOT)	SL-816-SA5D-SB-4.0-5.0 (TOT)			
SELENIUM	0.167	0.147	13	50.00	No Qualifiers Applied
SILVER	0.0648	0.0614	5	50.00	
STRONTIUM	27.8	25.9	7	50.00	
THALLIUM	0.301	0.289	4	50.00	

Method: 7199

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-516-SA5D-SB-4.0-5.0 (TOT)	SL-816-SA5D-SB-4.0-5.0 (TOT)			
HEXAVALENT CHROMIUM	0.44 U	0.16	200	50.00	J(all detects) UJ(all non-detects)

Method: 7471B

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-516-SA5D-SB-4.0-5.0 (TOT)	SL-816-SA5D-SB-4.0-5.0 (TOT)			
MERCURY	0.0337	0.0283	17	50.00	No Qualifiers Applied

Method: 9045M

Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-516-SA5D-SB-4.0-5.0	SL-816-SA5D-SB-4.0-5.0			
PH	7.79	8.32	7	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-516-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	0.330	5.31	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.0969	5.31	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.105	5.31	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0427	5.31	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.0747	5.31	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.271	5.31	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JQ	0.0549	5.31	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.421	5.31	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.677	5.31	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.102	5.31	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.168	5.31	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.118	5.31	PQL	ng/Kg	
	OCDD	JB	3.39	10.6	PQL	ng/Kg	
	OCDF	JB	0.365	10.6	PQL	ng/Kg	
SL-516-SA5D-SB-13.0-14.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.137	5.10	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0921	5.10	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0501	5.10	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0505	5.10	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.0758	5.10	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.0367	5.10	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	J	0.0515	5.10	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.0443	5.10	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.0384	5.10	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0548	5.10	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0757	5.10	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.0484	5.10	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0686	5.10	PQL	ng/Kg	
	OCDD	JB	1.40	10.2	PQL	ng/Kg	
OCDF	JBQ	0.156	10.2	PQL	ng/Kg		
SL-516-SA5D-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	0.313	5.44	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.112	5.44	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0665	5.44	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0899	5.44	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.122	5.44	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.0667	5.44	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JQ	0.0446	5.44	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.121	5.44	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.146	5.44	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0967	5.44	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.136	5.44	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0666	5.44	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.0691	5.44	PQL	ng/Kg	
	OCDD	JB	1.85	10.9	PQL	ng/Kg	
OCDF	JB	0.281	10.9	PQL	ng/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-516-SA5D-SB-9.0-10.0	1,2,3,4,6,7,8-HPCDD	JB	0.846	5.35	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.216	5.35	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.107	5.35	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0581	5.35	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.149	5.35	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.173	5.35	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JQ	0.122	5.35	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.216	5.35	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.205	5.35	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0896	5.35	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.303	5.35	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.133	5.35	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.122	5.35	PQL	ng/Kg	
	2,3,7,8-TCDF	JB	0.0992	1.07	PQL	ng/Kg	
	OCDD	JB	9.55	10.7	PQL	ng/Kg	
	OCDF	JBQ	0.601	10.7	PQL	ng/Kg	
SL-609-SA5D-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.580	5.17	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.544	5.17	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.439	5.17	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.452	5.17	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.393	5.17	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.397	5.17	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JQ	0.481	5.17	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.524	5.17	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.492	5.17	PQL	ng/Kg	
	1,2,3,7,8-PECDD	J	0.414	5.17	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.501	5.17	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.418	5.17	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.336	5.17	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.0640	1.03	PQL	ng/Kg	
	OCDD	JB	1.49	10.3	PQL	ng/Kg	
	OCDF	JBQ	1.05	10.3	PQL	ng/Kg	
SL-816-SA5D-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	0.280	5.44	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.236	5.44	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.205	5.44	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.195	5.44	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.213	5.44	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.228	5.44	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	J	0.195	5.44	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.224	5.44	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.222	5.44	PQL	ng/Kg	
	1,2,3,7,8-PECDD	J	0.172	5.44	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.214	5.44	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.224	5.44	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.223	5.44	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.111	1.09	PQL	ng/Kg	
	2,3,7,8-TCDF	JB	0.0787	1.09	PQL	ng/Kg	
	OCDD	JB	1.03	10.9	PQL	ng/Kg	
OCDF	JB	0.462	10.9	PQL	ng/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-516-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.899	1.05	PQL	mg/Kg	J (all detects)
	BORON	J	3.75	10.5	PQL	mg/Kg	
	MOLYBDENUM	J	0.366	2.10	PQL	mg/Kg	
	TIN	J	2.94	10.5	PQL	mg/Kg	
	Zirconium	J	4.90	5.24	PQL	mg/Kg	
SL-516-SA5D-SB-13.0-14.0	BERYLLIUM	J	0.650	0.999	PQL	mg/Kg	J (all detects)
	BORON	J	1.78	9.99	PQL	mg/Kg	
	MOLYBDENUM	J	0.509	2.00	PQL	mg/Kg	
	Zirconium	J	3.74	4.99	PQL	mg/Kg	
SL-516-SA5D-SB-4.0-5.0	BERYLLIUM	J	0.984	1.09	PQL	mg/Kg	J (all detects)
	BORON	J	2.58	10.9	PQL	mg/Kg	
	MOLYBDENUM	J	0.543	2.19	PQL	mg/Kg	
	TIN	J	3.31	10.9	PQL	mg/Kg	
	Zirconium	J	4.99	5.47	PQL	mg/Kg	
SL-516-SA5D-SB-9.0-10.0	BERYLLIUM	J	1.01	1.08	PQL	mg/Kg	J (all detects)
	BORON	J	1.67	10.8	PQL	mg/Kg	
	MOLYBDENUM	J	0.366	2.16	PQL	mg/Kg	
	TIN	J	3.07	10.8	PQL	mg/Kg	
	Zirconium	J	4.73	5.39	PQL	mg/Kg	
SL-609-SA5D-SB-0.0-0.5	BERYLLIUM	J	0.600	1.04	PQL	mg/Kg	J (all detects)
	BORON	J	1.27	10.4	PQL	mg/Kg	
	TIN	J	2.57	10.4	PQL	mg/Kg	
	Zirconium	J	4.77	5.21	PQL	mg/Kg	
SL-816-SA5D-SB-4.0-5.0	BERYLLIUM	J	1.01	1.08	PQL	mg/Kg	J (all detects)
	BORON	J	1.71	10.8	PQL	mg/Kg	
	MOLYBDENUM	J	0.294	2.16	PQL	mg/Kg	
	TIN	J	3.07	10.8	PQL	mg/Kg	
	Zirconium	J	5.10	5.41	PQL	mg/Kg	

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-516-SA5D-SB-0.0-0.5	SELENIUM	J	0.253	0.419	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0478	0.210	PQL	mg/Kg	
SL-516-SA5D-SB-13.0-14.0	SELENIUM	J	0.160	0.400	PQL	mg/Kg	J (all detects)
SL-516-SA5D-SB-4.0-5.0	SELENIUM	J	0.167	0.438	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0648	0.219	PQL	mg/Kg	
SL-516-SA5D-SB-9.0-10.0	SELENIUM	J	0.146	0.431	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0472	0.216	PQL	mg/Kg	
SL-609-SA5D-SB-0.0-0.5	SILVER	J	0.0283	0.208	PQL	mg/Kg	J (all detects)
SL-816-SA5D-SB-4.0-5.0	SELENIUM	J	0.147	0.432	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0614	0.216	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH130

Laboratory: LL

EDD Filename: PH130

eQAPP Name: CDM_SSFL_140113_Lan

Method: 7199

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-516-SA5D-SB-13.0-14.0	HEXAVALENT CHROMIUM	J	0.26	0.41	PQL	mg/Kg	J (all detects)
SL-816-SA5D-SB-4.0-5.0	HEXAVALENT CHROMIUM	J	0.16	0.45	PQL	mg/Kg	J (all detects)

Method: 7471B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-516-SA5D-SB-13.0-14.0	MERCURY	J	0.0103	0.0168	PQL	mg/Kg	J (all detects)

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-516-SA5D-SB-0.0-0.5	EFH (C21-C30)	J	3.9	5.3	PQL	mg/Kg	J (all detects)
	EFH (C30-C40)	J	6.2	11	PQL	mg/Kg	
SL-609-SA5D-SB-0.0-0.5	EFH (C21-C30)	J	2.6	5.2	PQL	mg/Kg	J (all detects)
	EFH (C30-C40)	J	5.5	10	PQL	mg/Kg	

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-516-SA5D-SB-13.0-14.0	2-METHYLNAPHTHALENE	J	0.91	1.7	PQL	ug/Kg	J (all detects)
	BENZO(B)FLUORANTHENE	J	0.71	1.7	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	1.3	1.7	PQL	ug/Kg	
	CHRYSENE	J	0.96	1.7	PQL	ug/Kg	
	FLUORANTHENE	J	1.1	1.7	PQL	ug/Kg	
	NAPHTHALENE	J	1.4	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	1.1	1.7	PQL	ug/Kg	
	PYRENE	J	0.87	1.7	PQL	ug/Kg	

LDC #: 31106A4

VALIDATION COMPLETENESS WORKSHEET

Date: 1/17/17

SDG #: PH130

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: CR

2nd Reviewer: SA

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	-	Sampling dates: 11/15/13
II.	ICP/MS Tune	-	
III.	Calibration	-	
IV.	Blanks	SW	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	SW	
VII.	Duplicate Sample Analysis	SW	
VIII.	Laboratory Control Samples (LCS)	A	
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	A	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	-	(2,3)
XV.	Field Blanks	SW	EB=EB1-111213 FB=FB-011613

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

(PT1127)
 D = Duplicate
 TB = Trip blank
 EB = Equipment blank

(PH032)

Validated Samples:

soil

1	SL-516-SA5D-SB-0.0-0.5	11		21		31	
2	SL-516-SA5D-SB-4.0-5.0	12		22		32	
3	SL-816-SA5D-SB-4.0-5.0	13		23		33	
4	SL-516-SA5D-SB-9.0-10.0	14		24		34	
5	SL-516-SA5D-SB-13.0-14.0	15		25		35	
6	SL-609-SA5D-SB-0.0-0.5	16		26		36	
7	SL-516-SA5D-SB-4.0-5.0MS	17		27		37	
8	SL-516-SA5D-SB-4.0-5.0MSD	18		28		38	
9	SL-516-SA5D-SB-4.0-5.0DUP	19		29		39	
10		20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET
PB/ICB/CCB QUALIFIED SAMPLES

METHOD: Trace metals (EPA SW 864 Method 6010B/6020/7000)

Soil preparation factor applied: 100x x 2xdil

Reason: B

Sample Concentration units, unless otherwise noted: mg/Kg

Associated Samples: All

					Sample Identification										
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/l)	Maximum ICB/CCB ^a (ug/l)	Action Level	1	2	3	4	5	6					
Ag			0.25	0.25	0.0478	0.0648	0.0614	0.0472		0.0283					
Tl			0.35	0.35	0.322	0.301	0.289	0.292	0.287	0.286					

Samples with analyte concentrations within five times the associated ICB, CCB or PB concentration are listed above with the identifications from the Validation Completeness Worksheet. These sample results were qualified as not detected, "U".

Note : a - The listed analyte concentration is the highest ICB, CCB, or PB detected in the analysis of each element.

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/16/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ **Associated Samples:** All Soil

Analyte	Blank ID	Sample Identification											
	FB-041613 (SDG: PH032)	Action Limit	1	2	3	4	5						
Mo	0.0132	6.60	0.366	0.543	0.294	0.366	0.509						
Sn	0.0029	1.45											

Sampling date: 9/25/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ **Associated Samples:** All Soil

Analyte	Blank ID	Sample Identification											
	EB1-111213 (SDG: PH127)	Action Limit	No Qualifiers										
Ba	0.00034	0.17											
Ca	0.0349	17.45											
Mg	0.0168	8.4											

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

Sr Post Spike = 351%

QUALITY ASSURANCE SUMMARY

FORM 5A(MS/MSD)

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

SDG No.: PH130

Matrix: SOIL

Level

(low/med):

LOW

74K

ALG RPD
O4(J/KS)

Background Lab Sample ID: 7280198BKG Matrix Spike Lab Sample ID: 7280199MS Matrix Spike Duplicate Lab Sample ID: 7280200MSD
Batch Id(s): P32937A, P32938A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit		M	
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q		%R
Aluminum		23697.1911		20259.1311		27981.5693		194.1748	198.0198	MG/KG	-1771		2164				20	P
Antimony		0.7327	U	15.6738		14.7802		48.5437	49.5050	MG/KG	32	N	30	N	6	75 - 125	20	P
Arsenic		6.7822		19.0243		20.6277		14.5631	14.8515	MG/KG	84		93		8	75 - 125	20	P
Barium		123.3040		246.2204		289.9851		194.1748	198.0198	MG/KG	63	N	84		16	75 - 125	20	P
Beryllium		0.8901	B	5.4767		5.9515		4.8544	4.9505	MG/KG	94		102		8	75 - 125	20	P
Boron		2.3347	B	182.0893		189.6356		194.1748	198.0198	MG/KG	93		95		4	75 - 125	20	P
Cadmium		0.0752	U	4.0922		4.1881		4.8544	4.9505	MG/KG	84		85		2	75 - 125	20	P
Calcium		3919.0495		2738.3893		3803.9267		388.3495	396.0396	MG/KG	-304		-29				20	P
Chromium		22.7931		39.7311		45.2564		19.4175	19.8020	MG/KG	87		113		13	75 - 125	20	P
Cobalt		6.3505		51.4612		53.3257		48.5437	49.5050	MG/KG	93		95		4	75 - 125	20	P
Copper		10.9505		32.6825		36.6663		24.2718	24.7525	MG/KG	90		104		11	75 - 125	20	P
Iron		23010.6000		21026.7767		24198.4574		97.0874	99.0099	MG/KG	-2043		1200		14		20	P
Lead		7.0941		19.3660		21.8911		14.5631	14.8515	MG/KG	84		100		12	75 - 125	20	P
Lithium		18.5752		110.5447		116.4564		97.0874	99.0099	MG/KG	95		99		5	75 - 125	20	P
Magnesium		4493.1970		4503.4990		4875.7832		194.1748	198.0198	MG/KG	5		193		8		20	P
Manganese		385.0317		279.8495		336.8871		48.5437	49.5050	MG/KG	-217		-97		18		20	P
Mercury		0.0305		0.1604		0.1634		0.1573	0.1581	MG/KG	83		84		2	65 - 135	20	CV
Molybdenum		0.4911	B	177.7204		182.1436		194.1748	198.0198	MG/KG	91		92		2	75 - 125	20	P
Nickel		12.9168		56.1350		61.3921		48.5437	49.5050	MG/KG	89		98		9	75 - 125	20	P
Phosphorus		119.7941		202.2670		232.2287		97.0874	99.0099	MG/KG	85		114		14	75 - 125	20	P
Potassium		2004.9950		2974.9796		3239.8099		970.8738	990.0990	MG/KG	100		125		9	75 - 125	20	P
Selenium	78	0.1507	B	2.1060		2.0099		1.9417	1.9802	MG/KG	101		94		5	75 - 125	20	MS
Silver	107	0.0586	B	10.4126		10.1723		9.7087	9.9010	MG/KG	107		102		2	75 - 125	20	MS
Sodium		435.1257		1390.2078		1401.5505		970.8738	990.0990	MG/KG	98		98		1	75 - 125	20	P
Strontium	88	25.1982		24.0800		32.6404		7.7670	7.9208	MG/KG	-14	N	94		30	* 75 - 125	20	MS
Thallium	203	0.2723		0.6431		0.6616		0.3883	0.3960	MG/KG	95		98		3	75 - 125	20	MS

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U = Below MDL, B = Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>
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74x

Background Lab Sample ID: 7280198BKG Matrix Spike Lab Sample ID: 7280199MS Matrix Spike Duplicate Lab Sample ID: 7280200MSD
 Batch Id(s): P32937A, P32938A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit				
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q	%R	RPD	M
Tin		2.9960	B	339.7379		352.5822		388.3495	396.0396	MG/KG	87		88		4	75 - 125	20	P	
Titanium		1034.6634		1240.9699		1205.4178		97.0874	99.0099	MG/KG	212		172		3			20	P
Vanadium		45.0119		86.7631		98.2257		48.5437	49.5050	MG/KG	86		107		12	75 - 125	20	P	
Zinc		43.5535		91.8078		95.1307		48.5437	49.5050	MG/KG	99		104		4	75 - 125	20	P	
Zirconium		4.5139	B	94.6136		98.2693		97.0874	99.0099	MG/KG	93		95		4	75 - 125	20	P	

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U = Below MDL, B = Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS
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Background Lab Sample ID: 7280198BKG
 Batch ID(s): P32937A, P32938A
 Concentration Units: MG/KG

Duplicate Lab Sample ID: 7280204DUP

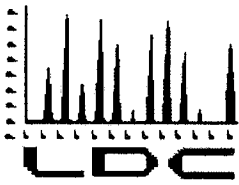
Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			23697.1911		24360.6725		3		P
Antimony			0.7327	U	0.7255	U			P
Arsenic		4.0	6.7822		6.9255		2		P
Barium			123.3040		110.2627		11		P
Beryllium			0.8901	B	0.9235	B	4		P
Boron			2.3347	B	1.8471	B	23	✓	P
Cadmium			-0.5574	B	-0.6255	B	-12		P
Calcium			3919.0495		3933.4922		0		P
Chromium			22.7931		24.2725		6		P
Cobalt			6.3505		5.3363		17		P
Copper			10.9505		11.6431		6		P
Iron			23010.6000		24548.6324		6		P
Lead		3.0	7.0941		8.2020		14		P
Lithium		4.0	18.5752		18.9069		2		P
Magnesium			4493.1970		4696.8225		4		P
Manganese			385.0317		200.2657		63	*✓	P
Mercury		0.0	0.0305		0.0181		51		CV
Molybdenum			0.4911	B	0.4059	B	19		P
Nickel			12.9168		12.1441		6		P
Phosphorus			119.7941		114.4520		5		P
Potassium			2004.9950		2006.1373		0		P
Selenium	78		0.1507	B	0.1425	B	6		MS
Silver	107		0.0586	B	0.0410	B	35		MS
Sodium		99.0	435.1257		448.0873		3		P
Strontium	88		25.1982		25.8012		2		MS
Thallium	203	0.2	0.2723		0.2500		9		MS
Tin			2.9960	B	2.8902	B	4		P
Titanium			1034.6634		1019.6863		1		P
Vanadium			45.0119		47.2412		5		P
Zinc			43.5535		45.0598		3		P
Zirconium		5.0	4.5139	B	5.0108		10		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).
 The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

ok by difference

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: = Duplicate Out of Spec
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CDM
555 17th Street, Suite 1100
Denver, CO 80202
ATTN: Mrs. Cherie Zakowski

September 10, 2013

SUBJECT: Santa Susana Field Laboratory, Data Validation

Dear Mrs. Zakowski,

Enclosed are the final validation reports for the fractions listed below. These SDGs were received on August 9, 2013. Attachment 1 is a summary of the samples that were reviewed for each analysis.

LDC Project # 30210:

<u>SDG #</u>	<u>Fraction</u>
PH062, PH063 PH064, PH065	Volatiles, Semivolatiles, Chlorinated Pesticides, Polychlorinated Biphenyls, Metals, Herbicides, Total Petroleum Hydrocarbons as Gasoline, Total Petroleum Hydrocarbons as Extractables, Dioxins/Dibenzofurans, Perchlorate

The data validation was performed under Standard Guidelines. The analyses were validated using the following documents, as applicable to each method:

- Quality Assurance Project Plan for Santa Susana Field Laboratory, RCRA Facility Investigation, Surficial Media Operable Unit, March 2009, Revision 4
- USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review, June 2008
- USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review, January 2010
- Polychlorinated Dioxins/Dibenzofurans Data Review, September 2005
- EPA SW 846, Third Edition, Test Methods for Evaluating Solid Waste, update 1, July 1992; update IIA, August 1993; update II, September 1994; update IIB, January 1995; update III, December 1996; update IIIA, April 1998; IIIB, November 2004; Update IV, February 2007

Please feel free to contact us if you have any questions.

Sincerely,

A handwritten signature in black ink, appearing to read 'S. McKellar', written in a cursive style.

Shauna McKellar
Project Manager/Senior Chemist

90/10 ADR/IV LDC #30210 (CDM Federal Programs-Chantilly VA / Santa Susana Field Laboratory, Subarea 8)

LDC	SDG#	DATE REC'D	(4) DATE DUE	VOA (8260B)		SVOA (8270D -SIM)		Pest. (8081A)		PCBs (8082A)		Metals & Hg (SW846)		Herbs (8151A)		TPH-G (8015M)		TPH-E (8015M)		Dioxins (1613B)		CLO ₄ (6850)																						
				W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S							
Matrix: Water/Soil																																												
A	PH062	08/09/13	09/06/13	-	-	0	4	-	-	0	4	0	4	-	-	1	1	0	4	0	4	0	3																					
B	PH063	08/09/13	09/06/13	-	-	0	5	-	-	0	5	0	5	-	-	1	2	0	5	0	5	-	-																					
C	PH064	08/09/13	09/06/13	3	2	2	11	2	0	2	11	2	11	2	0	3	8	2	11	2	8	2	0																					
D	PH065	08/09/13	09/06/13	1	2	0	7	-	-	0	7	0	7	-	-	1	5	0	7	0	4	-	-																					
Total	T/SM			4	4	2	27	2	0	2	27	2	27	2	0	6	16	2	27	2	21	2	3	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	0	178		

Shaded cells indicate Level IV validation (all other cells are ADR review). These sample counts do not include MS/MSD, and DUPs 30210ST-SA8.wpd

**Data Validation Report
Santa Susan Field Laboratory
Subarea 8**

SDG(s): PH062

Prepared for

CDM
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
7750 El Camino Real, Suite 2L
Carlsbad, California 92009

September 10, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on July 8, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory, RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles by Environmental Protection Agency (EPA) SW 846 Method 8270D-SIM
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A, and 7471B
Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M
Total Petroleum Hydrocarbons (TPH) as Extractables by EPA SW 846 Method 8015M
Dioxins/Dibenzofurans by EPA Method 1613B
Perchlorate by EPA Method 6850

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibrations blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks and field duplicate samples. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the calibration blanks and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of three method blanks for dioxins and metals. The associated sample results were qualified as non-detected (U) due to method blank contamination. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks with the following exceptions:

SDG/ Method	Method Blank ID	Analyte	Maximum Concentration	Associated Samples
PH062/ 6020A	ICB/CCB	Molybdenum	3.9 ug/L	SL-586-SA8-SB-0.0-0.5 SL-588-SA8-SB-0.0-0.5 SL-888-SA8-SB-0.0-0.5 SL-588-SA8-SB-2.0-3.0

Sample concentrations were compared to concentrations detected in the initial and continuing blanks. The sample concentrations were not detected or were significantly greater than the concentrations found in the associated blanks with the following exceptions:

SDG/Method	Sample	Compound	Reported Concentration	Modified Final Concentration
PH062/ 6020A	SL-586-SA8-SB-0.0-0.5	Molybdenum	0.65 mg/kg	0.65U mg/kg
PH062/ 6020A	SL-588-SA8-SB-0.0-0.5	Molybdenum	0.84 mg/kg	0.84U mg/kg
PH062/ 6020A	SL-888-SA8-SB-0.0-0.5	Molybdenum	0.56 mg/kg	0.56U mg/kg

SDG/Method	Sample	Compound	Reported Concentration	Modified Final Concentration
PH062/ 6020A	SL-588-SA8-SB-2.0-3.0	Molybdenum	0.51 mg/kg	0.51U mg/kg

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits with the exception of one sample for PCBs. The associated sample results were qualified as detected estimated (J) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data was not reviewed for Level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for dioxins, TPH as extractables and metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits with the exception of one DUP for metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH062	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pair was collected and analyzed for semivolatiles, PCBs, metals, perchlorate, TPH as extractables, and dioxins. All RPDs were within QC limits with the exception of several dioxins, semivolatiles, perchlorate and metals in the duplicate pair. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No contaminants were found in the trip blank.

One equipment blank (from SDG PH064) was collected and analyzed for semivolatiles, PCBs, metals, perchlorate, TPH as gasoline, TPH as extractables, and dioxins. No contaminants were found in the equipment blank with the exception of several dioxins, semivolatiles, and metals. The associated sample results were not detected or were significantly greater than the concentrations found in the associated blanks, therefore no data were qualified.

One field blank (from SDG PH029) was collected and analyzed for semivolatiles, PCBs, metals, perchlorate, TPH as gasoline, TPH as extractables and dioxins. No contaminants were found in the field blank with the exception of several dioxins, semivolatiles, and metals. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The samples results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
08-Jul-2013	TB-070813	7120508	TB	5030B	8015M	III
08-Jul-2013	SL-586-SA8-SB-0.0-0.5	7120501	N	3050B	6010C	III
08-Jul-2013	SL-586-SA8-SB-0.0-0.5	7120501	N	3050B	6020A	III
08-Jul-2013	SL-586-SA8-SB-0.0-0.5	7120501	N	3546	8015M	III
08-Jul-2013	SL-586-SA8-SB-0.0-0.5	7120501	N	3546	8082A	III
08-Jul-2013	SL-586-SA8-SB-0.0-0.5	7120501	N	3546	8270D SIM	III
08-Jul-2013	SL-586-SA8-SB-0.0-0.5	7120501	N	METHOD	1613B	III
08-Jul-2013	SL-586-SA8-SB-0.0-0.5	7120501	N	METHOD	7471B	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5	7120502	N	3050B	6010C	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5	7120502	N	3050B	6020A	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5	7120502	N	3546	8015M	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5	7120502	N	3546	8082A	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5	7120502	N	3546	8270D SIM	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5	7120502	N	METHOD	1613B	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5	7120502	N	METHOD	6850	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5	7120502	N	METHOD	7471B	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MS	7120503	MS	3050B	6010C	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MS	7120503	MS	3050B	6020A	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MS	7120503	MS	3546	8015M	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MS	7120503	MS	3546	8082A	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MS	7120503	MS	3546	8270D SIM	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MS	7120503	MS	METHOD	1613B	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MS	7120503	MS	METHOD	6850	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MS	7120503	MS	METHOD	7471B	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MSD	7120504	MSD	3050B	6010C	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MSD	7120504	MSD	3050B	6020A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MSD	7120504	MSD	3546	8015M	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MSD	7120504	MSD	3546	8270D SIM	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MSD	7120504	MSD	Gen Prep	8082A	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MSD	7120504	MSD	METHOD	1613B	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MSD	7120504	MSD	METHOD	6850	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5MSD	7120504	MSD	METHOD	7471B	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5DUP	7120505	DUP	3050B	6010C	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5DUP	7120505	DUP	3050B	6020A	III
08-Jul-2013	SL-588-SA8-SB-0.0-0.5DUP	7120505	DUP	METHOD	7471B	III
08-Jul-2013	SL-888-SA8-SB-0.0-0.5	7120506	FD	3050B	6010C	III
08-Jul-2013	SL-888-SA8-SB-0.0-0.5	7120506	FD	3050B	6020A	III
08-Jul-2013	SL-888-SA8-SB-0.0-0.5	7120506	FD	3546	8015M	III
08-Jul-2013	SL-888-SA8-SB-0.0-0.5	7120506	FD	3546	8082A	III
08-Jul-2013	SL-888-SA8-SB-0.0-0.5	7120506	FD	3546	8270D SIM	III
08-Jul-2013	SL-888-SA8-SB-0.0-0.5	7120506	FD	METHOD	1613B	III
08-Jul-2013	SL-888-SA8-SB-0.0-0.5	7120506	FD	METHOD	6850	III
08-Jul-2013	SL-888-SA8-SB-0.0-0.5	7120506	FD	METHOD	7471B	III
08-Jul-2013	SL-588-SA8-SB-2.0-3.0	7120507	N	3050B	6010C	III
08-Jul-2013	SL-588-SA8-SB-2.0-3.0	7120507	N	3050B	6020A	III
08-Jul-2013	SL-588-SA8-SB-2.0-3.0	7120507	N	3546	8015M	III
08-Jul-2013	SL-588-SA8-SB-2.0-3.0	7120507	N	3546	8082A	III
08-Jul-2013	SL-588-SA8-SB-2.0-3.0	7120507	N	3546	8270D SIM	III
08-Jul-2013	SL-588-SA8-SB-2.0-3.0	7120507	N	5035A	8015M	III
08-Jul-2013	SL-588-SA8-SB-2.0-3.0	7120507	N	METHOD	1613B	III
08-Jul-2013	SL-588-SA8-SB-2.0-3.0	7120507	N	METHOD	6850	III
08-Jul-2013	SL-588-SA8-SB-2.0-3.0	7120507	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-586-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:10:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.12	U	0.762	MDL	4.12	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.548	J	0.0690	MDL	1.03	PQL	mg/Kg	J	Z
BORON	4.37	J	0.865	MDL	10.3	PQL	mg/Kg	J	Z
CADMIUM	0.535	J	0.0782	MDL	1.03	PQL	mg/Kg	J	Z
CALCIUM	3660		3.44	MDL	20.6	PQL	mg/Kg	J	E
MOLYBDENUM	0.652	J	0.175	MDL	2.06	PQL	mg/Kg	U	B, F
POTASSIUM	3690		8.59	MDL	103	PQL	mg/Kg	J	Q, E
SODIUM	84.2	J	17.2	MDL	103	PQL	mg/Kg	J	Z
TIN	2.58	J	0.227	MDL	10.3	PQL	mg/Kg	U	B
ZINC	101		0.206	MDL	4.12	PQL	mg/Kg	J	E
Zirconium	3.99	J	0.865	MDL	5.15	PQL	mg/Kg	J	Z

Sample ID: SL-588-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:50:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	3.96	U	0.732	MDL	3.96	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.522	J	0.0663	MDL	0.990	PQL	mg/Kg	J	Z
BORON	2.63	J	0.831	MDL	9.90	PQL	mg/Kg	J	Z
CADMIUM	0.423	J	0.0752	MDL	0.990	PQL	mg/Kg	J	Z
CALCIUM	2210		3.31	MDL	19.8	PQL	mg/Kg	J	E
MOLYBDENUM	0.843	J	0.168	MDL	1.98	PQL	mg/Kg	U	B, F
POTASSIUM	3690		8.25	MDL	99.0	PQL	mg/Kg	J	Q, E
SODIUM	66.5	J	16.5	MDL	99.0	PQL	mg/Kg	J	Z
TIN	2.47	J	0.218	MDL	9.90	PQL	mg/Kg	U	B
ZINC	62.4		0.198	MDL	3.96	PQL	mg/Kg	J	E
Zirconium	3.62	J	0.831	MDL	4.95	PQL	mg/Kg	J	Z

Sample ID: SL-588-SA8-SB-2.0-3.0 Collected: 7/8/2013 2:35:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.08	U	0.755	MDL	4.08	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.604	J	0.0684	MDL	1.02	PQL	mg/Kg	J	Z
BORON	2.18	J	0.857	MDL	10.2	PQL	mg/Kg	J	Z
CADMIUM	0.393	J	0.0776	MDL	1.02	PQL	mg/Kg	J	Z
CALCIUM	2060		3.41	MDL	20.4	PQL	mg/Kg	J	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-588-SA8-SB-2.0-3.0 Collected: 7/8/2013 2:35:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MOLYBDENUM	0.513	J	0.173	MDL	2.04	PQL	mg/Kg	U	B, F
POTASSIUM	3500		8.51	MDL	102	PQL	mg/Kg	J	Q, E
SODIUM	74.3	J	17.0	MDL	102	PQL	mg/Kg	J	Z
TIN	2.39	J	0.224	MDL	10.2	PQL	mg/Kg	U	B
ZINC	61.9		0.204	MDL	4.08	PQL	mg/Kg	J	E
Zirconium	3.40	J	0.857	MDL	5.10	PQL	mg/Kg	J	Z

Sample ID: SL-888-SA8-SB-0.0-0.5 Collected: 7/8/2013 2:00:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	3.96	U	0.733	MDL	3.96	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.551	J	0.0663	MDL	0.990	PQL	mg/Kg	J	Z
BORON	2.49	J	0.832	MDL	9.90	PQL	mg/Kg	J	Z
CADMIUM	0.441	J	0.0752	MDL	0.990	PQL	mg/Kg	J	Z
CALCIUM	2320		3.31	MDL	19.8	PQL	mg/Kg	J	E
MOLYBDENUM	0.559	J	0.168	MDL	1.98	PQL	mg/Kg	U	B, F
POTASSIUM	3950		8.26	MDL	99.0	PQL	mg/Kg	J	Q, E
SODIUM	67.3	J	16.5	MDL	99.0	PQL	mg/Kg	J	Z
TIN	2.41	J	0.218	MDL	9.90	PQL	mg/Kg	U	B
ZINC	65.2		0.198	MDL	3.96	PQL	mg/Kg	J	E
Zirconium	3.65	J	0.832	MDL	4.95	PQL	mg/Kg	J	Z

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-586-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:10:00 PM Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.139	J	0.103	MDL	0.412	PQL	mg/Kg	J	Z

Sample ID: SL-586-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:10:00 PM Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0427	J	0.0268	MDL	0.206	PQL	mg/Kg	J	Z
STRONTIUM	25.7		0.0700	MDL	0.412	PQL	mg/Kg	J	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

<i>Sample ID:</i> SL-588-SA8-SB-0.0-0.5			<i>Collected:</i> 7/8/2013 1:50:00 PM			<i>Analysis Type:</i> REA		<i>Dilution:</i> 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.102	J	0.0990	MDL	0.396	PQL	mg/Kg	J	Z, FD

<i>Sample ID:</i> SL-588-SA8-SB-0.0-0.5			<i>Collected:</i> 7/8/2013 1:50:00 PM			<i>Analysis Type:</i> RES		<i>Dilution:</i> 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0326	J	0.0257	MDL	0.198	PQL	mg/Kg	J	Z
STRONTIUM	19.3		0.0673	MDL	0.396	PQL	mg/Kg	J	E

<i>Sample ID:</i> SL-588-SA8-SB-2.0-3.0			<i>Collected:</i> 7/8/2013 2:35:00 PM			<i>Analysis Type:</i> RES		<i>Dilution:</i> 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0476	J	0.0265	MDL	0.204	PQL	mg/Kg	J	Z
STRONTIUM	20.0		0.0694	MDL	0.408	PQL	mg/Kg	J	E

<i>Sample ID:</i> SL-888-SA8-SB-0.0-0.5			<i>Collected:</i> 7/8/2013 2:00:00 PM			<i>Analysis Type:</i> REA		<i>Dilution:</i> 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.181	J	0.0990	MDL	0.396	PQL	mg/Kg	J	Z, FD

<i>Sample ID:</i> SL-888-SA8-SB-0.0-0.5			<i>Collected:</i> 7/8/2013 2:00:00 PM			<i>Analysis Type:</i> RES		<i>Dilution:</i> 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0312	J	0.0257	MDL	0.198	PQL	mg/Kg	J	Z
STRONTIUM	20.8		0.0673	MDL	0.396	PQL	mg/Kg	J	E

Method Category:	METALS	
Method:	7471B	Matrix: SO

<i>Sample ID:</i> SL-588-SA8-SB-0.0-0.5			<i>Collected:</i> 7/8/2013 1:50:00 PM			<i>Analysis Type:</i> RES		<i>Dilution:</i> 1	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0120	J	0.0101	MDL	0.0168	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	7471B	Matrix: SO

Sample ID: SL-888-SA8-SB-0.0-0.5 Collected: 7/8/2013 2:00:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0099	J	0.0094	MDL	0.0157	PQL	mg/Kg	J	Z

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-586-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:10:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	2.14	JB	0.0258	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.221	JB	0.0319	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.249	JB	0.0335	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.378	JB	0.0212	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.807	JB	0.0362	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.383	JB	0.0211	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.711	JB	0.0343	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.345	JBQ	0.0213	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.397	JB	0.0356	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.914	JB	0.0257	MDL	5.05	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.337	JB	0.0197	MDL	5.05	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.566	JB	0.0239	MDL	5.05	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0950	J	0.0216	MDL	1.01	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.538	J	0.0529	MDL	1.01	PQL	ng/Kg	J	Z
OCDF	5.49	JB	0.0248	MDL	10.1	PQL	ng/Kg	J	Z

Sample ID: SL-588-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:50:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	0.972	JB	0.0144	MDL	4.91	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.118	JBQ	0.0205	MDL	4.91	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.130	JBQ	0.0388	MDL	4.91	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.270	JB	0.0224	MDL	4.91	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.460	JB	0.0428	MDL	4.91	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.223	JB	0.0227	MDL	4.91	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.467	JB	0.0409	MDL	4.91	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA		
Method:	1613B	Matrix:	SO

Sample ID: SL-588-SA8-SB-0.0-0.5	Collected: 7/8/2013 1:50:00 PM	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,7,8,9-HXCDF	0.309	JB	0.0235	MDL	4.91	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.178	JB	0.0285	MDL	4.91	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.397	JBQ	0.0249	MDL	4.91	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.243	JB	0.0222	MDL	4.91	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.518	JB	0.0231	MDL	4.91	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0397	JQ	0.0216	MDL	0.982	PQL	ng/Kg	J	Z, FD
2,3,7,8-TCDF	0.345	J	0.0504	MDL	0.982	PQL	ng/Kg	J	Z
OCDD	70.3	B	0.0211	MDL	9.82	PQL	ng/Kg	J	Q
OCDF	1.80	JB	0.0230	MDL	9.82	PQL	ng/Kg	J	Z

Sample ID: SL-588-SA8-SB-2.0-3.0	Collected: 7/8/2013 2:35:00 PM	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	0.508	JB	0.0381	MDL	4.97	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0811	JBQ	0.0471	MDL	4.97	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0750	JB	0.0342	MDL	4.97	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0905	JBQ	0.0150	MDL	4.97	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.257	JB	0.0365	MDL	4.97	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0816	JBQ	0.0149	MDL	4.97	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.266	JB	0.0379	MDL	4.97	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.163	JB	0.0159	MDL	4.97	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0492	JBQ	0.0283	MDL	4.97	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.170	JB	0.0202	MDL	4.97	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.105	JB	0.0163	MDL	4.97	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.165	JBQ	0.0173	MDL	4.97	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.0981	JQ	0.0334	MDL	0.995	PQL	ng/Kg	J	Z
OCDF	1.03	JB	0.0270	MDL	9.95	PQL	ng/Kg	J	Z

Sample ID: SL-888-SA8-SB-0.0-0.5	Collected: 7/8/2013 2:00:00 PM	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	0.910	JB	0.0165	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.103	JB	0.0239	MDL	5.03	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.152	JB	0.0345	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.190	JB	0.0207	MDL	5.03	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-888-SA8-SB-0.0-0.5 Collected: 7/8/2013 2:00:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,6,7,8-HXCDD	0.410	JBQ	0.0378	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.206	JB	0.0208	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.484	JB	0.0354	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.304	JB	0.0213	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.148	JBQ	0.0255	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.307	JB	0.0220	MDL	5.03	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.184	JB	0.0204	MDL	5.03	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.330	JB	0.0194	MDL	5.03	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0222	JQ	0.0191	MDL	1.01	PQL	ng/Kg	J	Z, FD
2,3,7,8-TCDF	0.265	JQ	0.0421	MDL	1.01	PQL	ng/Kg	J	Z
OCDF	1.73	JB	0.0194	MDL	10.1	PQL	ng/Kg	J	Z

Method Category:	SVOA	
Method:	6850	Matrix: SO

Sample ID: SL-588-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:50:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
PERCHLORATE	3.0	J	2.1	MDL	5.0	PQL	ug/Kg	J	Z, FD

Sample ID: SL-888-SA8-SB-0.0-0.5 Collected: 7/8/2013 2:00:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
PERCHLORATE	5.0	U	2.1	MDL	5.0	PQL	ug/Kg	UJ	FD

Method Category:	SVOA	
Method:	8015M	Matrix: SO

Sample ID: SL-586-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:10:00 PM Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	7.5	J	4.2	MDL	10	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8015M	Matrix: SO

Sample ID: SL-588-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:50:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	9.3		2.0	MDL	5.0	PQL	mg/Kg	J	Q, Q

Method Category:	SVOA	
Method:	8082A	Matrix: SO

Sample ID: SL-586-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:10:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
Aroclor 5460	12	J	11	MDL	35	PQL	ug/Kg	J	Z

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-586-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:10:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	1.2	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	1.3	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	0.73	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
Di-n-butylphthalate	9.3	J	6.3	MDL	19	PQL	ug/Kg	J	Z
Di-n-octylphthalate	7.8	J	6.3	MDL	19	PQL	ug/Kg	J	Z

Sample ID: SL-588-SA8-SB-0.0-0.5 Collected: 7/8/2013 1:50:00 PM Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.82	J	0.66	MDL	1.7	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	10	J	6.0	MDL	18	PQL	ug/Kg	J	Z, FD
CHRYSENE	1.1	J	0.33	MDL	1.7	PQL	ug/Kg	J	Z
FLUORANTHENE	0.87	J	0.66	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	0.73	J	0.66	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-588-SA8-SB-2.0-3.0 Collected: 7/8/2013 2:35:00 PM Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	0.41	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-888-SA8-SB-0.0-0.5

Collected: 7/8/2013 2:00:00 PM Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.71	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	18	U	6.0	MDL	18	PQL	ug/Kg	UJ	FD
CHRYSENE	0.94	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
FLUORANTHENE	0.90	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	0.67	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Calibration Blank Contamination
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Field Blank Contamination
FD	Field Duplicate Precision
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Enclosure I
Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH062

Method Blank Outlier Report

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK1920B370624	7/13/2013 6:24:00 AM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF OCDD OCDF	0.0814 ng/Kg 0.0265 ng/Kg 0.0357 ng/Kg 0.0239 ng/Kg 0.0351 ng/Kg 0.0206 ng/Kg 0.0220 ng/Kg 0.0274 ng/Kg 0.0184 ng/Kg 0.0216 ng/Kg 0.0503 ng/Kg 0.0269 ng/Kg 0.0333 ng/Kg 0.214 ng/Kg 0.0705 ng/Kg	SL-586-SA8-SB-0.0-0.5 SL-588-SA8-SB-0.0-0.5 SL-588-SA8-SB-2.0-3.0 SL-888-SA8-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-588-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.118 ng/Kg	0.118U ng/Kg
SL-588-SA8-SB-2.0-3.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0811 ng/Kg	0.0811U ng/Kg
SL-588-SA8-SB-2.0-3.0(RES)	1,2,3,4,7,8-HxCDD	0.0750 ng/Kg	0.0750U ng/Kg
SL-588-SA8-SB-2.0-3.0(RES)	1,2,3,4,7,8-HxCDF	0.0905 ng/Kg	0.0905U ng/Kg
SL-588-SA8-SB-2.0-3.0(RES)	1,2,3,6,7,8-HXCDF	0.0816 ng/Kg	0.0816U ng/Kg
SL-588-SA8-SB-2.0-3.0(RES)	1,2,3,7,8-PECDD	0.0492 ng/Kg	0.0492U ng/Kg
SL-588-SA8-SB-2.0-3.0(RES)	1,2,3,7,8-PECDF	0.170 ng/Kg	0.170U ng/Kg
SL-588-SA8-SB-2.0-3.0(RES)	2,3,4,6,7,8-HXCDF	0.105 ng/Kg	0.105U ng/Kg
SL-588-SA8-SB-2.0-3.0(RES)	2,3,4,7,8-PECDF	0.165 ng/Kg	0.165U ng/Kg
SL-888-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.103 ng/Kg	0.103U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19137AB221846	7/12/2013 6:46:00 PM	ALUMINUM CALCIUM IRON MAGNESIUM TIN TITANIUM ZINC	11.0 mg/Kg 12.8 mg/Kg 6.78 mg/Kg 5.68 mg/Kg 1.32 mg/Kg 0.177 mg/Kg 0.422 mg/Kg	SL-586-SA8-SB-0.0-0.5 SL-588-SA8-SB-0.0-0.5 SL-588-SA8-SB-2.0-3.0 SL-888-SA8-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-586-SA8-SB-0.0-0.5(RES)	TIN	2.58 mg/Kg	2.58U mg/Kg
SL-588-SA8-SB-0.0-0.5(RES)	TIN	2.47 mg/Kg	2.47U mg/Kg
SL-588-SA8-SB-2.0-3.0(RES)	TIN	2.39 mg/Kg	2.39U mg/Kg
SL-888-SA8-SB-0.0-0.5(RES)	TIN	2.41 mg/Kg	2.41U mg/Kg

Method Blank Outlier Report

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A				
Matrix: SO				
Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19137AB221713A	7/18/2013 5:13:00 PM	STRONTIUM	0.0774 mg/Kg	SL-586-SA8-SB-0.0-0.5 SL-588-SA8-SB-0.0-0.5 SL-588-SA8-SB-2.0-3.0 SL-888-SA8-SB-0.0-0.5

Field Blank Outlier Report

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(REA2)	4/11/2013 3:00:00 PM	COPPER MOLYBDENUM	0.0036 mg/L 0.0036 mg/L	SL-586-SA8-SB-0.0-0.5 SL-588-SA8-SB-0.0-0.5 SL-588-SA8-SB-2.0-3.0 SL-888-SA8-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-586-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.652 mg/Kg	0.652U mg/Kg
SL-588-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.843 mg/Kg	0.843U mg/Kg
SL-588-SA8-SB-2.0-3.0(RES)	MOLYBDENUM	0.513 mg/Kg	0.513U mg/Kg
SL-888-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.559 mg/Kg	0.559U mg/Kg

Surrogate Outlier Report

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8082A

Matrix: SO

<i>Sample ID (Analysis Type)</i>	<i>Surrogate</i>	<i>Sample % Recovery</i>	<i>% Recovery Limits</i>	<i>Affected Compounds</i>	<i>Flag</i>
SL-888-SA8-SB-0.0 -0.5	TETRACHLORO-M-XYLENE	121	45.00-120.00	All Target Analytes	J (all detects)

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-588-SA8-SB-0.0-0.5MS (SL-588-SA8-SB-0.0-0.5)	OCDD	138	-	40.00-135.00	-	OCDD	J (all detects)

Method: 8015M
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-588-SA8-SB-0.0-0.5MS	EFH (C15-C20)	173	132	49.00-123.00	27 (20.00)	EFH (C15-C20)	J(all detects)
SL-588-SA8-SB-0.0-0.5MSD	EFH (C21-C30)	162	-	49.00-123.00	24 (20.00)	EFH (C21-C30)	EFH (C30-C40), No Qual, >4x
(SL-588-SA8-SB-0.0-0.5)	EFH (C30-C40)	388	182	49.00-123.00	26 (20.00)	EFH (C30-C40)	

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-588-SA8-SB-0.0-0.5MS (TOT)	ALUMINUM	1971	2027	75.00-125.00	-	ALUMINUM	J(all detects) Al, Ca, Fe, Mg, Mn, Ti, No Qual, >4x
SL-588-SA8-SB-0.0-0.5MSD (TOT)	CALCIUM	228	188	75.00-125.00	-	CALCIUM	
(SL-586-SA8-SB-0.0-0.5)	IRON	1714	1495	75.00-125.00	-	IRON	
SL-588-SA8-SB-0.0-0.5	MAGNESIUM	296	298	75.00-125.00	-	MAGNESIUM	
SL-588-SA8-SB-0.0-0.5	MANGANESE	132	137	75.00-125.00	-	MANGANESE	
SL-588-SA8-SB-2.0-3.0	POTASSIUM	158	162	75.00-125.00	-	POTASSIUM	
SL-888-SA8-SB-0.0-0.5)	TITANIUM	371	392	75.00-125.00	-	TITANIUM	
SL-588-SA8-SB-0.0-0.5MS (TOT)	ANTIMONY	37	37	75.00-125.00	-	ANTIMONY	J(all detects) UU(all non-detects)
SL-588-SA8-SB-0.0-0.5MSD (TOT)							
(SL-586-SA8-SB-0.0-0.5)							
SL-588-SA8-SB-0.0-0.5							
SL-588-SA8-SB-2.0-3.0							
SL-888-SA8-SB-0.0-0.5)							

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-588-SA8-SB-0.0-0.5DUP (TOT)	BORON	25	20.00	J (all detects) UJ (all non-detects)
(SL-586-SA8-SB-0.0-0.5)	CADMIUM	72	20.00	
SL -588-SA8-SB-0.0-0.5	CALCIUM	41	20.00	B, Cd, Mo, Na, Zr, No Qual, OK by Difference
SL -588-SA8-SB-2.0-3.0	MOLYBDENUM	29	20.00	
SL -888-SA8-SB-0.0-0.5)	POTASSIUM	23	20.00	
	SODIUM	23	20.00	
	ZINC	60	20.00	
	Zirconium	29	20.00	

Method: 6020A
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-588-SA8-SB-0.0-0.5DUP (TOT)	SELENIUM	21	20.00	J(all detects) UJ(all non-detects)
(SL-586-SA8-SB-0.0-0.5)	STRONTIUM	26	20.00	
SL -588-SA8-SB-0.0-0.5				Se, No Qual, OK by Difference
SL -588-SA8-SB-2.0-3.0				
SL -888-SA8-SB-0.0-0.5)				

Method: 7471B
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-588-SA8-SB-0.0-0.5DUP (TOT)	MERCURY	80	20.00	No Qual, OK by Difference
(SL-586-SA8-SB-0.0-0.5)				
SL -588-SA8-SB-0.0-0.5				
SL -588-SA8-SB-2.0-3.0				
SL -888-SA8-SB-0.0-0.5)				

Field Duplicate RPD Report

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 160.3M
Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-588-SA8-SB-0.0-0.5	SL-888-SA8-SB-0.0-0.5			
MOISTURE	0.95	0.97	2		No Qualifiers Applied

Method: 1613B
Matrix: SO

Analyte	Concentration (ng/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-588-SA8-SB-0.0-0.5	SL-888-SA8-SB-0.0-0.5			
1,2,3,4,6,7,8-HPCDD	6.94	5.63	21	50.00	No Qualifiers Applied
1,2,3,4,6,7,8-HPCDF	0.972	0.910	7	50.00	
1,2,3,4,7,8,9-HPCDF	0.118	0.103	14	50.00	
1,2,3,4,7,8-HxCDD	0.130	0.152	16	50.00	
1,2,3,4,7,8-HxCDF	0.270	0.190	35	50.00	
1,2,3,6,7,8-HxCDD	0.460	0.410	11	50.00	
1,2,3,6,7,8-HxCDF	0.223	0.206	8	50.00	
1,2,3,7,8,9-HxCDD	0.467	0.484	4	50.00	
1,2,3,7,8,9-HxCDF	0.309	0.304	2	50.00	
1,2,3,7,8-PECDD	0.178	0.148	18	50.00	
1,2,3,7,8-PECDF	0.397	0.307	26	50.00	
2,3,4,6,7,8-HxCDF	0.243	0.184	28	50.00	
2,3,4,7,8-PECDF	0.518	0.330	44	50.00	
2,3,7,8-TCDF	0.345	0.265	26	50.00	
OCDD	70.3	55.8	23	50.00	
OCDF	1.80	1.73	4	50.00	
2,3,7,8-TCDD	0.0397	0.0222	57	50.00	

Field Duplicate RPD Report

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-588-SA8-SB-0.0-0.5 (TOT)	SL-888-SA8-SB-0.0-0.5 (TOT)			
ALUMINUM	16000	16600	4	50.00	No Qualifiers Applied
ARSENIC	4.15	4.22	2	50.00	
BARIUM	103	109	6	50.00	
BERYLLIUM	0.522	0.551	5	50.00	
BORON	2.63	2.49	5	50.00	
CADMIUM	0.423	0.441	4	50.00	
CALCIUM	2210	2320	5	50.00	
CHROMIUM	16.0	16.5	3	50.00	
COBALT	4.35	4.55	4	50.00	
COPPER	10.4	10.7	3	50.00	
IRON	17100	18000	5	50.00	
LEAD	9.00	8.79	2	50.00	
LITHIUM	20.1	21.1	5	50.00	
MAGNESIUM	3500	3680	5	50.00	
MANGANESE	325	341	5	50.00	
MOLYBDENUM	0.843	0.559	41	50.00	
NICKEL	10.3	10.5	2	50.00	
PHOSPHORUS	493	501	2	50.00	
POTASSIUM	3690	3950	7	50.00	
SODIUM	66.5	67.3	1	50.00	
TIN	2.47	2.41	2	50.00	
TITANIUM	1020	1050	3	50.00	
VANADIUM	28.9	30.7	6	50.00	
ZINC	62.4	65.2	4	50.00	
Zirconium	3.62	3.65	1	50.00	

Method: 6020A
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-588-SA8-SB-0.0-0.5 (TOT)	SL-888-SA8-SB-0.0-0.5 (TOT)			
SILVER	0.0326	0.0312	4	50.00	No Qualifiers Applied
STRONTIUM	19.3	20.8	7	50.00	
THALLIUM	0.240	0.249	4	50.00	
SELENIUM	0.102	0.181	56	50.00	J(all detects)

Method: 6850
Matrix: SO

Analyte	Concentration (ug/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-588-SA8-SB-0.0-0.5	SL-888-SA8-SB-0.0-0.5			
PERCHLORATE	3.0	5.0 U	200	50.00	J(all detects) UJ(all non-detects)

Method: 7471B
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-588-SA8-SB-0.0-0.5 (TOT)	SL-888-SA8-SB-0.0-0.5 (TOT)			
MERCURY	0.0120	0.0099	19	50.00	No Qualifiers Applied

Field Duplicate RPD Report

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8015M

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-588-SA8-SB-0.0-0.5	SL-888-SA8-SB-0.0-0.5			
EFH (C21-C30)	9.3	10	7	50.00	No Qualifiers Applied
EFH (C30-C40)	25	27	8	50.00	

Method: 8270D SIM

Matrix: SO

Analyte	Concentration (ug/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-588-SA8-SB-0.0-0.5	SL-888-SA8-SB-0.0-0.5			
BENZO(B)FLUORANTHENE	0.82	0.71	14	50.00	No Qualifiers Applied
BENZO(K)FLUORANTHENE	4.1	4.9	18	50.00	
CHRYSENE	1.1	0.94	16	50.00	
FLUORANTHENE	0.87	0.90	3	50.00	
PYRENE	0.73	0.67	9	50.00	
BIS(2-ETHYLHEXYL)PHTHALATE	10	18 U	200	50.00	J(all detects) UJ(all non-detects)

Method: 9045M

Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-588-SA8-SB-0.0-0.5	SL-888-SA8-SB-0.0-0.5			
PH	6.56	6.42	2	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-586-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	2.14	5.05	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JB	0.221	5.05	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.249	5.05	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.378	5.05	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.807	5.05	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.383	5.05	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.711	5.05	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.345	5.05	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.397	5.05	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.914	5.05	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.337	5.05	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.566	5.05	PQL	ng/Kg	
	2,3,7,8-TCDD	J	0.0950	1.01	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.538	1.01	PQL	ng/Kg	
	OCDF	JB	5.49	10.1	PQL	ng/Kg	
SL-588-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	0.972	4.91	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JBQ	0.118	4.91	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.130	4.91	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.270	4.91	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.460	4.91	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.223	4.91	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.467	4.91	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.309	4.91	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.178	4.91	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.397	4.91	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.243	4.91	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.518	4.91	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0397	0.982	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.345	0.982	PQL	ng/Kg	
OCDF	JB	1.80	9.82	PQL	ng/Kg		
SL-588-SA8-SB-2.0-3.0	1,2,3,4,6,7,8-HPCDF	JB	0.508	4.97	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0811	4.97	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0750	4.97	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0905	4.97	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.257	4.97	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0816	4.97	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.266	4.97	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.163	4.97	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.0492	4.97	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.170	4.97	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.105	4.97	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.165	4.97	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0981	0.995	PQL	ng/Kg	
	OCDF	JB	1.03	9.95	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-888-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	0.910	5.03	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JB	0.103	5.03	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.152	5.03	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.190	5.03	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.410	5.03	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.206	5.03	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.484	5.03	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.304	5.03	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.148	5.03	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.307	5.03	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.184	5.03	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.330	5.03	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0222	1.01	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.265	1.01	PQL	ng/Kg	
	OCDF	JB	1.73	10.1	PQL	ng/Kg	

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-586-SA8-SB-0.0-0.5	BERYLLIUM	J	0.548	1.03	PQL	mg/Kg	J (all detects)
	BORON	J	4.37	10.3	PQL	mg/Kg	
	CADMIUM	J	0.535	1.03	PQL	mg/Kg	
	MOLYBDENUM	J	0.652	2.06	PQL	mg/Kg	
	SODIUM	J	84.2	103	PQL	mg/Kg	
	TIN	J	2.58	10.3	PQL	mg/Kg	
	Zirconium	J	3.99	5.15	PQL	mg/Kg	
SL-588-SA8-SB-0.0-0.5	BERYLLIUM	J	0.522	0.990	PQL	mg/Kg	J (all detects)
	BORON	J	2.63	9.90	PQL	mg/Kg	
	CADMIUM	J	0.423	0.990	PQL	mg/Kg	
	MOLYBDENUM	J	0.843	1.98	PQL	mg/Kg	
	SODIUM	J	66.5	99.0	PQL	mg/Kg	
	TIN	J	2.47	9.90	PQL	mg/Kg	
	Zirconium	J	3.62	4.95	PQL	mg/Kg	
SL-588-SA8-SB-2.0-3.0	BERYLLIUM	J	0.604	1.02	PQL	mg/Kg	J (all detects)
	BORON	J	2.18	10.2	PQL	mg/Kg	
	CADMIUM	J	0.393	1.02	PQL	mg/Kg	
	MOLYBDENUM	J	0.513	2.04	PQL	mg/Kg	
	SODIUM	J	74.3	102	PQL	mg/Kg	
	TIN	J	2.39	10.2	PQL	mg/Kg	
	Zirconium	J	3.40	5.10	PQL	mg/Kg	
SL-888-SA8-SB-0.0-0.5	BERYLLIUM	J	0.551	0.990	PQL	mg/Kg	J (all detects)
	BORON	J	2.49	9.90	PQL	mg/Kg	
	CADMIUM	J	0.441	0.990	PQL	mg/Kg	
	MOLYBDENUM	J	0.559	1.98	PQL	mg/Kg	
	SODIUM	J	67.3	99.0	PQL	mg/Kg	
	TIN	J	2.41	9.90	PQL	mg/Kg	
	Zirconium	J	3.65	4.95	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-586-SA8-SB-0.0-0.5	SELENIUM	J	0.139	0.412	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0427	0.206	PQL	mg/Kg	
SL-588-SA8-SB-0.0-0.5	SELENIUM	J	0.102	0.396	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0326	0.198	PQL	mg/Kg	
SL-588-SA8-SB-2.0-3.0	SILVER	J	0.0476	0.204	PQL	mg/Kg	J (all detects)
SL-888-SA8-SB-0.0-0.5	SELENIUM	J	0.181	0.396	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0312	0.198	PQL	mg/Kg	

Method: 6850
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-588-SA8-SB-0.0-0.5	PERCHLORATE	J	3.0	5.0	PQL	ug/Kg	J (all detects)

Method: 7471B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-588-SA8-SB-0.0-0.5	MERCURY	J	0.0120	0.0168	PQL	mg/Kg	J (all detects)
SL-888-SA8-SB-0.0-0.5	MERCURY	J	0.0099	0.0157	PQL	mg/Kg	J (all detects)

Method: 8015M
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-586-SA8-SB-0.0-0.5	EFH (C15-C20)	J	7.5	10	PQL	mg/Kg	J (all detects)

Method: 8082A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-586-SA8-SB-0.0-0.5	Aroclor 5460	J	12	35	PQL	ug/Kg	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH062

Laboratory: LL

EDD Filename: PH062_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8270D SIM
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-586-SA8-SB-0.0-0.5	1-METHYLNAPHTHALENE	J	1.2	1.8	PQL	ug/Kg	J (all detects)
	BENZO(A)PYRENE	J	1.3	1.8	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	0.73	1.8	PQL	ug/Kg	
	Di-n-butylphthalate	J	9.3	19	PQL	ug/Kg	
	Di-n-octylphthalate	J	7.8	19	PQL	ug/Kg	
SL-588-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	0.82	1.7	PQL	ug/Kg	J (all detects)
	BIS(2-ETHYLHEXYL)PHTHALATE	J	10	18	PQL	ug/Kg	
	CHRYSENE	J	1.1	1.7	PQL	ug/Kg	
	FLUORANTHENE	J	0.87	1.7	PQL	ug/Kg	
	PYRENE	J	0.73	1.7	PQL	ug/Kg	
SL-588-SA8-SB-2.0-3.0	CHRYSENE	J	0.41	1.7	PQL	ug/Kg	J (all detects)
SL-888-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	0.71	1.7	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	0.94	1.7	PQL	ug/Kg	
	FLUORANTHENE	J	0.90	1.7	PQL	ug/Kg	
	PYRENE	J	0.67	1.7	PQL	ug/Kg	

LDC #: 30210A4

VALIDATION COMPLETENESS WORKSHEET

Date: 8/28/13

SDG #: PH062

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: CR

2nd Reviewer: W

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	—	Sampling dates: 7/8/13
II.	ICP/MS Tune	—	
III.	Calibration	—	
IV.	Blanks	SW	
V.	ICP Interference Check Sample (ICS) Analysis	—	
VI.	Matrix Spike Analysis	SW	MSIP
VII.	Duplicate Sample Analysis	SW	DUP
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	A	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	—	
XV.	Field Blanks	SW	EB=EB2-071013 (PH064)

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

FB=FB-041113
 (PH029)

Validated Samples:

soil

1	SL-586-SA8-SB-0.0-0.5	11		21		31	
2	SL-588-SA8-SB-0.0-0.5	12		22		32	
3	SL-888-SA8-SB-0.0-0.5	13		23		33	
4	SL-588-SA8-SB-2.0-3.0	14		24		34	
5	SL-588-SA8-SB-0.0-0.5MS	15		25		35	
6	SL-588-SA8-SB-0.0-0.5MSD	16		26		36	
7	SL-588-SA8-SB-0.0-0.5DUP	17		27		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes: _____

**VALIDATION FINDINGS WORKSHEET
PB/ICB/CCB QUALIFIED SAMPLES**

METHOD: Trace metals (EPA SW 864 Method 6010B/6020/7000)

Soil preparation factor applied: 100x

Reason: B

Sample Concentration units, unless otherwise noted: ug/L mg/kg

Associated Samples: All

Analyte	Maximum PB* (mg/Kg)	Maximum PB* (ug/L)	Maximum ICB/CCB* (ug/L)	Action Level	Sample Identification										
					1	2	3	4							
Mo			3.9	1.95	0.65	0.84	0.56	0.51							

Samples with analyte concentrations within five times the associated ICB, CCB or PB concentration are listed above with the identifications from the Validation Completeness Worksheet. These sample results were qualified as not detected, "U".

Note : a - The listed analyte concentration is the highest ICB, CCB, or PB detected in the analysis of each element.

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification										
	FB-041113 (SDG: PH029)	Action Limit	1	2	3	4						
Cu	0.0036	1.8										
Mo	0.0036	1.8	0.65	0.84	0.56	0.51						

Sampling date: 7/10/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification										
	EB2-071013 (SDG: PH064)	Action Limit	No Qualifiers									
Cu	0.0030	1.5										

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".



Background Lab Sample ID: 7120502BKG Matrix Spike Lab Sample ID: 7120503MS Matrix Spike Duplicate Lab Sample ID: 7120504MSD
Batch Id(s): P19137A, P19138A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Control Limit			
		Result	C	Result	C	Result	C				%R	Q	%R	Q		%R	RPD	M	
Aluminum		15805.5029		19747.6560		19780.0500		200.0000	196.0784	MG/KG	197		2027		0			20	P
Antimony		0.7255	U	18.6030		18.0853		50.0000	49.0196	MG/KG	37	N	37	N	3	75 - 125		20	P
Arsenic		4.1069		19.7570		19.2451		15.0000	14.7059	MG/KG	104		103		3	75 - 125		20	P
Barium		102.4804		313.7070		309.5902		200.0000	196.0784	MG/KG	106		106		1	75 - 125		20	P
Beryllium		0.5167	B	5.7610		5.6775		5.0000	4.9020	MG/KG	105		105		1	75 - 125		20	P
Boron		2.6059	B	204.8300		200.7216		200.0000	196.0784	MG/KG	101		101		2	75 - 125		20	P
Cadmium		0.4186	B	5.4390		5.2735		5.0000	4.9020	MG/KG	100		99		3	75 - 125		20	P
Calcium		2190.9853		3104.0020		2928.4078		400.0000	392.1569	MG/KG	228		188		6			20	P
Chromium		15.8127		39.8230		37.7255		20.0000	19.6078	MG/KG	120		112		5	75 - 125		20	P
Cobalt		4.3098		53.7250		52.3353		50.0000	49.0196	MG/KG	99		98		3	75 - 125		20	P
Copper		10.3059		37.2340		36.5784		25.0000	24.5098	MG/KG	108		107		2	75 - 125		20	P
Iron		16896.1314		18609.6560		18362.1882		100.0000	98.0392	MG/KG	1714		1495		1			20	P
Lead		8.9167		24.4460		23.7941		15.0000	14.7059	MG/KG	104		101		3	75 - 125		20	P
Lithium		19.9392		124.9810		122.4069		100.0000	98.0392	MG/KG	105		105		2	75 - 125		20	P
Magnesium		3466.4882		4058.9570		4051.0843		200.0000	196.0784	MG/KG	296		298		0			20	P
Manganese		321.6892		387.7610		389.0029		50.0000	49.0196	MG/KG	132		137		0			20	P
Mercury		0.0119	B	0.1931		0.2075		0.1652	0.1655	MG/KG	110		118		7	65 - 135		20	CV
Molybdenum		0.8353	B	198.8710		195.0304		200.0000	196.0784	MG/KG	99		99		2	75 - 125		20	P
Nickel		10.2500		61.4360		59.7853		50.0000	49.0196	MG/KG	102		101		3	75 - 125		20	P
Phosphorus		488.5598		608.8550		608.2971		100.0000	98.0392	MG/KG	120		122		0			20	P
Potassium		3654.5431		5232.0070		5243.0422		1000.0000	980.3922	MG/KG	158	N	162	N	0	75 - 125		20	P
Selenium	78	0.1012	B	1.8504		1.9647		2.0000	1.9608	MG/KG	87		95		6	75 - 125		20	MS
Silver	107	0.0323	B	10.3100		9.7078		10.0000	9.8039	MG/KG	103		99		6	75 - 125		20	MS
Sodium		65.8725	B	1069.6280		1039.3000		1000.0000	980.3922	MG/KG	100		99		3	75 - 125		20	P
Strontium	88	19.0863		28.3200		27.3333		8.0000	7.8431	MG/KG	115		105		4	75 - 125		20	MS
Thallium	203	0.2373		0.6384		0.6345		0.4000	0.3922	MG/KG	100		101		1	75 - 125		20	MS
Tin		2.4471	B	368.6250		361.4343		400.0000	392.1569	MG/KG	92		92		2	75 - 125		20	P
Titanium		1006.1961		1377.4590		1390.1735		100.0000	98.0392	MG/KG	371		392		1			20	P
Vanadium		28.6529		84.7950		82.9422		50.0000	49.0196	MG/KG	112		111		2	75 - 125		20	P
Zinc		61.7814		121.8310		116.8833		50.0000	49.0196	MG/KG	120		112		4	75 - 125		20	P
Zirconium		3.5882	B	101.0480		99.4794		100.0000	98.0392	MG/KG	97		98		2	75 - 125		20	P

Note: Results shown are reported on an as-received basis.

<p>METHODS: P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>		<p>CONCENTRATION QUALIFIERS: U= Below MDL, B= Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS</p>	
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Background Lab Sample ID: 7120502BKG

Duplicate Lab Sample ID: 7120505DUP

Batch ID(s): P19137A, P19138A

Concentration Units: MG/KG

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			15805.5029		16890.0900		7		P
Antimony			0.7255	U	0.8430	B	200		P
Arsenic		3.9	4.1069		4.4610		8		P
Barium			102.4804		109.3240		6		P
Beryllium			0.5167	B	0.5300	B	3		P
Boron			2.6059	B	3.3450	B	25		P
Cadmium			0.4186	B	0.8860	B	72		P
Calcium			2190.9853		3308.0810		41	*	P
Chromium			15.8127		18.4020		15		P
Cobalt		1.0	4.3098		4.8120		11		P
Copper			10.3059		11.3550		10		P
Iron			16896.1314		18563.3180		9		P
Lead		2.9	8.9167		10.3420		15		P
Lithium			19.9392		22.2850		11		P
Magnesium			3466.4882		3912.7770		12		P
Manganese			321.6892		345.0400		7		P
Mercury		0.0	0.0119	B	0.0278		80		CV
Molybdenum			0.8353	B	0.6260	B	29		P
Nickel			10.2500		11.4710		11		P
Phosphorus			488.5598		566.0640		15		P
Potassium			3654.5431		4623.5210		23	*	P
Selenium	78		0.1012	B	0.1254	B	21		MS
Silver	107		0.0323	B	0.0295	B	9		MS
Sodium			65.8725	B	82.7290	B	23		P
Strontium	88		19.0863		24.9000		26	*	MS
Thallium	203	0.2	0.2373		0.2486		5		MS
Tin			2.4471	B	2.5500	B	4		P
Titanium			1006.1961		1024.9500		2		P
Vanadium			28.6529		32.9080		14		P
Zinc			61.7814		114.4030		60	*	P
Zirconium			3.5882	B	4.8200	B	29		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).

The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

ok by difference

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer</p> <p>MS = ICP Mass Spectrometry</p> <p>CV = Cold Vapor</p> <p>AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U= Below MDL</p> <p>B= Below LOQ</p> <p>FLAGS:</p> <p>= Duplicate Out of Spec</p>
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**Data Validation Report
Santa Susan Field Laboratory
Subarea 8**

SDG(s): PH063

Prepared for

CDM
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
7750 El Camino Real, Suite 2L
Carlsbad, California 92009

September 10, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on July 9, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory, RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles by Environmental Protection Agency (EPA) SW 846 Method 8270D-SIM
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A, and 7471B
Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M
Total Petroleum Hydrocarbons (TPH) as Extractables by EPA SW 846 Method 8015M
Dioxins/Dibenzofurans by EPA Method 1613B

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibrations blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, field blanks and equipment blanks. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the calibration blanks and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of three method blanks for dioxins and metals. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The samples results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks with the following exceptions:

SDG/ Method	Method Blank ID	Analyte	Maximum Concentration	Associated Samples
PH063/ 6020A	ICB/CCB	Molybdenum	3.9 ug/L	SL-561-SA8-SB-0.0-0.5 SL-561-SA8-SB-4.0-5.0 SL-562-SA8-SB-0.0-0.5 SL-563-SA8-SB-0.0-0.5 SL-563-SA8-SB-2.0-3.0

Sample concentrations were compared to concentrations detected in the initial and continuing blanks. The sample concentrations were not detected or were significantly greater than the concentrations found in the associated blanks with the following exceptions:

SDG/Method	Sample	Compound	Reported Concentration	Modified Final Concentration
PH063/ 6020A	SL-561-SA8-SB-0.0-0.5	Molybdenum	0.51 mg/kg	0.51U mg/kg
PH063/ 6020A	SL-561-SA8-SB-4.0-5.0	Molybdenum	0.28 mg/kg	0.28U mg/kg
PH063/ 6020A	SL-562-SA8-SB-0.0-0.5	Molybdenum	0.47 mg/kg	0.47U mg/kg

SDG/Method	Sample	Compound	Reported Concentration	Modified Final Concentration
PH063/ 6020A	SL-563-SA8-SB-0.0-0.5	Molybdenum	0.39 mg/kg	0.39U mg/kg
PH063/ 6020A	SL-563-SA8-SB-2.0-3.0	Molybdenum	0.36 mg/kg	0.36U mg/kg

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for metals performed in SDG PH062. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. Details regarding the qualification of data are provided in Enclosure I of the DVR for SDG PH062.

VIII. Laboratory Duplicates Sample

Laboratory duplicate (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits with the exception of one DUP for metals performed in SDG PH062. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. Details regarding the qualification of data are provided in the Enclosure I of the DVR for SDG PH062.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified

requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH063	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

No field duplicates were identified in this SDG.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH as gasoline. No contaminants were found in the trip blank.

One equipment blank (from SDG PH064) was collected and analyzed for semivolatiles, PCBs, metals, TPH as gasoline, TPH as extractables, and dioxins. No contaminants were found in the equipment blank with the exception of several dioxins, semivolatiles, and metals. The associated sample results were not detected or were significantly greater than the concentrations found in the associated blanks, therefore no data were qualified.

One field blank (from SDG PH029) was collected and analyzed for semivolatiles, PCBs, metals, TPH as gasoline, TPH as extractables and dioxins. No contaminants were found in the field blank with the exception of several dioxins, semivolatiles, and metals. Associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The samples results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
09-Jul-2013	TB-070913	7122287	TB	5030B	8015M	III
09-Jul-2013	SL-562-SA8-SB-0.0-0.5	7122290	N	3050B	6010C	III
09-Jul-2013	SL-562-SA8-SB-0.0-0.5	7122290	N	3050B	6020A	III
09-Jul-2013	SL-562-SA8-SB-0.0-0.5	7122290	N	3546	8015M	III
09-Jul-2013	SL-562-SA8-SB-0.0-0.5	7122290	N	3546	8082A	III
09-Jul-2013	SL-562-SA8-SB-0.0-0.5	7122290	N	3546	8270D SIM	III
09-Jul-2013	SL-562-SA8-SB-0.0-0.5	7122290	N	METHOD	1613B	III
09-Jul-2013	SL-562-SA8-SB-0.0-0.5	7122290	N	METHOD	7471B	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5	7122288	N	3050B	6010C	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5	7122288	N	3050B	6020A	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5	7122288	N	3546	8015M	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5	7122288	N	3546	8082A	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5	7122288	N	3546	8270D SIM	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5	7122288	N	METHOD	1613B	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5	7122288	N	METHOD	7471B	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5MSD	P122288M240210A	MSD	3546	8082A	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5MSD	P122288M260632	MSD	3546	8270D SIM	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5MS	P122288R240151A	MS	3546	8082A	III
09-Jul-2013	SL-561-SA8-SB-0.0-0.5MS	P122288R260600	MS	3546	8270D SIM	III
09-Jul-2013	SL-561-SA8-SB-4.0-5.0	7122289	N	3050B	6010C	III
09-Jul-2013	SL-561-SA8-SB-4.0-5.0	7122289	N	3050B	6020A	III
09-Jul-2013	SL-561-SA8-SB-4.0-5.0	7122289	N	3546	8015M	III
09-Jul-2013	SL-561-SA8-SB-4.0-5.0	7122289	N	3546	8082A	III
09-Jul-2013	SL-561-SA8-SB-4.0-5.0	7122289	N	3546	8270D SIM	III
09-Jul-2013	SL-561-SA8-SB-4.0-5.0	7122289	N	5035A	8015M	III
09-Jul-2013	SL-561-SA8-SB-4.0-5.0	7122289	N	METHOD	1613B	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
09-Jul-2013	SL-561-SA8-SB-4.0-5.0	7122289	N	METHOD	7471B	III
09-Jul-2013	SL-563-SA8-SB-0.0-0.5	7122291	N	3050B	6010C	III
09-Jul-2013	SL-563-SA8-SB-0.0-0.5	7122291	N	3050B	6020A	III
09-Jul-2013	SL-563-SA8-SB-0.0-0.5	7122291	N	3546	8015M	III
09-Jul-2013	SL-563-SA8-SB-0.0-0.5	7122291	N	3546	8082A	III
09-Jul-2013	SL-563-SA8-SB-0.0-0.5	7122291	N	3546	8270D SIM	III
09-Jul-2013	SL-563-SA8-SB-0.0-0.5	7122291	N	METHOD	1613B	III
09-Jul-2013	SL-563-SA8-SB-0.0-0.5	7122291	N	METHOD	7471B	III
09-Jul-2013	SL-563-SA8-SB-2.0-3.0	7122292	N	3050B	6010C	III
09-Jul-2013	SL-563-SA8-SB-2.0-3.0	7122292	N	3050B	6020A	III
09-Jul-2013	SL-563-SA8-SB-2.0-3.0	7122292	N	3546	8015M	III
09-Jul-2013	SL-563-SA8-SB-2.0-3.0	7122292	N	3546	8082A	III
09-Jul-2013	SL-563-SA8-SB-2.0-3.0	7122292	N	3546	8270D SIM	III
09-Jul-2013	SL-563-SA8-SB-2.0-3.0	7122292	N	5035A	8015M	III
09-Jul-2013	SL-563-SA8-SB-2.0-3.0	7122292	N	METHOD	1613B	III
09-Jul-2013	SL-563-SA8-SB-2.0-3.0	7122292	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-561-SA8-SB-0.0-0.5

Collected: 7/9/2013 9:45:00 AM Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.08	U	0.755	MDL	4.08	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.611	J	0.0684	MDL	1.02	PQL	mg/Kg	J	Z
BORON	4.63	J	0.857	MDL	10.2	PQL	mg/Kg	J	Z
CADMIUM	0.477	J	0.0776	MDL	1.02	PQL	mg/Kg	J	Z
CALCIUM	3500		3.41	MDL	20.4	PQL	mg/Kg	J	E
MOLYBDENUM	0.508	J	0.174	MDL	2.04	PQL	mg/Kg	U	B, F
POTASSIUM	4740		8.51	MDL	102	PQL	mg/Kg	J	Q, E
SODIUM	76.3	J	17.0	MDL	102	PQL	mg/Kg	J	Z
TIN	2.46	J	0.225	MDL	10.2	PQL	mg/Kg	U	B
ZINC	58.6		0.204	MDL	4.08	PQL	mg/Kg	J	E
Zirconium	4.50	J	0.857	MDL	5.10	PQL	mg/Kg	J	Z

Sample ID: SL-561-SA8-SB-4.0-5.0

Collected: 7/9/2013 11:40:00 Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.31	U	0.797	MDL	4.31	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.975	J	0.0722	MDL	1.08	PQL	mg/Kg	J	Z
BORON	3.82	J	0.905	MDL	10.8	PQL	mg/Kg	J	Z
CADMIUM	0.400	J	0.0819	MDL	1.08	PQL	mg/Kg	J	Z
CALCIUM	6820		3.60	MDL	21.5	PQL	mg/Kg	J	E
MOLYBDENUM	0.280	J	0.183	MDL	2.15	PQL	mg/Kg	U	B, F
POTASSIUM	2910		8.99	MDL	108	PQL	mg/Kg	J	Q, E
TIN	2.80	J	0.237	MDL	10.8	PQL	mg/Kg	U	B
ZINC	62.1		0.215	MDL	4.31	PQL	mg/Kg	J	E

Sample ID: SL-562-SA8-SB-0.0-0.5

Collected: 7/9/2013 8:45:00 AM Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	3.96	U	0.733	MDL	3.96	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.618	J	0.0664	MDL	0.991	PQL	mg/Kg	J	Z
BORON	2.75	J	0.832	MDL	9.91	PQL	mg/Kg	J	Z
CADMIUM	0.405	J	0.0753	MDL	0.991	PQL	mg/Kg	J	Z
CALCIUM	2520		3.31	MDL	19.8	PQL	mg/Kg	J	E
MOLYBDENUM	0.467	J	0.168	MDL	1.98	PQL	mg/Kg	U	B, F
POTASSIUM	3550		8.26	MDL	99.1	PQL	mg/Kg	J	Q, E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-562-SA8-SB-0.0-0.5 Collected: 7/9/2013 8:45:00 AM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SODIUM	80.1	J	16.5	MDL	99.1	PQL	mg/Kg	J	Z
TIN	2.38	J	0.218	MDL	9.91	PQL	mg/Kg	U	B
ZINC	50.1		0.198	MDL	3.96	PQL	mg/Kg	J	E
Zirconium	4.13	J	0.832	MDL	4.95	PQL	mg/Kg	J	Z

Sample ID: SL-563-SA8-SB-0.0-0.5 Collected: 7/9/2013 1:50:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.02	U	0.744	MDL	4.02	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.573	J	0.0673	MDL	1.01	PQL	mg/Kg	J	Z
BORON	2.72	J	0.844	MDL	10.1	PQL	mg/Kg	J	Z
CADMIUM	0.431	J	0.0764	MDL	1.01	PQL	mg/Kg	J	Z
CALCIUM	2780		3.36	MDL	20.1	PQL	mg/Kg	J	E
MOLYBDENUM	0.390	J	0.171	MDL	2.01	PQL	mg/Kg	U	B, F
POTASSIUM	3510		8.38	MDL	101	PQL	mg/Kg	J	Q, E
SODIUM	73.0	J	16.8	MDL	101	PQL	mg/Kg	J	Z
TIN	2.30	J	0.221	MDL	10.1	PQL	mg/Kg	U	B
ZINC	52.8		0.201	MDL	4.02	PQL	mg/Kg	J	E
Zirconium	3.77	J	0.844	MDL	5.03	PQL	mg/Kg	J	Z

Sample ID: SL-563-SA8-SB-2.0-3.0 Collected: 7/9/2013 2:30:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.08	U	0.755	MDL	4.08	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.755	J	0.0684	MDL	1.02	PQL	mg/Kg	J	Z
BORON	2.67	J	0.857	MDL	10.2	PQL	mg/Kg	J	Z
CADMIUM	0.434	J	0.0776	MDL	1.02	PQL	mg/Kg	J	Z
CALCIUM	2940		3.41	MDL	20.4	PQL	mg/Kg	J	E
MOLYBDENUM	0.364	J	0.174	MDL	2.04	PQL	mg/Kg	U	B, F
POTASSIUM	3160		8.51	MDL	102	PQL	mg/Kg	J	Q, E
SODIUM	100	J	17.0	MDL	102	PQL	mg/Kg	J	Z
TIN	2.63	J	0.225	MDL	10.2	PQL	mg/Kg	U	B
ZINC	56.5		0.204	MDL	4.08	PQL	mg/Kg	J	E
Zirconium	4.55	J	0.857	MDL	5.10	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-561-SA8-SB-0.0-0.5 Collected: 7/9/2013 9:45:00 AM Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.135	J	0.102	MDL	0.408	PQL	mg/Kg	J	Z

Sample ID: SL-561-SA8-SB-0.0-0.5 Collected: 7/9/2013 9:45:00 AM Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	24.6		0.0694	MDL	0.408	PQL	mg/Kg	J	E

Sample ID: SL-561-SA8-SB-4.0-5.0 Collected: 7/9/2013 11:40:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0465	J	0.0280	MDL	0.215	PQL	mg/Kg	J	Z
STRONTIUM	36.6		0.0733	MDL	0.431	PQL	mg/Kg	J	E

Sample ID: SL-562-SA8-SB-0.0-0.5 Collected: 7/9/2013 8:45:00 AM Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.130	J	0.0991	MDL	0.396	PQL	mg/Kg	J	Z

Sample ID: SL-562-SA8-SB-0.0-0.5 Collected: 7/9/2013 8:45:00 AM Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0399	J	0.0258	MDL	0.198	PQL	mg/Kg	J	Z
STRONTIUM	22.7		0.0674	MDL	0.396	PQL	mg/Kg	J	E

Sample ID: SL-563-SA8-SB-0.0-0.5 Collected: 7/9/2013 1:50:00 PM Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.151	J	0.101	MDL	0.402	PQL	mg/Kg	J	Z

Sample ID: SL-563-SA8-SB-0.0-0.5 Collected: 7/9/2013 1:50:00 PM Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0280	J	0.0261	MDL	0.201	PQL	mg/Kg	J	Z
STRONTIUM	21.9		0.0684	MDL	0.402	PQL	mg/Kg	J	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-563-SA8-SB-2.0-3.0 Collected: 7/9/2013 2:30:00 PM Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.158	J	0.102	MDL	0.408	PQL	mg/Kg	J	Z

Sample ID: SL-563-SA8-SB-2.0-3.0 Collected: 7/9/2013 2:30:00 PM Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0400	J	0.0265	MDL	0.204	PQL	mg/Kg	J	Z
STRONTIUM	25.0		0.0694	MDL	0.408	PQL	mg/Kg	J	E

Method Category:	METALS	
Method:	7471B	Matrix: SO

Sample ID: SL-562-SA8-SB-0.0-0.5 Collected: 7/9/2013 8:45:00 AM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0148	J	0.0099	MDL	0.0165	PQL	mg/Kg	J	Z

Sample ID: SL-563-SA8-SB-0.0-0.5 Collected: 7/9/2013 1:50:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0116	J	0.0096	MDL	0.0160	PQL	mg/Kg	J	Z

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-561-SA8-SB-0.0-0.5 Collected: 7/9/2013 9:45:00 AM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	1.81	JB	0.0234	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.187	JBQ	0.0284	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.185	JB	0.0458	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDF	0.229	JB	0.0260	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.945	JB	0.0474	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.284	JB	0.0242	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	1.14	JB	0.0482	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDF	0.844	JB	0.0272	MDL	5.11	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-561-SA8-SB-0.0-0.5 Collected: 7/9/2013 9:45:00 AM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,7,8-PECDD	0.205	JB	0.0362	MDL	5.11	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.484	JB	0.0252	MDL	5.11	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.281	JB	0.0242	MDL	5.11	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.475	JB	0.0238	MDL	5.11	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.349	J	0.0546	MDL	1.02	PQL	ng/Kg	J	Z
OCDF	3.70	JB	0.0182	MDL	10.2	PQL	ng/Kg	J	Z

Sample ID: SL-561-SA8-SB-4.0-5.0 Collected: 7/9/2013 11:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.26	JB	0.0298	MDL	5.42	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.248	JBQ	0.0115	MDL	5.42	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0995	JBQ	0.0149	MDL	5.42	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.162	JB	0.0284	MDL	5.42	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.182	JBQ	0.0139	MDL	5.42	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.213	JBQ	0.0284	MDL	5.42	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.181	JBQ	0.0132	MDL	5.42	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.199	JB	0.0296	MDL	5.42	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.193	JB	0.0157	MDL	5.42	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.255	JBQ	0.0246	MDL	5.42	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.419	JB	0.0169	MDL	5.42	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.128	JB	0.0133	MDL	5.42	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.259	JB	0.0151	MDL	5.42	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0640	JQ	0.0256	MDL	1.08	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.124	JQ	0.0284	MDL	1.08	PQL	ng/Kg	J	Z
OCDD	9.19	JB	0.0189	MDL	10.8	PQL	ng/Kg	J	Z
OCDF	0.457	JB	0.0224	MDL	10.8	PQL	ng/Kg	J	Z

Sample ID: SL-562-SA8-SB-0.0-0.5 Collected: 7/9/2013 8:45:00 AM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	3.38	JB	0.0287	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.812	JB	0.0153	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.127	JB	0.0196	MDL	5.09	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.165	JB	0.0305	MDL	5.09	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-562-SA8-SB-0.0-0.5

Collected: 7/9/2013 8:45:00 AM Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HXCDF	0.229	JB	0.0179	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.835	JB	0.0319	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.299	JB	0.0167	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	1.08	JB	0.0314	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.716	JB	0.0195	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.343	JB	0.0312	MDL	5.09	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.561	JB	0.0177	MDL	5.09	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.200	JB	0.0173	MDL	5.09	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.496	JBQ	0.0161	MDL	5.09	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.177	J	0.0330	MDL	1.02	PQL	ng/Kg	J	Z
OCDF	1.40	JB	0.0170	MDL	10.2	PQL	ng/Kg	J	Z

Sample ID: SL-563-SA8-SB-0.0-0.5

Collected: 7/9/2013 1:50:00 PM Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	4.37	JB	0.0309	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	1.03	JB	0.0139	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.146	JBQ	0.0160	MDL	5.03	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.184	JBQ	0.0329	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.236	JB	0.0156	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.827	JB	0.0350	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.260	JB	0.0149	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.982	JB	0.0357	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.729	JB	0.0168	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.315	JB	0.0240	MDL	5.03	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.498	JB	0.0151	MDL	5.03	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.190	JBQ	0.0153	MDL	5.03	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.380	JB	0.0141	MDL	5.03	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0540	JQ	0.0190	MDL	1.01	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.110	J	0.0309	MDL	1.01	PQL	ng/Kg	J	Z
OCDF	2.97	JB	0.0150	MDL	10.1	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-563-SA8-SB-2.0-3.0 Collected: 7/9/2013 2:30:00 PM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.454	JB	0.0277	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.152	JB	0.0131	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0678	JBQ	0.0160	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.116	JB	0.0346	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.141	JBQ	0.0118	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.449	JBQ	0.0363	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.157	JBQ	0.0121	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.658	JB	0.0359	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDF	0.732	JBQ	0.0131	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.211	JBQ	0.0281	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.342	JB	0.0143	MDL	5.22	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HxCDF	0.0936	JB	0.0120	MDL	5.22	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.194	JB	0.0123	MDL	5.22	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0293	J	0.0242	MDL	1.04	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0655	J	0.0235	MDL	1.04	PQL	ng/Kg	J	Z
OCDD	4.24	JB	0.0210	MDL	10.4	PQL	ng/Kg	J	Z
OCDF	0.310	JB	0.0273	MDL	10.4	PQL	ng/Kg	U	B

Method Category:	SVOA	
Method:	8015M	Matrix: SO

Sample ID: SL-561-SA8-SB-0.0-0.5 Collected: 7/9/2013 9:45:00 AM Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	3.1	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z

Sample ID: SL-563-SA8-SB-2.0-3.0 Collected: 7/9/2013 2:30:00 PM Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	4.9	J	2.1	MDL	5.3	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-561-SA8-SB-0.0-0.5 Collected: 7/9/2013 9:45:00 AM Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	1.2	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(A)ANTHRACENE	0.79	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	0.94	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	0.82	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	8.8	J	6.2	MDL	18	PQL	ug/Kg	J	Z
FLUORENE	0.87	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
INDENO(1,2,3-CD)PYRENE	0.71	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
NAPHTHALENE	1.3	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-562-SA8-SB-0.0-0.5 Collected: 7/9/2013 8:45:00 AM Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.70	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
CHRYSENE	0.80	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-563-SA8-SB-0.0-0.5 Collected: 7/9/2013 1:50:00 PM Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	0.81	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z
FLUORANTHENE	0.76	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-563-SA8-SB-2.0-3.0 Collected: 7/9/2013 2:30:00 PM Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(K)FLUORANTHENE	0.88	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	0.37	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Calibration Blank Contamination
B	Method Blank Contamination
E	Laboratory Duplicate Precision
F	Field Blank Contamination
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

9/4/2013 8:45:25 AM

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Enclosure I
Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH063

Method Blank Outlier Report

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK1920B370624	7/13/2013 6:24:00 AM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF OCDD OCDF	0.0814 ng/Kg 0.0265 ng/Kg 0.0357 ng/Kg 0.0239 ng/Kg 0.0351 ng/Kg 0.0206 ng/Kg 0.0220 ng/Kg 0.0274 ng/Kg 0.0184 ng/Kg 0.0216 ng/Kg 0.0503 ng/Kg 0.0269 ng/Kg 0.0333 ng/Kg 0.214 ng/Kg 0.0705 ng/Kg	SL-561-SA8-SB-0.0-0.5 SL-561-SA8-SB-4.0-5.0 SL-562-SA8-SB-0.0-0.5 SL-563-SA8-SB-0.0-0.5 SL-563-SA8-SB-2.0-3.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-561-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0995 ng/Kg	0.0995U ng/Kg
SL-561-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.128 ng/Kg	0.128U ng/Kg
SL-562-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.127 ng/Kg	0.127U ng/Kg
SL-563-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.146 ng/Kg	0.146U ng/Kg
SL-563-SA8-SB-2.0-3.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0678 ng/Kg	0.0678U ng/Kg
SL-563-SA8-SB-2.0-3.0(RES)	1,2,3,4,7,8-HxCDD	0.116 ng/Kg	0.116U ng/Kg
SL-563-SA8-SB-2.0-3.0(RES)	1,2,3,4,7,8-HxCDF	0.141 ng/Kg	0.141U ng/Kg
SL-563-SA8-SB-2.0-3.0(RES)	2,3,4,6,7,8-HXCDF	0.0936 ng/Kg	0.0936U ng/Kg
SL-563-SA8-SB-2.0-3.0(RES)	OCDF	0.310 ng/Kg	0.310U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19137AB221846	7/12/2013 6:46:00 PM	ALUMINUM CALCIUM IRON MAGNESIUM TIN TITANIUM ZINC	11.0 mg/Kg 12.8 mg/Kg 6.78 mg/Kg 5.68 mg/Kg 1.32 mg/Kg 0.177 mg/Kg 0.422 mg/Kg	SL-561-SA8-SB-0.0-0.5 SL-561-SA8-SB-4.0-5.0 SL-562-SA8-SB-0.0-0.5 SL-563-SA8-SB-0.0-0.5 SL-563-SA8-SB-2.0-3.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-561-SA8-SB-0.0-0.5(RES)	TIN	2.46 mg/Kg	2.46U mg/Kg
SL-561-SA8-SB-4.0-5.0(RES)	TIN	2.80 mg/Kg	2.80U mg/Kg
SL-562-SA8-SB-0.0-0.5(RES)	TIN	2.38 mg/Kg	2.38U mg/Kg
SL-563-SA8-SB-0.0-0.5(RES)	TIN	2.30 mg/Kg	2.30U mg/Kg
SL-563-SA8-SB-2.0-3.0(RES)	TIN	2.63 mg/Kg	2.63U mg/Kg

Method Blank Outlier Report

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19137AB221713A	7/18/2013 5:13:00 PM	STRONTIUM	0.0774 mg/Kg	SL-561-SA8-SB-0.0-0.5 SL-561-SA8-SB-4.0-5.0 SL-562-SA8-SB-0.0-0.5 SL-563-SA8-SB-0.0-0.5 SL-563-SA8-SB-2.0-3.0

Field Blank Outlier Report

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(REA2)	4/11/2013 3:00:00 PM	COPPER MOLYBDENUM	0.0036 mg/L 0.0036 mg/L	SL-561-SA8-SB-0.0-0.5 SL-561-SA8-SB-4.0-5.0 SL-562-SA8-SB-0.0-0.5 SL-563-SA8-SB-0.0-0.5 SL-563-SA8-SB-2.0-3.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-561-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.508 mg/Kg	0.508U mg/Kg
SL-561-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.280 mg/Kg	0.280U mg/Kg
SL-562-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.467 mg/Kg	0.467U mg/Kg
SL-563-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.390 mg/Kg	0.390U mg/Kg
SL-563-SA8-SB-2.0-3.0(RES)	MOLYBDENUM	0.364 mg/Kg	0.364U mg/Kg

Reporting Limit Outliers

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	1.81	5.11	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JBQ	0.187	5.11	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.185	5.11	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.229	5.11	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.945	5.11	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.284	5.11	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	1.14	5.11	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.844	5.11	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.205	5.11	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.484	5.11	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.281	5.11	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.475	5.11	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.349	1.02	PQL	ng/Kg	
	OCDF	JB	3.70	10.2	PQL	ng/Kg	
SL-561-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	1.26	5.42	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.248	5.42	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0995	5.42	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.162	5.42	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.182	5.42	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.213	5.42	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.181	5.42	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.199	5.42	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.193	5.42	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.255	5.42	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.419	5.42	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.128	5.42	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.259	5.42	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0640	1.08	PQL	ng/Kg	
2,3,7,8-TCDF	JQ	0.124	1.08	PQL	ng/Kg		
OCDD	JB	9.19	10.8	PQL	ng/Kg		
OCDF	JB	0.457	10.8	PQL	ng/Kg		
SL-562-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	3.38	5.09	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.812	5.09	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.127	5.09	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.165	5.09	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.229	5.09	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.835	5.09	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.299	5.09	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	1.08	5.09	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.716	5.09	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.343	5.09	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.561	5.09	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.200	5.09	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.496	5.09	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.177	1.02	PQL	ng/Kg	
OCDF	JB	1.40	10.2	PQL	ng/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-563-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	4.37	5.03	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	1.03	5.03	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.146	5.03	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.184	5.03	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.236	5.03	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.827	5.03	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.260	5.03	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.982	5.03	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.729	5.03	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.315	5.03	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.498	5.03	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.190	5.03	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.380	5.03	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0540	1.01	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.110	1.01	PQL	ng/Kg	
OCDF	JB	2.97	10.1	PQL	ng/Kg		
SL-563-SA8-SB-2.0-3.0	1,2,3,4,6,7,8-HPCDD	JB	0.454	5.22	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.152	5.22	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0678	5.22	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.116	5.22	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.141	5.22	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.449	5.22	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.157	5.22	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.658	5.22	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JBQ	0.732	5.22	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.211	5.22	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.342	5.22	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.0936	5.22	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.194	5.22	PQL	ng/Kg	
	2,3,7,8-TCDD	J	0.0293	1.04	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.0655	1.04	PQL	ng/Kg	
OCDD	JB	4.24	10.4	PQL	ng/Kg		
OCDF	JB	0.310	10.4	PQL	ng/Kg		

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA8-SB-0.0-0.5	BERYLLIUM	J	0.611	1.02	PQL	mg/Kg	J (all detects)
	BORON	J	4.63	10.2	PQL	mg/Kg	
	CADMIUM	J	0.477	1.02	PQL	mg/Kg	
	MOLYBDENUM	J	0.508	2.04	PQL	mg/Kg	
	SODIUM	J	76.3	102	PQL	mg/Kg	
	TIN	J	2.46	10.2	PQL	mg/Kg	
	Zirconium	J	4.50	5.10	PQL	mg/Kg	
SL-561-SA8-SB-4.0-5.0	BERYLLIUM	J	0.975	1.08	PQL	mg/Kg	J (all detects)
	BORON	J	3.82	10.8	PQL	mg/Kg	
	CADMIUM	J	0.400	1.08	PQL	mg/Kg	
	MOLYBDENUM	J	0.280	2.15	PQL	mg/Kg	
	TIN	J	2.80	10.8	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-562-SA8-SB-0.0-0.5	BERYLLIUM	J	0.618	0.991	PQL	mg/Kg	J (all detects)
	BORON	J	2.75	9.91	PQL	mg/Kg	
	CADMIUM	J	0.405	0.991	PQL	mg/Kg	
	MOLYBDENUM	J	0.467	1.98	PQL	mg/Kg	
	SODIUM	J	80.1	99.1	PQL	mg/Kg	
	TIN	J	2.38	9.91	PQL	mg/Kg	
	Zirconium	J	4.13	4.95	PQL	mg/Kg	
SL-563-SA8-SB-0.0-0.5	BERYLLIUM	J	0.573	1.01	PQL	mg/Kg	J (all detects)
	BORON	J	2.72	10.1	PQL	mg/Kg	
	CADMIUM	J	0.431	1.01	PQL	mg/Kg	
	MOLYBDENUM	J	0.390	2.01	PQL	mg/Kg	
	SODIUM	J	73.0	101	PQL	mg/Kg	
	TIN	J	2.30	10.1	PQL	mg/Kg	
	Zirconium	J	3.77	5.03	PQL	mg/Kg	
SL-563-SA8-SB-2.0-3.0	BERYLLIUM	J	0.755	1.02	PQL	mg/Kg	J (all detects)
	BORON	J	2.67	10.2	PQL	mg/Kg	
	CADMIUM	J	0.434	1.02	PQL	mg/Kg	
	MOLYBDENUM	J	0.364	2.04	PQL	mg/Kg	
	SODIUM	J	100	102	PQL	mg/Kg	
	TIN	J	2.63	10.2	PQL	mg/Kg	
	Zirconium	J	4.55	5.10	PQL	mg/Kg	

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA8-SB-0.0-0.5	SELENIUM	J	0.135	0.408	PQL	mg/Kg	J (all detects)
SL-561-SA8-SB-4.0-5.0	SILVER	J	0.0465	0.215	PQL	mg/Kg	J (all detects)
SL-562-SA8-SB-0.0-0.5	SELENIUM	J	0.130	0.396	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0399	0.198	PQL	mg/Kg	
SL-563-SA8-SB-0.0-0.5	SELENIUM	J	0.151	0.402	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0280	0.201	PQL	mg/Kg	
SL-563-SA8-SB-2.0-3.0	SELENIUM	J	0.158	0.408	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0400	0.204	PQL	mg/Kg	

Method: 7471B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-562-SA8-SB-0.0-0.5	MERCURY	J	0.0148	0.0165	PQL	mg/Kg	J (all detects)
SL-563-SA8-SB-0.0-0.5	MERCURY	J	0.0116	0.0160	PQL	mg/Kg	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH063

Laboratory: LL

EDD Filename: PH063_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8015M
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA8-SB-0.0-0.5	EFH (C15-C20)	J	3.1	5.2	PQL	mg/Kg	J (all detects)
SL-563-SA8-SB-2.0-3.0	EFH (C21-C30)	J	4.9	5.3	PQL	mg/Kg	J (all detects)

Method: 8270D SIM
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-561-SA8-SB-0.0-0.5	2-METHYLNAPHTHALENE	J	1.2	1.7	PQL	ug/Kg	J (all detects)
	BENZO(A)ANTHRACENE	J	0.79	1.7	PQL	ug/Kg	
	BENZO(A)PYRENE	J	0.94	1.7	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	0.82	1.7	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	8.8	18	PQL	ug/Kg	
	FLUORENE	J	0.87	1.7	PQL	ug/Kg	
	INDENO(1,2,3-CD)PYRENE	J	0.71	1.7	PQL	ug/Kg	
NAPHTHALENE	J	1.3	1.7	PQL	ug/Kg		
SL-562-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	0.70	1.7	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	0.80	1.7	PQL	ug/Kg	
SL-563-SA8-SB-0.0-0.5	CHRYSENE	J	0.81	1.7	PQL	ug/Kg	J (all detects)
	FLUORANTHENE	J	0.76	1.7	PQL	ug/Kg	
SL-563-SA8-SB-2.0-3.0	BENZO(K)FLUORANTHENE	J	0.88	1.8	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	0.37	1.8	PQL	ug/Kg	

LDC #: 30210B4

VALIDATION COMPLETENESS WORKSHEET

Date: 8/29/13

SDG #: PH063

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: a

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	-	Sampling dates: 7/9/13
II.	ICP/MS Tune	-	
III.	Calibration	-	
IV.	Blanks	SW	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	SW	MS/D (from PH062)
VII.	Duplicate Sample Analysis	SW	Dup ↓
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	A	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	-	
XV.	Field Blanks	SW	EB=EB2-071013 (PH064) FB=FB01113 (PH029)

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: 501

1	SL-561-SA8-SB-0.0-0.5	11	21	31
2	SL-561-SA8-SB-4.0-5.0	12	22	32
3	SL-562-SA8-SB-0.0-0.5	13	23	33
4	SL-563-SA8-SB-0.0-0.5	14	24	34
5	SL-563-SA8-SB-2.0-3.0	15	25	35
6		16	26	36
7		17	27	37
8		18	28	38
9		19	29	39
10		20	30	40

Notes: _____

VALIDATION FINDINGS WORKSHEET
PB/ICB/CCB QUALIFIED SAMPLES

METHOD: Trace metals (EPA SW 864 Method 6010B/6020/7000)

Soil preparation factor applied: 100x

Reason: B

Sample Concentration units, unless otherwise noted: mg/Kg

Associated Samples: All

Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/L)	Maximum ICB/CCB ^a (ug/L)	Action Level	Sample Identification											
					1	2	3	4	5							
Mo			3.9	1.95	0.51	0.28	0.47	0.39	0.36							

Samples with analyte concentrations within five times the associated ICB, CCB or PB concentration are listed above with the identifications from the Validation Completeness Worksheet. These sample results were qualified as not detected, "U".

Note : a - The listed analyte concentration is the highest ICB, CCB, or PB detected in the analysis of each element.

VALIDATION FINDINGS WORKSHEET Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification									
	FB-041113 (SDG: PH029)	Action Limit	1	2	3	4	5				
Cu	0.0036	1.8									
Mo	0.0036	1.8	0.51	0.28	0.47	0.39	0.36				

Sampling date: 7/10/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification									
	EB2-071013 (SDG: PH064)	Action Limit	No Qualifiers								
Cu	0.0030	1.5									

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

**Data Validation Report
Santa Susan Field Laboratory
Subarea 8**

SDG(s): PH064

Prepared for

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Prepared by

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September 10, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on July 10, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory, RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Volatile Organic Compounds (VOCs) by Environmental Protection Agency (EPA) SW 846 Method 8260B

Semivolatiles by EPA SW 846 Method 8270D-SIM

Pesticides by EPA SW 846 Method 8081A

Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A

Metals by EPA SW 846 Method 6010C, 6020A, 7470A and 7471B

Herbicides by EPA SW 846 8151A

Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M

Total Petroleum Hydrocarbons (TPH) as Extractables by EPA SW 846 Method 8015M

Dioxins/Dibenzofurans by EPA Method 1613B

Perchlorate by EPA Method 6850

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibrations blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, field blanks and equipment blanks. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the calibration blanks and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met with the exception of one sample for herbicides. The associated sample results were qualified as estimated. The details regarding the qualification of data are provided in Enclosure I.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of method blanks for VOCs, semivolatiles, herbicides, metals, and dioxins. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks with the following exceptions:

SDG/ Method	Method Blank ID	Analyte	Maximum Concentration	Associated Samples
PH064/ 6020A	ICB/CCB	Copper	2.9 ug/L	EB1-071013 EB2-071013

Sample concentrations were compared to concentrations detected in the initial and continuing blanks. The sample concentrations were not detected or were significantly greater than the concentrations found in the associated blanks with the following exceptions:

SDG/Method	Sample	Compound	Reported Concentration	Modified Final Concentration
PH064/ 6020A	EB1-071013	Copper	3.1 ug/L	3.1 U ug/L
PH064/ 6020A	EB2-071013	Copper	3.0 ug/L	3.0U ug/L

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of several LCS/LCSD pairs for VOAs, semivolatiles, and herbicides. The associated sample results were qualified as detected estimated (J) as applicable. The details regarding the qualification of data are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH064	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

No field duplicates were identified in this SDG.

XIV. Field Blank Samples

One trip blank was collected and analyzed for VOCs and TPH as gasoline. No contaminants were found in the trip blank.

Two equipment blanks were collected and analyzed for VOCs, semivolatiles, pesticides, PCBs, metals, herbicides, perchlorate, TPH as gasoline, TPH as extractables and dioxins. No contaminants were found in the equipment blanks with the exception of several VOAs, herbicides, dioxins, semivolatiles, and metals. The associated samples were qualified as non-detected (U) due to equipment blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The equipment blank outlier reports are presented in Enclosure I.

One field blank (from SDG PH029) was collected and analyzed for VOCs, semivolatiles, pesticides, PCBs, metals, herbicides, perchlorate, TPH as gasoline, TPH as extractables and dioxins. No contaminants were found in the field blank with the exception of several VOAs, dioxins, semivolatiles, and metals. Associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
10-Jul-2013	TB-071013	7123477	TB	5030B	8015M	III
10-Jul-2013	TB-071013	7123477	TB	5030B	8260B	III
10-Jul-2013	SL-573-SA8-SB-0.0-0.5	7123474	N	3050B	6010C	III
10-Jul-2013	SL-573-SA8-SB-0.0-0.5	7123474	N	3050B	6020A	III
10-Jul-2013	SL-573-SA8-SB-0.0-0.5	7123474	N	3546	8015M	III
10-Jul-2013	SL-573-SA8-SB-0.0-0.5	7123474	N	3546	8082A	III
10-Jul-2013	SL-573-SA8-SB-0.0-0.5	7123474	N	3546	8270D SIM	III
10-Jul-2013	SL-573-SA8-SB-0.0-0.5	7123474	N	METHOD	1613B	III
10-Jul-2013	SL-573-SA8-SB-0.0-0.5	7123474	N	METHOD	7471B	III
10-Jul-2013	SL-573-SA8-SB-4.0-5.0	7123475	N	3050B	6010C	III
10-Jul-2013	SL-573-SA8-SB-4.0-5.0	7123475	N	3050B	6020A	III
10-Jul-2013	SL-573-SA8-SB-4.0-5.0	7123475	N	3546	8015M	III
10-Jul-2013	SL-573-SA8-SB-4.0-5.0	7123475	N	3546	8082A	III
10-Jul-2013	SL-573-SA8-SB-4.0-5.0	7123475	N	3546	8270D SIM	III
10-Jul-2013	SL-573-SA8-SB-4.0-5.0	7123475	N	5035A	8015M	III
10-Jul-2013	SL-573-SA8-SB-4.0-5.0	7123475	N	METHOD	1613B	III
10-Jul-2013	SL-573-SA8-SB-4.0-5.0	7123475	N	METHOD	7471B	III
10-Jul-2013	SL-573-SA8-SB-9.5-10.5	7123476	N	3050B	6010C	III
10-Jul-2013	SL-573-SA8-SB-9.5-10.5	7123476	N	3050B	6020A	III
10-Jul-2013	SL-573-SA8-SB-9.5-10.5	7123476	N	3546	8015M	III
10-Jul-2013	SL-573-SA8-SB-9.5-10.5	7123476	N	3546	8082A	III
10-Jul-2013	SL-573-SA8-SB-9.5-10.5	7123476	N	3546	8270D SIM	III
10-Jul-2013	SL-573-SA8-SB-9.5-10.5	7123476	N	5035A	8015M	III
10-Jul-2013	SL-573-SA8-SB-9.5-10.5	7123476	N	5035A	8260B	III
10-Jul-2013	SL-573-SA8-SB-9.5-10.5	7123476	N	METHOD	7471B	III
10-Jul-2013	SL-572-SA8-SB-0.0-0.5	7123469	N	3050B	6010C	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
10-Jul-2013	SL-572-SA8-SB-0.0-0.5	7123469	N	3050B	6020A	III
10-Jul-2013	SL-572-SA8-SB-0.0-0.5	7123469	N	3546	8015M	III
10-Jul-2013	SL-572-SA8-SB-0.0-0.5	7123469	N	3546	8082A	III
10-Jul-2013	SL-572-SA8-SB-0.0-0.5	7123469	N	3546	8270D SIM	III
10-Jul-2013	SL-572-SA8-SB-0.0-0.5	7123469	N	METHOD	1613B	III
10-Jul-2013	SL-572-SA8-SB-0.0-0.5	7123469	N	METHOD	7471B	III
10-Jul-2013	SL-572-SA8-SB-0.0-0.5DUP	P123469D220949	DUP	METHOD	7471B	III
10-Jul-2013	SL-572-SA8-SB-0.0-0.5MSD	P123469M220953	MSD	METHOD	7471B	III
10-Jul-2013	SL-572-SA8-SB-0.0-0.5MS	P123469R220951	MS	METHOD	7471B	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0	7123470	N	3050B	6010C	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0	7123470	N	3050B	6020A	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0	7123470	N	3546	8015M	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0	7123470	N	3546	8082A	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0	7123470	N	3546	8270D SIM	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0	7123470	N	5035A	8015M	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0	7123470	N	METHOD	1613B	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0	7123470	N	METHOD	7471B	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0DUP	P123470D220843	DUP	3050B	6010C	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0DUP	P123470D221910A	DUP	3050B	6020A	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0DUP	P123470D221910B	DUP	3050B	6020A	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0MSD	P123470M220851	MSD	3050B	6010C	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0MSD	P123470M221915A	MSD	3050B	6020A	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0MSD	P123470M221915B	MSD	3050B	6020A	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0MS	P123470R220847	MS	3050B	6010C	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0MS	P123470R221912A	MS	3050B	6020A	III
10-Jul-2013	SL-572-SA8-SB-4.0-5.0MS	P123470R221912B	MS	3050B	6020A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
10-Jul-2013	SL-572-SA8-SB-9.0-10.0	7123471	N	3050B	6010C	III
10-Jul-2013	SL-572-SA8-SB-9.0-10.0	7123471	N	3050B	6020A	III
10-Jul-2013	SL-572-SA8-SB-9.0-10.0	7123471	N	3546	8015M	III
10-Jul-2013	SL-572-SA8-SB-9.0-10.0	7123471	N	3546	8082A	III
10-Jul-2013	SL-572-SA8-SB-9.0-10.0	7123471	N	3546	8270D SIM	III
10-Jul-2013	SL-572-SA8-SB-9.0-10.0	7123471	N	5035A	8015M	III
10-Jul-2013	SL-572-SA8-SB-9.0-10.0	7123471	N	METHOD	1613B	III
10-Jul-2013	SL-572-SA8-SB-9.0-10.0	7123471	N	METHOD	7471B	III
10-Jul-2013	SL-572-SA8-SB-14.0-15.0	7123472	N	3050B	6010C	III
10-Jul-2013	SL-572-SA8-SB-14.0-15.0	7123472	N	3050B	6020A	III
10-Jul-2013	SL-572-SA8-SB-14.0-15.0	7123472	N	3546	8015M	III
10-Jul-2013	SL-572-SA8-SB-14.0-15.0	7123472	N	3546	8082A	III
10-Jul-2013	SL-572-SA8-SB-14.0-15.0	7123472	N	3546	8270D SIM	III
10-Jul-2013	SL-572-SA8-SB-14.0-15.0	7123472	N	5035A	8015M	III
10-Jul-2013	SL-572-SA8-SB-14.0-15.0	7123472	N	METHOD	7471B	III
10-Jul-2013	SL-572-SA8-SB-18.5-19.5	7123473	N	3050B	6010C	III
10-Jul-2013	SL-572-SA8-SB-18.5-19.5	7123473	N	3050B	6020A	III
10-Jul-2013	SL-572-SA8-SB-18.5-19.5	7123473	N	3546	8015M	III
10-Jul-2013	SL-572-SA8-SB-18.5-19.5	7123473	N	3546	8082A	III
10-Jul-2013	SL-572-SA8-SB-18.5-19.5	7123473	N	3546	8270D SIM	III
10-Jul-2013	SL-572-SA8-SB-18.5-19.5	7123473	N	5035A	8015M	III
10-Jul-2013	SL-572-SA8-SB-18.5-19.5	7123473	N	5035A	8260B	III
10-Jul-2013	SL-572-SA8-SB-18.5-19.5	7123473	N	METHOD	7471B	III
10-Jul-2013	SL-571-SA8-SB-0.0-0.5	7123478	N	3050B	6010C	III
10-Jul-2013	SL-571-SA8-SB-0.0-0.5	7123478	N	3050B	6020A	III
10-Jul-2013	SL-571-SA8-SB-0.0-0.5	7123478	N	3546	8015M	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
10-Jul-2013	SL-571-SA8-SB-0.0-0.5	7123478	N	3546	8082A	III
10-Jul-2013	SL-571-SA8-SB-0.0-0.5	7123478	N	3546	8270D SIM	III
10-Jul-2013	SL-571-SA8-SB-0.0-0.5	7123478	N	METHOD	1613B	III
10-Jul-2013	SL-571-SA8-SB-0.0-0.5	7123478	N	METHOD	7471B	III
10-Jul-2013	SL-571-SA8-SB-6.5-7.5	7123479	N	3050B	6010C	III
10-Jul-2013	SL-571-SA8-SB-6.5-7.5	7123479	N	3050B	6020A	III
10-Jul-2013	SL-571-SA8-SB-6.5-7.5	7123479	N	3546	8015M	III
10-Jul-2013	SL-571-SA8-SB-6.5-7.5	7123479	N	3546	8082A	III
10-Jul-2013	SL-571-SA8-SB-6.5-7.5	7123479	N	3546	8270D SIM	III
10-Jul-2013	SL-571-SA8-SB-6.5-7.5	7123479	N	5035A	8015M	III
10-Jul-2013	SL-571-SA8-SB-6.5-7.5	7123479	N	METHOD	1613B	III
10-Jul-2013	SL-571-SA8-SB-6.5-7.5	7123479	N	METHOD	7471B	III
10-Jul-2013	EB1-071013	7123467	EB	3005A	6010C	III
10-Jul-2013	EB1-071013	7123467	EB	3510C	8015M	III
10-Jul-2013	EB1-071013	7123467	EB	3510C	8081B	III
10-Jul-2013	EB1-071013	7123467	EB	3510C	8082A	III
10-Jul-2013	EB1-071013	7123467	EB	3510C	8270D SIM	III
10-Jul-2013	EB1-071013	7123467	EB	5030B	8015M	III
10-Jul-2013	EB1-071013	7123467	EB	5030B	8260B	III
10-Jul-2013	EB1-071013	7123467	EB	Gen Prep	6850	III
10-Jul-2013	EB1-071013	7123467	EB	M3010A	6020A	III
10-Jul-2013	EB1-071013	7123467	EB	METHOD	1613B	III
10-Jul-2013	EB1-071013	7123467	EB	METHOD	7470A	III
10-Jul-2013	EB1-071013	7123467	EB	METHOD	8151A	III
10-Jul-2013	EB1-071013MSD	P123467M241827A	MSD	Gen Prep	6850	III
10-Jul-2013	EB1-071013MS	P123467R241814A	MS	Gen Prep	6850	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
10-Jul-2013	SL-571-SA8-SB-12.5-13.5	7123480	N	3050B	6010C	III
10-Jul-2013	SL-571-SA8-SB-12.5-13.5	7123480	N	3050B	6020A	III
10-Jul-2013	SL-571-SA8-SB-12.5-13.5	7123480	N	3546	8015M	III
10-Jul-2013	SL-571-SA8-SB-12.5-13.5	7123480	N	3546	8082A	III
10-Jul-2013	SL-571-SA8-SB-12.5-13.5	7123480	N	3546	8270D SIM	III
10-Jul-2013	SL-571-SA8-SB-12.5-13.5	7123480	N	5035A	8015M	III
10-Jul-2013	SL-571-SA8-SB-12.5-13.5	7123480	N	METHOD	1613B	III
10-Jul-2013	SL-571-SA8-SB-12.5-13.5	7123480	N	METHOD	7471B	III
10-Jul-2013	EB2-071013	7123468	EB	3005A	6010C	III
10-Jul-2013	EB2-071013	7123468	EB	3510C	8015M	III
10-Jul-2013	EB2-071013	7123468	EB	3510C	8081B	III
10-Jul-2013	EB2-071013	7123468	EB	3510C	8082A	III
10-Jul-2013	EB2-071013	7123468	EB	3510C	8270D SIM	III
10-Jul-2013	EB2-071013	7123468	EB	5030B	8015M	III
10-Jul-2013	EB2-071013	7123468	EB	5030B	8260B	III
10-Jul-2013	EB2-071013	7123468	EB	Gen Prep	6850	III
10-Jul-2013	EB2-071013	7123468	EB	M3010A	6020A	III
10-Jul-2013	EB2-071013	7123468	EB	METHOD	1613B	III
10-Jul-2013	EB2-071013	7123468	EB	METHOD	7470A	III
10-Jul-2013	EB2-071013	7123468	EB	METHOD	8151A	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: AQ

Sample ID: EB1-071013	Collected: 7/10/2013 3:00:00	Analysis Type: REA2	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
COPPER	0.0031	J	0.0027	MDL	0.0200	PQL	mg/L	U	B, B
MOLYBDENUM	0.0035	J	0.0017	MDL	0.0200	PQL	mg/L	J	Z

Sample ID: EB2-071013	Collected: 7/10/2013 3:30:00	Analysis Type: REA2	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
COPPER	0.0030	J	0.0027	MDL	0.0200	PQL	mg/L	U	B, B

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-571-SA8-SB-0.0-0.5	Collected: 7/10/2013 2:15:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.49	U	0.831	MDL	4.49	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.524	J	0.0752	MDL	1.12	PQL	mg/Kg	J	Z
BORON	5.51	J	0.943	MDL	11.2	PQL	mg/Kg	J	Z
CADMIUM	0.407	J	0.0853	MDL	1.12	PQL	mg/Kg	J	Z
MOLYBDENUM	0.611	J	0.191	MDL	2.25	PQL	mg/Kg	U	F, F
PHOSPHORUS	361		3.24	MDL	11.2	PQL	mg/Kg	J	Q
POTASSIUM	2800		9.36	MDL	112	PQL	mg/Kg	J	Q
TIN	2.79	J	0.247	MDL	11.2	PQL	mg/Kg	U	B

Sample ID: SL-571-SA8-SB-12.5-13.5	Collected: 7/10/2013 3:05:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.20	U	0.778	MDL	4.20	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.525	J	0.0704	MDL	1.05	PQL	mg/Kg	J	Z
BORON	3.81	J	0.883	MDL	10.5	PQL	mg/Kg	J	Z
CADMIUM	0.314	J	0.0799	MDL	1.05	PQL	mg/Kg	J	Z
MOLYBDENUM	0.200	J	0.179	MDL	2.10	PQL	mg/Kg	U	F, F
PHOSPHORUS	447		3.04	MDL	10.5	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-571-SA8-SB-12.5-13.5 Collected: 7/10/2013 3:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
POTASSIUM	3560		8.76	MDL	105	PQL	mg/Kg	J	Q
TIN	2.93	J	0.231	MDL	10.5	PQL	mg/Kg	U	B
Zirconium	4.13	J	0.883	MDL	5.25	PQL	mg/Kg	J	Z

Sample ID: SL-571-SA8-SB-6.5-7.5 Collected: 7/10/2013 2:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.31	U	0.797	MDL	4.31	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.590	J	0.0722	MDL	1.08	PQL	mg/Kg	J	Z
BORON	4.64	J	0.905	MDL	10.8	PQL	mg/Kg	J	Z
CADMIUM	0.288	J	0.0819	MDL	1.08	PQL	mg/Kg	J	Z
MOLYBDENUM	0.538	J	0.183	MDL	2.16	PQL	mg/Kg	U	F, F
PHOSPHORUS	383		3.11	MDL	10.8	PQL	mg/Kg	J	Q
POTASSIUM	3240		8.99	MDL	108	PQL	mg/Kg	J	Q
TIN	2.96	J	0.237	MDL	10.8	PQL	mg/Kg	U	B
Zirconium	4.77	J	0.905	MDL	5.39	PQL	mg/Kg	J	Z

Sample ID: SL-572-SA8-SB-0.0-0.5 Collected: 7/10/2013 10:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	0.944	J	0.863	MDL	4.66	PQL	mg/Kg	J	Z, Q
ARSENIC	4.37	J	0.816	MDL	4.66	PQL	mg/Kg	J	Z
BERYLLIUM	1.03	J	0.0781	MDL	1.17	PQL	mg/Kg	J	Z
BORON	9.52	J	0.979	MDL	11.7	PQL	mg/Kg	J	Z
CADMIUM	0.422	J	0.0886	MDL	1.17	PQL	mg/Kg	J	Z
MOLYBDENUM	0.516	J	0.198	MDL	2.33	PQL	mg/Kg	U	F, F
PHOSPHORUS	213		3.37	MDL	11.7	PQL	mg/Kg	J	Q
POTASSIUM	4060		9.72	MDL	117	PQL	mg/Kg	J	Q
TIN	3.45	J	0.256	MDL	11.7	PQL	mg/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-572-SA8-SB-14.0-15.0 Collected: 7/10/2013 11:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.37	U	0.808	MDL	4.37	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.643	J	0.0731	MDL	1.09	PQL	mg/Kg	J	Z
BORON	5.60	J	0.917	MDL	10.9	PQL	mg/Kg	J	Z
CADMIUM	0.400	J	0.0830	MDL	1.09	PQL	mg/Kg	J	Z
MOLYBDENUM	0.224	J	0.186	MDL	2.18	PQL	mg/Kg	U	F, F
PHOSPHORUS	401		3.15	MDL	10.9	PQL	mg/Kg	J	Q
POTASSIUM	2980		9.10	MDL	109	PQL	mg/Kg	J	Q
TIN	3.17	J	0.240	MDL	10.9	PQL	mg/Kg	U	B
Zirconium	5.43	J	0.917	MDL	5.46	PQL	mg/Kg	J	Z

Sample ID: SL-572-SA8-SB-18.5-19.5 Collected: 7/10/2013 12:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.38	U	0.811	MDL	4.38	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.584	J	0.0734	MDL	1.10	PQL	mg/Kg	J	Z
BORON	5.18	J	0.920	MDL	11.0	PQL	mg/Kg	J	Z
CADMIUM	0.289	J	0.0833	MDL	1.10	PQL	mg/Kg	J	Z
PHOSPHORUS	387		3.17	MDL	11.0	PQL	mg/Kg	J	Q
POTASSIUM	3020		9.14	MDL	110	PQL	mg/Kg	J	Q
TIN	3.07	J	0.241	MDL	11.0	PQL	mg/Kg	U	B
Zirconium	4.53	J	0.920	MDL	5.48	PQL	mg/Kg	J	Z

Sample ID: SL-572-SA8-SB-4.0-5.0 Collected: 7/10/2013 11:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.41	U	0.815	MDL	4.41	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.808	J	0.0738	MDL	1.10	PQL	mg/Kg	J	Z
BORON	7.40	J	0.926	MDL	11.0	PQL	mg/Kg	J	Z
CADMIUM	0.358	J	0.0838	MDL	1.10	PQL	mg/Kg	J	Z
MOLYBDENUM	0.346	J	0.187	MDL	2.20	PQL	mg/Kg	U	F, F
PHOSPHORUS	330		3.18	MDL	11.0	PQL	mg/Kg	J	Q
POTASSIUM	3660		9.19	MDL	110	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-572-SA8-SB-4.0-5.0 Collected: 7/10/2013 11:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
TIN	3.29	J	0.242	MDL	11.0	PQL	mg/Kg	U	B

Sample ID: SL-572-SA8-SB-9.0-10.0 Collected: 7/10/2013 11:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.35	U	0.804	MDL	4.35	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.573	J	0.0728	MDL	1.09	PQL	mg/Kg	J	Z
BORON	5.77	J	0.913	MDL	10.9	PQL	mg/Kg	J	Z
CADMIUM	0.408	J	0.0826	MDL	1.09	PQL	mg/Kg	J	Z
MOLYBDENUM	0.211	J	0.185	MDL	2.17	PQL	mg/Kg	U	F, F
PHOSPHORUS	356		3.14	MDL	10.9	PQL	mg/Kg	J	Q
POTASSIUM	2860		9.06	MDL	109	PQL	mg/Kg	J	Q
TIN	2.92	J	0.239	MDL	10.9	PQL	mg/Kg	U	B
Zirconium	5.18	J	0.913	MDL	5.43	PQL	mg/Kg	J	Z

Sample ID: SL-573-SA8-SB-0.0-0.5 Collected: 7/10/2013 9:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	1.27	J	0.847	MDL	4.58	PQL	mg/Kg	J	Z, Q
ARSENIC	2.91	J	0.801	MDL	4.58	PQL	mg/Kg	J	Z
BERYLLIUM	0.583	J	0.0767	MDL	1.14	PQL	mg/Kg	J	Z
BORON	5.75	J	0.962	MDL	11.4	PQL	mg/Kg	J	Z
CADMIUM	0.688	J	0.0870	MDL	1.14	PQL	mg/Kg	J	Z
MOLYBDENUM	0.723	J	0.195	MDL	2.29	PQL	mg/Kg	U	F, F
PHOSPHORUS	767		3.31	MDL	11.4	PQL	mg/Kg	J	Q
POTASSIUM	3270		9.55	MDL	114	PQL	mg/Kg	J	Q
TIN	2.99	J	0.252	MDL	11.4	PQL	mg/Kg	U	B

Sample ID: SL-573-SA8-SB-4.0-5.0 Collected: 7/10/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.39	U	0.812	MDL	4.39	PQL	mg/Kg	UJ	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-573-SA8-SB-4.0-5.0 Collected: 7/10/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.751	J	0.0735	MDL	1.10	PQL	mg/Kg	J	Z
BORON	6.38	J	0.922	MDL	11.0	PQL	mg/Kg	J	Z
CADMIUM	0.368	J	0.0834	MDL	1.10	PQL	mg/Kg	J	Z
MOLYBDENUM	0.285	J	0.186	MDL	2.19	PQL	mg/Kg	U	F, F
PHOSPHORUS	238		3.17	MDL	11.0	PQL	mg/Kg	J	Q
POTASSIUM	3020		9.15	MDL	110	PQL	mg/Kg	J	Q
TIN	3.08	J	0.241	MDL	11.0	PQL	mg/Kg	U	B

Sample ID: SL-573-SA8-SB-9.5-10.5 Collected: 7/10/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.14	U	0.765	MDL	4.14	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.422	J	0.0693	MDL	1.03	PQL	mg/Kg	J	Z
BORON	3.25	J	0.869	MDL	10.3	PQL	mg/Kg	J	Z
CADMIUM	0.256	J	0.0786	MDL	1.03	PQL	mg/Kg	J	Z
MOLYBDENUM	0.194	J	0.176	MDL	2.07	PQL	mg/Kg	U	F, F
PHOSPHORUS	332		2.99	MDL	10.3	PQL	mg/Kg	J	Q
POTASSIUM	2700		8.62	MDL	103	PQL	mg/Kg	J	Q
TIN	2.70	J	0.227	MDL	10.3	PQL	mg/Kg	U	B
Zirconium	3.15	J	0.869	MDL	5.17	PQL	mg/Kg	J	Z

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-571-SA8-SB-0.0-0.5 Collected: 7/10/2013 2:15:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0750	J	0.0292	MDL	0.225	PQL	mg/Kg	J	Z
STRONTIUM	62.8		0.0763	MDL	0.449	PQL	mg/Kg	J	E
THALLIUM	0.206	J	0.0337	MDL	0.225	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

<i>Sample ID:</i> SL-571-SA8-SB-12.5-13.5		<i>Collected:</i> 7/10/2013 3:05:00		<i>Analysis Type:</i> RES		<i>Dilution:</i> 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	31.4		0.0715	MDL	0.420	PQL	mg/Kg	J	E

<i>Sample ID:</i> SL-571-SA8-SB-6.5-7.5		<i>Collected:</i> 7/10/2013 2:50:00		<i>Analysis Type:</i> RES		<i>Dilution:</i> 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	26.3		0.0733	MDL	0.431	PQL	mg/Kg	J	E

<i>Sample ID:</i> SL-572-SA8-SB-0.0-0.5		<i>Collected:</i> 7/10/2013 10:45:00		<i>Analysis Type:</i> RES		<i>Dilution:</i> 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0410	J	0.0303	MDL	0.233	PQL	mg/Kg	J	Z
STRONTIUM	47.1		0.0793	MDL	0.466	PQL	mg/Kg	J	E

<i>Sample ID:</i> SL-572-SA8-SB-14.0-15.0		<i>Collected:</i> 7/10/2013 11:45:00		<i>Analysis Type:</i> RES		<i>Dilution:</i> 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0331	J	0.0284	MDL	0.218	PQL	mg/Kg	J	Z
STRONTIUM	34.8		0.0742	MDL	0.437	PQL	mg/Kg	J	E

<i>Sample ID:</i> SL-572-SA8-SB-18.5-19.5		<i>Collected:</i> 7/10/2013 12:00:00		<i>Analysis Type:</i> RES		<i>Dilution:</i> 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0509	J	0.0285	MDL	0.219	PQL	mg/Kg	J	Z
STRONTIUM	33.2		0.0745	MDL	0.438	PQL	mg/Kg	J	E

<i>Sample ID:</i> SL-572-SA8-SB-4.0-5.0		<i>Collected:</i> 7/10/2013 11:10:00		<i>Analysis Type:</i> RES		<i>Dilution:</i> 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0463	J	0.0287	MDL	0.220	PQL	mg/Kg	J	Z
STRONTIUM	42.4		0.0749	MDL	0.441	PQL	mg/Kg	J	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-572-SA8-SB-9.0-10.0	Collected: 7/10/2013 11:30:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	34.7		0.0739	MDL	0.435	PQL	mg/Kg	J	E

Sample ID: SL-573-SA8-SB-0.0-0.5	Collected: 7/10/2013 9:10:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	67.1		0.0778	MDL	0.458	PQL	mg/Kg	J	E
THALLIUM	0.181	J	0.0343	MDL	0.229	PQL	mg/Kg	J	Z

Sample ID: SL-573-SA8-SB-4.0-5.0	Collected: 7/10/2013 9:30:00	Analysis Type: REA	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.175	J	0.110	MDL	0.439	PQL	mg/Kg	J	Z

Sample ID: SL-573-SA8-SB-4.0-5.0	Collected: 7/10/2013 9:30:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0576	J	0.0285	MDL	0.219	PQL	mg/Kg	J	Z
STRONTIUM	53.3		0.0746	MDL	0.439	PQL	mg/Kg	J	E

Sample ID: SL-573-SA8-SB-9.5-10.5	Collected: 7/10/2013 9:45:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	17.3		0.0703	MDL	0.414	PQL	mg/Kg	J	E

Method Category:	SVOA	
Method:	1613B	Matrix: AQ

Sample ID: EB1-071013	Collected: 7/10/2013 3:00:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.309	JBQ	0.193	MDL	9.64	PQL	pg/L	U	B
1,2,3,4,6,7,8-HPCDF	0.282	JBQ	0.0837	MDL	9.64	PQL	pg/L	U	B
1,2,3,4,7,8,9-HPCDF	0.203	JBQ	0.0997	MDL	9.64	PQL	pg/L	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: AQ

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HxCDD	0.224	JBQ	0.159	MDL	9.64	PQL	pg/L	U	B
1,2,3,4,7,8-HxCDF	0.412	JBQ	0.0847	MDL	9.64	PQL	pg/L	U	B
1,2,3,6,7,8-HxCDD	0.241	JBQ	0.170	MDL	9.64	PQL	pg/L	U	B
1,2,3,6,7,8-HxCDF	0.201	JBQ	0.0855	MDL	9.64	PQL	pg/L	U	B
1,2,3,7,8,9-HxCDD	0.393	JBQ	0.164	MDL	9.64	PQL	pg/L	U	B
1,2,3,7,8,9-HxCDF	0.336	JB	0.0935	MDL	9.64	PQL	pg/L	U	B
1,2,3,7,8-PECDF	0.405	JBQ	0.152	MDL	9.64	PQL	pg/L	U	B
2,3,4,6,7,8-HxCDF	0.235	JBQ	0.0799	MDL	9.64	PQL	pg/L	U	B
OCDD	1.04	JBQ	0.187	MDL	19.3	PQL	pg/L	U	B
OCDF	0.838	JBQ	0.204	MDL	19.3	PQL	pg/L	U	B

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.532	JB	0.200	MDL	10.2	PQL	pg/L	U	B
1,2,3,4,6,7,8-HPCDF	0.248	JBQ	0.0753	MDL	10.2	PQL	pg/L	U	B
1,2,3,4,7,8,9-HPCDF	0.333	JBQ	0.0903	MDL	10.2	PQL	pg/L	U	B
1,2,3,4,7,8-HxCDF	0.344	JB	0.0967	MDL	10.2	PQL	pg/L	U	B
1,2,3,6,7,8-HxCDD	0.375	JBQ	0.181	MDL	10.2	PQL	pg/L	U	B
1,2,3,6,7,8-HxCDF	0.187	JBQ	0.101	MDL	10.2	PQL	pg/L	U	B
1,2,3,7,8,9-HxCDF	0.410	JBQ	0.112	MDL	10.2	PQL	pg/L	U	B
1,2,3,7,8-PECDD	0.268	JQ	0.262	MDL	10.2	PQL	pg/L	J	Z
1,2,3,7,8-PECDF	0.450	JB	0.150	MDL	10.2	PQL	pg/L	U	B
2,3,4,6,7,8-HxCDF	0.206	JBQ	0.0930	MDL	10.2	PQL	pg/L	U	B
2,3,4,7,8-PECDF	0.209	JBQ	0.132	MDL	10.2	PQL	pg/L	U	B
OCDD	0.674	JBQ	0.174	MDL	20.4	PQL	pg/L	U	B
OCDF	0.599	JB	0.240	MDL	20.4	PQL	pg/L	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-571-SA8-SB-0.0-0.5 Collected: 7/10/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	2.93	JB	0.0220	MDL	5.60	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.425	JB	0.0170	MDL	5.60	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0682	JBQ	0.0209	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0645	JBQ	0.0230	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.135	JBQ	0.0163	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.184	JB	0.0246	MDL	5.60	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.108	JB	0.0158	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.141	JBQ	0.0237	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0961	JB	0.0167	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.141	JB	0.0222	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.335	JB	0.0204	MDL	5.60	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0901	JB	0.0147	MDL	5.60	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.101	JBQ	0.0187	MDL	5.60	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.0577	JQ	0.0223	MDL	1.12	PQL	ng/Kg	J	Z
OCDF	0.728	JB	0.0197	MDL	11.2	PQL	ng/Kg	J	Z

Sample ID: SL-571-SA8-SB-12.5-13.5 Collected: 7/10/2013 3:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.616	JB	0.0202	MDL	5.37	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.130	JB	0.00656	MDL	5.37	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0463	JB	0.00984	MDL	5.37	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0378	JB	0.0158	MDL	5.37	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0558	JBQ	0.00929	MDL	5.37	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.0630	JBQ	0.0167	MDL	5.37	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0439	JBQ	0.00907	MDL	5.37	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0685	JBQ	0.0165	MDL	5.37	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0800	JBQ	0.0105	MDL	5.37	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0920	JBQ	0.0128	MDL	5.37	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0367	JBQ	0.00885	MDL	5.37	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0530	JBQ	0.0127	MDL	5.37	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0410	JBQ	0.0210	MDL	1.07	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-571-SA8-SB-12.5-13.5	Collected: 7/10/2013 3:05:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,3,7,8-TCDF	0.0241	JQ	0.0170	MDL	1.07	PQL	ng/Kg	J	Z
OCDD	8.87	JB	0.0183	MDL	10.7	PQL	ng/Kg	J	Z
OCDF	0.367	JB	0.0179	MDL	10.7	PQL	ng/Kg	U	B

Sample ID: SL-571-SA8-SB-6.5-7.5	Collected: 7/10/2013 2:50:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.296	JB	0.0216	MDL	5.32	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.109	JBQ	0.00744	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0437	JB	0.0134	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0624	JB	0.0164	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.131	JB	0.0107	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.0627	JBQ	0.0176	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDF	0.101	JBQ	0.0101	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0682	JB	0.0168	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDF	0.0970	JB	0.0131	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.118	JBQ	0.0214	MDL	5.32	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.234	JB	0.0138	MDL	5.32	PQL	ng/Kg	U	B
2,3,4,6,7,8-HxCDF	0.0596	JB	0.0101	MDL	5.32	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.155	JBQ	0.0133	MDL	5.32	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0441	JBQ	0.0198	MDL	1.06	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.0589	JQ	0.0169	MDL	1.06	PQL	ng/Kg	J	Z
OCDD	3.66	JB	0.0273	MDL	10.6	PQL	ng/Kg	J	Z
OCDF	0.218	JB	0.0336	MDL	10.6	PQL	ng/Kg	U	B

Sample ID: SL-572-SA8-SB-0.0-0.5	Collected: 7/10/2013 10:45:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	2.58	JB	0.0279	MDL	5.82	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.362	JB	0.0108	MDL	5.82	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0525	JBQ	0.0163	MDL	5.82	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0392	JB	0.0247	MDL	5.82	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	1613B	Matrix:	SO
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Sample ID: SL-572-SA8-SB-0.0-0.5 Collected: 7/10/2013 10:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HXCDF	0.0721	JBQ	0.0185	MDL	5.82	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.195	JB	0.0257	MDL	5.82	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0638	JB	0.0170	MDL	5.82	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.169	JB	0.0251	MDL	5.82	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.162	JBQ	0.0209	MDL	5.82	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0684	JBQ	0.0264	MDL	5.82	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.119	JB	0.0256	MDL	5.82	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0548	JBQ	0.0174	MDL	5.82	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0542	JBQ	0.0233	MDL	5.82	PQL	ng/Kg	U	B
OCDF	0.630	JB	0.0291	MDL	11.6	PQL	ng/Kg	J	Z

Sample ID: SL-572-SA8-SB-4.0-5.0 Collected: 7/10/2013 11:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.895	JB	0.0217	MDL	5.53	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.139	JB	0.00988	MDL	5.53	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0284	JB	0.0116	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0307	JBQ	0.0198	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0499	JBQ	0.0117	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0747	JBQ	0.0208	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0394	JBQ	0.0115	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0450	JBQ	0.0202	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0285	JBQ	0.0121	MDL	5.53	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0477	JB	0.0177	MDL	5.53	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0382	JBQ	0.0108	MDL	5.53	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0424	JB	0.0157	MDL	5.53	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0261	JBQ	0.0235	MDL	1.11	PQL	ng/Kg	U	B
OCDD	6.56	JB	0.0242	MDL	11.1	PQL	ng/Kg	J	Z
OCDF	0.204	JB	0.0213	MDL	11.1	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA		
Method:	1613B	Matrix:	SO

Sample ID: SL-572-SA8-SB-9.0-10.0

Collected: 7/10/2013 11:30:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.04	JB	0.0187	MDL	5.34	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.208	JBQ	0.00931	MDL	5.34	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0291	JB	0.0118	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0450	JBQ	0.00998	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0572	JB	0.0177	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0214	JBQ	0.00965	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0400	JBQ	0.0171	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0345	JBQ	0.00998	MDL	5.34	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0612	JBQ	0.0151	MDL	5.34	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0482	JBQ	0.00931	MDL	5.34	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0273	JBQ	0.0134	MDL	5.34	PQL	ng/Kg	U	B
OCDD	8.94	JB	0.0220	MDL	10.7	PQL	ng/Kg	J	Z
OCDF	0.332	JBQ	0.0183	MDL	10.7	PQL	ng/Kg	U	B

Sample ID: SL-573-SA8-SB-0.0-0.5

Collected: 7/10/2013 9:10:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	5.29	JB	0.0320	MDL	5.80	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.492	JBQ	0.0173	MDL	5.80	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0658	JBQ	0.0248	MDL	5.80	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0521	JBQ	0.0158	MDL	5.80	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.258	JBQ	0.0296	MDL	5.80	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0576	JB	0.0156	MDL	5.80	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.234	JB	0.0276	MDL	5.80	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.247	JB	0.0196	MDL	5.80	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0552	JBQ	0.0287	MDL	5.80	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.117	JBQ	0.0185	MDL	5.80	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0778	JB	0.0146	MDL	5.80	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0381	JBQ	0.0169	MDL	5.80	PQL	ng/Kg	U	B
OCDF	1.04	JB	0.0379	MDL	11.6	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-573-SA8-SB-4.0-5.0 Collected: 7/10/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.682	JB	0.0235	MDL	5.51	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.0894	JBQ	0.0108	MDL	5.51	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0197	JBQ	0.0124	MDL	5.51	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0315	JBQ	0.0200	MDL	5.51	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0426	JBQ	0.0111	MDL	5.51	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.0450	JB	0.0210	MDL	5.51	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0394	JBQ	0.0105	MDL	5.51	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0557	JBQ	0.0200	MDL	5.51	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0595	JBQ	0.0112	MDL	5.51	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0392	JBQ	0.0233	MDL	5.51	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0614	JBQ	0.0181	MDL	5.51	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0497	JB	0.0104	MDL	5.51	PQL	ng/Kg	U	B
OCDD	5.37	JB	0.0333	MDL	11.0	PQL	ng/Kg	J	Z
OCDF	0.259	JBQ	0.0221	MDL	11.0	PQL	ng/Kg	U	B

Method Category: SVOA
Method: 8015M **Matrix:** SO

Sample ID: SL-571-SA8-SB-12.5-13.5 Collected: 7/10/2013 3:05:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	5.1	J	2.2	MDL	5.4	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 8151A **Matrix:** AQ

Sample ID: EB2-071013 Collected: 7/10/2013 3:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,4,5-T	0.051	U	0.015	MDL	0.051	PQL	ug/L	UJ	H
2,4,5-TP (Silvex)	0.051	U	0.010	MDL	0.051	PQL	ug/L	UJ	H
2,4-D	0.51	U	0.16	MDL	0.51	PQL	ug/L	UJ	H

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8151A	Matrix: AQ

Sample ID: EB2-071013 Collected: 7/10/2013 3:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,4-DB	1.0	U	0.31	MDL	1.0	PQL	ug/L	UJ	H
DALAPON	1.3	U	0.26	MDL	1.3	PQL	ug/L	UJ	H
DICAMBA	0.31	U	0.082	MDL	0.31	PQL	ug/L	UJ	H
DICHLOROPROP	0.51	U	0.16	MDL	0.51	PQL	ug/L	UJ	H
DINOSEB	0.51	U	0.12	MDL	0.51	PQL	ug/L	UJ	H
MCPA	210	U	51	MDL	210	PQL	ug/L	UJ	H
MCPP	210	U	51	MDL	210	PQL	ug/L	UJ	H

Method Category:	SVOA	
Method:	8270D SIM	Matrix: AQ

Sample ID: EB1-071013 Collected: 7/10/2013 3:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BIS(2-ETHYLHEXYL)PHTHALATE	0.43	J	0.052	MDL	1.0	PQL	ug/L	U	B
NAPHTHALENE	0.034	J	0.031	MDL	0.052	PQL	ug/L	J	Z

Sample ID: EB2-071013 Collected: 7/10/2013 3:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	0.032	J	0.030	MDL	0.051	PQL	ug/L	J	Z

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-571-SA8-SB-0.0-0.5 Collected: 7/10/2013 2:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	13	J	7.6	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	13	J	3.8	MDL	19	PQL	ug/Kg	J	Z
PHENANTHRENE	7.8	J	7.6	MDL	19	PQL	ug/Kg	J	Z
PYRENE	8.8	J	7.6	MDL	19	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-571-SA8-SB-12.5-13.5 Collected: 7/10/2013 3:05:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	0.70	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-571-SA8-SB-6.5-7.5 Collected: 7/10/2013 2:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	1.4	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z
2-METHYLNAPHTHALENE	1.3	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(B)FLUORANTHENE	1.1	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	1.1	J	0.37	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	0.93	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z
PHENANTHRENE	1.4	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z
PYRENE	0.85	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-572-SA8-SB-0.0-0.5 Collected: 7/10/2013 10:45:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.80	J	0.79	MDL	2.0	PQL	ug/Kg	J	Z
ANTHRACENE	0.87	J	0.39	MDL	2.0	PQL	ug/Kg	J	Z
BENZO(E)PYRENE	5.2	J	3.9	MDL	20	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	17	J	7.1	MDL	21	PQL	ug/Kg	J	Z
DIBENZO(A,H)ANTHRACENE	1.0	J	0.79	MDL	2.0	PQL	ug/Kg	J	Z
Di-n-octylphthalate	12	J	7.1	MDL	21	PQL	ug/Kg	J	Z

Sample ID: SL-572-SA8-SB-4.0-5.0 Collected: 7/10/2013 11:10:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	16	J	7.5	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	16	J	3.8	MDL	19	PQL	ug/Kg	J	Z
FLUORANTHENE	11	J	7.5	MDL	19	PQL	ug/Kg	J	Z
PHENANTHRENE	14	J	7.5	MDL	19	PQL	ug/Kg	J	Z
PYRENE	13	J	7.5	MDL	19	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-572-SA8-SB-9.0-10.0 Collected: 7/10/2013 11:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTHRACENE	0.39	J	0.37	MDL	1.8	PQL	ug/Kg	J	Z
INDENO(1,2,3-CD)PYRENE	1.3	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-573-SA8-SB-0.0-0.5 Collected: 7/10/2013 9:10:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(A)ANTHRACENE	13	J	7.9	MDL	20	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	16	J	7.9	MDL	20	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	11	J	7.9	MDL	20	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	11	J	7.9	MDL	20	PQL	ug/Kg	J	Z
PHENANTHRENE	19	J	7.9	MDL	20	PQL	ug/Kg	J	Z

Sample ID: SL-573-SA8-SB-4.0-5.0 Collected: 7/10/2013 9:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	16	J	7.5	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	15	J	3.7	MDL	19	PQL	ug/Kg	J	Z
PHENANTHRENE	9.8	J	7.5	MDL	19	PQL	ug/Kg	J	Z
PYRENE	8.8	J	7.5	MDL	19	PQL	ug/Kg	J	Z

Sample ID: SL-573-SA8-SB-9.5-10.5 Collected: 7/10/2013 9:45:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(A)ANTHRACENE	1.0	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	1.0	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	0.92	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	VOA	
Method:	8260B	Matrix: AQ

Sample ID: EB1-071013	Collected: 7/10/2013 3:00:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ACETONE	8	J	6	MDL	20	PQL	ug/L	J	Z

Sample ID: EB2-071013	Collected: 7/10/2013 3:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ACETONE	7	J	6	MDL	20	PQL	ug/L	J	Z

Method Category:	VOA	
Method:	8260B	Matrix: SO

Sample ID: SL-572-SA8-SB-18.5-19.5	Collected: 7/10/2013 12:00:00	Analysis Type: RES	Dilution: 0.93						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ACETONE	9	J	7	MDL	21	PQL	ug/Kg	U	F

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Calibration Blank Contamination
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Equipment Blank Contamination
F	Field Blank Contamination
H	Sampling to Extraction Estimation
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Lower Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH064

QC Outlier Report: HoldingTimes

Lab Reporting Batch ID: PH064
EDD Filename: PH064_v1

Laboratory: LL
eQAPP Name: CDM_SSFL_130808_Lan

Method: 8151A Preparation Method: 3546
Matrix: AQ

<i>Sample ID</i>	<i>Type</i>	<i>Actual</i>	<i>Criteria</i>	<i>Units</i>	<i>Flag</i>
EB2-071013 (RES)	Sampling To Extraction	8.00	7.00	DAYS	J(all detects) UJ(all non-detects)

Method Blank Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2000B372114	7/19/2013 9:14:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF OCDD OCDF	0.373 pg/L 0.413 pg/L 0.325 pg/L 0.368 pg/L 0.527 pg/L 0.364 pg/L 0.476 pg/L 0.327 pg/L 0.586 pg/L 0.465 pg/L 0.238 pg/L 0.511 pg/L 1.18 pg/L 0.838 pg/L	EB1-071013 EB2-071013

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB1-071013(RES)	1,2,3,4,6,7,8-HPCDD	0.309 pg/L	0.309U pg/L
EB1-071013(RES)	1,2,3,4,6,7,8-HPCDF	0.282 pg/L	0.282U pg/L
EB1-071013(RES)	1,2,3,4,7,8,9-HPCDF	0.203 pg/L	0.203U pg/L
EB1-071013(RES)	1,2,3,4,7,8-HxCDD	0.224 pg/L	0.224U pg/L
EB1-071013(RES)	1,2,3,4,7,8-HxCDF	0.412 pg/L	0.412U pg/L
EB1-071013(RES)	1,2,3,6,7,8-HXCDD	0.241 pg/L	0.241U pg/L
EB1-071013(RES)	1,2,3,6,7,8-HXCDF	0.201 pg/L	0.201U pg/L
EB1-071013(RES)	1,2,3,7,8,9-HXCDD	0.393 pg/L	0.393U pg/L
EB1-071013(RES)	1,2,3,7,8,9-HXCDF	0.336 pg/L	0.336U pg/L
EB1-071013(RES)	1,2,3,7,8-PECDF	0.405 pg/L	0.405U pg/L
EB1-071013(RES)	2,3,4,6,7,8-HXCDF	0.235 pg/L	0.235U pg/L
EB1-071013(RES)	OCDD	1.04 pg/L	1.04U pg/L
EB1-071013(RES)	OCDF	0.838 pg/L	0.838U pg/L
EB2-071013(RES)	1,2,3,4,6,7,8-HPCDD	0.532 pg/L	0.532U pg/L
EB2-071013(RES)	1,2,3,4,6,7,8-HPCDF	0.248 pg/L	0.248U pg/L
EB2-071013(RES)	1,2,3,4,7,8,9-HPCDF	0.333 pg/L	0.333U pg/L
EB2-071013(RES)	1,2,3,4,7,8-HxCDF	0.344 pg/L	0.344U pg/L
EB2-071013(RES)	1,2,3,6,7,8-HXCDD	0.375 pg/L	0.375U pg/L
EB2-071013(RES)	1,2,3,6,7,8-HXCDF	0.187 pg/L	0.187U pg/L
EB2-071013(RES)	1,2,3,7,8,9-HXCDF	0.410 pg/L	0.410U pg/L
EB2-071013(RES)	1,2,3,7,8-PECDF	0.450 pg/L	0.450U pg/L
EB2-071013(RES)	2,3,4,6,7,8-HXCDF	0.206 pg/L	0.206U pg/L
EB2-071013(RES)	2,3,4,7,8-PECDF	0.209 pg/L	0.209U pg/L
EB2-071013(RES)	OCDD	0.674 pg/L	0.674U pg/L
EB2-071013(RES)	OCDF	0.599 pg/L	0.599U pg/L

Method Blank Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK1960B371832	7/16/2013 6:32:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDD 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDD 1,2,3,7,8,9-HxCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PCDF 2,3,4,6,7,8-HxCDF 2,3,4,7,8-PCDF 2,3,7,8-TCDD OCDD OCDF	0.0478 ng/Kg 0.0258 ng/Kg 0.0609 ng/Kg 0.0220 ng/Kg 0.0412 ng/Kg 0.0258 ng/Kg 0.0216 ng/Kg 0.0474 ng/Kg 0.0394 ng/Kg 0.0613 ng/Kg 0.0646 ng/Kg 0.0356 ng/Kg 0.0232 ng/Kg 0.0400 ng/Kg 0.263 ng/Kg 0.106 ng/Kg	SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-571-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0682 ng/Kg	0.0682U ng/Kg
SL-571-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0645 ng/Kg	0.0645U ng/Kg
SL-571-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.135 ng/Kg	0.135U ng/Kg
SL-571-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HxCDF	0.108 ng/Kg	0.108U ng/Kg
SL-571-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HxCDD	0.141 ng/Kg	0.141U ng/Kg
SL-571-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HxCDF	0.0961 ng/Kg	0.0961U ng/Kg
SL-571-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.141 ng/Kg	0.141U ng/Kg
SL-571-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HxCDF	0.0901 ng/Kg	0.0901U ng/Kg
SL-571-SA8-SB-0.0-0.5(RES)	2,3,4,7,8-PCDF	0.101 ng/Kg	0.101U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0463 ng/Kg	0.0463U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	1,2,3,4,7,8-HxCDD	0.0378 ng/Kg	0.0378U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	1,2,3,4,7,8-HxCDF	0.0558 ng/Kg	0.0558U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	1,2,3,6,7,8-HxCDD	0.0630 ng/Kg	0.0630U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	1,2,3,6,7,8-HxCDF	0.0439 ng/Kg	0.0439U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	1,2,3,7,8,9-HxCDD	0.0685 ng/Kg	0.0685U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	1,2,3,7,8,9-HxCDF	0.0800 ng/Kg	0.0800U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	1,2,3,7,8-PCDF	0.0920 ng/Kg	0.0920U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	2,3,4,6,7,8-HxCDF	0.0367 ng/Kg	0.0367U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	2,3,4,7,8-PCDF	0.0530 ng/Kg	0.0530U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	2,3,7,8-TCDD	0.0410 ng/Kg	0.0410U ng/Kg
SL-571-SA8-SB-12.5-13.5(RES)	OCDF	0.367 ng/Kg	0.367U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	1,2,3,4,6,7,8-HPCDF	0.109 ng/Kg	0.109U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0437 ng/Kg	0.0437U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	1,2,3,4,7,8-HxCDD	0.0624 ng/Kg	0.0624U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	1,2,3,4,7,8-HxCDF	0.131 ng/Kg	0.131U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	1,2,3,6,7,8-HxCDD	0.0627 ng/Kg	0.0627U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	1,2,3,6,7,8-HxCDF	0.101 ng/Kg	0.101U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	1,2,3,7,8,9-HxCDD	0.0682 ng/Kg	0.0682U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	1,2,3,7,8,9-HxCDF	0.0970 ng/Kg	0.0970U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-571-SA8-SB-6.5-7.5(RES)	1,2,3,7,8-PECDD	0.118 ng/Kg	0.118U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	1,2,3,7,8-PECDF	0.234 ng/Kg	0.234U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	2,3,4,6,7,8-HXCDF	0.0596 ng/Kg	0.0596U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	2,3,7,8-TCDD	0.0441 ng/Kg	0.0441U ng/Kg
SL-571-SA8-SB-6.5-7.5(RES)	OCDF	0.218 ng/Kg	0.218U ng/Kg
SL-572-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0525 ng/Kg	0.0525U ng/Kg
SL-572-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0392 ng/Kg	0.0392U ng/Kg
SL-572-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0721 ng/Kg	0.0721U ng/Kg
SL-572-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0638 ng/Kg	0.0638U ng/Kg
SL-572-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.169 ng/Kg	0.169U ng/Kg
SL-572-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.162 ng/Kg	0.162U ng/Kg
SL-572-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.0684 ng/Kg	0.0684U ng/Kg
SL-572-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.119 ng/Kg	0.119U ng/Kg
SL-572-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0548 ng/Kg	0.0548U ng/Kg
SL-572-SA8-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.0542 ng/Kg	0.0542U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0284 ng/Kg	0.0284U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDD	0.0307 ng/Kg	0.0307U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HXCDF	0.0499 ng/Kg	0.0499U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDD	0.0747 ng/Kg	0.0747U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDF	0.0394 ng/Kg	0.0394U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDD	0.0450 ng/Kg	0.0450U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDF	0.0285 ng/Kg	0.0285U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.0477 ng/Kg	0.0477U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0382 ng/Kg	0.0382U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	2,3,4,7,8-PECDF	0.0424 ng/Kg	0.0424U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	2,3,7,8-TCDD	0.0261 ng/Kg	0.0261U ng/Kg
SL-572-SA8-SB-4.0-5.0(RES)	OCDF	0.204 ng/Kg	0.204U ng/Kg
SL-572-SA8-SB-9.0-10.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0291 ng/Kg	0.0291U ng/Kg
SL-572-SA8-SB-9.0-10.0(RES)	1,2,3,4,7,8-HXCDF	0.0450 ng/Kg	0.0450U ng/Kg
SL-572-SA8-SB-9.0-10.0(RES)	1,2,3,6,7,8-HXCDD	0.0572 ng/Kg	0.0572U ng/Kg
SL-572-SA8-SB-9.0-10.0(RES)	1,2,3,6,7,8-HXCDF	0.0214 ng/Kg	0.0214U ng/Kg
SL-572-SA8-SB-9.0-10.0(RES)	1,2,3,7,8,9-HXCDD	0.0400 ng/Kg	0.0400U ng/Kg
SL-572-SA8-SB-9.0-10.0(RES)	1,2,3,7,8,9-HXCDF	0.0345 ng/Kg	0.0345U ng/Kg
SL-572-SA8-SB-9.0-10.0(RES)	1,2,3,7,8-PECDF	0.0612 ng/Kg	0.0612U ng/Kg
SL-572-SA8-SB-9.0-10.0(RES)	2,3,4,6,7,8-HXCDF	0.0482 ng/Kg	0.0482U ng/Kg
SL-572-SA8-SB-9.0-10.0(RES)	2,3,4,7,8-PECDF	0.0273 ng/Kg	0.0273U ng/Kg
SL-572-SA8-SB-9.0-10.0(RES)	OCDF	0.332 ng/Kg	0.332U ng/Kg
SL-573-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0658 ng/Kg	0.0658U ng/Kg
SL-573-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0521 ng/Kg	0.0521U ng/Kg
SL-573-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0576 ng/Kg	0.0576U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-573-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.234 ng/Kg	0.234U ng/Kg
SL-573-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.0552 ng/Kg	0.0552U ng/Kg
SL-573-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.117 ng/Kg	0.117U ng/Kg
SL-573-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0778 ng/Kg	0.0778U ng/Kg
SL-573-SA8-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.0381 ng/Kg	0.0381U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0894 ng/Kg	0.0894U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0197 ng/Kg	0.0197U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDD	0.0315 ng/Kg	0.0315U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HXCDF	0.0426 ng/Kg	0.0426U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDD	0.0450 ng/Kg	0.0450U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDF	0.0394 ng/Kg	0.0394U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDD	0.0557 ng/Kg	0.0557U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDF	0.0595 ng/Kg	0.0595U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDD	0.0392 ng/Kg	0.0392U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.0614 ng/Kg	0.0614U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0497 ng/Kg	0.0497U ng/Kg
SL-573-SA8-SB-4.0-5.0(RES)	OCDF	0.259 ng/Kg	0.259U ng/Kg

Method: 6010C
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19735AB222028	7/18/2013 8:28:00 PM	COPPER	0.0034 mg/L	EB1-071013 EB2-071013

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB1-071013(REA2)	COPPER	0.0031 mg/L	0.0031U mg/L
EB2-071013(REA2)	COPPER	0.0030 mg/L	0.0030U mg/L

Method Blank Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19237AB222027	7/18/2013 8:27:00 PM	CALCIUM MAGNESIUM TIN ZINC	8.96 mg/Kg 2.47 mg/Kg 1.45 mg/Kg 0.430 mg/Kg	SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-571-SA8-SB-0.0-0.5(RES)	TIN	2.79 mg/Kg	2.79U mg/Kg
SL-571-SA8-SB-12.5-13.5(RES)	TIN	2.93 mg/Kg	2.93U mg/Kg
SL-571-SA8-SB-6.5-7.5(RES)	TIN	2.96 mg/Kg	2.96U mg/Kg
SL-572-SA8-SB-0.0-0.5(RES)	TIN	3.45 mg/Kg	3.45U mg/Kg
SL-572-SA8-SB-14.0-15.0(RES)	TIN	3.17 mg/Kg	3.17U mg/Kg
SL-572-SA8-SB-18.5-19.5(RES)	TIN	3.07 mg/Kg	3.07U mg/Kg
SL-572-SA8-SB-4.0-5.0(RES)	TIN	3.29 mg/Kg	3.29U mg/Kg
SL-572-SA8-SB-9.0-10.0(RES)	TIN	2.92 mg/Kg	2.92U mg/Kg
SL-573-SA8-SB-0.0-0.5(RES)	TIN	2.99 mg/Kg	2.99U mg/Kg
SL-573-SA8-SB-4.0-5.0(RES)	TIN	3.08 mg/Kg	3.08U mg/Kg
SL-573-SA8-SB-9.5-10.5(RES)	TIN	2.70 mg/Kg	2.70U mg/Kg

Method: 8151A
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P99991AB241002A	7/19/2013 10:02:00 AM	2,4,5-T	0.099 ug/L	EB2-071013

Method: 8270D SIM
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
PLKWC19B262100	7/19/2013 9:00:00 PM	BIS(2-ETHYLHEXYL)PHTHALATE	0.31 ug/L	EB1-071013 EB2-071013

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB1-071013(RES)	BIS(2-ETHYLHEXYL)PHTHALATE	0.43 ug/L	1.0U ug/L

Equipment Rinsate Blank Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Equipment Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
EB1-071013(REA2)	7/10/2013 3:00:00 PM	COPPER MOLYBDENUM	0.0031 mg/L 0.0035 mg/L	SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-571-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.611 mg/Kg	0.611U mg/Kg
SL-571-SA8-SB-12.5-13.5(RES)	MOLYBDENUM	0.200 mg/Kg	0.200U mg/Kg
SL-571-SA8-SB-6.5-7.5(RES)	MOLYBDENUM	0.538 mg/Kg	0.538U mg/Kg
SL-572-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.516 mg/Kg	0.516U mg/Kg
SL-572-SA8-SB-14.0-15.0(RES)	MOLYBDENUM	0.224 mg/Kg	0.224U mg/Kg
SL-572-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.346 mg/Kg	0.346U mg/Kg
SL-572-SA8-SB-9.0-10.0(RES)	MOLYBDENUM	0.211 mg/Kg	0.211U mg/Kg
SL-573-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.723 mg/Kg	0.723U mg/Kg
SL-573-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.285 mg/Kg	0.285U mg/Kg
SL-573-SA8-SB-9.5-10.5(RES)	MOLYBDENUM	0.194 mg/Kg	0.194U mg/Kg

Method: 8260B
Matrix: SO

Equipment Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
EB1-071013(RES)	7/10/2013 3:00:00 PM	ACETONE	8 ug/L	SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-572-SA8-SB-18.5-19.5(RES)	ACETONE	9 ug/Kg	21U ug/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PrepPH064

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(REA2)	4/11/2013 3:00:00 PM	COPPER MOLYBDENUM	0.0036 mg/L 0.0036 mg/L	SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-571-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.611 mg/Kg	0.611U mg/Kg
SL-571-SA8-SB-12.5-13.5(RES)	MOLYBDENUM	0.200 mg/Kg	0.200U mg/Kg
SL-571-SA8-SB-6.5-7.5(RES)	MOLYBDENUM	0.538 mg/Kg	0.538U mg/Kg
SL-572-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.516 mg/Kg	0.516U mg/Kg
SL-572-SA8-SB-14.0-15.0(RES)	MOLYBDENUM	0.224 mg/Kg	0.224U mg/Kg
SL-572-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.346 mg/Kg	0.346U mg/Kg
SL-572-SA8-SB-9.0-10.0(RES)	MOLYBDENUM	0.211 mg/Kg	0.211U mg/Kg
SL-573-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.723 mg/Kg	0.723U mg/Kg
SL-573-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.285 mg/Kg	0.285U mg/Kg
SL-573-SA8-SB-9.5-10.5(RES)	MOLYBDENUM	0.194 mg/Kg	0.194U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-572-SA8-SB-4.0-5.0MS (TOT) SL-572-SA8-SB-4.0-5.0MSD (TOT) (SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5)	ALUMINUM IRON MAGNESIUM POTASSIUM TITANIUM	1834 1182 229 135 274	1927 1485 277 148 360	75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00	- - - - -	ALUMINUM IRON MAGNESIUM POTASSIUM TITANIUM	J (all detects) Al, Fe, Mg, Ti No Qual, >4x
SL-572-SA8-SB-4.0-5.0MS (TOT) SL-572-SA8-SB-4.0-5.0MSD (TOT) (SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5)	CALCIUM	-365	-585	75.00-125.00	-	CALCIUM	No Qual, >4x
SL-572-SA8-SB-4.0-5.0MS (TOT) SL-572-SA8-SB-4.0-5.0MSD (TOT) (SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5)	ANTIMONY PHOSPHORUS	60 64	58 -	75.00-125.00 75.00-125.00	- -	ANTIMONY PHOSPHORUS	J(all detects) UJ(all non-detects)

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-572-SA8-SB-4.0-5.0MS (TOT) SL-572-SA8-SB-4.0-5.0MSD (TOT) (SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5)	STRONTIUM	65	197	75.00-125.00	22 (20.00)	STRONTIUM	J(all detects) UJ(all non-detects) Sr, No Qual %R >4x

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-572-SA8-SB-4.0-5.0DUP (TOT) (SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5)	ANTIMONY	200	20.00	No Qual, OK by Difference

Method: 6020A
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-572-SA8-SB-4.0-5.0DUP (TOT) (SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5)	SILVER	200	20.00	No Qual, OK by Difference

Method: 7471B
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-572-SA8-SB-0.0-0.5DUP (TOT) (SL-571-SA8-SB-0.0-0.5 SL-571-SA8-SB-12.5-13.5 SL-571-SA8-SB-6.5-7.5 SL-572-SA8-SB-0.0-0.5 SL-572-SA8-SB-14.0-15.0 SL-572-SA8-SB-18.5-19.5 SL-572-SA8-SB-4.0-5.0 SL-572-SA8-SB-9.0-10.0 SL-573-SA8-SB-0.0-0.5 SL-573-SA8-SB-4.0-5.0 SL-573-SA8-SB-9.5-10.5)	MERCURY	200	20.00	No Qual, OK by Difference

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8151A
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P31991AQ241029A	2,4,5-TP (Silvex)	-	161	58.00-155.00	-	2,4,5-TP (Silvex)	J (all detects)
P31991AY241055A	2,4-D	-	161	68.00-155.00	-	2,4-D	
(EB2-071013)	DICHLOROPROP	182	182	89.00-162.00	-	DICHLOROPROP	

Method: 8270D SIM
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P7WCLCSQ262127 P7WCLCSY262155 (EB1-071013 EB2-071013)	INDENO(1,2,3-CD)PYRENE	140	138	66.00-122.00	-	INDENO(1,2,3-CD)PYRENE	J(all detects)

Method: 8260B
Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
LCSB54Q212231A LCSB55Q212253A LCSB55Y212316A (SL-572-SA8-SB-18.5-19.5 SL-573-SA8-SB-9.5-10.5)	4-METHYL-2-PENTANONE (MIB VINYL ACETATE	129 135	- 127	52.00-125.00 29.00-111.00	- -	4-METHYL-2-PENTANONE (MI VINYL ACETATE	J(all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-071013	1,2,3,4,6,7,8-HPCDD	JBQ	0.309	9.64	PQL	pg/L	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.282	9.64	PQL	pg/L	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.203	9.64	PQL	pg/L	
	1,2,3,4,7,8-HxCDD	JBQ	0.224	9.64	PQL	pg/L	
	1,2,3,4,7,8-HxCDF	JBQ	0.412	9.64	PQL	pg/L	
	1,2,3,6,7,8-HxCDD	JBQ	0.241	9.64	PQL	pg/L	
	1,2,3,6,7,8-HxCDF	JBQ	0.201	9.64	PQL	pg/L	
	1,2,3,7,8,9-HxCDD	JBQ	0.393	9.64	PQL	pg/L	
	1,2,3,7,8,9-HxCDF	JB	0.336	9.64	PQL	pg/L	
	1,2,3,7,8-PECDF	JBQ	0.405	9.64	PQL	pg/L	
	2,3,4,6,7,8-HxCDF	JBQ	0.235	9.64	PQL	pg/L	
	OCDD	JBQ	1.04	19.3	PQL	pg/L	
	OCDF	JBQ	0.838	19.3	PQL	pg/L	
EB2-071013	1,2,3,4,6,7,8-HPCDD	JB	0.532	10.2	PQL	pg/L	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.248	10.2	PQL	pg/L	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.333	10.2	PQL	pg/L	
	1,2,3,4,7,8-HxCDF	JB	0.344	10.2	PQL	pg/L	
	1,2,3,6,7,8-HxCDD	JBQ	0.375	10.2	PQL	pg/L	
	1,2,3,6,7,8-HxCDF	JBQ	0.187	10.2	PQL	pg/L	
	1,2,3,7,8,9-HxCDF	JBQ	0.410	10.2	PQL	pg/L	
	1,2,3,7,8-PECDD	JQ	0.268	10.2	PQL	pg/L	
	1,2,3,7,8-PECDF	JB	0.450	10.2	PQL	pg/L	
	2,3,4,6,7,8-HxCDF	JBQ	0.206	10.2	PQL	pg/L	
	2,3,4,7,8-PECDF	JBQ	0.209	10.2	PQL	pg/L	
	OCDD	JBQ	0.674	20.4	PQL	pg/L	
	OCDF	JB	0.599	20.4	PQL	pg/L	

Method: 6010C
Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-071013	COPPER	J	0.0031	0.0200	PQL	mg/L	J (all detects)
	MOLYBDENUM	J	0.0035	0.0200	PQL	mg/L	
EB2-071013	COPPER	J	0.0030	0.0200	PQL	mg/L	J (all detects)

Method: 8260B
Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-071013	ACETONE	J	8	20	PQL	ug/L	J (all detects)
EB2-071013	ACETONE	J	7	20	PQL	ug/L	J (all detects)

Method: 8270D SIM
Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-071013	BIS(2-ETHYLHEXYL)PHTHALATE	J	0.43	1.0	PQL	ug/L	J (all detects)
	NAPHTHALENE	J	0.034	0.052	PQL	ug/L	

Reporting Limit Outliers

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8270D SIM
Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB2-071013	NAPHTHALENE	J	0.032	0.051	PQL	ug/L	J (all detects)

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-571-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	2.93	5.60	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.425	5.60	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0682	5.60	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0645	5.60	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.135	5.60	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.184	5.60	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.108	5.60	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JBQ	0.141	5.60	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.0961	5.60	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.141	5.60	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.335	5.60	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.0901	5.60	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.101	5.60	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0577	1.12	PQL	ng/Kg	
OCDF	JB	0.728	11.2	PQL	ng/Kg		
SL-571-SA8-SB-12.5-13.5	1,2,3,4,6,7,8-HPCDD	JB	0.616	5.37	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.130	5.37	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0463	5.37	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0378	5.37	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.0558	5.37	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.0630	5.37	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.0439	5.37	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JBQ	0.0685	5.37	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JBQ	0.0800	5.37	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0920	5.37	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.0367	5.37	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0530	5.37	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.0410	1.07	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0241	1.07	PQL	ng/Kg	
OCDD	JB	8.87	10.7	PQL	ng/Kg		
OCDF	JB	0.367	10.7	PQL	ng/Kg		
SL-571-SA8-SB-6.5-7.5	1,2,3,4,6,7,8-HPCDD	JB	0.296	5.32	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.109	5.32	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0437	5.32	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0624	5.32	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.131	5.32	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.0627	5.32	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.101	5.32	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.0682	5.32	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.0970	5.32	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.118	5.32	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.234	5.32	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.0596	5.32	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.155	5.32	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.0441	1.06	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0589	1.06	PQL	ng/Kg	
	OCDD	JB	3.66	10.6	PQL	ng/Kg	
OCDF	JB	0.218	10.6	PQL	ng/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-572-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	2.58	5.82	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.362	5.82	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0525	5.82	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0392	5.82	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.0721	5.82	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.195	5.82	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.0638	5.82	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.169	5.82	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JBQ	0.162	5.82	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.0684	5.82	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.119	5.82	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.0548	5.82	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0542	5.82	PQL	ng/Kg	
	OCDF	JB	0.630	11.6	PQL	ng/Kg	
SL-572-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	0.895	5.53	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.139	5.53	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0284	5.53	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0307	5.53	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.0499	5.53	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.0747	5.53	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.0394	5.53	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JBQ	0.0450	5.53	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JBQ	0.0285	5.53	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.0477	5.53	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.0382	5.53	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.0424	5.53	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.0261	1.11	PQL	ng/Kg	
	OCDD	JB	6.56	11.1	PQL	ng/Kg	
OCDF	JB	0.204	11.1	PQL	ng/Kg		
SL-572-SA8-SB-9.0-10.0	1,2,3,4,6,7,8-HPCDD	JB	1.04	5.34	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.208	5.34	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0291	5.34	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.0450	5.34	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.0572	5.34	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.0214	5.34	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JBQ	0.0400	5.34	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JBQ	0.0345	5.34	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0612	5.34	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.0482	5.34	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0273	5.34	PQL	ng/Kg	
	OCDD	JB	8.94	10.7	PQL	ng/Kg	
	OCDF	JBQ	0.332	10.7	PQL	ng/Kg	
	SL-573-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	5.29	5.80	PQL	
1,2,3,4,6,7,8-HPCDF		JBQ	0.492	5.80	PQL	ng/Kg	
1,2,3,4,7,8,9-HPCDF		JBQ	0.0658	5.80	PQL	ng/Kg	
1,2,3,4,7,8-HxCDF		JBQ	0.0521	5.80	PQL	ng/Kg	
1,2,3,6,7,8-HxCDD		JBQ	0.258	5.80	PQL	ng/Kg	
1,2,3,6,7,8-HxCDF		JB	0.0576	5.80	PQL	ng/Kg	
1,2,3,7,8,9-HxCDD		JB	0.234	5.80	PQL	ng/Kg	
1,2,3,7,8,9-HxCDF		JB	0.247	5.80	PQL	ng/Kg	
1,2,3,7,8-PECDD		JBQ	0.0552	5.80	PQL	ng/Kg	
1,2,3,7,8-PECDF		JBQ	0.117	5.80	PQL	ng/Kg	
2,3,4,6,7,8-HxCDF		JB	0.0778	5.80	PQL	ng/Kg	
2,3,4,7,8-PECDF		JBQ	0.0381	5.80	PQL	ng/Kg	
OCDF		JB	1.04	11.6	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-573-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	0.682	5.51	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0894	5.51	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0197	5.51	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0315	5.51	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.0426	5.51	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.0450	5.51	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.0394	5.51	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JBQ	0.0557	5.51	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JBQ	0.0595	5.51	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.0392	5.51	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0614	5.51	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.0497	5.51	PQL	ng/Kg	
	OCDD	JB	5.37	11.0	PQL	ng/Kg	
	OCDF	JBQ	0.259	11.0	PQL	ng/Kg	

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-571-SA8-SB-0.0-0.5	BERYLLIUM	J	0.524	1.12	PQL	mg/Kg	J (all detects)
	BORON	J	5.51	11.2	PQL	mg/Kg	
	CADMIUM	J	0.407	1.12	PQL	mg/Kg	
	MOLYBDENUM	J	0.611	2.25	PQL	mg/Kg	
	TIN	J	2.79	11.2	PQL	mg/Kg	
SL-571-SA8-SB-12.5-13.5	BERYLLIUM	J	0.525	1.05	PQL	mg/Kg	J (all detects)
	BORON	J	3.81	10.5	PQL	mg/Kg	
	CADMIUM	J	0.314	1.05	PQL	mg/Kg	
	MOLYBDENUM	J	0.200	2.10	PQL	mg/Kg	
	TIN	J	2.93	10.5	PQL	mg/Kg	
	Zirconium	J	4.13	5.25	PQL	mg/Kg	
SL-571-SA8-SB-6.5-7.5	BERYLLIUM	J	0.590	1.08	PQL	mg/Kg	J (all detects)
	BORON	J	4.64	10.8	PQL	mg/Kg	
	CADMIUM	J	0.288	1.08	PQL	mg/Kg	
	MOLYBDENUM	J	0.538	2.16	PQL	mg/Kg	
	TIN	J	2.96	10.8	PQL	mg/Kg	
	Zirconium	J	4.77	5.39	PQL	mg/Kg	
SL-572-SA8-SB-0.0-0.5	ANTIMONY	J	0.944	4.66	PQL	mg/Kg	J (all detects)
	ARSENIC	J	4.37	4.66	PQL	mg/Kg	
	BERYLLIUM	J	1.03	1.17	PQL	mg/Kg	
	BORON	J	9.52	11.7	PQL	mg/Kg	
	CADMIUM	J	0.422	1.17	PQL	mg/Kg	
	MOLYBDENUM	J	0.516	2.33	PQL	mg/Kg	
	TIN	J	3.45	11.7	PQL	mg/Kg	
SL-572-SA8-SB-14.0-15.0	BERYLLIUM	J	0.643	1.09	PQL	mg/Kg	J (all detects)
	BORON	J	5.60	10.9	PQL	mg/Kg	
	CADMIUM	J	0.400	1.09	PQL	mg/Kg	
	MOLYBDENUM	J	0.224	2.18	PQL	mg/Kg	
	TIN	J	3.17	10.9	PQL	mg/Kg	
	Zirconium	J	5.43	5.46	PQL	mg/Kg	
SL-572-SA8-SB-18.5-19.5	BERYLLIUM	J	0.584	1.10	PQL	mg/Kg	J (all detects)
	BORON	J	5.18	11.0	PQL	mg/Kg	
	CADMIUM	J	0.289	1.10	PQL	mg/Kg	
	TIN	J	3.07	11.0	PQL	mg/Kg	
	Zirconium	J	4.53	5.48	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-572-SA8-SB-4.0-5.0	BERYLLIUM	J	0.808	1.10	PQL	mg/Kg	J (all detects)
	BORON	J	7.40	11.0	PQL	mg/Kg	
	CADMIUM	J	0.358	1.10	PQL	mg/Kg	
	MOLYBDENUM	J	0.346	2.20	PQL	mg/Kg	
	TIN	J	3.29	11.0	PQL	mg/Kg	
SL-572-SA8-SB-9.0-10.0	BERYLLIUM	J	0.573	1.09	PQL	mg/Kg	J (all detects)
	BORON	J	5.77	10.9	PQL	mg/Kg	
	CADMIUM	J	0.408	1.09	PQL	mg/Kg	
	MOLYBDENUM	J	0.211	2.17	PQL	mg/Kg	
	TIN	J	2.92	10.9	PQL	mg/Kg	
	Zirconium	J	5.18	5.43	PQL	mg/Kg	
SL-573-SA8-SB-0.0-0.5	ANTIMONY	J	1.27	4.58	PQL	mg/Kg	J (all detects)
	ARSENIC	J	2.91	4.58	PQL	mg/Kg	
	BERYLLIUM	J	0.583	1.14	PQL	mg/Kg	
	BORON	J	5.75	11.4	PQL	mg/Kg	
	CADMIUM	J	0.688	1.14	PQL	mg/Kg	
	MOLYBDENUM	J	0.723	2.29	PQL	mg/Kg	
	TIN	J	2.99	11.4	PQL	mg/Kg	
SL-573-SA8-SB-4.0-5.0	BERYLLIUM	J	0.751	1.10	PQL	mg/Kg	J (all detects)
	BORON	J	6.38	11.0	PQL	mg/Kg	
	CADMIUM	J	0.368	1.10	PQL	mg/Kg	
	MOLYBDENUM	J	0.285	2.19	PQL	mg/Kg	
	TIN	J	3.08	11.0	PQL	mg/Kg	
SL-573-SA8-SB-9.5-10.5	BERYLLIUM	J	0.422	1.03	PQL	mg/Kg	J (all detects)
	BORON	J	3.25	10.3	PQL	mg/Kg	
	CADMIUM	J	0.256	1.03	PQL	mg/Kg	
	MOLYBDENUM	J	0.194	2.07	PQL	mg/Kg	
	TIN	J	2.70	10.3	PQL	mg/Kg	
	Zirconium	J	3.15	5.17	PQL	mg/Kg	

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-571-SA8-SB-0.0-0.5	SILVER	J	0.0750	0.225	PQL	mg/Kg	J (all detects)
	THALLIUM	J	0.206	0.225	PQL	mg/Kg	
SL-572-SA8-SB-0.0-0.5	SILVER	J	0.0410	0.233	PQL	mg/Kg	J (all detects)
SL-572-SA8-SB-14.0-15.0	SILVER	J	0.0331	0.218	PQL	mg/Kg	J (all detects)
SL-572-SA8-SB-18.5-19.5	SILVER	J	0.0509	0.219	PQL	mg/Kg	J (all detects)
SL-572-SA8-SB-4.0-5.0	SILVER	J	0.0463	0.220	PQL	mg/Kg	J (all detects)
SL-573-SA8-SB-0.0-0.5	THALLIUM	J	0.181	0.229	PQL	mg/Kg	J (all detects)
SL-573-SA8-SB-4.0-5.0	SELENIUM	J	0.175	0.439	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0576	0.219	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH064

Laboratory: LL

EDD Filename: PH064_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8015M
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-571-SA8-SB-12.5-13.5	EFH (C21-C30)	J	5.1	5.4	PQL	mg/Kg	J (all detects)

Method: 8260B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-572-SA8-SB-18.5-19.5	ACETONE	J	9	21	PQL	ug/Kg	J (all detects)

Method: 8270D SIM
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-571-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	13	19	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	13	19	PQL	ug/Kg	
	PHENANTHRENE	J	7.8	19	PQL	ug/Kg	
	PYRENE	J	8.8	19	PQL	ug/Kg	
SL-571-SA8-SB-12.5-13.5	CHRYSENE	J	0.70	1.8	PQL	ug/Kg	J (all detects)
SL-571-SA8-SB-6.5-7.5	1-METHYLNAPHTHALENE	J	1.4	1.8	PQL	ug/Kg	J (all detects)
	2-METHYLNAPHTHALENE	J	1.3	1.8	PQL	ug/Kg	
	BENZO(B)FLUORANTHENE	J	1.1	1.8	PQL	ug/Kg	
	CHRYSENE	J	1.1	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	0.93	1.8	PQL	ug/Kg	
	PHENANTHRENE	J	1.4	1.8	PQL	ug/Kg	
SL-572-SA8-SB-0.0-0.5	2-METHYLNAPHTHALENE	J	0.80	2.0	PQL	ug/Kg	J (all detects)
	ANTHRACENE	J	0.87	2.0	PQL	ug/Kg	
	BENZO(E)PYRENE	J	5.2	20	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	17	21	PQL	ug/Kg	
	DIBENZO(A,H)ANTHRACENE	J	1.0	2.0	PQL	ug/Kg	
	Di-n-octylphthalate	J	12	21	PQL	ug/Kg	
SL-572-SA8-SB-4.0-5.0	BENZO(B)FLUORANTHENE	J	16	19	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	16	19	PQL	ug/Kg	
	FLUORANTHENE	J	11	19	PQL	ug/Kg	
	PHENANTHRENE	J	14	19	PQL	ug/Kg	
	PYRENE	J	13	19	PQL	ug/Kg	
SL-572-SA8-SB-9.0-10.0	ANTHRACENE	J	0.39	1.8	PQL	ug/Kg	J (all detects)
	INDENO(1,2,3-CD)PYRENE	J	1.3	1.8	PQL	ug/Kg	
SL-573-SA8-SB-0.0-0.5	BENZO(A)ANTHRACENE	J	13	20	PQL	ug/Kg	J (all detects)
	BENZO(A)PYRENE	J	16	20	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	11	20	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	11	20	PQL	ug/Kg	
	PHENANTHRENE	J	19	20	PQL	ug/Kg	
SL-573-SA8-SB-4.0-5.0	BENZO(B)FLUORANTHENE	J	16	19	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	15	19	PQL	ug/Kg	
	PHENANTHRENE	J	9.8	19	PQL	ug/Kg	
	PYRENE	J	8.8	19	PQL	ug/Kg	
SL-573-SA8-SB-9.5-10.5	BENZO(A)ANTHRACENE	J	1.0	1.8	PQL	ug/Kg	J (all detects)
	BENZO(G,H,I)PERYLENE	J	1.0	1.8	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	0.92	1.8	PQL	ug/Kg	

LDC #: 30210C4

VALIDATION COMPLETENESS WORKSHEET

Date: 8/28/13

SDG #: PH064

ADR

Page: 2 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: [Signature]

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	—	Sampling dates: 7/10/13
II.	ICP/MS Tune	—	
III.	Calibration	—	
IV.	Blanks	SW	
V.	ICP Interference Check Sample (ICS) Analysis	—	
VI.	Matrix Spike Analysis	SW	MS/D
VII.	Duplicate Sample Analysis	SW	Dp
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	A	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	—	
XV.	Field Blanks	SW	EB=1,2 FB=FB-041113

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

(PH029)

Validated Samples:

soil/water

1	EB1-071013	w	11	SL-571-SA8-SB-0.0-0.5	21	31
2	EB2-071013	✓	12	SL-571-SA8-SB-6.5-7.5	22	32
3	SL-572-SA8-SB-0.0-0.5		13	SL-571-SA8-SB-12.5-13.5	23	33
4	SL-572-SA8-SB-4.0-5.0		14	A3 MS (Hg)	24	34
5	SL-572-SA8-SB-9.0-10.0		15	MSD ↓	25	35
6	SL-572-SA8-SB-14.0-15.0		16	Dp ↓	26	36
7	SL-572-SA8-SB-18.5-19.5		17	A4 MS	27	37
8	SL-573-SA8-SB-0.0-0.5		18	Dp	28	38
9	SL-573-SA8-SB-4.0-5.0		19	MS	29	39
10	SL-573-SA8-SB-9.5-10.5		20		30	40

Notes: _____

LDC #: 30210C4

VALIDATION FINDINGS WORKSHEET
PB/ICB/CCB QUALIFIED SAMPLES

Page: 1 of 1

Reviewer: *a*

2nd Reviewer: *h*

METHOD: Trace metals (EPA SW 864 Method 6010B/6020/7000)

Soil preparation factor applied: _____

Reason: B

Sample Concentration units, unless otherwise noted: ug/L

Associated Samples: All Water

					Sample Identification											
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/l)	Maximum ICB/CCB ^a (ug/l)	Action Level	1	2										
Cu			2.9	14.5	3.1	3.0										

Samples with analyte concentrations within five times the associated ICB, CCB or PB concentration are listed above with the identifications from the Validation Completeness Worksheet. These sample results were qualified as not detected, "U".

Note : a - The listed analyte concentration is the highest ICB, CCB, or PB detected in the analysis of each element.

VALIDATION FINDINGS WORKSHEET Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All Soil

Analyte	Blank ID	Sample Identification										
	FB-041113 (SDG: PH029)	Action Limit	3	4	5	6	8	9	10	11	12	13
Cu	0.0036	1.8										
Mo	0.0036	1.8	0.52	0.35	0.21	0.22	0.72	0.29	0.19	0.64	0.21	0.20

Sampling date: 7/10/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All Soil

Analyte	Blank ID	Sample Identification										
	EB1-071013 (SDG: PH064)	Action Limit	See FB									
Cu	0.0031	1.55										
Mo	0.0035	1.75										

Sampling date: 7/10/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: None

Analyte	Blank ID	Sample Identification										
	EB2-071013 (SDG: PH064)	Action Limit	No Qualifiers									
Cu	0.0030	1.5										

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

7/11/06 qual CSR qualified from RPD

Background Lab Sample ID: 7123470BKG Matrix Spike Lab Sample ID: 7123470MS Matrix Spike Duplicate Lab Sample ID: 7123470MSD
Batch Id(s): P19237A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Control Limit			
		Result	C	Result	C	Result	C				%R	Q	%R	Q		%R	RPD	M	
Aluminum		23391.4621		26953.0427		27133.0252		194.1748	194.1748	MG/KG	1834		1927		1			20	P
Antimony		0.7184	U	29.2136		28.3951		48.5437	48.5437	MG/KG	60	N	58	N	3	75 - 125		20	P
Arsenic		5.0757		19.6214		19.6097		14.5631	14.5631	MG/KG	100		100		0	75 - 125		20	P
Barium		76.9359		316.9078		280.6427		194.1748	194.1748	MG/KG	124		105		12	75 - 125		20	P
Beryllium		0.7117	B	5.7485		5.6816		4.8544	4.8544	MG/KG	104		102		1	75 - 125		20	P
Boron		6.5233	B	200.9621		199.1728		194.1748	194.1748	MG/KG	100		99		1	75 - 125		20	P
Cadmium		0.3155	B	4.7612		4.7816		4.8544	4.8544	MG/KG	92		92		0	75 - 125		20	P
Calcium		12994.3019		11577.3097		10721.0272		388.3495	388.3495	MG/KG	-365		-585		8			20	P
Chromium		25.6942		47.2233		46.6951		19.4175	19.4175	MG/KG	111		108		1	75 - 125		20	P
Cobalt		6.8398		52.6563		52.8320		48.5437	48.5437	MG/KG	94		95		0	75 - 125		20	P
Copper		11.7311		37.6126		37.8408		24.2718	24.2718	MG/KG	107		108		1	75 - 125		20	P
Iron		24840.0583		25987.7427		26281.6942		97.0874	97.0874	MG/KG	1182		1485		1			20	P
Lead		7.1000		19.6553		19.8505		14.5631	14.5631	MG/KG	86		88		1	75 - 125		20	P
Lithium		20.5650		120.4408		120.6126		97.0874	97.0874	MG/KG	103		103		0	75 - 125		20	P
Magnesium		5436.3796		5881.2340		5974.7563		194.1748	194.1748	MG/KG	229		277		2			20	P
Manganese		322.3951		368.3738		370.2243		48.5437	48.5437	MG/KG	95		99		1			20	P
Molybdenum		0.3049	B	181.5175		181.2689		194.1748	194.1748	MG/KG	93		93		0	75 - 125		20	P
Nickel		14.6184		60.4806		60.9699		48.5437	48.5437	MG/KG	94		95		1	75 - 125		20	P
Phosphorus		290.4971		352.8932		383.7000		97.0874	97.0874	MG/KG	64	N	96		8	75 - 125		20	P
Potassium		3220.3204		4530.9660		4656.8282		970.8738	970.8738	MG/KG	135	N	148	N	3	75 - 125		20	P
Selenium	78	0.0971	U	1.8289		1.9748		1.9417	1.9417	MG/KG	94		102		8	75 - 125		20	MS
Silver	107	0.0408	B	9.6427		9.8913		9.7087	9.7087	MG/KG	99		101		3	75 - 125		20	MS
Sodium		511.6466		1489.2680		1500.4592		970.8738	970.8738	MG/KG	101		102		1	75 - 125		20	P
Strontium	88	37.3398		42.3883		52.6214		7.7670	7.7670	MG/KG	65		197		22 *			20	MS
Thallium	203	0.2540		0.6258		0.6575		0.3883	0.3883	MG/KG	96		104		5	75 - 125		20	MS

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor</p> <p>MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>		<p>CONCENTRATION QUALIFIERS:</p> <p>U= Below MDL, B= Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>	
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Background Lab Sample ID: 7123470BKG Matrix Spike Lab Sample ID: 7123470MS Matrix Spike Duplicate Lab Sample ID: 7123470MSD
 Batch Id(s): P19237A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit				
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q	%R	RPD	M
Tin		2.8971	B	348.7757		347.5243		388.3495	388.3495	MG/KG	89		89		0	75 - 125	20	P	
Titanium		1342.8107		1608.9417		1692.4320		97.0874	97.0874	MG/KG	274		360		5			20	P
Vanadium		46.3039		100.4806		100.8155		48.5437	48.5437	MG/KG	112		112		0	75 - 125	20	P	
Zinc		55.7602		104.2175		104.7592		48.5437	48.5437	MG/KG	100		101		1	75 - 125	20	P	
Zirconium		5.6204		101.1019		101.2485		97.0874	97.0874	MG/KG	98		98		0	75 - 125	20	P	

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL, B= Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS
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Background Lab Sample ID: 7123470BKG
 Batch ID(s): P19237A
 Concentration Units: MG/KG

Duplicate Lab Sample ID: 7123470DUP

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			23391.4621		22124.6767		6		P
Antimony	107		0.7184	U	0.9476	B	200	*	P
Arsenic		3.9	5.0757		5.2117		3		P
Barium			76.9359		72.8718		5		P
Beryllium			0.7117	B	0.6961	B	2		P
Boron			6.5233	B	6.1767	B	5		P
Cadmium			0.3155	B	0.3398	B	7		P
Calcium			12994.3019		13316.8000		2		P
Chromium			25.6942		24.9699		3		P
Cobalt			6.8398		6.5893		4		P
Copper			11.7311		11.4204		3		P
Iron			24840.0583		25659.6650		3		P
Lead		2.9	7.1000		6.8379		4		P
Lithium			20.5650		20.0272		3		P
Magnesium			5436.3796		5207.5155		4		P
Manganese			322.3951		309.9816		4		P
Molybdenum			0.3049	B	0.3000	B	2		P
Nickel			14.6184		14.1155		4		P
Phosphorus			290.4971		256.8825		12		P
Potassium			3220.3204		3043.9942		6		P
Selenium	78		0.0971	U	0.0971	U			MS
Silver	107		0.0408	B	0.0252	U	200		MS
Sodium			511.6466		491.4311		4		P
Strontium	88		37.3398		34.9320		7		MS
Thallium	203	0.2	0.2540		0.2579		2		MS
Tin			2.8971	B	2.7476	B	5		P
Titanium			1342.8107		1249.3107		7		P
Vanadium			46.3039		45.0272		3		P
Zinc			55.7602		54.6981		2		P
Zirconium		4.9	5.6204		6.1592		9		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).
 The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.



Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: Duplicate Out of Spec
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Background Lab Sample ID: 7123469BKG
 Batch ID(s): P19238A
 Concentration Units: MG/KG

Duplicate Lab Sample ID: 7123469DUP

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Mercury			0.0095	U	0.0101	B	200	*	CV

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).
 The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

1.0% difference

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: * = Duplicate Out of Spec
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**Data Validation Report
Santa Susan Field Laboratory
Subarea 8**

SDG(s): PH065

Prepared for

CDM
555 17th Street, Suite 1100
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Prepared by

Laboratory Data Consultants, Inc
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September 10, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on July 11, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory, RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Volatile Organic Compounds (VOCs) by Environmental Protection Agency (EPA) SW 846 Method 8260B

Semivolatiles by EPA SW 846 Method 8270D-SIM

Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A

Metals by EPA SW 846 Method 6010C, 6020A, and 7471B

Total Petroleum Hydrocarbons (TPH) as Gasoline by EPA SW 846 Method 8015M

Total Petroleum Hydrocarbons (TPH) as Extractables by EPA SW 846 Method 8015M

Dioxins/Dibenzofurans by EPA Method 1613B

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibrations blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilution, method blanks, trip blanks, equipment blanks, field blanks and field duplicates. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the calibration blanks and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method blanks or initial, continuing, and preparation blanks with the exception of two method blanks for dioxins and metals. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicate (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits with the exception of one DUP for metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of three VOCs and semivolatiles. Associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met with the following exceptions:

Diluted Sample	Analyte	%D (Limits)	Associated Samples	Flag	A or P
SL-574-SA8-SB-14.0-15.0	Iron Lithium	12 (≤ 10) 11 (≤ 10)	SL-574-SA8-SB-0.0-0.5 SL-574-SA8-SB-4.0-5.0 SL-574-SA8-SB-9.0-10.0 SL-574-SA8-SB-14.0-15.0 SL-874-SA8-SB-14.0-15.0 SL-570-SA8-SB-0.0-0.5 SL-570-SA8-SB-4.0-5.0	J(all detects) UJ(all non-detects)	A

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH065	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pair was collected and analyzed for VOCs, semivolatiles, PCBs, metals, TPH as gasoline, TPH as extractables and dioxins. All RPDs were within QC criteria with the exception of two metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for VOCs and TPH as gasoline. No contaminants were found in the trip blank.

One equipment blank (from SDG PH064) was collected and analyzed for VOCs, semivolatiles, PCBs, metals, TPH as gasoline, TPH as extractables and dioxins. No contaminants were found in the equipment blank with the exception of several VOAs, semivolatiles, dioxins and metals. The associated sample results were not detected or were significantly greater than the concentrations found in the equipment blank, therefore no data were qualified.

One field blank (from SDG PH029) was collected and analyzed for VOCs, semivolatiles, PCBs, metals, TPH as gasoline, TPH as extractables and dioxins. No contaminants were found in the field blank with the exception of several VOAs, semivolatiles, dioxins and metals. The associated sample results were not detected or were significantly greater than the concentrations found in the field blank, therefore no data were qualified.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
11-Jul-2013	TB-071113	7125066	TB	5030B	8015M	III
11-Jul-2013	TB-071113	7125066	TB	5030B	8260B	III
11-Jul-2013	SL-574-SA8-SB-0.0-0.5	7125058	N	3050B	6010C	III
11-Jul-2013	SL-574-SA8-SB-0.0-0.5	7125058	N	3050B	6020A	III
11-Jul-2013	SL-574-SA8-SB-0.0-0.5	7125058	N	3546	8015M	III
11-Jul-2013	SL-574-SA8-SB-0.0-0.5	7125058	N	3546	8082A	III
11-Jul-2013	SL-574-SA8-SB-0.0-0.5	7125058	N	3546	8270D SIM	III
11-Jul-2013	SL-574-SA8-SB-0.0-0.5	7125058	N	METHOD	1613B	III
11-Jul-2013	SL-574-SA8-SB-0.0-0.5	7125058	N	METHOD	7471B	III
11-Jul-2013	SL-574-SA8-SB-4.0-5.0	7125059	N	3050B	6010C	III
11-Jul-2013	SL-574-SA8-SB-4.0-5.0	7125059	N	3050B	6020A	III
11-Jul-2013	SL-574-SA8-SB-4.0-5.0	7125059	N	3546	8015M	III
11-Jul-2013	SL-574-SA8-SB-4.0-5.0	7125059	N	3546	8082A	III
11-Jul-2013	SL-574-SA8-SB-4.0-5.0	7125059	N	3546	8270D SIM	III
11-Jul-2013	SL-574-SA8-SB-4.0-5.0	7125059	N	5035A	8015M	III
11-Jul-2013	SL-574-SA8-SB-4.0-5.0	7125059	N	METHOD	1613B	III
11-Jul-2013	SL-574-SA8-SB-4.0-5.0	7125059	N	METHOD	7471B	III
11-Jul-2013	SL-574-SA8-SB-9.0-10.0	7125060	N	3050B	6010C	III
11-Jul-2013	SL-574-SA8-SB-9.0-10.0	7125060	N	3050B	6020A	III
11-Jul-2013	SL-574-SA8-SB-9.0-10.0	7125060	N	3546	8015M	III
11-Jul-2013	SL-574-SA8-SB-9.0-10.0	7125060	N	3546	8082A	III
11-Jul-2013	SL-574-SA8-SB-9.0-10.0	7125060	N	3546	8270D SIM	III
11-Jul-2013	SL-574-SA8-SB-9.0-10.0	7125060	N	5035A	8015M	III
11-Jul-2013	SL-574-SA8-SB-9.0-10.0	7125060	N	METHOD	7471B	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0	7125061	N	3050B	6010C	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0	7125061	N	3050B	6020A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
11-Jul-2013	SL-574-SA8-SB-14.0-15.0	7125061	N	3546	8015M	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0	7125061	N	3546	8082A	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0	7125061	N	3546	8270D SIM	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0	7125061	N	5035A	8015M	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0	7125061	N	5035A	8260B	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0	7125061	N	METHOD	7471B	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0MS	7125062	MS	3050B	6010C	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0MS	7125062	MS	3050B	6020A	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0MS	7125062	MS	3546	8015M	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0MS	7125062	MS	3546	8082A	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0MS	7125062	MS	3546	8270D SIM	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0MS	7125062	MS	5035A	8015M	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0MS	7125062	MS	5035A	8260B	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0MS	7125062	MS	METHOD	7471B	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0DU	7125064	DUP	3050B	6010C	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0DU	7125064	DUP	3050B	6020A	III
11-Jul-2013	SL-574-SA8-SB-14.0-15.0DU	7125064	DUP	METHOD	7471B	III
11-Jul-2013	SL-874-SA8-SB-14.0-15.0	7125065	FD	3050B	6010C	III
11-Jul-2013	SL-874-SA8-SB-14.0-15.0	7125065	FD	3050B	6020A	III
11-Jul-2013	SL-874-SA8-SB-14.0-15.0	7125065	FD	3546	8015M	III
11-Jul-2013	SL-874-SA8-SB-14.0-15.0	7125065	FD	3546	8082A	III
11-Jul-2013	SL-874-SA8-SB-14.0-15.0	7125065	FD	3546	8270D SIM	III
11-Jul-2013	SL-874-SA8-SB-14.0-15.0	7125065	FD	5035A	8015M	III
11-Jul-2013	SL-874-SA8-SB-14.0-15.0	7125065	FD	5035A	8260B	III
11-Jul-2013	SL-874-SA8-SB-14.0-15.0	7125065	FD	METHOD	7471B	III
11-Jul-2013	SL-570-SA8-SB-0.0-0.5	7125067	N	3050B	6010C	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
11-Jul-2013	SL-570-SA8-SB-0.0-0.5	7125067	N	3050B	6020A	III
11-Jul-2013	SL-570-SA8-SB-0.0-0.5	7125067	N	3546	8015M	III
11-Jul-2013	SL-570-SA8-SB-0.0-0.5	7125067	N	3546	8082A	III
11-Jul-2013	SL-570-SA8-SB-0.0-0.5	7125067	N	3546	8270D SIM	III
11-Jul-2013	SL-570-SA8-SB-0.0-0.5	7125067	N	METHOD	1613B	III
11-Jul-2013	SL-570-SA8-SB-0.0-0.5	7125067	N	METHOD	7471B	III
11-Jul-2013	SL-570-SA8-SB-4.0-5.0	7125068	N	3050B	6010C	III
11-Jul-2013	SL-570-SA8-SB-4.0-5.0	7125068	N	3050B	6020A	III
11-Jul-2013	SL-570-SA8-SB-4.0-5.0	7125068	N	3546	8015M	III
11-Jul-2013	SL-570-SA8-SB-4.0-5.0	7125068	N	3546	8082A	III
11-Jul-2013	SL-570-SA8-SB-4.0-5.0	7125068	N	3546	8270D SIM	III
11-Jul-2013	SL-570-SA8-SB-4.0-5.0	7125068	N	5035A	8015M	III
11-Jul-2013	SL-570-SA8-SB-4.0-5.0	7125068	N	METHOD	1613B	III
11-Jul-2013	SL-570-SA8-SB-4.0-5.0	7125068	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-570-SA8-SB-0.0-0.5		Collected: 7/11/2013 1:50:00		Analysis Type: REA2		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	69200		7.10	MDL	42.5	PQL	mg/Kg	J	E

Sample ID: SL-570-SA8-SB-0.0-0.5		Collected: 7/11/2013 1:50:00		Analysis Type: REA3		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	14000		3.81	MDL	42.1	PQL	mg/Kg	J	A

Sample ID: SL-570-SA8-SB-0.0-0.5		Collected: 7/11/2013 1:50:00		Analysis Type: RES		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	1.18	J	0.787	MDL	4.25	PQL	mg/Kg	J	Z, Q
BERYLLIUM	0.140	J	0.0712	MDL	1.06	PQL	mg/Kg	J	Z
BORON	9.64	J	0.893	MDL	10.6	PQL	mg/Kg	J	Z
CADMIUM	0.548	J	0.0808	MDL	1.06	PQL	mg/Kg	J	Z
CHROMIUM	18.4		0.170	MDL	3.19	PQL	mg/Kg	J	Q
LITHIUM	8.9		0.36	MDL	4.3	PQL	mg/Kg	J	A
MANGANESE	225		0.0883	MDL	1.06	PQL	mg/Kg	J	E
POTASSIUM	1910		8.87	MDL	106	PQL	mg/Kg	J	Q
TIN	2.06	J	0.234	MDL	10.6	PQL	mg/Kg	U	B
ZINC	44.0		0.213	MDL	4.25	PQL	mg/Kg	J	Q

Sample ID: SL-570-SA8-SB-4.0-5.0		Collected: 7/11/2013 2:15:00		Analysis Type: REA3		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	25900		4.10	MDL	45.3	PQL	mg/Kg	J	A

Sample ID: SL-570-SA8-SB-4.0-5.0		Collected: 7/11/2013 2:15:00		Analysis Type: RES		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.53	U	0.838	MDL	4.53	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.714	J	0.0759	MDL	1.13	PQL	mg/Kg	J	Z
BORON	8.50	J	0.952	MDL	11.3	PQL	mg/Kg	J	Z
CADMIUM	0.497	J	0.0861	MDL	1.13	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-570-SA8-SB-4.0-5.0

Collected: 7/11/2013 2:15:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	7720		3.78	MDL	22.7	PQL	mg/Kg	J	E
CHROMIUM	26.7		0.181	MDL	3.40	PQL	mg/Kg	J	Q
LITHIUM	21.3		0.39	MDL	4.5	PQL	mg/Kg	J	A
MANGANESE	188		0.0940	MDL	1.13	PQL	mg/Kg	J	E
POTASSIUM	2490		9.45	MDL	113	PQL	mg/Kg	J	Q
TIN	3.01	J	0.249	MDL	11.3	PQL	mg/Kg	U	B
ZINC	58.4		0.227	MDL	4.53	PQL	mg/Kg	J	Q
Zirconium	2.90	J	0.952	MDL	5.66	PQL	mg/Kg	J	Z

Sample ID: SL-574-SA8-SB-0.0-0.5

Collected: 7/11/2013 8:45:00

Analysis Type: REA3

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	26500		4.13	MDL	45.6	PQL	mg/Kg	J	A

Sample ID: SL-574-SA8-SB-0.0-0.5

Collected: 7/11/2013 8:45:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.51	U	0.835	MDL	4.51	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.368	J	0.0756	MDL	1.13	PQL	mg/Kg	J	Z
BORON	9.30	J	0.948	MDL	11.3	PQL	mg/Kg	J	Z
CADMIUM	0.764	J	0.0858	MDL	1.13	PQL	mg/Kg	J	Z
CALCIUM	27900		3.77	MDL	22.6	PQL	mg/Kg	J	E
CHROMIUM	19.5		0.181	MDL	3.39	PQL	mg/Kg	J	Q
LITHIUM	12.2		0.38	MDL	4.5	PQL	mg/Kg	J	A
MANGANESE	251		0.0937	MDL	1.13	PQL	mg/Kg	J	E
POTASSIUM	2210		9.41	MDL	113	PQL	mg/Kg	J	Q
TIN	2.25	J	0.248	MDL	11.3	PQL	mg/Kg	U	B
ZINC	77.7		0.226	MDL	4.51	PQL	mg/Kg	J	Q
Zirconium	4.46	J	0.948	MDL	5.64	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-574-SA8-SB-14.0-15.0	Collected: 7/11/2013 10:00:00	Analysis Type: REA3	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	27600		3.81	MDL	42.1	PQL	mg/Kg	J	A

Sample ID: SL-574-SA8-SB-14.0-15.0	Collected: 7/11/2013 10:00:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.38	U	0.810	MDL	4.38	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.681	J	0.0734	MDL	1.10	PQL	mg/Kg	J	Z
BORON	5.22	J	0.920	MDL	11.0	PQL	mg/Kg	J	Z
CADMIUM	0.508	J	0.0832	MDL	1.10	PQL	mg/Kg	J	Z
CALCIUM	9730		3.66	MDL	21.9	PQL	mg/Kg	J	E, FD
CHROMIUM	31.9		0.175	MDL	3.29	PQL	mg/Kg	J	Q
LITHIUM	26.1		0.37	MDL	4.4	PQL	mg/Kg	J	A
MANGANESE	328		0.0909	MDL	1.10	PQL	mg/Kg	J	E
POTASSIUM	2720		9.13	MDL	110	PQL	mg/Kg	J	Q
TIN	3.20	J	0.241	MDL	11.0	PQL	mg/Kg	U	B
ZINC	75.1		0.219	MDL	4.38	PQL	mg/Kg	J	Q
Zirconium	5.48	U	0.920	MDL	5.48	PQL	mg/Kg	UJ	FD

Sample ID: SL-574-SA8-SB-4.0-5.0	Collected: 7/11/2013 9:30:00	Analysis Type: REA3	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	27200		3.65	MDL	40.3	PQL	mg/Kg	J	A

Sample ID: SL-574-SA8-SB-4.0-5.0	Collected: 7/11/2013 9:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.11	U	0.761	MDL	4.11	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.854	J	0.0689	MDL	1.03	PQL	mg/Kg	J	Z
CADMIUM	0.439	J	0.0781	MDL	1.03	PQL	mg/Kg	J	Z
CALCIUM	7560		3.43	MDL	20.6	PQL	mg/Kg	J	E
CHROMIUM	33.7		0.165	MDL	3.08	PQL	mg/Kg	J	Q
LITHIUM	27.7		0.35	MDL	4.1	PQL	mg/Kg	J	A

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-574-SA8-SB-4.0-5.0 Collected: 7/11/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MANGANESE	341		0.0853	MDL	1.03	PQL	mg/Kg	J	E
POTASSIUM	4040		8.57	MDL	103	PQL	mg/Kg	J	Q
TIN	2.58	J	0.226	MDL	10.3	PQL	mg/Kg	U	B
ZINC	62.4		0.206	MDL	4.11	PQL	mg/Kg	J	Q
Zirconium	4.00	J	0.864	MDL	5.14	PQL	mg/Kg	J	Z

Sample ID: SL-574-SA8-SB-9.0-10.0 Collected: 7/11/2013 9:45:00 Analysis Type: REA3 Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	20100		3.95	MDL	43.6	PQL	mg/Kg	J	A

Sample ID: SL-574-SA8-SB-9.0-10.0 Collected: 7/11/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.45	U	0.823	MDL	4.45	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.652	J	0.0745	MDL	1.11	PQL	mg/Kg	J	Z
BORON	5.92	J	0.934	MDL	11.1	PQL	mg/Kg	J	Z
CADMIUM	0.345	J	0.0845	MDL	1.11	PQL	mg/Kg	J	Z
CALCIUM	11900		3.72	MDL	22.2	PQL	mg/Kg	J	E
CHROMIUM	21.2		0.178	MDL	3.34	PQL	mg/Kg	J	Q
LITHIUM	19.9		0.38	MDL	4.4	PQL	mg/Kg	J	A
MANGANESE	145		0.0923	MDL	1.11	PQL	mg/Kg	J	E
POTASSIUM	2710		9.28	MDL	111	PQL	mg/Kg	J	Q
TIN	2.68	J	0.245	MDL	11.1	PQL	mg/Kg	U	B
ZINC	45.7		0.222	MDL	4.45	PQL	mg/Kg	J	Q
Zirconium	1.89	J	0.934	MDL	5.56	PQL	mg/Kg	J	Z

Sample ID: SL-874-SA8-SB-14.0-15.0 Collected: 7/11/2013 10:15:00 Analysis Type: REA3 Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
IRON	21200		3.85	MDL	42.5	PQL	mg/Kg	J	A

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-874-SA8-SB-14.0-15.0 Collected: 7/11/2013 10:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.17	U	0.771	MDL	4.17	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.493	J	0.0698	MDL	1.04	PQL	mg/Kg	J	Z
BORON	4.00	J	0.875	MDL	10.4	PQL	mg/Kg	J	Z
CADMIUM	0.447	J	0.0792	MDL	1.04	PQL	mg/Kg	J	Z
CALCIUM	5690		3.48	MDL	20.8	PQL	mg/Kg	J	E, FD
CHROMIUM	19.4		0.167	MDL	3.13	PQL	mg/Kg	J	Q
LITHIUM	19.6		0.35	MDL	4.2	PQL	mg/Kg	J	A
MANGANESE	267		0.0865	MDL	1.04	PQL	mg/Kg	J	E
POTASSIUM	2590		8.69	MDL	104	PQL	mg/Kg	J	Q
TIN	2.73	J	0.229	MDL	10.4	PQL	mg/Kg	U	B
ZINC	52.6		0.208	MDL	4.17	PQL	mg/Kg	J	Q
Zirconium	1.50	J	0.875	MDL	5.21	PQL	mg/Kg	J	Z, FD

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-570-SA8-SB-0.0-0.5 Collected: 7/11/2013 1:50:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.280	J	0.106	MDL	0.425	PQL	mg/Kg	J	Z

Sample ID: SL-570-SA8-SB-0.0-0.5 Collected: 7/11/2013 1:50:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0560	J	0.0276	MDL	0.213	PQL	mg/Kg	J	Z
THALLIUM	0.110	J	0.0319	MDL	0.213	PQL	mg/Kg	J	Z

Sample ID: SL-570-SA8-SB-4.0-5.0 Collected: 7/11/2013 2:15:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0354	J	0.0295	MDL	0.227	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-574-SA8-SB-0.0-0.5 Collected: 7/11/2013 8:45:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0390	J	0.0293	MDL	0.226	PQL	mg/Kg	J	Z
THALLIUM	0.155	J	0.0339	MDL	0.226	PQL	mg/Kg	J	Z

Sample ID: SL-574-SA8-SB-4.0-5.0 Collected: 7/11/2013 9:30:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0400	J	0.0267	MDL	0.206	PQL	mg/Kg	J	Z

Method Category: METALS
Method: 7471B **Matrix:** SO

Sample ID: SL-574-SA8-SB-9.0-10.0 Collected: 7/11/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0163	J	0.0106	MDL	0.0177	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-570-SA8-SB-0.0-0.5 Collected: 7/11/2013 1:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8,9-HPCDF	0.542	JB	0.0780	MDL	5.45	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.431	JBQ	0.0588	MDL	5.45	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.763	JB	0.0472	MDL	5.45	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	2.08	JB	0.0597	MDL	5.45	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.464	JBQ	0.0468	MDL	5.45	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.861	JBQ	0.0587	MDL	5.45	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.271	JB	0.0769	MDL	5.45	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.241	JB	0.0456	MDL	5.45	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.292	JB	0.0564	MDL	5.45	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.660	JB	0.0469	MDL	5.45	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-570-SA8-SB-0.0-0.5 Collected: 7/11/2013 1:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,3,4,7,8-PECDF	1.08	JB	0.0536	MDL	5.45	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0427	JBQ	0.0335	MDL	1.09	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.201	JQ	0.0620	MDL	1.09	PQL	ng/Kg	J	Z
OCDF	9.42	JB	0.111	MDL	10.9	PQL	ng/Kg	J	Z

Sample ID: SL-570-SA8-SB-4.0-5.0 Collected: 7/11/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.20	JB	0.0469	MDL	5.60	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.265	JBQ	0.0190	MDL	5.60	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0594	JB	0.0241	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0374	JB	0.0232	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.0553	JB	0.0137	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.0606	JB	0.0243	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDF	0.0420	JBQ	0.0130	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0480	JB	0.0235	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDF	0.0497	JBQ	0.0151	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0308	JBQ	0.0262	MDL	5.60	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0599	JBQ	0.0167	MDL	5.60	PQL	ng/Kg	U	B
2,3,4,6,7,8-HxCDF	0.0476	JBQ	0.0133	MDL	5.60	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0318	JBQ	0.0158	MDL	5.60	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0237	JB	0.0184	MDL	1.12	PQL	ng/Kg	U	B
OCDD	9.62	JB	0.0695	MDL	11.2	PQL	ng/Kg	J	Z
OCDF	0.419	JBQ	0.0636	MDL	11.2	PQL	ng/Kg	U	B

Sample ID: SL-574-SA8-SB-0.0-0.5 Collected: 7/11/2013 8:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	1.47	JB	0.0189	MDL	5.76	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.122	JB	0.0293	MDL	5.76	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.136	JBQ	0.0396	MDL	5.76	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDF	0.171	JB	0.0300	MDL	5.76	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-574-SA8-SB-0.0-0.5 Collected: 7/11/2013 8:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,6,7,8-HXCDD	0.893	JB	0.0431	MDL	5.76	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.178	JBQ	0.0299	MDL	5.76	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.533	JBQ	0.0410	MDL	5.76	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.265	JB	0.0362	MDL	5.76	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.174	JB	0.0407	MDL	5.76	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.298	JB	0.0339	MDL	5.76	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.262	JBQ	0.0288	MDL	5.76	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.476	JB	0.0296	MDL	5.76	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.105	JBQ	0.0269	MDL	1.15	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.192	J	0.0501	MDL	1.15	PQL	ng/Kg	J	Z
OCDF	2.47	JB	0.0370	MDL	11.5	PQL	ng/Kg	J	Z

Sample ID: SL-574-SA8-SB-4.0-5.0 Collected: 7/11/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.17	JB	0.0272	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.143	JBQ	0.0105	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0518	JBQ	0.0134	MDL	5.19	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0465	JBQ	0.0226	MDL	5.19	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0884	JB	0.0130	MDL	5.19	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.157	JB	0.0248	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.108	JBQ	0.0132	MDL	5.19	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.186	JB	0.0229	MDL	5.19	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.195	JB	0.0143	MDL	5.19	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.102	JBQ	0.0213	MDL	5.19	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.196	JBQ	0.0170	MDL	5.19	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0606	JB	0.0124	MDL	5.19	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.109	JBQ	0.0151	MDL	5.19	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0333	JBQ	0.0194	MDL	1.04	PQL	ng/Kg	U	B
OCDD	5.70	JB	0.0325	MDL	10.4	PQL	ng/Kg	J	Z
OCDF	0.217	JB	0.0280	MDL	10.4	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8082A	Matrix: SO

Sample ID: SL-570-SA8-SB-0.0-0.5 Collected: 7/11/2013 1:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
Aroclor 5460	12	J	11	MDL	36	PQL	ug/Kg	J	Z

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-570-SA8-SB-0.0-0.5 Collected: 7/11/2013 1:50:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.99	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	0.94	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	11	J	6.6	MDL	20	PQL	ug/Kg	J	Z
CHRYSENE	0.59	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
PYRENE	0.76	J	0.73	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-570-SA8-SB-4.0-5.0 Collected: 7/11/2013 2:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	6.0	J	3.8	MDL	19	PQL	ug/Kg	J	Z

Sample ID: SL-574-SA8-SB-4.0-5.0 Collected: 7/11/2013 9:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	4.7	J	3.4	MDL	17	PQL	ug/Kg	J	Z

Sample ID: SL-574-SA8-SB-9.0-10.0 Collected: 7/11/2013 9:45:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	6.4	J	3.7	MDL	19	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PrepPH065

eQAPP Name: CDM_SSFL_130808_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
*#	Professional Judgment
A	ICP Serial Dilution
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
FD	Field Duplicate Precision
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH065

Method Blank Outlier Report

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK1960B371832	7/16/2013 6:32:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDD 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDD 1,2,3,7,8,9-HxCDF 1,2,3,7,8-PECDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HxCDF 2,3,4,7,8-PECDF 2,3,7,8-TCDD OCDD OCDF	0.0478 ng/Kg 0.0258 ng/Kg 0.0609 ng/Kg 0.0220 ng/Kg 0.0412 ng/Kg 0.0258 ng/Kg 0.0216 ng/Kg 0.0474 ng/Kg 0.0394 ng/Kg 0.0613 ng/Kg 0.0646 ng/Kg 0.0356 ng/Kg 0.0232 ng/Kg 0.0400 ng/Kg 0.263 ng/Kg 0.106 ng/Kg	SL-570-SA8-SB-0.0-0.5 SL-570-SA8-SB-4.0-5.0 SL-574-SA8-SB-0.0-0.5 SL-574-SA8-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-570-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.241 ng/Kg	0.241U ng/Kg
SL-570-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.292 ng/Kg	0.292U ng/Kg
SL-570-SA8-SB-0.0-0.5(RES)	2,3,7,8-TCDD	0.0427 ng/Kg	0.0427U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0594 ng/Kg	0.0594U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDD	0.0374 ng/Kg	0.0374U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDF	0.0553 ng/Kg	0.0553U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HxCDD	0.0606 ng/Kg	0.0606U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HxCDF	0.0420 ng/Kg	0.0420U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDD	0.0480 ng/Kg	0.0480U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDF	0.0497 ng/Kg	0.0497U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDD	0.0308 ng/Kg	0.0308U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.0599 ng/Kg	0.0599U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HxCDF	0.0476 ng/Kg	0.0476U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	2,3,4,7,8-PECDF	0.0318 ng/Kg	0.0318U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	2,3,7,8-TCDD	0.0237 ng/Kg	0.0237U ng/Kg
SL-570-SA8-SB-4.0-5.0(RES)	OCDF	0.419 ng/Kg	0.419U ng/Kg
SL-574-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.122 ng/Kg	0.122U ng/Kg
SL-574-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.171 ng/Kg	0.171U ng/Kg
SL-574-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDD	0.174 ng/Kg	0.174U ng/Kg
SL-574-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.298 ng/Kg	0.298U ng/Kg
SL-574-SA8-SB-0.0-0.5(RES)	2,3,7,8-TCDD	0.105 ng/Kg	0.105U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0518 ng/Kg	0.0518U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDD	0.0465 ng/Kg	0.0465U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDF	0.0884 ng/Kg	0.0884U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HxCDF	0.108 ng/Kg	0.108U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDD	0.186 ng/Kg	0.186U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDF	0.195 ng/Kg	0.195U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDD	0.102 ng/Kg	0.102U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.196 ng/Kg	0.196U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-574-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0606 ng/Kg	0.0606U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	2,3,4,7,8-PECDF	0.109 ng/Kg	0.109U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	2,3,7,8-TCDD	0.0333 ng/Kg	0.0333U ng/Kg
SL-574-SA8-SB-4.0-5.0(RES)	OCDF	0.217 ng/Kg	0.217U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19737AB220512	7/18/2013 5:12:00 AM	CALCIUM TIN	5.84 mg/Kg 1.28 mg/Kg	SL-570-SA8-SB-0.0-0.5 SL-570-SA8-SB-4.0-5.0 SL-574-SA8-SB-0.0-0.5 SL-574-SA8-SB-14.0-15.0 SL-574-SA8-SB-4.0-5.0 SL-574-SA8-SB-9.0-10.0 SL-874-SA8-SB-14.0-15.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-570-SA8-SB-0.0-0.5(RES)	TIN	2.06 mg/Kg	2.06U mg/Kg
SL-570-SA8-SB-4.0-5.0(RES)	TIN	3.01 mg/Kg	3.01U mg/Kg
SL-574-SA8-SB-0.0-0.5(RES)	TIN	2.25 mg/Kg	2.25U mg/Kg
SL-574-SA8-SB-14.0-15.0(RES)	TIN	3.20 mg/Kg	3.20U mg/Kg
SL-574-SA8-SB-4.0-5.0(RES)	TIN	2.58 mg/Kg	2.58U mg/Kg
SL-574-SA8-SB-9.0-10.0(RES)	TIN	2.68 mg/Kg	2.68U mg/Kg
SL-874-SA8-SB-14.0-15.0(RES)	TIN	2.73 mg/Kg	2.73U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-574-SA8-SB-14.0-15.0MS (TOT) SL-574-SA8-SB-14.0-15.0MSD (TOT) (SL-570-SA8-SB-0.0-0.5 SL-570-SA8-SB-4.0-5.0 SL-574-SA8-SB-0.0-0.5 SL-574-SA8-SB-14.0-15.0 SL-574-SA8-SB-4.0-5.0 SL-574-SA8-SB-9.0-10.0 SL-874-SA8-SB-14.0-15.0)	ALUMINUM POTASSIUM TITANIUM	604 136 293	190 - -	75.00-125.00 75.00-125.00 75.00-125.00	- - -	ALUMINUM POTASSIUM TITANIUM	J (all detects) Al, Ti, No Qual, >4x
SL-574-SA8-SB-14.0-15.0MS (TOT) SL-574-SA8-SB-14.0-15.0MSD (TOT) (SL-570-SA8-SB-0.0-0.5 SL-570-SA8-SB-4.0-5.0 SL-574-SA8-SB-0.0-0.5 SL-574-SA8-SB-14.0-15.0 SL-574-SA8-SB-4.0-5.0 SL-574-SA8-SB-9.0-10.0 SL-874-SA8-SB-14.0-15.0)	CALCIUM MAGNESIUM PHOSPHORUS	-709 -74 -14	-964 -558 -90	75.00-125.00 75.00-125.00 75.00-125.00	- - -	CALCIUM MAGNESIUM PHOSPHORUS	No Qual, >4x
SL-574-SA8-SB-14.0-15.0MS (TOT) SL-574-SA8-SB-14.0-15.0MSD (TOT) (SL-570-SA8-SB-0.0-0.5 SL-570-SA8-SB-4.0-5.0 SL-574-SA8-SB-0.0-0.5 SL-574-SA8-SB-14.0-15.0 SL-574-SA8-SB-4.0-5.0 SL-574-SA8-SB-9.0-10.0 SL-874-SA8-SB-14.0-15.0)	ANTIMONY CHROMIUM MANGANESE ZINC	57 - 346 -	55 54 36 56	75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00	- - 39 (20.00) -	ANTIMONY CHROMIUM MANGANESE ZINC	J(all detects) UJ(all non-detects) Mn, No Qual %R >4x

Method: 6020A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-574-SA8-SB-14.0-15.0MS (TOT) SL-574-SA8-SB-14.0-15.0MSD (TOT) (SL-570-SA8-SB-0.0-0.5 SL-570-SA8-SB-4.0-5.0 SL-574-SA8-SB-0.0-0.5 SL-574-SA8-SB-14.0-15.0 SL-574-SA8-SB-4.0-5.0 SL-574-SA8-SB-9.0-10.0 SL-874-SA8-SB-14.0-15.0)	STRONTIUM	0	21	75.00-125.00	-	STRONTIUM	No Qual, >4x

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-574-SA8-SB-14.0-15.0MS (TOT) SL-574-SA8-SB-14.0-15.0MSD (TOT) (SL-570-SA8-SB-0.0-0.5 SL-570-SA8-SB-4.0-5.0 SL-574-SA8-SB-0.0-0.5 SL-574-SA8-SB-14.0-15.0 SL-574-SA8-SB-4.0-5.0 SL-574-SA8-SB-9.0-10.0 SL-874-SA8-SB-14.0-15.0)	IRON	-1653	-2598	75.00-125.00	-	IRON	No Qual, >4x

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-574-SA8-SB-14.0-15.0DUP (TOT)	ARSENIC	32	20.00	J (all detects) UJ (all non-detects) As, Pb, Zr, No Qual, OK by Difference
(SL-570-SA8-SB-0.0-0.5	CALCIUM	29	20.00	
SL -570-SA8-SB-4.0-5.0	LEAD	26	20.00	
SL -574-SA8-SB-0.0-0.5	Zirconium	200	20.00	
SL -574-SA8-SB-14.0-15.0				
SL -574-SA8-SB-4.0-5.0				
SL -574-SA8-SB-9.0-10.0				
SL -874-SA8-SB-14.0-15.0)				

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8270D SIM

Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P7LCLCSQ260839 (SL-570-SA8-SB-0.0-0.5 SL-570-SA8-SB-4.0-5.0 SL-574-SA8-SB-0.0-0.5 SL-574-SA8-SB-14.0-15.0 SL-574-SA8-SB-4.0-5.0 SL-574-SA8-SB-9.0-10.0 SL-874-SA8-SB-14.0-15.0)	Diethylphthalate	128	-	68.00-125.00	-	Diethylphthalate	J (all detects)

Method: 8260B

Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
LCSB54Q212231A LCSB55Q212253A LCSB55Y212316A (SL-574-SA8-SB-14.0-15.0 SL-874-SA8-SB-14.0-15.0)	4-METHYL-2-PENTANONE (MIB VINYL ACETATE	129 135	- 127	52.00-125.00 29.00-111.00	- -	4-METHYL-2-PENTANONE (MI VINYL ACETATE	J(all detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 160.3M

Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-574-SA8-SB-14.0-15.0	SL-874-SA8-SB-14.0-15.0			
MOISTURE	9.6	6.8	34		No Qualifiers Applied

Method: 6010C

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-574-SA8-SB-14.0-15.0 (TOT)	SL-874-SA8-SB-14.0-15.0 (TOT)			
ALUMINUM	18200	13600	29	50.00	No Qualifiers Applied
ARSENIC	6.86	5.89	15	50.00	
BARIUM	89.1	65.5	31	50.00	
BERYLLIUM	0.681	0.493	32	50.00	
BORON	5.22	4.00	26	50.00	
CADMIUM	0.508	0.447	13	50.00	
CHROMIUM	31.9	19.4	49	50.00	
COBALT	9.05	6.20	37	50.00	
COPPER	16.8	11.0	42	50.00	
IRON	27600	21200	26	50.00	
LEAD	6.13	4.56	29	50.00	
LITHIUM	26.1	19.6	28	50.00	
MAGNESIUM	6550	4510	37	50.00	
MANGANESE	328	267	21	50.00	
NICKEL	16.5	10.2	47	50.00	
PHOSPHORUS	683	417	48	50.00	
POTASSIUM	2720	2590	5	50.00	
SODIUM	448	300	40	50.00	
TIN	3.20	2.73	16	50.00	
TITANIUM	1670	1390	18	50.00	
VANADIUM	54.4	41.4	27	50.00	
ZINC	75.1	52.6	35	50.00	
CALCIUM	9730	5690	52	50.00	J(all detects)
Zirconium	5.48 U	1.50	200	50.00	UJ(all non-detects)

Method: 6020A

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-574-SA8-SB-14.0-15.0 (TOT)	SL-874-SA8-SB-14.0-15.0 (TOT)			
STRONTIUM	38.7	24.5	45	50.00	No Qualifiers Applied
THALLIUM	0.408	0.303	30	50.00	

Method: 9045M

Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-574-SA8-SB-14.0-15.0	SL-874-SA8-SB-14.0-15.0			
PH	8.10	7.96	2	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-570-SA8-SB-0.0-0.5	1,2,3,4,7,8,9-HPCDF	JB	0.542	5.45	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8-HxCDD	JBQ	0.431	5.45	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.763	5.45	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	2.08	5.45	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.464	5.45	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.861	5.45	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.271	5.45	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.241	5.45	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.292	5.45	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.660	5.45	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	1.08	5.45	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.0427	1.09	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.201	1.09	PQL	ng/Kg	
	OCDF	JB	9.42	10.9	PQL	ng/Kg	
SL-570-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	1.20	5.60	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.265	5.60	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0594	5.60	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.0374	5.60	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.0553	5.60	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.0606	5.60	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0420	5.60	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.0480	5.60	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.0497	5.60	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.0308	5.60	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0599	5.60	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0476	5.60	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0318	5.60	PQL	ng/Kg	
	2,3,7,8-TCDD	JB	0.0237	1.12	PQL	ng/Kg	
OCDD	JB	9.62	11.2	PQL	ng/Kg		
OCDF	JBQ	0.419	11.2	PQL	ng/Kg		
SL-574-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	1.47	5.76	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JB	0.122	5.76	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.136	5.76	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.171	5.76	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.893	5.76	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.178	5.76	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.533	5.76	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.265	5.76	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JB	0.174	5.76	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.298	5.76	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.262	5.76	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JB	0.476	5.76	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.105	1.15	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.192	1.15	PQL	ng/Kg	
OCDF	JB	2.47	11.5	PQL	ng/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-574-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	1.17	5.19	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.143	5.19	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0518	5.19	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0465	5.19	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.0884	5.19	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JB	0.157	5.19	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.108	5.19	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.186	5.19	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.195	5.19	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JBQ	0.102	5.19	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.196	5.19	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.0606	5.19	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.109	5.19	PQL	ng/Kg	
	2,3,7,8-TCDD	JBQ	0.0333	1.04	PQL	ng/Kg	
	OCDD	JB	5.70	10.4	PQL	ng/Kg	
OCDF	JB	0.217	10.4	PQL	ng/Kg		

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-570-SA8-SB-0.0-0.5	ANTIMONY	J	1.18	4.25	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.140	1.06	PQL	mg/Kg	
	BORON	J	9.64	10.6	PQL	mg/Kg	
	CADMIUM	J	0.548	1.06	PQL	mg/Kg	
SL-570-SA8-SB-4.0-5.0	TIN	J	2.06	10.6	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.714	1.13	PQL	mg/Kg	
	BORON	J	8.50	11.3	PQL	mg/Kg	
	CADMIUM	J	0.497	1.13	PQL	mg/Kg	
	TIN	J	3.01	11.3	PQL	mg/Kg	
SL-574-SA8-SB-0.0-0.5	Zirconium	J	2.90	5.66	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.368	1.13	PQL	mg/Kg	
	BORON	J	9.30	11.3	PQL	mg/Kg	
	CADMIUM	J	0.764	1.13	PQL	mg/Kg	
	TIN	J	2.25	11.3	PQL	mg/Kg	
SL-574-SA8-SB-14.0-15.0	Zirconium	J	4.46	5.64	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.681	1.10	PQL	mg/Kg	
	BORON	J	5.22	11.0	PQL	mg/Kg	
	CADMIUM	J	0.508	1.10	PQL	mg/Kg	
SL-574-SA8-SB-4.0-5.0	TIN	J	3.20	11.0	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.854	1.03	PQL	mg/Kg	
	CADMIUM	J	0.439	1.03	PQL	mg/Kg	
	TIN	J	2.58	10.3	PQL	mg/Kg	
SL-574-SA8-SB-9.0-10.0	Zirconium	J	4.00	5.14	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.652	1.11	PQL	mg/Kg	
	BORON	J	5.92	11.1	PQL	mg/Kg	
	CADMIUM	J	0.345	1.11	PQL	mg/Kg	
	TIN	J	2.68	11.1	PQL	mg/Kg	
SL-874-SA8-SB-14.0-15.0	Zirconium	J	1.89	5.56	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.493	1.04	PQL	mg/Kg	
	BORON	J	4.00	10.4	PQL	mg/Kg	
	CADMIUM	J	0.447	1.04	PQL	mg/Kg	
	TIN	J	2.73	10.4	PQL	mg/Kg	
Zirconium	J	1.50	5.21	PQL	mg/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH065

Laboratory: LL

EDD Filename: PH065_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-570-SA8-SB-0.0-0.5	SELENIUM	J	0.280	0.425	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0560	0.213	PQL	mg/Kg	
	THALLIUM	J	0.110	0.213	PQL	mg/Kg	
SL-570-SA8-SB-4.0-5.0	SILVER	J	0.0354	0.227	PQL	mg/Kg	J (all detects)
SL-574-SA8-SB-0.0-0.5	SILVER	J	0.0390	0.226	PQL	mg/Kg	J (all detects)
	THALLIUM	J	0.155	0.226	PQL	mg/Kg	
SL-574-SA8-SB-4.0-5.0	SILVER	J	0.0400	0.206	PQL	mg/Kg	J (all detects)

Method: 7471B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-574-SA8-SB-9.0-10.0	MERCURY	J	0.0163	0.0177	PQL	mg/Kg	J (all detects)

Method: 8082A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-570-SA8-SB-0.0-0.5	Aroclor 5460	J	12	36	PQL	ug/Kg	J (all detects)

Method: 8270D SIM
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-570-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	0.99	1.8	PQL	ug/Kg	J (all detects)
	BENZO(G,H,I)PERYLENE	J	0.94	1.8	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	11	20	PQL	ug/Kg	
	CHRYSENE	J	0.59	1.8	PQL	ug/Kg	
	PYRENE	J	0.76	1.8	PQL	ug/Kg	
SL-570-SA8-SB-4.0-5.0	CHRYSENE	J	6.0	19	PQL	ug/Kg	J (all detects)
SL-574-SA8-SB-4.0-5.0	CHRYSENE	J	4.7	17	PQL	ug/Kg	J (all detects)
SL-574-SA8-SB-9.0-10.0	CHRYSENE	J	6.4	19	PQL	ug/Kg	J (all detects)

LDC #: 30210D4

VALIDATION COMPLETENESS WORKSHEET

Date: 8/29/13

SDG #: PH065

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: al

2nd Reviewer: W

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	—	Sampling dates: 7/11/13
II.	ICP/MS Tune	—	
III.	Calibration	—	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	—	
VI.	Matrix Spike Analysis	SW	MS/D
VII.	Duplicate Sample Analysis	SW	DUP
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	SW	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	—	(4,5)
XV.	Field Blanks	SW	FB = FB-041113 EB = EB1-071013

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

(PH029)

(PH064)

Validated Samples: soil

1	SL-574-SA8-SB-0.0-0.5	11		21		31	
2	SL-574-SA8-SB-4.0-5.0	12		22		32	
3	SL-574-SA8-SB-9.0-10.0	13		23		33	
4	SL-574-SA8-SB-14.0-15.0	14		24		34	
5	SL-874-SA8-SB-14.0-15.0	15		25		35	
6	SL-570-SA8-SB-0.0-0.5	16		26		36	
7	SL-570-SA8-SB-4.0-5.0	17		27		37	
8	SL-574-SA8-SB-14.0-15.0MS	18		28		38	
9	SL-574-SA8-SB-14.0-15.0MSD	19		29		39	
10	SL-574-SA8-SB-14.0-15.0DUP	20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification											
	FB-041113 (SDG: PH029)	Action Limit	No Qualifiers										
Cu	0.0036	1.8											
Mo	0.0036	1.8											

Sampling date: 7/10/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification											
	EB1-071013 (SDG: PH064)	Action Limit	No Qualifiers										
Cu	0.0031	1.55											
Mo	0.0035	1.75											

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:

Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

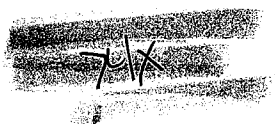
24K (Mn qualified off RPD) (8145)

Background Lab Sample ID: 7125061BKG Matrix Spike Lab Sample ID: 7125062MS Matrix Spike Duplicate Lab Sample ID: 7125063MSD
 Batch Id(s): P19737A, P20337C, P19738A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Control Limit		M
		Result	C	Result	C	Result	C				%R	Q	%R	Q		%R	RPD	
Aluminum		16452.8287		17661.2270		16833.8270		200.0000	200.0000	MG/KG	604		190		5		20	P
Antimony		0.7327	U	28.4360		27.6340		50.0000	50.0000	MG/KG	57	N	55	N	3	75 - 125	20	P
Arsenic		6.2020		20.1320		20.5150		15.0000	15.0000	MG/KG	93		95		2	75 - 125	20	P
Barium		80.5545		275.8390		251.6190		200.0000	200.0000	MG/KG	98		86		9	75 - 125	20	P
Beryllium		0.6158	B	5.4550		5.3280		5.0000	5.0000	MG/KG	97		94		2	75 - 125	20	P
Boron		4.7168	B	194.5330		189.3040		200.0000	200.0000	MG/KG	95		92		3	75 - 125	20	P
Cadmium		0.4594	B	5.1790		5.0620		5.0000	5.0000	MG/KG	94		92		2	75 - 125	20	P
Calcium		8798.4990		5961.7170		4944.0980		400.0000	400.0000	MG/KG	709		964		19			P
Chromium		28.8703		47.1680		39.7440		20.0000	20.0000	MG/KG	91		54	N	17	75 - 125	20	P
Cobalt		8.1842		55.3250		52.7760		50.0000	50.0000	MG/KG	94		89		5	75 - 125	20	P
Copper		15.2010		39.1010		35.4280		25.0000	25.0000	MG/KG	96		81		10	75 - 125	20	P
Iron		24921.5476		23316.9757		22399.2816		97.0874	97.0874	MG/KG	1653		2598		4			P
Lead		5.5416		18.1510		19.1270		15.0000	15.0000	MG/KG	84		91		5	75 - 125	20	P
Lithium		23.6000		121.7930		116.5740		100.0000	100.0000	MG/KG	98		93		4	75 - 125	20	P
Magnesium		5917.5525		5768.8950		4801.9540		200.0000	200.0000	MG/KG	74		558		18			P
Manganese		296.3139		469.1770		314.5130		50.0000	50.0000	MG/KG	346		36		39*			P
Mercury		0.0094	U	0.1917		0.1879		0.1661	0.1655	MG/KG	115		114		2	65 - 135	20	CV
Molybdenum		0.1683	U	190.1370		186.1480		200.0000	200.0000	MG/KG	95		93		2	75 - 125	20	P
Nickel		14.8772		64.4700		56.7430		50.0000	50.0000	MG/KG	99		84		13	75 - 125	20	P
Phosphorus		617.4337		603.4620		527.0730		100.0000	100.0000	MG/KG	14		90		14			P
Potassium		2458.2040		3815.3380		3474.3990		1000.0000	1000.0000	MG/KG	136	N	102		9	75 - 125	20	P
Selenium	78	0.0990	U	2.0980		1.8492		2.0000	2.0000	MG/KG	105		92		13	75 - 125	20	MS
Silver	107	0.0257	U	10.6160		10.5340		10.0000	10.0000	MG/KG	106		105		1	75 - 125	20	MS
Sodium		404.6277		1316.2020		1253.0350		1000.0000	1000.0000	MG/KG	91		85		5	75 - 125	20	P
Strontium	88	34.9703		35.0000		36.6600		8.0000	8.0000	MG/KG	0		21		5			MS
Thallium	203	0.3689		0.7988		0.6898		0.4000	0.4000	MG/KG	107		80		15	75 - 125	20	MS

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor</p> <p>MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U= Below MDL, B= Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>
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Background Lab Sample ID: 7125061BKG Matrix Spike Lab Sample ID: 7125062MS Matrix Spike Duplicate Lab Sample ID: 7125063MSD

Batch Id(s): P19737A, P20337C, P19738A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit		M		
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q		%R	RPD
Tin		2.8950	B	350.8090		346.5550		400.0000	400.0000	MG/KG	87		86		1	75 - 125	20	P	
Titanium		1505.7337		1799.1200		1581.1720		100.0000	100.0000	MG/KG	293		75		13			20	P
Vanadium		49.2129		96.8260		92.5530		50.0000	50.0000	MG/KG	95		87		5	75 - 125	20	P	
Zinc		67.8762		108.1970		96.0920		50.0000	50.0000	MG/KG	81		56	N	12	75 - 125	20	P	
Zirconium		0.8317	U	101.3780		99.3320		100.0000	100.0000	MG/KG	101		99		2	75 - 125	20	P	

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor</p> <p>MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U= Below MDL, B= Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>
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Background Lab Sample ID: 7125061BKG
 Batch ID(s): P19737A, P20337C, P19738A
 Concentration Units: MG/KG

Duplicate Lab Sample ID: 7125064DUP

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			16452.8287		17448.0120		6		P
Antimony			-1.5089	B	-1.3370	B	-12		P
Arsenic		4.0	6.2020		8.5450		32	*	P
Barium			80.5545		77.5640		4		P
Beryllium			0.6158	B	0.7390	B	18		P
Boron			4.7168	B	4.3340	B	8		P
Cadmium			0.4594	B	0.4840	B	5		P
Calcium			8798.4990		6562.5670		29	*	P
Chromium			28.8703		26.1500		10		P
Cobalt			8.1842		8.3310		2		P
Copper			15.2010		16.3060		7		P
Iron			24921.5476		25118.9080		1		P
Lead		3.0	5.5416		7.1820		26	*	P
Lithium			23.6000		25.0490		6		P
Magnesium			5917.5525		5521.0420		7		P
Manganese			296.3139		334.8200		12		P
Mercury			0.0094	U	0.0100	U			CV
Molybdenum			0.1683	U	0.1700	U			P
Nickel			14.8772		14.1210		5		P
Phosphorus			617.4337		573.4900		7		P
Potassium			2458.2040		2330.8220		5		P
Selenium	78		0.0990	U	0.1000	U			MS
Silver	107		0.0257	U	0.0260	U			MS
Sodium		99.0	404.6277		407.1170		1		P
Strontium	88		34.9703		31.5600		10		MS
Thallium	203	0.2	0.3689		0.3874		5		MS
Tin			2.8950	B	3.1200	B	7		P
Titanium			1505.7337		1386.3700		8		P
Vanadium			49.2129		53.2620		8		P
Zinc			67.8762		62.2810		9		P
Zirconium			0.8317	U	1.3700	B	200		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x IOQ).
 The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

ok by difference

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: = Duplicate Out of Spec
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QUALITY ASSURANCE SUMMARY
FORM 9

SERIAL DILUTIONS

SDG No.: PH065

Matrix: SOIL

Level

LOW

(low/med):

Background Lab Sample ID: 7125061BKG

Serial Dilution Lab Sample ID: 7125061L

Batch ID(s): P19737A, P20337C

Concentration Units: UG/L

Analyte	Mass	Initial Sample Result (I)	C	Serial Dilution Result (S)	C	% Diff.	Q	M
Aluminum		166173.5700		171492.2500		3		P
Antimony		7.4000	U	37.0000	U			P
Arsenic		62.6400		92.1000	B	47		P
Barium		813.6000		855.7000		5		P
Beryllium		6.2200	B	6.6500	B	7		P
Boron		47.6400	B	86.1500	B	81		P
Cadmium		4.6400	B	6.8500	B	48		P
Calcium		88864.8400		92560.5500		4		P
Chromium		291.5900		302.9500		4		P
Cobalt		82.6600		87.4500		6		P
Copper		153.5300		162.7500		6		P
Iron		261676.2500		292991.9500		12		P
Lead		55.9700		72.0500	B	29		P
Lithium		238.3600		265.5500		11		P
Magnesium		59767.2800		62261.1000		4		P
Manganese		2992.7700		3181.6000		6		P
Molybdenum		1.7000	U	8.5000	U			P
Nickel		150.2600		160.6000		7		P
Phosphorus		6236.0800		6339.9000		2		P
Potassium		24827.8600		25556.0000		3		P
Selenium	78	0.5000	U	2.5000	U			MS
Silver	107	0.1300	U	0.6500	U			MS
Sodium		4086.7400		4157.5500	B	2		P
Strontium	88	176.6000		177.4000		0		MS
Thallium	203	1.8630		2.0540	B	10		MS
Tin		29.2400	B	27.6500	B	5		P
Titanium		15207.9100		15760.6500		4		P
Vanadium		497.0500		512.9500		3		P
Zinc		685.5500		683.6500		0		P
Zirconium		8.4000	U	42.0000	U			P

NOTE: An E in column Q indicates the presence of a chemical or physical interference in the matrix when the % difference is greater than 10%. This applies only when (I) is greater than or equal to 50x MDL for ICP, 100x MDL for ICP-MS (6020), 50x MDL for ICP-MS (200.8), or 25x MDL for GFAA.

~~PH065 (A) All~~

METHODS:

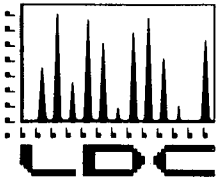
P = ICP Atomic Emission Spectrometer
MS = ICP Mass Spectrometry

CONCENTRATION QUALIFIERS:

U = Below MDL
B = Below LOQ

FLAGS:

E = Matrix Effects exist as proven by



LABORATORY DATA CONSULTANTS, INC.

2701 Loker Ave. West, Suite 220, Carlsbad, CA 92010 Bus: 760-827-1100 Fax: 760-827-1099

CDM
555 17th Street, Suite 1100
Denver, CO 80202
ATTN: Mrs. Cherie Zakowski

October 8, 2013

SUBJECT: Santa Susana Field Laboratory, Subarea 8, Data Validation

Dear Mrs. Zakowski,

Enclosed are the final validation reports for the fractions listed below. These SDGs were received on August 22, 2013. Attachment 1 is a summary of the samples that were reviewed for each analysis.

LDC Project # 30289:

<u>SDG #</u>	<u>Fraction</u>
PH066, PH067 PH068, PH069 PH070, PH071 PH072, PH073	Volatiles, Semivolatiles, Chlorinated Pesticides, Polychlorinated Biphenyls, Metals, Herbicides, Total Petroleum Hydrocarbons as Gasoline, Total Petroleum Hydrocarbons as Extractables, Dioxins/Dibenzofurans, Perchlorate, Wet Chemistry

The data validation was performed under Level III & IV guidelines. The analyses were validated using the following documents, as applicable to each method:

- Quality Assurance Project Plan for Santa Susana Field Laboratory, RCRA Facility Investigation, Surficial Media Operable Unit, March 2009, Revision 4
- USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review, June 2008
- USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review, January 2010
- Polychlorinated Dioxins/Dibenzofurans Data Review, September 2005
- EPA SW 846, Third Edition, Test Methods for Evaluating Solid Waste, update 1, July 1992; update IIA, August 1993; update II, September 1994; update IIB, January 1995; update III, December 1996; update IIIA, April 1998; IIIB, November 2004; Update IV, February 2007

Please feel free to contact us if you have any questions.

Sincerely,

A handwritten signature in black ink, appearing to read 'SMcKellar', written in a cursive style.

Shauna McKellar
Project Manager/Chemist

90/10 ADR/IV LDC #30289 (CDM Federal Programs-Chantilly VA / Santa Susana Field Laboratory, Subarea 8)

LDC	SDG#	DATE REC'D	(4) DATE DUE	VOA (8260B)		SVOA (8270D)		SVOA (8270D -SIM)		Pest. (8081B)		PCBs (8082)		Metals & Hg (SW846)		Herbs (8151A)		TPH-G (8015M)		TPH-E (8015M)		Dioxins (1613B)		CLO ₄ (6850)		Cr(VI) (7199)													
				W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S	W	S		
Matrix: Water/Soil																																							
A	PH066	08/22/13	09/20/13	-	-	0	7	0	11	-	-	0	11	0	11	-	-	1	7	0	11	0	11	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
B	PH067	08/22/13	09/20/13	-	-	-	-	0	12	0	5	0	12	0	12	0	5	1	7	0	12	0	12	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-	-
C	PH068	08/22/13	09/20/13	1	3	0	6	0	9	-	-	0	9	0	9	-	-	1	4	0	9	0	9	0	6	0	6	0	6	0	6	0	6	0	6	0	6	0	6
D	PH069	08/22/13	09/20/13	2	1	1	3	2	11	2	3	2	8	2	11	2	3	3	7	2	11	2	10	1	7	2	3												
E	PH070	08/22/13	09/20/13	1	2	0	4	0	11	-	-	0	11	0	11	-	-	1	5	0	11	0	11	0	4	0	4												
F	PH071	08/22/13	09/20/13	-	-	-	-	0	10	-	-	0	10	0	10	-	-	1	4	0	10	0	10	-	-	0	3												
G	PH072	08/22/13	09/20/13	-	-	-	-	0	9	-	-	0	9	0	9	-	-	1	6	0	9	0	6	-	-	-	-												
H	PH073	08/22/13	09/20/13	-	-	-	-	0	8	-	-	0	12	0	8	-	-	1	6	0	8	0	5	-	-	-	-												
Total	T/SM			4	6	1	20	2	81	2	8	2	82	2	81	2	8	10	46	2	81	2	74	1	17	2	16	0	0	0	0	0	0	0	0	0	0	0	552

Shaded cells indicate Level IV validation (all other cells are ADR review). These sample counts do not include MS/MSD, and DUPs

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 8

SDG: PH066

Prepared for

CDM
555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc
7750 El Camino Real, Suite 2L
Carlsbad, California 92009

September 24, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on July 12th, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOAs) by EPA SW 846 Method 8270D
SVOAs by EPA SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A, and 7471B
Total Petroleum Hydrocarbon as Gasoline (TPH-G) by EPA SW 846 Method 8015M
Total Petroleum Hydrocarbons as Extractable (TPH-E) by EPA SW 846 Method 8015M
Dioxins/Dibenzofurans by EPA Method 1613B

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks, and field duplicate samples. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for dioxins and metals. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for SVOAs, TPH-E, and metals. The benzidine result in sample SL-578-SA8-SB-4.0-5.0 was qualified as rejected (R) due to MS/MSD %Rs grossly outside QC limits (i.e., $\leq 0\%$). The remainder of the associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ), as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits with the exception of one DUP for metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of two LCS/LCSD pairs for SVOA. Since the percent recoveries were high and the results in the associated samples were non-detect, no data were qualified due to %Rs. The details regarding the qualification of data are provided in Enclosure I.

The lab noted that due to a lab error, 2,6-dichlorophenol was not spiked into the associated QC for samples analyzed for SVOAs by method 8270D. The lab stated that the sample was re-extracted outside the method required holding time and the QC was within acceptable limits; however, all results reported are from the original run. Due to lack of required QC in the original run, all 2,6-dichlorophenol results were qualified as detected estimated (J) or non-detected estimated (UJ) in the associated samples.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met with the following exceptions:

Diluted Sample	Analyte	%D (Limits)	Associated Samples	Flag	A or P
SL-578-SA8-SB-4.0-5.0	Barium Copper Manganese	11 (≤10) 24 (≤10) 21 (≤10)	All soil samples in SDG PH066	J (all detects) UJ (all non-detects)	A

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH066	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pair was collected and analyzed for SVOAs, PCBs, metals, TPH-G, TPH-E, and dioxins. All RPDs were within QC limits with the exception of several SVOAs, dioxins, and metals. In these duplicate pairs, the associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The field duplicate result

comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH-G. No contaminant concentrations were found in the trip blanks.

One equipment blank (from SDG PH064) was collected and analyzed for SVOAs, PCBs, metals, TPH-G, TPH-E, and dioxins. The equipment blanks had detections for several SVOAs, metals, and dioxins. The associated sample results were qualified as non-detected (U) due to equipment blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the equipment blanks were not qualified. The equipment blank outlier reports are presented in Enclosure I.

One field blank (from SDG PH029) was collected and analyzed for SVOAs, PCBs, metals, TPH-G, TPH-E, and dioxins. The field blank had detections for several semivolatiles, metals and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

One SVOC result was rejected due to MS/MSD %Rs grossly outside of QC limits. These results are not useable for all purposes.

The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. The remainder of the data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
12-Jul-2013	TB-071213	7126358	TB	5030B	8015M	III
12-Jul-2013	SL-578-SA8-SB-0.0-0.5	7126350	N	3050B	6010C	III
12-Jul-2013	SL-578-SA8-SB-0.0-0.5	7126350	N	3050B	6020A	III
12-Jul-2013	SL-578-SA8-SB-0.0-0.5	7126350	N	3546	8015M	III
12-Jul-2013	SL-578-SA8-SB-0.0-0.5	7126350	N	3546	8082A	III
12-Jul-2013	SL-578-SA8-SB-0.0-0.5	7126350	N	3546	8270D	III
12-Jul-2013	SL-578-SA8-SB-0.0-0.5	7126350	N	3546	8270D SIM	III
12-Jul-2013	SL-578-SA8-SB-0.0-0.5	7126350	N	METHOD	1613B	III
12-Jul-2013	SL-578-SA8-SB-0.0-0.5	7126350	N	METHOD	7471B	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0	7126351	N	3050B	6010C	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0	7126351	N	3050B	6020A	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0	7126351	N	3546	8015M	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0	7126351	N	3546	8082A	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0	7126351	N	3546	8270D	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0	7126351	N	3546	8270D SIM	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0	7126351	N	5035A	8015M	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0	7126351	N	METHOD	1613B	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0	7126351	N	METHOD	7471B	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MS	7126352	MS	3050B	6010C	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MS	7126352	MS	3050B	6020A	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MS	7126352	MS	3546	8015M	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MS	7126352	MS	3546	8082A	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MS	7126352	MS	3546	8270D	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MS	7126352	MS	3546	8270D SIM	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MS	7126352	MS	5035A	8015M	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MS	7126352	MS	METHOD	1613B	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MS	7126352	MS	METHOD	7471B	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MSD	7126353	MSD	3050B	6010C	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MSD	7126353	MSD	3050B	6020A	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MSD	7126353	MSD	3546	8015M	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MSD	7126353	MSD	3546	8082A	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MSD	7126353	MSD	3546	8270D	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MSD	7126353	MSD	3546	8270D SIM	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MSD	7126353	MSD	5035A	8015M	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MSD	7126353	MSD	METHOD	1613B	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0MSD	7126353	MSD	METHOD	7471B	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0DUP	7126354	DUP	3050B	6010C	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0DUP	7126354	DUP	3050B	6020A	III
12-Jul-2013	SL-578-SA8-SB-4.0-5.0DUP	7126354	DUP	METHOD	7471B	III
12-Jul-2013	SL-878-SA8-SB-4.0-5.0	7126355	N	3050B	6010C	III
12-Jul-2013	SL-878-SA8-SB-4.0-5.0	7126355	N	3050B	6020A	III
12-Jul-2013	SL-878-SA8-SB-4.0-5.0	7126355	N	3546	8015M	III
12-Jul-2013	SL-878-SA8-SB-4.0-5.0	7126355	N	3546	8082A	III
12-Jul-2013	SL-878-SA8-SB-4.0-5.0	7126355	N	3546	8270D	III
12-Jul-2013	SL-878-SA8-SB-4.0-5.0	7126355	N	3546	8270D SIM	III
12-Jul-2013	SL-878-SA8-SB-4.0-5.0	7126355	N	5035A	8015M	III
12-Jul-2013	SL-878-SA8-SB-4.0-5.0	7126355	N	METHOD	1613B	III
12-Jul-2013	SL-878-SA8-SB-4.0-5.0	7126355	N	METHOD	7471B	III
12-Jul-2013	SL-578-SA8-SB-9.0-10.0	7126356	N	3050B	6010C	III
12-Jul-2013	SL-578-SA8-SB-9.0-10.0	7126356	N	3050B	6020A	III
12-Jul-2013	SL-578-SA8-SB-9.0-10.0	7126356	N	3546	8015M	III
12-Jul-2013	SL-578-SA8-SB-9.0-10.0	7126356	N	3546	8082A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
12-Jul-2013	SL-578-SA8-SB-9.0-10.0	7126356	N	3546	8270D	III
12-Jul-2013	SL-578-SA8-SB-9.0-10.0	7126356	N	3546	8270D SIM	III
12-Jul-2013	SL-578-SA8-SB-9.0-10.0	7126356	N	5035A	8015M	III
12-Jul-2013	SL-578-SA8-SB-9.0-10.0	7126356	N	METHOD	1613B	III
12-Jul-2013	SL-578-SA8-SB-9.0-10.0	7126356	N	METHOD	7471B	III
12-Jul-2013	SL-580-SA8-SB-0.0-0.5	7126357	N	3050B	6010C	III
12-Jul-2013	SL-580-SA8-SB-0.0-0.5	7126357	N	3050B	6020A	III
12-Jul-2013	SL-580-SA8-SB-0.0-0.5	7126357	N	3546	8015M	III
12-Jul-2013	SL-580-SA8-SB-0.0-0.5	7126357	N	3546	8082A	III
12-Jul-2013	SL-580-SA8-SB-0.0-0.5	7126357	N	3546	8270D	III
12-Jul-2013	SL-580-SA8-SB-0.0-0.5	7126357	N	3546	8270D SIM	III
12-Jul-2013	SL-580-SA8-SB-0.0-0.5	7126357	N	METHOD	1613B	III
12-Jul-2013	SL-580-SA8-SB-0.0-0.5	7126357	N	METHOD	7471B	III
12-Jul-2013	SL-580-SA8-SB-4.0-5.0	7126359	N	3050B	6010C	III
12-Jul-2013	SL-580-SA8-SB-4.0-5.0	7126359	N	3050B	6020A	III
12-Jul-2013	SL-580-SA8-SB-4.0-5.0	7126359	N	3546	8015M	III
12-Jul-2013	SL-580-SA8-SB-4.0-5.0	7126359	N	3546	8082A	III
12-Jul-2013	SL-580-SA8-SB-4.0-5.0	7126359	N	3546	8270D	III
12-Jul-2013	SL-580-SA8-SB-4.0-5.0	7126359	N	3546	8270D SIM	III
12-Jul-2013	SL-580-SA8-SB-4.0-5.0	7126359	N	5035A	8015M	III
12-Jul-2013	SL-580-SA8-SB-4.0-5.0	7126359	N	METHOD	1613B	III
12-Jul-2013	SL-580-SA8-SB-4.0-5.0	7126359	N	METHOD	7471B	III
12-Jul-2013	SL-580-SA8-SB-10.5-11.5	7126360	N	3050B	6010C	III
12-Jul-2013	SL-580-SA8-SB-10.5-11.5	7126360	N	3050B	6020A	III
12-Jul-2013	SL-580-SA8-SB-10.5-11.5	7126360	N	3546	8015M	III
12-Jul-2013	SL-580-SA8-SB-10.5-11.5	7126360	N	3546	8082A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
12-Jul-2013	SL-580-SA8-SB-10.5-11.5	7126360	N	3546	8270D	III
12-Jul-2013	SL-580-SA8-SB-10.5-11.5	7126360	N	3546	8270D SIM	III
12-Jul-2013	SL-580-SA8-SB-10.5-11.5	7126360	N	5035A	8015M	III
12-Jul-2013	SL-580-SA8-SB-10.5-11.5	7126360	N	METHOD	1613B	III
12-Jul-2013	SL-580-SA8-SB-10.5-11.5	7126360	N	METHOD	7471B	III
12-Jul-2013	SL-581-SA8-SB-0.0-0.5	7126361	N	3050B	6010C	III
12-Jul-2013	SL-581-SA8-SB-0.0-0.5	7126361	N	3050B	6020A	III
12-Jul-2013	SL-581-SA8-SB-0.0-0.5	7126361	N	3546	8015M	III
12-Jul-2013	SL-581-SA8-SB-0.0-0.5	7126361	N	3546	8082A	III
12-Jul-2013	SL-581-SA8-SB-0.0-0.5	7126361	N	3546	8270D SIM	III
12-Jul-2013	SL-581-SA8-SB-0.0-0.5	7126361	N	METHOD	1613B	III
12-Jul-2013	SL-581-SA8-SB-0.0-0.5	7126361	N	METHOD	7471B	III
12-Jul-2013	SL-581-SA8-SB-4.5-5.5	7126362	N	3050B	6010C	III
12-Jul-2013	SL-581-SA8-SB-4.5-5.5	7126362	N	3050B	6020A	III
12-Jul-2013	SL-581-SA8-SB-4.5-5.5	7126362	N	3546	8015M	III
12-Jul-2013	SL-581-SA8-SB-4.5-5.5	7126362	N	3546	8082A	III
12-Jul-2013	SL-581-SA8-SB-4.5-5.5	7126362	N	3546	8270D SIM	III
12-Jul-2013	SL-581-SA8-SB-4.5-5.5	7126362	N	5035A	8015M	III
12-Jul-2013	SL-581-SA8-SB-4.5-5.5	7126362	N	METHOD	1613B	III
12-Jul-2013	SL-581-SA8-SB-4.5-5.5	7126362	N	METHOD	7471B	III
12-Jul-2013	SL-587-SA8-SB-0.0-0.5	7126363	N	3050B	6010C	III
12-Jul-2013	SL-587-SA8-SB-0.0-0.5	7126363	N	3050B	6020A	III
12-Jul-2013	SL-587-SA8-SB-0.0-0.5	7126363	N	3546	8015M	III
12-Jul-2013	SL-587-SA8-SB-0.0-0.5	7126363	N	3546	8082A	III
12-Jul-2013	SL-587-SA8-SB-0.0-0.5	7126363	N	3546	8270D SIM	III
12-Jul-2013	SL-587-SA8-SB-0.0-0.5	7126363	N	METHOD	1613B	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
12-Jul-2013	SL-587-SA8-SB-0.0-0.5	7126363	N	METHOD	7471B	III
12-Jul-2013	SL-587-SA8-SB-6.5-7.5	7126364	N	3050B	6010C	III
12-Jul-2013	SL-587-SA8-SB-6.5-7.5	7126364	N	3050B	6020A	III
12-Jul-2013	SL-587-SA8-SB-6.5-7.5	7126364	N	3546	8015M	III
12-Jul-2013	SL-587-SA8-SB-6.5-7.5	7126364	N	3546	8082A	III
12-Jul-2013	SL-587-SA8-SB-6.5-7.5	7126364	N	3546	8270D SIM	III
12-Jul-2013	SL-587-SA8-SB-6.5-7.5	7126364	N	5035A	8015M	III
12-Jul-2013	SL-587-SA8-SB-6.5-7.5	7126364	N	METHOD	1613B	III
12-Jul-2013	SL-587-SA8-SB-6.5-7.5	7126364	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS								
Method:	6010C	Matrix:		SO					

Sample ID: SL-578-SA8-SB-0.0-0.5 Collected: 7/12/2013 8:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.25	U	0.787	MDL	4.25	PQL	mg/Kg	UJ	Q, E
BARIUM	118		0.0351	MDL	1.06	PQL	mg/Kg	J	A
BERYLLIUM	0.735	J	0.0712	MDL	1.06	PQL	mg/Kg	J	Z
BORON	1.97	J	0.893	MDL	10.6	PQL	mg/Kg	J	Z
CADMIUM	0.371	J	0.0808	MDL	1.06	PQL	mg/Kg	J	Z
CHROMIUM	29.9		0.170	MDL	3.19	PQL	mg/Kg	J	Q
COPPER	23.4		0.308	MDL	2.13	PQL	mg/Kg	J	A
MANGANESE	534		0.0883	MDL	1.06	PQL	mg/Kg	J	E, A
MOLYBDENUM	0.308	J	0.181	MDL	2.13	PQL	mg/Kg	U	F, F
PHOSPHORUS	365		3.07	MDL	10.6	PQL	mg/Kg	J	Q
TIN	3.15	J	0.234	MDL	10.6	PQL	mg/Kg	U	B
VANADIUM	52.2		0.138	MDL	1.06	PQL	mg/Kg	J	Q

Sample ID: SL-578-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.44	U	0.821	MDL	4.44	PQL	mg/Kg	UJ	Q, E
BARIUM	116		0.0366	MDL	1.11	PQL	mg/Kg	J	A
BERYLLIUM	0.830	J	0.0743	MDL	1.11	PQL	mg/Kg	J	Z
BORON	2.75	J	0.932	MDL	11.1	PQL	mg/Kg	J	Z
CADMIUM	0.533	J	0.0843	MDL	1.11	PQL	mg/Kg	J	Z
CHROMIUM	29.7		0.177	MDL	3.33	PQL	mg/Kg	J	Q
COPPER	16.9		0.322	MDL	2.22	PQL	mg/Kg	J	A
MANGANESE	373		0.0921	MDL	1.11	PQL	mg/Kg	J	E, A
MOLYBDENUM	0.416	J	0.189	MDL	2.22	PQL	mg/Kg	U	F, F
PHOSPHORUS	286		3.21	MDL	11.1	PQL	mg/Kg	J	Q
TIN	3.04	J	0.244	MDL	11.1	PQL	mg/Kg	U	B
VANADIUM	55.0		0.144	MDL	1.11	PQL	mg/Kg	J	Q

Sample ID: SL-578-SA8-SB-9.0-10.0 Collected: 7/12/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.18	U	0.774	MDL	4.18	PQL	mg/Kg	UJ	Q, E
BARIUM	87.1		0.0345	MDL	1.05	PQL	mg/Kg	J	A
BERYLLIUM	0.440	J	0.0700	MDL	1.05	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS		
Method:	6010C	Matrix:	SO

Sample ID: SL-578-SA8-SB-9.0-10.0 Collected: 7/12/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CADMIUM	0.351	J	0.0795	MDL	1.05	PQL	mg/Kg	J	Z
CHROMIUM	21.2		0.167	MDL	3.14	PQL	mg/Kg	J	Q
COPPER	7.74		0.303	MDL	2.09	PQL	mg/Kg	J	A
MANGANESE	419		0.0868	MDL	1.05	PQL	mg/Kg	J	E, A
PHOSPHORUS	499		3.02	MDL	10.5	PQL	mg/Kg	J	Q
TIN	3.04	J	0.230	MDL	10.5	PQL	mg/Kg	U	B
VANADIUM	44.2		0.136	MDL	1.05	PQL	mg/Kg	J	Q
Zirconium	3.63	J	0.878	MDL	5.23	PQL	mg/Kg	J	Z

Sample ID: SL-580-SA8-SB-0.0-0.5 Collected: 7/12/2013 10:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.37	U	0.809	MDL	4.37	PQL	mg/Kg	UJ	Q, E
BARIUM	101		0.0361	MDL	1.09	PQL	mg/Kg	J	A
BERYLLIUM	0.754	J	0.0732	MDL	1.09	PQL	mg/Kg	J	Z
BORON	2.49	J	0.918	MDL	10.9	PQL	mg/Kg	J	Z
CADMIUM	0.303	J	0.0831	MDL	1.09	PQL	mg/Kg	J	Z
CHROMIUM	26.6		0.175	MDL	3.28	PQL	mg/Kg	J	Q
COPPER	11.1		0.317	MDL	2.19	PQL	mg/Kg	J	A
MANGANESE	306		0.0907	MDL	1.09	PQL	mg/Kg	J	E, A
MOLYBDENUM	0.238	J	0.186	MDL	2.19	PQL	mg/Kg	U	F, F
PHOSPHORUS	300		3.16	MDL	10.9	PQL	mg/Kg	J	Q
SODIUM	99.2	J	18.3	MDL	109	PQL	mg/Kg	J	Z
TIN	3.08	J	0.240	MDL	10.9	PQL	mg/Kg	U	B
VANADIUM	49.6		0.142	MDL	1.09	PQL	mg/Kg	J	Q
Zirconium	5.13	J	0.918	MDL	5.46	PQL	mg/Kg	J	Z

Sample ID: SL-580-SA8-SB-10.5-11.5 Collected: 7/12/2013 11:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.17	U	0.772	MDL	4.17	PQL	mg/Kg	UJ	Q, E
BARIUM	69.8		0.0344	MDL	1.04	PQL	mg/Kg	J	A
BERYLLIUM	0.609	J	0.0699	MDL	1.04	PQL	mg/Kg	J	Z
CADMIUM	0.321	J	0.0792	MDL	1.04	PQL	mg/Kg	J	Z
CHROMIUM	19.8		0.167	MDL	3.13	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS								
Method:	6010C		Matrix:	SO					

Sample ID: SL-580-SA8-SB-10.5-11.5 Collected: 7/12/2013 11:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
COPPER	9.10		0.302	MDL	2.09	PQL	mg/Kg	J	A
MANGANESE	245		0.0865	MDL	1.04	PQL	mg/Kg	J	E, A
MOLYBDENUM	1.08	J	0.177	MDL	2.09	PQL	mg/Kg	U	F, F
PHOSPHORUS	326		3.01	MDL	10.4	PQL	mg/Kg	J	Q
TIN	2.85	J	0.229	MDL	10.4	PQL	mg/Kg	U	B
VANADIUM	42.4		0.136	MDL	1.04	PQL	mg/Kg	J	Q
Zirconium	4.17	J	0.876	MDL	5.21	PQL	mg/Kg	J	Z

Sample ID: SL-580-SA8-SB-4.0-5.0 Collected: 7/12/2013 10:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.21	U	0.778	MDL	4.21	PQL	mg/Kg	UJ	Q, E
BARIUM	125		0.0347	MDL	1.05	PQL	mg/Kg	J	A
BERYLLIUM	0.769	J	0.0705	MDL	1.05	PQL	mg/Kg	J	Z
BORON	4.29	J	0.884	MDL	10.5	PQL	mg/Kg	J	Z
CADMIUM	0.449	J	0.0799	MDL	1.05	PQL	mg/Kg	J	Z
CHROMIUM	26.8		0.168	MDL	3.16	PQL	mg/Kg	J	Q
COPPER	15.3		0.305	MDL	2.10	PQL	mg/Kg	J	A
MANGANESE	435		0.0873	MDL	1.05	PQL	mg/Kg	J	E, A
MOLYBDENUM	0.332	J	0.179	MDL	2.10	PQL	mg/Kg	U	F, F
PHOSPHORUS	306		3.04	MDL	10.5	PQL	mg/Kg	J	Q
TIN	2.94	J	0.231	MDL	10.5	PQL	mg/Kg	U	B
VANADIUM	50.0		0.137	MDL	1.05	PQL	mg/Kg	J	Q

Sample ID: SL-581-SA8-SB-0.0-0.5 Collected: 7/12/2013 12:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.16	U	0.770	MDL	4.16	PQL	mg/Kg	UJ	Q, E
BARIUM	119		0.0343	MDL	1.04	PQL	mg/Kg	J	A
BERYLLIUM	0.810	J	0.0697	MDL	1.04	PQL	mg/Kg	J	Z
BORON	2.82	J	0.874	MDL	10.4	PQL	mg/Kg	J	Z
CADMIUM	0.496	J	0.0791	MDL	1.04	PQL	mg/Kg	J	Z
CHROMIUM	28.8		0.166	MDL	3.12	PQL	mg/Kg	J	Q
COPPER	16.1		0.302	MDL	2.08	PQL	mg/Kg	J	A
MANGANESE	438		0.0864	MDL	1.04	PQL	mg/Kg	J	E, A

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-581-SA8-SB-0.0-0.5	Collected: 7/12/2013 12:40:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MOLYBDENUM	0.314	J	0.177	MDL	2.08	PQL	mg/Kg	U	F, F
PHOSPHORUS	248		3.01	MDL	10.4	PQL	mg/Kg	J	Q
SODIUM	77.6	J	17.4	MDL	104	PQL	mg/Kg	J	Z
TIN	2.85	J	0.229	MDL	10.4	PQL	mg/Kg	U	B
VANADIUM	54.2		0.135	MDL	1.04	PQL	mg/Kg	J	Q

Sample ID: SL-581-SA8-SB-4.5-5.5	Collected: 7/12/2013 12:50:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	3.92	U	0.726	MDL	3.92	PQL	mg/Kg	UJ	Q, E
BARIUM	71.1		0.0324	MDL	0.981	PQL	mg/Kg	J	A
BERYLLIUM	0.418	J	0.0657	MDL	0.981	PQL	mg/Kg	J	Z
CADMIUM	0.294	J	0.0745	MDL	0.981	PQL	mg/Kg	J	Z
CHROMIUM	18.1		0.157	MDL	2.94	PQL	mg/Kg	J	Q
COPPER	7.24		0.284	MDL	1.96	PQL	mg/Kg	J	A
MANGANESE	251		0.0814	MDL	0.981	PQL	mg/Kg	J	E, A
MOLYBDENUM	0.204	J	0.167	MDL	1.96	PQL	mg/Kg	U	F, F
PHOSPHORUS	384		2.83	MDL	9.81	PQL	mg/Kg	J	Q
SODIUM	86.7	J	16.4	MDL	98.1	PQL	mg/Kg	J	Z
TIN	2.97	J	0.216	MDL	9.81	PQL	mg/Kg	U	B
VANADIUM	37.4		0.128	MDL	0.981	PQL	mg/Kg	J	Q
Zirconium	3.05	J	0.824	MDL	4.90	PQL	mg/Kg	J	Z

Sample ID: SL-587-SA8-SB-0.0-0.5	Collected: 7/12/2013 1:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	3.88	U	0.717	MDL	3.88	PQL	mg/Kg	UJ	Q, E
ARSENIC	3.51	J	0.678	MDL	3.88	PQL	mg/Kg	J	Z
BARIUM	112		0.0320	MDL	0.969	PQL	mg/Kg	J	A
BERYLLIUM	0.542	J	0.0649	MDL	0.969	PQL	mg/Kg	J	Z
BORON	0.814	J	0.814	MDL	9.69	PQL	mg/Kg	J	Z
CADMIUM	0.360	J	0.0736	MDL	0.969	PQL	mg/Kg	J	Z
CHROMIUM	18.3		0.155	MDL	2.91	PQL	mg/Kg	J	Q
COPPER	9.63		0.281	MDL	1.94	PQL	mg/Kg	J	A
MANGANESE	322		0.0804	MDL	0.969	PQL	mg/Kg	J	E, A

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-587-SA8-SB-0.0-0.5 Collected: 7/12/2013 1:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MOLYBDENUM	0.453	J	0.165	MDL	1.94	PQL	mg/Kg	U	F, F
PHOSPHORUS	574		2.80	MDL	9.69	PQL	mg/Kg	J	Q
SODIUM	67.7	J	16.2	MDL	96.9	PQL	mg/Kg	J	Z
TIN	2.50	J	0.213	MDL	9.69	PQL	mg/Kg	U	B
VANADIUM	31.7		0.126	MDL	0.969	PQL	mg/Kg	J	Q
Zirconium	3.20	J	0.814	MDL	4.84	PQL	mg/Kg	J	Z

Sample ID: SL-587-SA8-SB-6.5-7.5 Collected: 7/12/2013 1:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	3.98	U	0.736	MDL	3.98	PQL	mg/Kg	UJ	Q, E
ARSENIC	3.25	J	0.696	MDL	3.98	PQL	mg/Kg	J	Z
BARIUM	66.0		0.0328	MDL	0.994	PQL	mg/Kg	J	A
BERYLLIUM	0.483	J	0.0666	MDL	0.994	PQL	mg/Kg	J	Z
CADMIUM	0.203	J	0.0756	MDL	0.994	PQL	mg/Kg	J	Z
CHROMIUM	14.7		0.159	MDL	2.98	PQL	mg/Kg	J	Q
COPPER	5.33		0.288	MDL	1.99	PQL	mg/Kg	J	A
MANGANESE	144		0.0825	MDL	0.994	PQL	mg/Kg	J	E, A
PHOSPHORUS	166		2.87	MDL	9.94	PQL	mg/Kg	J	Q
TIN	2.70	J	0.219	MDL	9.94	PQL	mg/Kg	U	B
VANADIUM	26.9		0.129	MDL	0.994	PQL	mg/Kg	J	Q
Zirconium	2.46	J	0.835	MDL	4.97	PQL	mg/Kg	J	Z

Sample ID: SL-878-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.30	U	0.795	MDL	4.30	PQL	mg/Kg	UJ	Q, E
BARIUM	135		0.0355	MDL	1.07	PQL	mg/Kg	J	A
BERYLLIUM	1.02	J	0.0720	MDL	1.07	PQL	mg/Kg	J	Z
BORON	3.04	J	0.903	MDL	10.7	PQL	mg/Kg	J	Z
CADMIUM	0.408	J	0.0817	MDL	1.07	PQL	mg/Kg	J	Z
CHROMIUM	36.2		0.172	MDL	3.22	PQL	mg/Kg	J	Q
COPPER	17.7		0.312	MDL	2.15	PQL	mg/Kg	J	A
MANGANESE	524		0.0892	MDL	1.07	PQL	mg/Kg	J	E, A
MOLYBDENUM	0.229	J	0.183	MDL	2.15	PQL	mg/Kg	U	F, F

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-878-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
PHOSPHORUS	242		3.11	MDL	10.7	PQL	mg/Kg	J	Q
TIN	3.10	J	0.236	MDL	10.7	PQL	mg/Kg	U	B
VANADIUM	66.8		0.140	MDL	1.07	PQL	mg/Kg	J	Q

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-578-SA8-SB-0.0-5.0 Collected: 7/12/2013 8:35:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	31.5		0.0723	MDL	0.425	PQL	mg/Kg	J	Q

Sample ID: SL-578-SA8-SB-0.0-5.0 Collected: 7/12/2013 8:35:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.425	U	0.106	MDL	0.425	PQL	mg/Kg	UJ	E

Sample ID: SL-578-SA8-SB-0.0-5.0 Collected: 7/12/2013 8:35:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0890	J	0.0276	MDL	0.213	PQL	mg/Kg	J	Z, Q
THALLIUM	0.368		0.0319	MDL	0.213	PQL	mg/Kg	J	Q

Sample ID: SL-578-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:15:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	33.4		0.0754	MDL	0.444	PQL	mg/Kg	J	Q

Sample ID: SL-578-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:15:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.277	J	0.111	MDL	0.444	PQL	mg/Kg	J	Z, Q, E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-578-SA8-SB-4.0-5.0			Collected: 7/12/2013 9:15:00			Analysis Type: RES		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0557	J	0.0288	MDL	0.222	PQL	mg/Kg	J	Z, Q
THALLIUM	0.402		0.0333	MDL	0.222	PQL	mg/Kg	J	Q

Sample ID: SL-578-SA8-SB-9.0-10.0			Collected: 7/12/2013 9:45:00			Analysis Type: REA		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	20.8		0.0711	MDL	0.418	PQL	mg/Kg	J	Q

Sample ID: SL-578-SA8-SB-9.0-10.0			Collected: 7/12/2013 9:45:00			Analysis Type: REA2		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.146	J	0.105	MDL	0.418	PQL	mg/Kg	J	Z, Q, E

Sample ID: SL-578-SA8-SB-9.0-10.0			Collected: 7/12/2013 9:45:00			Analysis Type: RES		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
THALLIUM	0.415		0.0314	MDL	0.209	PQL	mg/Kg	J	Q

Sample ID: SL-580-SA8-SB-0.0-0.5			Collected: 7/12/2013 10:30:00			Analysis Type: REA		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	29.5		0.0743	MDL	0.437	PQL	mg/Kg	J	Q

Sample ID: SL-580-SA8-SB-0.0-0.5			Collected: 7/12/2013 10:30:00			Analysis Type: REA2		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.123	J	0.109	MDL	0.437	PQL	mg/Kg	J	Z, Q, E

Sample ID: SL-580-SA8-SB-0.0-0.5			Collected: 7/12/2013 10:30:00			Analysis Type: RES		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
THALLIUM	0.364		0.0328	MDL	0.219	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-580-SA8-SB-10.5-11.5		Collected: 7/12/2013 11:00:00		Analysis Type: REA		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	20.4		0.0709	MDL	0.417	PQL	mg/Kg	J	Q

Sample ID: SL-580-SA8-SB-10.5-11.5		Collected: 7/12/2013 11:00:00		Analysis Type: REA2		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.105	J	0.104	MDL	0.417	PQL	mg/Kg	J	Z, Q, E

Sample ID: SL-580-SA8-SB-10.5-11.5		Collected: 7/12/2013 11:00:00		Analysis Type: RES		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
THALLIUM	0.320		0.0313	MDL	0.209	PQL	mg/Kg	J	Q

Sample ID: SL-580-SA8-SB-4.0-5.0		Collected: 7/12/2013 10:45:00		Analysis Type: REA		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	32.9		0.0715	MDL	0.421	PQL	mg/Kg	J	Q

Sample ID: SL-580-SA8-SB-4.0-5.0		Collected: 7/12/2013 10:45:00		Analysis Type: REA2		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.247	J	0.105	MDL	0.421	PQL	mg/Kg	J	Z, Q, E

Sample ID: SL-580-SA8-SB-4.0-5.0		Collected: 7/12/2013 10:45:00		Analysis Type: RES		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0404	J	0.0273	MDL	0.210	PQL	mg/Kg	J	Z, Q
THALLIUM	0.441		0.0316	MDL	0.210	PQL	mg/Kg	J	Q

Sample ID: SL-581-SA8-SB-0.0-0.5		Collected: 7/12/2013 12:40:00		Analysis Type: REA		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	28.7		0.0708	MDL	0.416	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-581-SA8-SB-0.0-0.5			Collected: 7/12/2013 12:40:00			Analysis Type: REA2		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.207	J	0.104	MDL	0.416	PQL	mg/Kg	J	Z, Q, E

Sample ID: SL-581-SA8-SB-0.0-0.5			Collected: 7/12/2013 12:40:00			Analysis Type: RES		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0503	J	0.0271	MDL	0.208	PQL	mg/Kg	J	Z, Q
THALLIUM	0.457		0.0312	MDL	0.208	PQL	mg/Kg	J	Q

Sample ID: SL-581-SA8-SB-4.5-5.5			Collected: 7/12/2013 12:50:00			Analysis Type: REA		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	15.6		0.0667	MDL	0.392	PQL	mg/Kg	J	Q

Sample ID: SL-581-SA8-SB-4.5-5.5			Collected: 7/12/2013 12:50:00			Analysis Type: REA2		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.392	U	0.0981	MDL	0.392	PQL	mg/Kg	UJ	E

Sample ID: SL-581-SA8-SB-4.5-5.5			Collected: 7/12/2013 12:50:00			Analysis Type: RES		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
THALLIUM	0.298		0.0294	MDL	0.196	PQL	mg/Kg	J	Q

Sample ID: SL-587-SA8-SB-0.0-0.5			Collected: 7/12/2013 1:30:00			Analysis Type: REA		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	17.8		0.0659	MDL	0.388	PQL	mg/Kg	J	Q

Sample ID: SL-587-SA8-SB-0.0-0.5			Collected: 7/12/2013 1:30:00			Analysis Type: REA2		Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.107	J	0.0969	MDL	0.388	PQL	mg/Kg	J	Z, Q, E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-587-SA8-SB-0.0-0.5		Collected: 7/12/2013 1:30:00		Analysis Type: RES				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
THALLIUM	0.285		0.0291	MDL	0.194	PQL	mg/Kg	J	Q

Sample ID: SL-587-SA8-SB-6.5-7.5		Collected: 7/12/2013 1:40:00		Analysis Type: REA				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	17.3		0.0676	MDL	0.398	PQL	mg/Kg	J	Q

Sample ID: SL-587-SA8-SB-6.5-7.5		Collected: 7/12/2013 1:40:00		Analysis Type: REA2				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.398	U	0.0994	MDL	0.398	PQL	mg/Kg	UJ	E

Sample ID: SL-587-SA8-SB-6.5-7.5		Collected: 7/12/2013 1:40:00		Analysis Type: RES				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0650	J	0.0259	MDL	0.199	PQL	mg/Kg	J	Z, Q
THALLIUM	0.269		0.0298	MDL	0.199	PQL	mg/Kg	J	Q

Sample ID: SL-878-SA8-SB-4.0-5.0		Collected: 7/12/2013 9:30:00		Analysis Type: REA				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
STRONTIUM	34.1		0.0731	MDL	0.430	PQL	mg/Kg	J	Q

Sample ID: SL-878-SA8-SB-4.0-5.0		Collected: 7/12/2013 9:30:00		Analysis Type: REA2				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.326	J	0.107	MDL	0.430	PQL	mg/Kg	J	Z, Q, E

Sample ID: SL-878-SA8-SB-4.0-5.0		Collected: 7/12/2013 9:30:00		Analysis Type: RES				Dilution: 2	
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0645	J	0.0279	MDL	0.215	PQL	mg/Kg	J	Z, Q
THALLIUM	0.588		0.0322	MDL	0.215	PQL	mg/Kg	J	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA								
Method:	1613B	Matrix:	SO						

Sample ID: SL-578-SA8-SB-0.0-0.5 Collected: 7/12/2013 8:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.41	JB	0.0497	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.194	JB	0.0163	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0406	JBQ	0.0247	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0256	JB	0.0153	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0938	JQ	0.0307	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0418	JBQ	0.0136	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.131	J	0.0329	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.168	J	0.0161	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.0515	JBQ	0.0204	MDL	5.30	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0272	JBQ	0.0139	MDL	5.30	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0735	JQ	0.0197	MDL	5.30	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0326	JQ	0.0325	MDL	1.06	PQL	ng/Kg	J	Z
OCDF	0.478	J	0.0359	MDL	10.6	PQL	ng/Kg	J	Z

Sample ID: SL-578-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.41	JB	0.0528	MDL	5.49	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.204	JBQ	0.0880	MDL	5.49	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.0838	J	0.0657	MDL	5.49	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0311	JBQ	0.0296	MDL	5.49	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.520	JQ	0.0699	MDL	5.49	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0590	JBQ	0.0279	MDL	5.49	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.800	J	0.0635	MDL	5.49	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.901	J	0.0334	MDL	5.49	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.127	JBQ	0.0372	MDL	5.49	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0870	JB	0.0273	MDL	5.49	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.131	JQ	0.0331	MDL	5.49	PQL	ng/Kg	J	Z
OCDF	0.414	J	0.0456	MDL	11.0	PQL	ng/Kg	J	Z

Sample ID: SL-578-SA8-SB-9.0-10.0 Collected: 7/12/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.163	JBQ	0.0270	MDL	5.05	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0450	JBQ	0.0120	MDL	5.05	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA		
Method:	1613B	Matrix:	SO

Sample ID: SL-578-SA8-SB-9.0-10.0 Collected: 7/12/2013 9:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8,9-HPCDF	0.0234	JBQ	0.0182	MDL	5.05	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0298	JB	0.00993	MDL	5.05	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0353	JQ	0.0206	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.0288	JQ	0.0109	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.0281	JB	0.0168	MDL	5.05	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0155	JQ	0.0151	MDL	5.05	PQL	ng/Kg	J	Z
OCDD	0.988	JBQ	0.0254	MDL	10.1	PQL	ng/Kg	U	B
OCDF	0.0542	JQ	0.0261	MDL	10.1	PQL	ng/Kg	J	Z

Sample ID: SL-580-SA8-SB-0.0-0.5 Collected: 7/12/2013 10:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	2.09	JB	0.0439	MDL	5.43	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.123	JBQ	0.0110	MDL	5.43	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.100	JQ	0.0295	MDL	5.43	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0390	JBQ	0.0164	MDL	5.43	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.144	JQ	0.0281	MDL	5.43	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.216	J	0.0191	MDL	5.43	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.0781	JBQ	0.0192	MDL	5.43	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0200	JBQ	0.0162	MDL	5.43	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0531	JQ	0.0178	MDL	5.43	PQL	ng/Kg	J	Z
OCDF	0.300	JQ	0.0360	MDL	10.9	PQL	ng/Kg	J	Z

Sample ID: SL-580-SA8-SB-10.5-11.5 Collected: 7/12/2013 11:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.316	JB	0.0399	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.0890	JBQ	0.0130	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0340	JBQ	0.0187	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0653	JQ	0.0264	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.122	JB	0.0141	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.115	JQ	0.0283	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.111	JBQ	0.0143	MDL	5.00	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0783	JQ	0.0277	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.0682	J	0.0161	MDL	5.00	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA									
Method:	1613B	Matrix:			SO					

Sample ID: SL-580-SA8-SB-10.5-11.5 Collected: 7/12/2013 11:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,7,8-PECDD	0.198	J	0.0511	MDL	5.00	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.167	JB	0.0230	MDL	5.00	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0612	JBQ	0.0143	MDL	5.00	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.230	JQ	0.0213	MDL	5.00	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0524	JQ	0.0310	MDL	1.00	PQL	ng/Kg	J	Z
OCDD	2.19	JB	0.0426	MDL	10.0	PQL	ng/Kg	J	Z
OCDF	0.136	JQ	0.0437	MDL	10.0	PQL	ng/Kg	J	Z

Sample ID: SL-580-SA8-SB-4.0-5.0 Collected: 7/12/2013 10:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.560	JBQ	0.0272	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.0549	JBQ	0.0105	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0349	JBQ	0.0119	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0489	J	0.0257	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0745	JB	0.0137	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.332	JQ	0.0269	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0749	JBQ	0.0144	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.473	J	0.0250	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.705	J	0.0146	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0910	JQ	0.0397	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.223	JBQ	0.0208	MDL	5.29	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0519	JBQ	0.0135	MDL	5.29	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.137	JQ	0.0180	MDL	5.29	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0295	JQ	0.0275	MDL	1.06	PQL	ng/Kg	J	Z
OCDD	5.04	JB	0.0222	MDL	10.6	PQL	ng/Kg	J	Z
OCDF	0.195	JQ	0.0216	MDL	10.6	PQL	ng/Kg	J	Z

Sample ID: SL-581-SA8-SB-0.0-0.5 Collected: 7/12/2013 12:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.63	JB	0.0371	MDL	5.27	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.345	JB	0.0126	MDL	5.27	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0554	JB	0.0254	MDL	5.27	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0682	JQ	0.0309	MDL	5.27	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA								
Method:	1613B	Matrix:	SO						

Sample ID: SL-581-SA8-SB-0.0-0.5 Collected: 7/12/2013 12:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HXCDF	0.101	JBQ	0.0176	MDL	5.27	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.758	J	0.0335	MDL	5.27	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.120	JBQ	0.0155	MDL	5.27	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	1.02	J	0.0315	MDL	5.27	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	1.36	J	0.0210	MDL	5.27	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.147	JQ	0.0462	MDL	5.27	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.495	JB	0.0191	MDL	5.27	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.102	JBQ	0.0173	MDL	5.27	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.145	JQ	0.0210	MDL	5.27	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0438	J	0.0259	MDL	1.05	PQL	ng/Kg	J	Z
OCDF	0.720	J	0.0387	MDL	10.5	PQL	ng/Kg	J	Z

Sample ID: SL-581-SA8-SB-4.5-5.5 Collected: 7/12/2013 12:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0525	JB	0.0270	MDL	4.91	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0193	JBQ	0.00968	MDL	4.91	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0482	JBQ	0.0196	MDL	4.91	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0516	JBQ	0.0113	MDL	4.91	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0291	JB	0.0103	MDL	4.91	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0290	JQ	0.0219	MDL	4.91	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.0210	JQ	0.0157	MDL	4.91	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0469	JQ	0.0444	MDL	4.91	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0203	JBQ	0.0112	MDL	4.91	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0294	JQ	0.0174	MDL	4.91	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0283	JQ	0.0279	MDL	0.983	PQL	ng/Kg	J	Z
OCDD	0.180	JBQ	0.0314	MDL	9.83	PQL	ng/Kg	U	B
OCDF	0.0513	JQ	0.0384	MDL	9.83	PQL	ng/Kg	J	Z

Sample ID: SL-587-SA8-SB-0.0-0.5 Collected: 7/12/2013 1:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.88	JB	0.0447	MDL	4.89	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.209	JBQ	0.0123	MDL	4.89	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0386	JBQ	0.0215	MDL	4.89	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	1613B	Matrix:	SO
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Sample ID: SL-587-SA8-SB-0.0-0.5 Collected: 7/12/2013 1:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HXCDF	0.0163	JBQ	0.0156	MDL	4.89	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.352	JQ	0.0465	MDL	4.89	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0670	JB	0.0149	MDL	4.89	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.556	J	0.0456	MDL	4.89	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.374	JQ	0.0183	MDL	4.89	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0615	JQ	0.0553	MDL	4.89	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.114	JB	0.0229	MDL	4.89	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0344	JBQ	0.0151	MDL	4.89	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.0401	JQ	0.0389	MDL	0.978	PQL	ng/Kg	J	Z
OCDF	0.372	J	0.0355	MDL	9.78	PQL	ng/Kg	J	Z

Sample ID: SL-587-SA8-SB-6.5-7.5 Collected: 7/12/2013 1:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0629	JB	0.0277	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0255	JBQ	0.00734	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0177	JB	0.0147	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.00854	JBQ	0.00796	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0176	JB	0.00765	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0155	JQ	0.0116	MDL	4.98	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.0307	JBQ	0.0146	MDL	4.98	PQL	ng/Kg	U	B
OCDD	0.227	JBQ	0.0241	MDL	9.96	PQL	ng/Kg	U	B
OCDF	0.110	J	0.0365	MDL	9.96	PQL	ng/Kg	J	Z

Sample ID: SL-878-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.273	JBQ	0.0362	MDL	5.40	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0814	JBQ	0.0142	MDL	5.40	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0496	JBQ	0.0181	MDL	5.40	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0627	J	0.0405	MDL	5.40	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0716	JBQ	0.0193	MDL	5.40	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.430	J	0.0413	MDL	5.40	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.114	JBQ	0.0194	MDL	5.40	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.735	J	0.0418	MDL	5.40	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-878-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,7,8,9-HXCDF	1.06	J	0.0235	MDL	5.40	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.144	JQ	0.0658	MDL	5.40	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.248	JBQ	0.0257	MDL	5.40	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0605	JBQ	0.0193	MDL	5.40	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0415	JQ	0.0249	MDL	5.40	PQL	ng/Kg	J	Z
OCDD	1.52	JB	0.0406	MDL	10.8	PQL	ng/Kg	J	Z
OCDF	0.147	J	0.0349	MDL	10.8	PQL	ng/Kg	J	Z

Method Category:	SVOA	
Method:	8015M	Matrix: SO

Sample ID: SL-578-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:15:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	5.7		2.2	MDL	5.6	PQL	mg/Kg	J	Q, Q
EFH (C30-C40)	28		4.5	MDL	11	PQL	mg/Kg	J	Q

Sample ID: SL-578-SA8-SB-9.0-10.0 Collected: 7/12/2013 9:45:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	5.0	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z

Sample ID: SL-587-SA8-SB-0.0-0.5 Collected: 7/12/2013 1:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	2.8	J	2.0	MDL	5.1	PQL	mg/Kg	J	Z
EFH (C30-C40)	6.6	J	4.1	MDL	10	PQL	mg/Kg	J	Z

Sample ID: SL-587-SA8-SB-6.5-7.5 Collected: 7/12/2013 1:40:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C30-C40)	4.6	J	4.1	MDL	10	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8270D	Matrix: SO

Sample ID: SL-578-SA8-SB-0.0-0.5 Collected: 7/12/2013 8:35:00 Analysis Type: RES-ACID Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,6-DICHLOROPHENOL	36	U	18	MDL	36	PQL	ug/Kg	UJ	L

Sample ID: SL-578-SA8-SB-4.0-0.5 Collected: 7/12/2013 9:15:00 Analysis Type: RES-ACID Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,6-DICHLOROPHENOL	38	U	19	MDL	38	PQL	ug/Kg	UJ	L
BENZIDINE	1900	U	790	MDL	1900	PQL	ug/Kg	R	Q
BENZO(A)ANTHRACENE	8	J	4	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	9	J	4	MDL	19	PQL	ug/Kg	J	Z
FLUORANTHENE	8	J	4	MDL	19	PQL	ug/Kg	J	Z
PHENANTHRENE	5	J	4	MDL	19	PQL	ug/Kg	J	Z
PYRENE	11	J	4	MDL	19	PQL	ug/Kg	J	Z

Sample ID: SL-578-SA8-SB-9.0-10.0 Collected: 7/12/2013 9:45:00 Analysis Type: RES-ACID Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,6-DICHLOROPHENOL	35	U	18	MDL	35	PQL	ug/Kg	UJ	L

Sample ID: SL-580-SA8-SB-0.0-0.5 Collected: 7/12/2013 10:30:00 Analysis Type: RES-ACID Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,6-DICHLOROPHENOL	37	U	18	MDL	37	PQL	ug/Kg	UJ	L

Sample ID: SL-580-SA8-SB-10.5-11.5 Collected: 7/12/2013 11:00:00 Analysis Type: RES-ACID Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,6-DICHLOROPHENOL	35	U	17	MDL	35	PQL	ug/Kg	UJ	L

Sample ID: SL-580-SA8-SB-4.0-5.0 Collected: 7/12/2013 10:45:00 Analysis Type: RES-ACID Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,6-DICHLOROPHENOL	36	U	18	MDL	36	PQL	ug/Kg	UJ	L
FLUORANTHENE	7	J	4	MDL	18	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA									
Method:	8270D	Matrix:			SO					

Sample ID: SL-878-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:30:00 Analysis Type: RES-ACID Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,6-DICHLOROPHENOL	38	U	19	MDL	38	PQL	ug/Kg	UJ	L

Method Category:	SVOA									
Method:	8270D SIM	Matrix:			SO					

Sample ID: SL-578-SA8-SB-0.0-5.0 Collected: 7/12/2013 8:35:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.79	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	9.3	J	6.5	MDL	20	PQL	ug/Kg	J	Z
CHRYSENE	0.71	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-578-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	0.89	J	0.75	MDL	1.9	PQL	ug/Kg	J	Z
2-METHYLNAPHTHALENE	0.92	J	0.75	MDL	1.9	PQL	ug/Kg	J	Z
ANTHRACENE	0.77	J	0.38	MDL	1.9	PQL	ug/Kg	J	Z
BENZO(E)PYRENE	5.0	J	3.8	MDL	19	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	7.8	J	6.8	MDL	20	PQL	ug/Kg	J	Z
DIBENZO(A,H)ANTHRACENE	1.2	J	0.75	MDL	1.9	PQL	ug/Kg	J	Z
NAPHTHALENE	0.78	J	0.75	MDL	1.9	PQL	ug/Kg	U	F

Sample ID: SL-578-SA8-SB-9.0-10.0 Collected: 7/12/2013 9:45:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(A)ANTHRACENE	1.3	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	1.7	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	0.82	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	1.0	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	9.7	J	6.3	MDL	19	PQL	ug/Kg	J	Z
PHENANTHRENE	1.4	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	8270D SIM	Matrix:	SO
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Sample ID: SL-580-SA8-SB-0.0-0.5 Collected: 7/12/2013 10:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(G,H,I)PERYLENE	1.5	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z
Di-n-octylphthalate	14	J	6.6	MDL	20	PQL	ug/Kg	J	Z
INDENO(1,2,3-CD)PYRENE	1.0	J	0.74	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-580-SA8-SB-10.5-11.5 Collected: 7/12/2013 11:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTHRACENE	0.59	J	0.35	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(E)PYRENE	5.3	J	3.5	MDL	18	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	9.6	J	6.3	MDL	19	PQL	ug/Kg	J	Z
DIBENZO(A,H)ANTHRACENE	1.1	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z
PHENANTHRENE	1.5	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-580-SA8-SB-4.0-5.0 Collected: 7/12/2013 10:45:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTHRACENE	0.46	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(E)PYRENE	4.7	J	3.6	MDL	18	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	7.8	J	6.5	MDL	20	PQL	ug/Kg	J	Z
DIBENZO(A,H)ANTHRACENE	1.0	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-581-SA8-SB-0.0-0.5 Collected: 7/12/2013 12:40:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.84	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	0.96	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	7.3	J	6.4	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	1.3	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
Di-n-octylphthalate	13	J	6.4	MDL	19	PQL	ug/Kg	J	Z
FLUORANTHENE	0.85	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-581-SA8-SB-4.5-5.5 Collected: 7/12/2013 12:50:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	0.44	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-581-SA8-SB-4.5-5.5 Collected: 7/12/2013 12:50:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
FLUORANTHENE	0.83	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
PHENANTHRENE	0.80	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	0.78	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-878-SA8-SB-4.0-5.0 Collected: 7/12/2013 9:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(A)ANTHRACENE	0.88	J	0.75	MDL	1.9	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	1.1	J	0.75	MDL	1.9	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	1.1	J	0.75	MDL	1.9	PQL	ug/Kg	J	Z
FLUORANTHENE	1.8	J	0.75	MDL	1.9	PQL	ug/Kg	J	Z
NAPHTHALENE	0.78	J	0.75	MDL	1.9	PQL	ug/Kg	U	F
PHENANTHRENE	1.5	J	0.75	MDL	1.9	PQL	ug/Kg	J	Z
PYRENE	1.6	J	0.75	MDL	1.9	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH066
EDD Filename: PrepPH066_v1

Laboratory: LL
eQAPP Name: CDM_SSFL_130808_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
*#	Professional Judgment
A	ICP Serial Dilution
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Equipment Blank Contamination
F	Field Blank Contamination
L	Compound Not Spiked in QC
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH066

Method Blank Outlier Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK1980B371635	7/19/2013 4:35:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDF 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF OCDD	0.0588 ng/Kg 0.0270 ng/Kg 0.0381 ng/Kg 0.0344 ng/Kg 0.0260 ng/Kg 0.0697 ng/Kg 0.0292 ng/Kg 0.262 ng/Kg	SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-581-SA8-SB-0.0-0.5 SL-581-SA8-SB-4.5-5.5 SL-587-SA8-SB-0.0-0.5 SL-587-SA8-SB-6.5-7.5 SL-878-SA8-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-578-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0406 ng/Kg	0.0406U ng/Kg
SL-578-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0256 ng/Kg	0.0256U ng/Kg
SL-578-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0418 ng/Kg	0.0418U ng/Kg
SL-578-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.0515 ng/Kg	0.0515U ng/Kg
SL-578-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0272 ng/Kg	0.0272U ng/Kg
SL-578-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HXCDF	0.0311 ng/Kg	0.0311U ng/Kg
SL-578-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDF	0.0590 ng/Kg	0.0590U ng/Kg
SL-578-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.127 ng/Kg	0.127U ng/Kg
SL-578-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0870 ng/Kg	0.0870U ng/Kg
SL-578-SA8-SB-9.0-10.0(RES)	1,2,3,4,6,7,8-HPCDD	0.163 ng/Kg	0.163U ng/Kg
SL-578-SA8-SB-9.0-10.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0450 ng/Kg	0.0450U ng/Kg
SL-578-SA8-SB-9.0-10.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0234 ng/Kg	0.0234U ng/Kg
SL-578-SA8-SB-9.0-10.0(RES)	1,2,3,6,7,8-HXCDF	0.0298 ng/Kg	0.0298U ng/Kg
SL-578-SA8-SB-9.0-10.0(RES)	1,2,3,7,8-PECDF	0.0281 ng/Kg	0.0281U ng/Kg
SL-578-SA8-SB-9.0-10.0(RES)	OCDD	0.988 ng/Kg	0.988U ng/Kg
SL-580-SA8-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.123 ng/Kg	0.123U ng/Kg
SL-580-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0390 ng/Kg	0.0390U ng/Kg
SL-580-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.0781 ng/Kg	0.0781U ng/Kg
SL-580-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0200 ng/Kg	0.0200U ng/Kg
SL-580-SA8-SB-10.5-11.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0890 ng/Kg	0.0890U ng/Kg
SL-580-SA8-SB-10.5-11.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0340 ng/Kg	0.0340U ng/Kg
SL-580-SA8-SB-10.5-11.5(RES)	1,2,3,4,7,8-HXCDF	0.122 ng/Kg	0.122U ng/Kg
SL-580-SA8-SB-10.5-11.5(RES)	1,2,3,6,7,8-HXCDF	0.111 ng/Kg	0.111U ng/Kg
SL-580-SA8-SB-10.5-11.5(RES)	1,2,3,7,8-PECDF	0.167 ng/Kg	0.167U ng/Kg
SL-580-SA8-SB-10.5-11.5(RES)	2,3,4,6,7,8-HXCDF	0.0612 ng/Kg	0.0612U ng/Kg
SL-580-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0549 ng/Kg	0.0549U ng/Kg
SL-580-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0349 ng/Kg	0.0349U ng/Kg
SL-580-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HXCDF	0.0745 ng/Kg	0.0745U ng/Kg
SL-580-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDF	0.0749 ng/Kg	0.0749U ng/Kg
SL-580-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.223 ng/Kg	0.223U ng/Kg
SL-580-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0519 ng/Kg	0.0519U ng/Kg
SL-581-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0554 ng/Kg	0.0554U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-581-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.101 ng/Kg	0.101U ng/Kg
SL-581-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.120 ng/Kg	0.120U ng/Kg
SL-581-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.102 ng/Kg	0.102U ng/Kg
SL-581-SA8-SB-4.5-5.5(RES)	1,2,3,4,6,7,8-HPCDD	0.0525 ng/Kg	0.0525U ng/Kg
SL-581-SA8-SB-4.5-5.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0193 ng/Kg	0.0193U ng/Kg
SL-581-SA8-SB-4.5-5.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0482 ng/Kg	0.0482U ng/Kg
SL-581-SA8-SB-4.5-5.5(RES)	1,2,3,4,7,8-HXCDF	0.0516 ng/Kg	0.0516U ng/Kg
SL-581-SA8-SB-4.5-5.5(RES)	1,2,3,6,7,8-HXCDF	0.0291 ng/Kg	0.0291U ng/Kg
SL-581-SA8-SB-4.5-5.5(RES)	2,3,4,6,7,8-HXCDF	0.0203 ng/Kg	0.0203U ng/Kg
SL-581-SA8-SB-4.5-5.5(RES)	OCDD	0.180 ng/Kg	0.180U ng/Kg
SL-587-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0386 ng/Kg	0.0386U ng/Kg
SL-587-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0163 ng/Kg	0.0163U ng/Kg
SL-587-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0670 ng/Kg	0.0670U ng/Kg
SL-587-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.114 ng/Kg	0.114U ng/Kg
SL-587-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0344 ng/Kg	0.0344U ng/Kg
SL-587-SA8-SB-6.5-7.5(RES)	1,2,3,4,6,7,8-HPCDD	0.0629 ng/Kg	0.0629U ng/Kg
SL-587-SA8-SB-6.5-7.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0255 ng/Kg	0.0255U ng/Kg
SL-587-SA8-SB-6.5-7.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0177 ng/Kg	0.0177U ng/Kg
SL-587-SA8-SB-6.5-7.5(RES)	1,2,3,4,7,8-HXCDF	0.00854 ng/Kg	0.00854U ng/Kg
SL-587-SA8-SB-6.5-7.5(RES)	1,2,3,6,7,8-HXCDF	0.0176 ng/Kg	0.0176U ng/Kg
SL-587-SA8-SB-6.5-7.5(RES)	1,2,3,7,8-PECDF	0.0307 ng/Kg	0.0307U ng/Kg
SL-587-SA8-SB-6.5-7.5(RES)	OCDD	0.227 ng/Kg	0.227U ng/Kg
SL-878-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDD	0.273 ng/Kg	0.273U ng/Kg
SL-878-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0814 ng/Kg	0.0814U ng/Kg
SL-878-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0496 ng/Kg	0.0496U ng/Kg
SL-878-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HXCDF	0.0716 ng/Kg	0.0716U ng/Kg
SL-878-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDF	0.114 ng/Kg	0.114U ng/Kg
SL-878-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.248 ng/Kg	0.248U ng/Kg
SL-878-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0605 ng/Kg	0.0605U ng/Kg

Method Blank Outlier Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19937AB221015	7/24/2013 10:15:00 AM	CALCIUM TIN ZINC	6.97 mg/Kg 1.67 mg/Kg 0.455 mg/Kg	SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-581-SA8-SB-0.0-0.5 SL-581-SA8-SB-4.5-5.5 SL-587-SA8-SB-0.0-0.5 SL-587-SA8-SB-6.5-7.5 SL-878-SA8-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-578-SA8-SB-0.0-0.5(RES)	TIN	3.15 mg/Kg	3.15U mg/Kg
SL-578-SA8-SB-4.0-5.0(RES)	TIN	3.04 mg/Kg	3.04U mg/Kg
SL-578-SA8-SB-9.0-10.0(RES)	TIN	3.04 mg/Kg	3.04U mg/Kg
SL-580-SA8-SB-0.0-0.5(RES)	TIN	3.08 mg/Kg	3.08U mg/Kg
SL-580-SA8-SB-10.5-11.5(RES)	TIN	2.85 mg/Kg	2.85U mg/Kg
SL-580-SA8-SB-4.0-5.0(RES)	TIN	2.94 mg/Kg	2.94U mg/Kg
SL-581-SA8-SB-0.0-0.5(RES)	TIN	2.85 mg/Kg	2.85U mg/Kg
SL-581-SA8-SB-4.5-5.5(RES)	TIN	2.97 mg/Kg	2.97U mg/Kg
SL-587-SA8-SB-0.0-0.5(RES)	TIN	2.50 mg/Kg	2.50U mg/Kg
SL-587-SA8-SB-6.5-7.5(RES)	TIN	2.70 mg/Kg	2.70U mg/Kg
SL-878-SA8-SB-4.0-5.0(RES)	TIN	3.10 mg/Kg	3.10U mg/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(REA2)	4/11/2013 3:00:00 PM	COPPER MOLYBDENUM	0.0036 mg/L 0.0036 mg/L	SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-581-SA8-SB-0.0-0.5 SL-581-SA8-SB-4.5-5.5 SL-587-SA8-SB-0.0-0.5 SL-587-SA8-SB-6.5-7.5 SL-878-SA8-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-578-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.308 mg/Kg	0.308U mg/Kg
SL-578-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.416 mg/Kg	0.416U mg/Kg
SL-580-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.238 mg/Kg	0.238U mg/Kg
SL-580-SA8-SB-10.5-11.5(RES)	MOLYBDENUM	1.08 mg/Kg	1.08U mg/Kg
SL-580-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.332 mg/Kg	0.332U mg/Kg
SL-581-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.314 mg/Kg	0.314U mg/Kg
SL-581-SA8-SB-4.5-5.5(RES)	MOLYBDENUM	0.204 mg/Kg	0.204U mg/Kg
SL-587-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.453 mg/Kg	0.453U mg/Kg
SL-878-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.229 mg/Kg	0.229U mg/Kg

Method: 8270D SIM
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(RES)	4/11/2013 3:00:00 PM	1-METHYLNAPHTHALENE 2-METHYLNAPHTHALENE BIS(2-ETHYLHEXYL)PHTHALATE Diethylphthalate Di-n-butylphthalate NAPHTHALENE	0.019 ug/L 0.024 ug/L 0.082 ug/L 0.18 ug/L 0.17 ug/L 0.17 ug/L	SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-581-SA8-SB-0.0-0.5 SL-581-SA8-SB-4.5-5.5 SL-587-SA8-SB-0.0-0.5 SL-587-SA8-SB-6.5-7.5 SL-878-SA8-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-578-SA8-SB-4.0-5.0(RES)	NAPHTHALENE	0.78 ug/Kg	1.9U ug/Kg
SL-878-SA8-SB-4.0-5.0(RES)	NAPHTHALENE	0.78 ug/Kg	1.9U ug/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

9/9/2013 11:18:09 AM

ADR version 1.7.0.207

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Equipment Rinsate Blank Outlier Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Equipment Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
EB1-071013(REA2)	7/10/2013 3:00:00 PM	COPPER MOLYBDENUM	0.0031 mg/L 0.0035 mg/L	SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-581-SA8-SB-0.0-0.5 SL-581-SA8-SB-4.5-5.5 SL-587-SA8-SB-0.0-0.5 SL-587-SA8-SB-6.5-7.5 SL-878-SA8-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-578-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.308 mg/Kg	0.308U mg/Kg
SL-578-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.416 mg/Kg	0.416U mg/Kg
SL-580-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.238 mg/Kg	0.238U mg/Kg
SL-580-SA8-SB-10.5-11.5(RES)	MOLYBDENUM	1.08 mg/Kg	1.08U mg/Kg
SL-580-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.332 mg/Kg	0.332U mg/Kg
SL-581-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.314 mg/Kg	0.314U mg/Kg
SL-581-SA8-SB-4.5-5.5(RES)	MOLYBDENUM	0.204 mg/Kg	0.204U mg/Kg
SL-587-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.453 mg/Kg	0.453U mg/Kg
SL-878-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.229 mg/Kg	0.229U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8015M
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-578-SA8-SB-4.0-5.0MS SL-578-SA8-SB-4.0-5.0MSD (SL-578-SA8-SB-4.0-5.0)	EFH (C15-C20) EFH (C21-C30)	154 273	125 -	49.00-123.00 49.00-123.00	21 (20.00) 52 (20.00)	EFH (C15-C20) EFH (C21-C30)	J (all detects)
SL-578-SA8-SB-4.0-5.0MS SL-578-SA8-SB-4.0-5.0MSD (SL-578-SA8-SB-4.0-5.0)	EFH (C30-C40)	364	26	49.00-123.00	49 (20.00)	EFH (C30-C40)	J(all detects) No Qual %R, >4X

Method: 8270D SIM
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-578-SA8-SB-4.0-5.0MS SL-578-SA8-SB-4.0-5.0MSD (SL-578-SA8-SB-4.0-5.0)	Diethylphthalate Di-n-octylphthalate FLUORANTHENE PYRENE	- 172 - -	135 178 - -	76.00-127.00 52.00-162.00 47.00-135.00 26.00-143.00	- - 33 (30.00) 33 (30.00)	Diethylphthalate Di-n-octylphthalate FLUORANTHENE PYRENE	J(all detects)

Method: 8270D
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-578-SA8-SB-4.0-5.0MS SL-578-SA8-SB-4.0-5.0MSD (SL-578-SA8-SB-4.0-5.0)	BENZIDINE	0	0	35.00-141.00	-	BENZIDINE	J(all detects) R(all non-detects)

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-578-SA8-SB-4.0-5.0MS (TOT)	ALUMINUM	4971	6312	75.00-125.00	-	ALUMINUM	J(all detects) Al, Ca, Fe, Mg, Mn, K, Ti No Qual, >4X
SL-578-SA8-SB-4.0-5.0MSD (TOT)	CALCIUM	162	200	75.00-125.00	-	CALCIUM	
(SL-578-SA8-SB-0.0-0.5)	CHROMIUM	135	145	75.00-125.00	-	CHROMIUM	
SL-578-SA8-SB-4.0-5.0	IRON	3655	5430	75.00-125.00	-	IRON	
SL-578-SA8-SB-9.0-10.0	MAGNESIUM	556	709	75.00-125.00	-	MAGNESIUM	
SL-580-SA8-SB-0.0-0.5	MANGANESE	126	196	75.00-125.00	-	MANGANESE	
SL-580-SA8-SB-10.5-11.5	POTASSIUM	151	-	75.00-125.00	-	POTASSIUM	
SL-580-SA8-SB-4.0-5.0	TITANIUM	502	480	75.00-125.00	-	TITANIUM	
SL-581-SA8-SB-0.0-0.5	VANADIUM	128	136	75.00-125.00	-	VANADIUM	
SL-581-SA8-SB-4.5-5.5							
SL-587-SA8-SB-0.0-0.5							
SL-587-SA8-SB-6.5-7.5							
SL-878-SA8-SB-4.0-5.0)							

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-578-SA8-SB-4.0-5.0MS (TOT) SL-578-SA8-SB-4.0-5.0MSD (TOT) (SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-581-SA8-SB-0.0-0.5 SL-581-SA8-SB-4.5-5.5 SL-587-SA8-SB-0.0-0.5 SL-587-SA8-SB-6.5-7.5 SL-878-SA8-SB-4.0-5.0)	ANTIMONY	35	29	75.00-125.00	21 (20.00)	ANTIMONY	J(all detects) UJ(all non-detects) PS=102%
SL-578-SA8-SB-4.0-5.0MS (TOT) SL-578-SA8-SB-4.0-5.0MSD (TOT) (SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-581-SA8-SB-0.0-0.5 SL-581-SA8-SB-4.5-5.5 SL-587-SA8-SB-0.0-0.5 SL-587-SA8-SB-6.5-7.5 SL-878-SA8-SB-4.0-5.0)	PHOSPHORUS	67	49	75.00-125.00	-	PHOSPHORUS	J(all detects) UJ(all non-detects)

Method: 6020A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-578-SA8-SB-4.0-5.0MS (TOT) SL-578-SA8-SB-4.0-5.0MSD (TOT) (SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-581-SA8-SB-0.0-0.5 SL-581-SA8-SB-4.5-5.5 SL-587-SA8-SB-0.0-0.5 SL-587-SA8-SB-6.5-7.5 SL-878-SA8-SB-4.0-5.0)	SILVER STRONTIUM THALLIUM	129 - 145	137 132 170	75.00-125.00 75.00-125.00 75.00-125.00	- - -	SILVER STRONTIUM THALLIUM	J(all detects)

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH066
 EDD Filename: PH066_v1.

Laboratory: LL
 eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-578-SA8-SB-4.0-5.0MSD (TOT) (SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-581-SA8-SB-0.0-0.5 SL-581-SA8-SB-4.5-5.5 SL-587-SA8-SB-0.0-0.5 SL-587-SA8-SB-6.5-7.5 SL-878-SA8-SB-4.0-5.0)	SELENIUM	-	138	75.00-125.00	21 (20.00)	SELENIUM	J(all detects) UJ(all non-detects)

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-578-SA8-SB-4.0-5.0DUP (TOT)	MANGANESE	40	20.00	J (all detects) UJ (all non-detects) Mo, No Qual OK by difference
(SL-578-SA8-SB-0.0-0.5)	MOLYBDENUM	31	20.00	
SL -578-SA8-SB-4.0-5.0				
SL -578-SA8-SB-9.0-10.0				
SL -580-SA8-SB-0.0-0.5				
SL -580-SA8-SB-10.5-11.5				
SL -580-SA8-SB-4.0-5.0				
SL -581-SA8-SB-0.0-0.5				
SL -581-SA8-SB-4.5-5.5				
SL -587-SA8-SB-0.0-0.5				
SL -587-SA8-SB-6.5-7.5				
SL -878-SA8-SB-4.0-5.0)				

Method: 6020A
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-578-SA8-SB-4.0-5.0DUP (TOT)	SILVER	38	20.00	No Qual, OK by difference
(SL-578-SA8-SB-0.0-0.5)				
SL -578-SA8-SB-4.0-5.0				
SL -578-SA8-SB-9.0-10.0				
SL -580-SA8-SB-0.0-0.5				
SL -580-SA8-SB-10.5-11.5				
SL -580-SA8-SB-4.0-5.0				
SL -581-SA8-SB-0.0-0.5				
SL -581-SA8-SB-4.5-5.5				
SL -587-SA8-SB-0.0-0.5				
SL -587-SA8-SB-6.5-7.5				
SL -878-SA8-SB-4.0-5.0)				

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PrepPH066_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8270D SIM
Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P8LBLCSEQ260943 (SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-581-SA8-SB-0.0-0.5 SL-581-SA8-SB-4.5-5.5 SL-587-SA8-SB-0.0-0.5 SL-587-SA8-SB-6.5-7.5 SL-878-SA8-SB-4.0-5.0)	Diethylphthalate	130	-	68.00-125.00	-	Diethylphthalate	J (all detects)

Method: 8270D
Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P8LDLCSQ260031 (SL-578-SA8-SB-0.0-0.5 SL-578-SA8-SB-4.0-5.0 SL-578-SA8-SB-9.0-10.0 SL-580-SA8-SB-0.0-0.5 SL-580-SA8-SB-10.5-11.5 SL-580-SA8-SB-4.0-5.0 SL-878-SA8-SB-4.0-5.0)	1,2,4-TRICHLOROBENZENE 1,2-Diphenylhydrazine/Azobenzene	110 118	- -	73.00-108.00 77.00-111.00	- -	1,2,4-TRICHLOROBENZENE 1,2-Diphenylhydrazine/Azobenzene	J(all detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 160.3M
Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-578-SA8-SB-4.0-5.0	SL-878-SA8-SB-4.0-5.0			
MOISTURE	11.6	11.4	2		No Qualifiers Applied

Method: 1613B
Matrix: SO

Analyte	Concentration (ng/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-578-SA8-SB-4.0-5.0	SL-878-SA8-SB-4.0-5.0			
1,2,3,4,7,8-HxCDD	0.0838	0.0627	29	50.00	No Qualifiers Applied
1,2,3,6,7,8-HxCDD	0.520	0.430	19	50.00	
1,2,3,7,8,9-HxCDD	0.800	0.735	8	50.00	
1,2,3,7,8,9-HxCDF	0.901	1.06	16	50.00	
2,3,4,6,7,8-HxCDF	0.0870	0.0605	36	50.00	
1,2,3,4,6,7,8-HPCDD	1.41	0.273	135	50.00	J(all detects) UJ(all non-detects)
1,2,3,4,6,7,8-HPCDF	0.204	0.0814	86	50.00	
1,2,3,4,7,8,9-HPCDF	5.49 U	0.0496	200	50.00	
1,2,3,4,7,8-HxCDF	0.0311	0.0716	79	50.00	
1,2,3,6,7,8-HxCDF	0.0590	0.114	64	50.00	
1,2,3,7,8-PECDD	5.49 U	0.144	200	50.00	
1,2,3,7,8-PECDF	0.127	0.248	65	50.00	
2,3,4,7,8-PECDF	0.131	0.0415	104	50.00	
OCDD	13.2	1.52	159	50.00	
OCDF	0.414	0.147	95	50.00	

Method: 6010C
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-578-SA8-SB-4.0-5.0 (TOT)	SL-878-SA8-SB-4.0-5.0 (TOT)			
ALUMINUM	25300	31500	22	50.00	No Qualifiers Applied
ARSENIC	5.71	6.49	13	50.00	
BARIUM	116	135	15	50.00	
BERYLLIUM	0.830	1.02	21	50.00	
BORON	2.75	3.04	10	50.00	
CADMIUM	0.533	0.408	27	50.00	
CALCIUM	5110	5050	1	50.00	
CHROMIUM	29.7	36.2	20	50.00	
COBALT	8.52	10.1	17	50.00	
COPPER	16.9	17.7	5	50.00	
IRON	27500	32600	17	50.00	
LEAD	8.58	8.91	4	50.00	
LITHIUM	20.5	24.3	17	50.00	
MAGNESIUM	5560	6350	13	50.00	
MANGANESE	373	524	34	50.00	
NICKEL	17.9	20.8	15	50.00	
PHOSPHORUS	286	242	17	50.00	
POTASSIUM	5200	5300	2	50.00	
SODIUM	214	207	3	50.00	
TIN	3.04	3.10	2	50.00	
TITANIUM	1210	1340	10	50.00	
VANADIUM	55.0	66.8	19	50.00	
ZINC	65.4	70.2	7	50.00	
Zirconium	7.25	7.25	0	50.00	
MOLYBDENUM	0.416	0.229	58	50.00	J(all detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-578-SA8-SB-4.0-5.0 (TOT)	SL-878-SA8-SB-4.0-5.0 (TOT)			
SELENIUM	0.277	0.326	16	50.00	No Qualifiers Applied
SILVER	0.0557	0.0645	15	50.00	
STRONTIUM	33.4	34.1	2	50.00	
THALLIUM	0.402	0.588	38	50.00	

Method: 8015M

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-578-SA8-SB-4.0-5.0	SL-878-SA8-SB-4.0-5.0			
EFH (C21-C30)	5.7	5.6	2	50.00	No Qualifiers Applied
EFH (C30-C40)	28	18	43	50.00	

Method: 8270D SIM

Matrix: SO

Analyte	Concentration (ug/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-578-SA8-SB-4.0-5.0	SL-878-SA8-SB-4.0-5.0			
NAPHTHALENE	0.78	0.78	0	50.00	No Qualifiers Applied
1-METHYLNAPHTHALENE	0.89	1.9 U	200	50.00	J(all detects) UJ(all non-detects)
2-METHYLNAPHTHALENE	0.92	1.9 U	200	50.00	
ANTHRACENE	0.77	1.9 U	200	50.00	
BENZO(A)PYRENE	6.3	1.1	141	50.00	
BENZO(B)FLUORANTHENE	9.9	2.4	122	50.00	
BENZO(E)PYRENE	5.0	19 U	200	50.00	
BENZO(G,H,I)PERYLENE	3.1	1.9 U	200	50.00	
BENZO(K)FLUORANTHENE	4.6	1.1	123	50.00	
BIS(2-ETHYLHEXYL)PHTHALATE	7.8	20 U	200	50.00	
DIBENZO(A,H)ANTHRACENE	1.2	1.9 U	200	50.00	
INDENO(1,2,3-CD)PYRENE	2.2	1.9 U	200	50.00	

Method: 9045M

Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-578-SA8-SB-4.0-5.0	SL-878-SA8-SB-4.0-5.0			
PH	6.12	5.91	3	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-578-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	1.41	5.30	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.194	5.30	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0406	5.30	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.0256	5.30	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.0938	5.30	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0418	5.30	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	J	0.131	5.30	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.168	5.30	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0515	5.30	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0272	5.30	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0735	5.30	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0326	1.06	PQL	ng/Kg	
	OCDF	J	0.478	10.6	PQL	ng/Kg	
	SL-578-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	1.41	5.49	PQL	
1,2,3,4,6,7,8-HPCDF		JBQ	0.204	5.49	PQL	ng/Kg	
1,2,3,4,7,8-HxCDD		J	0.0838	5.49	PQL	ng/Kg	
1,2,3,4,7,8-HXCDF		JBQ	0.0311	5.49	PQL	ng/Kg	
1,2,3,6,7,8-HXCDD		JQ	0.520	5.49	PQL	ng/Kg	
1,2,3,6,7,8-HXCDF		JBQ	0.0590	5.49	PQL	ng/Kg	
1,2,3,7,8,9-HXCDD		J	0.800	5.49	PQL	ng/Kg	
1,2,3,7,8,9-HXCDF		J	0.901	5.49	PQL	ng/Kg	
1,2,3,7,8-PECDF		JBQ	0.127	5.49	PQL	ng/Kg	
2,3,4,6,7,8-HXCDF		JB	0.0870	5.49	PQL	ng/Kg	
2,3,4,7,8-PECDF		JQ	0.131	5.49	PQL	ng/Kg	
OCDF		J	0.414	11.0	PQL	ng/Kg	
SL-578-SA8-SB-9.0-10.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.163	5.05	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0450	5.05	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0234	5.05	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.0298	5.05	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JQ	0.0353	5.05	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.0288	5.05	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.0281	5.05	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0155	5.05	PQL	ng/Kg	
	OCDD	JBQ	0.988	10.1	PQL	ng/Kg	
	OCDF	JQ	0.0542	10.1	PQL	ng/Kg	
SL-580-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	2.09	5.43	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.123	5.43	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.100	5.43	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0390	5.43	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JQ	0.144	5.43	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.216	5.43	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0781	5.43	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0200	5.43	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0531	5.43	PQL	ng/Kg	
	OCDF	JQ	0.300	10.9	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-580-SA8-SB-10.5-11.5	1,2,3,4,6,7,8-HPCDD	JB	0.316	5.00	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0890	5.00	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0340	5.00	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0653	5.00	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.122	5.00	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JQ	0.115	5.00	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.111	5.00	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JQ	0.0783	5.00	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.0682	5.00	PQL	ng/Kg	
	1,2,3,7,8-PECDD	J	0.198	5.00	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.167	5.00	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0612	5.00	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.230	5.00	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0524	1.00	PQL	ng/Kg	
	OCDD	JB	2.19	10.0	PQL	ng/Kg	
OCDF	JQ	0.136	10.0	PQL	ng/Kg		
SL-580-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.560	5.29	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0549	5.29	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0349	5.29	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	J	0.0489	5.29	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.0745	5.29	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JQ	0.332	5.29	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0749	5.29	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	J	0.473	5.29	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.705	5.29	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0910	5.29	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.223	5.29	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0519	5.29	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.137	5.29	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0295	1.06	PQL	ng/Kg	
	OCDD	JB	5.04	10.6	PQL	ng/Kg	
OCDF	JQ	0.195	10.6	PQL	ng/Kg		
SL-581-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	1.63	5.27	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.345	5.27	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0554	5.27	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0682	5.27	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.101	5.27	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	J	0.758	5.27	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.120	5.27	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	J	1.02	5.27	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	1.36	5.27	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.147	5.27	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.495	5.27	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.102	5.27	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.145	5.27	PQL	ng/Kg	
	2,3,7,8-TCDD	J	0.0438	1.05	PQL	ng/Kg	
	OCDF	J	0.720	10.5	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-581-SA8-SB-4.5-5.5	1,2,3,4,6,7,8-HPCDD	JB	0.0525	4.91	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0193	4.91	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0482	4.91	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0516	4.91	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.0291	4.91	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JQ	0.0290	4.91	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.0210	4.91	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0469	4.91	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0203	4.91	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0294	4.91	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0283	0.983	PQL	ng/Kg	
	OCDD	JBQ	0.180	9.83	PQL	ng/Kg	
	OCDF	JQ	0.0513	9.83	PQL	ng/Kg	
SL-587-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	1.88	4.89	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.209	4.89	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0386	4.89	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0163	4.89	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.352	4.89	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.0670	4.89	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	J	0.556	4.89	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.374	4.89	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0615	4.89	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.114	4.89	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0344	4.89	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0401	0.978	PQL	ng/Kg	
	OCDF	J	0.372	9.78	PQL	ng/Kg	
SL-587-SA8-SB-6.5-7.5	1,2,3,4,6,7,8-HPCDD	JB	0.0629	4.98	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0255	4.98	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JB	0.0177	4.98	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.00854	4.98	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.0176	4.98	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.0155	4.98	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0307	4.98	PQL	ng/Kg	
	OCDD	JBQ	0.227	9.96	PQL	ng/Kg	
	OCDF	J	0.110	9.96	PQL	ng/Kg	
	SL-878-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.273	5.40	PQL	
1,2,3,4,6,7,8-HPCDF		JBQ	0.0814	5.40	PQL	ng/Kg	
1,2,3,4,7,8,9-HPCDF		JBQ	0.0496	5.40	PQL	ng/Kg	
1,2,3,4,7,8-HxCDD		J	0.0627	5.40	PQL	ng/Kg	
1,2,3,4,7,8-HXCDF		JBQ	0.0716	5.40	PQL	ng/Kg	
1,2,3,6,7,8-HXCDD		J	0.430	5.40	PQL	ng/Kg	
1,2,3,6,7,8-HXCDF		JBQ	0.114	5.40	PQL	ng/Kg	
1,2,3,7,8,9-HXCDD		J	0.735	5.40	PQL	ng/Kg	
1,2,3,7,8,9-HXCDF		J	1.06	5.40	PQL	ng/Kg	
1,2,3,7,8-PECDD		JQ	0.144	5.40	PQL	ng/Kg	
1,2,3,7,8-PECDF		JBQ	0.248	5.40	PQL	ng/Kg	
2,3,4,6,7,8-HXCDF		JBQ	0.0605	5.40	PQL	ng/Kg	
2,3,4,7,8-PECDF		JQ	0.0415	5.40	PQL	ng/Kg	
OCDD		JB	1.52	10.8	PQL	ng/Kg	
OCDF		J	0.147	10.8	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-578-SA8-SB-0.0-0.5	BERYLLIUM	J	0.735	1.06	PQL	mg/Kg	J (all detects)
	BORON	J	1.97	10.6	PQL	mg/Kg	
	CADMIUM	J	0.371	1.06	PQL	mg/Kg	
	MOLYBDENUM	J	0.308	2.13	PQL	mg/Kg	
	TIN	J	3.15	10.6	PQL	mg/Kg	
SL-578-SA8-SB-4.0-5.0	BERYLLIUM	J	0.830	1.11	PQL	mg/Kg	J (all detects)
	BORON	J	2.75	11.1	PQL	mg/Kg	
	CADMIUM	J	0.533	1.11	PQL	mg/Kg	
	MOLYBDENUM	J	0.416	2.22	PQL	mg/Kg	
	TIN	J	3.04	11.1	PQL	mg/Kg	
SL-578-SA8-SB-9.0-10.0	BERYLLIUM	J	0.440	1.05	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.351	1.05	PQL	mg/Kg	
	TIN	J	3.04	10.5	PQL	mg/Kg	
	Zirconium	J	3.63	5.23	PQL	mg/Kg	
SL-580-SA8-SB-0.0-0.5	BERYLLIUM	J	0.754	1.09	PQL	mg/Kg	J (all detects)
	BORON	J	2.49	10.9	PQL	mg/Kg	
	CADMIUM	J	0.303	1.09	PQL	mg/Kg	
	MOLYBDENUM	J	0.238	2.19	PQL	mg/Kg	
	SODIUM	J	99.2	109	PQL	mg/Kg	
	TIN	J	3.08	10.9	PQL	mg/Kg	
	Zirconium	J	5.13	5.46	PQL	mg/Kg	
SL-580-SA8-SB-10.5-11.5	BERYLLIUM	J	0.609	1.04	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.321	1.04	PQL	mg/Kg	
	MOLYBDENUM	J	1.08	2.09	PQL	mg/Kg	
	TIN	J	2.85	10.4	PQL	mg/Kg	
	Zirconium	J	4.17	5.21	PQL	mg/Kg	
SL-580-SA8-SB-4.0-5.0	BERYLLIUM	J	0.769	1.05	PQL	mg/Kg	J (all detects)
	BORON	J	4.29	10.5	PQL	mg/Kg	
	CADMIUM	J	0.449	1.05	PQL	mg/Kg	
	MOLYBDENUM	J	0.332	2.10	PQL	mg/Kg	
	TIN	J	2.94	10.5	PQL	mg/Kg	
SL-581-SA8-SB-0.0-0.5	BERYLLIUM	J	0.810	1.04	PQL	mg/Kg	J (all detects)
	BORON	J	2.82	10.4	PQL	mg/Kg	
	CADMIUM	J	0.496	1.04	PQL	mg/Kg	
	MOLYBDENUM	J	0.314	2.08	PQL	mg/Kg	
	SODIUM	J	77.6	104	PQL	mg/Kg	
	TIN	J	2.85	10.4	PQL	mg/Kg	
SL-581-SA8-SB-4.5-5.5	BERYLLIUM	J	0.418	0.981	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.294	0.981	PQL	mg/Kg	
	MOLYBDENUM	J	0.204	1.96	PQL	mg/Kg	
	SODIUM	J	86.7	98.1	PQL	mg/Kg	
	TIN	J	2.97	9.81	PQL	mg/Kg	
	Zirconium	J	3.05	4.90	PQL	mg/Kg	
SL-587-SA8-SB-0.0-0.5	ARSENIC	J	3.51	3.88	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.542	0.969	PQL	mg/Kg	
	BORON	J	0.814	9.69	PQL	mg/Kg	
	CADMIUM	J	0.360	0.969	PQL	mg/Kg	
	MOLYBDENUM	J	0.453	1.94	PQL	mg/Kg	
	SODIUM	J	67.7	96.9	PQL	mg/Kg	
	TIN	J	2.50	9.69	PQL	mg/Kg	
	Zirconium	J	3.20	4.84	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-587-SA8-SB-6.5-7.5	ARSENIC	J	3.25	3.98	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.483	0.994	PQL	mg/Kg	
	CADMIUM	J	0.203	0.994	PQL	mg/Kg	
	TIN	J	2.70	9.94	PQL	mg/Kg	
	Zirconium	J	2.46	4.97	PQL	mg/Kg	
SL-878-SA8-SB-4.0-5.0	BERYLLIUM	J	1.02	1.07	PQL	mg/Kg	J (all detects)
	BORON	J	3.04	10.7	PQL	mg/Kg	
	CADMIUM	J	0.408	1.07	PQL	mg/Kg	
	MOLYBDENUM	J	0.229	2.15	PQL	mg/Kg	
	TIN	J	3.10	10.7	PQL	mg/Kg	

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-578-SA8-SB-0.0-0.5	SILVER	J	0.0890	0.213	PQL	mg/Kg	J (all detects)
SL-578-SA8-SB-4.0-5.0	SELENIUM	J	0.277	0.444	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0557	0.222	PQL	mg/Kg	
SL-578-SA8-SB-9.0-10.0	SELENIUM	J	0.146	0.418	PQL	mg/Kg	J (all detects)
SL-580-SA8-SB-0.0-0.5	SELENIUM	J	0.123	0.437	PQL	mg/Kg	J (all detects)
SL-580-SA8-SB-10.5-11.5	SELENIUM	J	0.105	0.417	PQL	mg/Kg	J (all detects)
SL-580-SA8-SB-4.0-5.0	SELENIUM	J	0.247	0.421	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0404	0.210	PQL	mg/Kg	
SL-581-SA8-SB-0.0-0.5	SELENIUM	J	0.207	0.416	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0503	0.208	PQL	mg/Kg	
SL-587-SA8-SB-0.0-0.5	SELENIUM	J	0.107	0.388	PQL	mg/Kg	J (all detects)
SL-587-SA8-SB-6.5-7.5	SILVER	J	0.0650	0.199	PQL	mg/Kg	J (all detects)
SL-878-SA8-SB-4.0-5.0	SELENIUM	J	0.326	0.430	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0645	0.215	PQL	mg/Kg	

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-578-SA8-SB-9.0-10.0	EFH (C21-C30)	J	5.0	5.2	PQL	mg/Kg	J (all detects)
SL-587-SA8-SB-0.0-0.5	EFH (C21-C30)	J	2.8	5.1	PQL	mg/Kg	J (all detects)
	EFH (C30-C40)	J	6.6	10	PQL	mg/Kg	
SL-587-SA8-SB-6.5-7.5	EFH (C30-C40)	J	4.6	10	PQL	mg/Kg	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8270D
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-578-SA8-SB-4.0-5.0	BENZO(A)ANTHRACENE	J	8	19	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	9	19	PQL	ug/Kg	
	FLUORANTHENE	J	8	19	PQL	ug/Kg	
	PHENANTHRENE	J	5	19	PQL	ug/Kg	
	PYRENE	J	11	19	PQL	ug/Kg	
SL-580-SA8-SB-4.0-5.0	FLUORANTHENE	J	7	18	PQL	ug/Kg	J (all detects)

Method: 8270D SIM
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-578-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	0.79	1.8	PQL	ug/Kg	J (all detects)
	BIS(2-ETHYLHEXYL)PHTHALATE	J	9.3	20	PQL	ug/Kg	
	CHRYSENE	J	0.71	1.8	PQL	ug/Kg	
SL-578-SA8-SB-4.0-5.0	1-METHYLNAPHTHALENE	J	0.89	1.9	PQL	ug/Kg	J (all detects)
	2-METHYLNAPHTHALENE	J	0.92	1.9	PQL	ug/Kg	
	ANTHRACENE	J	0.77	1.9	PQL	ug/Kg	
	BENZO(E)PYRENE	J	5.0	19	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	7.8	20	PQL	ug/Kg	
	DIBENZO(A,H)ANTHRACENE	J	1.2	1.9	PQL	ug/Kg	
	NAPHTHALENE	J	0.78	1.9	PQL	ug/Kg	
SL-578-SA8-SB-9.0-10.0	BENZO(A)ANTHRACENE	J	1.3	1.8	PQL	ug/Kg	J (all detects)
	BENZO(A)PYRENE	J	1.7	1.8	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	0.82	1.8	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	1.0	1.8	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	9.7	19	PQL	ug/Kg	
	PHENANTHRENE	J	1.4	1.8	PQL	ug/Kg	
SL-580-SA8-SB-0.0-0.5	BENZO(G,H,I)PERYLENE	J	1.5	1.8	PQL	ug/Kg	J (all detects)
	Di-n-octylphthalate	J	14	20	PQL	ug/Kg	
	INDENO(1,2,3-CD)PYRENE	J	1.0	1.8	PQL	ug/Kg	
SL-580-SA8-SB-10.5-11.5	ANTHRACENE	J	0.59	1.7	PQL	ug/Kg	J (all detects)
	BENZO(E)PYRENE	J	5.3	18	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	9.6	19	PQL	ug/Kg	
	DIBENZO(A,H)ANTHRACENE	J	1.1	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	1.5	1.7	PQL	ug/Kg	
SL-580-SA8-SB-4.0-5.0	ANTHRACENE	J	0.46	1.8	PQL	ug/Kg	J (all detects)
	BENZO(E)PYRENE	J	4.7	18	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	7.8	20	PQL	ug/Kg	
	DIBENZO(A,H)ANTHRACENE	J	1.0	1.8	PQL	ug/Kg	
SL-581-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	0.84	1.8	PQL	ug/Kg	J (all detects)
	BENZO(K)FLUORANTHENE	J	0.96	1.8	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	7.3	19	PQL	ug/Kg	
	CHRYSENE	J	1.3	1.8	PQL	ug/Kg	
	Di-n-octylphthalate	J	13	19	PQL	ug/Kg	
	FLUORANTHENE	J	0.85	1.8	PQL	ug/Kg	
SL-581-SA8-SB-4.5-5.5	CHRYSENE	J	0.44	1.7	PQL	ug/Kg	J (all detects)
	FLUORANTHENE	J	0.83	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	0.80	1.7	PQL	ug/Kg	
	PYRENE	J	0.78	1.7	PQL	ug/Kg	

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

9/9/2013 11:18:30 AM

ADR version 1.7.0.207

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Reporting Limit Outliers

Lab Reporting Batch ID: PH066

Laboratory: LL

EDD Filename: PH066_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8270D SIM
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-878-SA8-SB-4.0-5.0	BENZO(A)ANTHRACENE	J	0.88	1.9	PQL	ug/Kg	J (all detects)
	BENZO(A)PYRENE	J	1.1	1.9	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	1.1	1.9	PQL	ug/Kg	
	FLUORANTHENE	J	1.8	1.9	PQL	ug/Kg	
	NAPHTHALENE	J	0.78	1.9	PQL	ug/Kg	
	PHENANTHRENE	J	1.5	1.9	PQL	ug/Kg	
	PYRENE	J	1.6	1.9	PQL	ug/Kg	

LDC #: 30289A4

VALIDATION COMPLETENESS WORKSHEET

Date: 9/11/13

SDG #: PH066

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: CR

2nd Reviewer: W

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	-	Sampling dates: 7/12/13
II.	ICP/MS Tune	-	
III.	Calibration	-	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	SW MSD	
VII.	Duplicate Sample Analysis	SW DUP	
VIII.	Laboratory Control Samples (LCS)	A LCS	
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	SW	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	-	(2,3)
XV.	Field Blanks	SW	EB = EB1-071013 FB = FB-041113

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

(PH064)
D = Duplicate
TB = Trip blank
EB = Equipment blank

(PH029)

Validated Samples:

Soil

1	SL-578-SA8-SB-0.0-0.5	11	SL-587-SA8-SB-6.5-7.5	21		31	
2	SL-578-SA8-SB-4.0-5.0	12	SL-578-SA8-SB-4.0-5.0MS	22		32	
3	SL-878-SA8-SB-4.0-5.0	13	SL-578-SA8-SB-4.0-5.0MSD	23		33	
4	SL-578-SA8-SB-9.0-10.0	14	SL-578-SA8-SB-4.0-5.0DUP	24		34	
5	SL-580-SA8-SB-0.0-0.5	15		25		35	
6	SL-580-SA8-SB-4.0-5.0	16		26		36	
7	SL-580-SA8-SB-10.5-11.5	17		27		37	
8	SL-581-SA8-SB-0.0-0.5	18		28		38	
9	SL-581-SA8-SB-4.5-5.5	19		29		39	
10	SL-587-SA8-SB-0.0-0.5	20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 **Soil factor applied:** 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ **Associated Samples:** Soil

Analyte	Blank ID	Sample Identification										
	FB-041113 (SDG: PH029)	Action Limit	1	2	3	5	6	7	8	9	10	
Cu	0.0036	1.8										
Mo	0.0036	1.8	0.31	0.42	0.23	0.24	0.33	1.1	0.31	0.20	0.45	

Sampling date: 7/10/13 **Soil factor applied:** 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ **Associated Samples:** Soil

Analyte	Blank ID	Sample Identification										
	EB1-071013 (SDG: PH064)	Action Limit	See FB									
Cu	0.0031	1.55										
Mo	0.0035	1.75										

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:

Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

2H no qual based on %R

PS = 102% (J/US/A)

Background Lab Sample ID: 7126351BKG Matrix Spike Lab Sample ID: 7126352MS Matrix Spike Duplicate Lab Sample ID: 7126353MSD
Batch Id(s): P19937A, P20338A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Control Limit		M	
		Result	C	Result	C	Result	C				%R	Q	%R	Q		%R	RPD		
Aluminum		22338.0059		32279.6970		34836.1218		200.0000	198.0198	MG/KG	4971		6312		8		20	P	
Antimony		0.7255	U	17.6230		14.2238		50.0000	49.5050	MG/KG	35	N	29	N	21	*	75 - 125	20	P
Arsenic		5.0451		20.3780		20.4663		15.0000	14.8515	MG/KG	102		104		0		75 - 125	20	P
Barium		102.4892		325.1630		308.0812		200.0000	198.0198	MG/KG	111		104		5		75 - 125	20	P
Beryllium		0.7333	B	5.9440		5.9584		5.0000	4.9505	MG/KG	104		106		0		75 - 125	20	P
Boron		2.4324	B	194.7070		192.7089		200.0000	198.0198	MG/KG	96		96		1		75 - 125	20	P
Cadmium		0.4716	B	5.0390		4.9396		5.0000	4.9505	MG/KG	91		90		2		75 - 125	20	P
Calcium		4514.8951		5164.8580		5307.4376		400.0000	396.0396	MG/KG	162		200		3				P
Chromium		26.2716		53.2480		54.9436		20.0000	19.8020	MG/KG	135	N	145	N	3		75 - 125	20	P
Cobalt		7.5324		54.3460		53.6950		50.0000	49.5050	MG/KG	94		93		1		75 - 125	20	P
Copper		14.9814		40.6750		39.4020		25.0000	24.7525	MG/KG	103		99		3		75 - 125	20	P
Iron		24293.8755		27948.8520		29669.9723		100.0000	99.0099	MG/KG	3655		5430		6				P
Lead		7.5814		22.1480		21.5525		15.0000	14.8515	MG/KG	97		94		3		75 - 125	20	P
Lithium		18.1314		122.1080		122.4208		100.0000	99.0099	MG/KG	104		105		0		75 - 125	20	P
Magnesium		4918.2892		6030.9930		6323.1901		200.0000	198.0198	MG/KG	556		709		5				P
Manganese		330.0039		392.9510		426.8941		50.0000	49.5050	MG/KG	126		196		8				P
Mercury		0.0095	U	0.1863		0.1850		0.1609	0.1567	MG/KG	116		118		1		65 - 135	20	CV
Molybdenum		0.3676	B	183.9710		177.0139		200.0000	198.0198	MG/KG	92		89		4		75 - 125	20	P
Nickel		15.7833		64.4710		64.4307		50.0000	49.5050	MG/KG	97		98		0		75 - 125	20	P
Phosphorus		252.6676		319.9870		301.5416		100.0000	99.0099	MG/KG	67	N	49	N	6		75 - 125	20	P
Potassium		4596.2324		6110.8490		5619.6911		1000.0000	990.0990	MG/KG	151		103		8				P
Selenium	78	0.2451	B	2.4060		2.9842		2.0000	1.9802	MG/KG	108		138	N	21	*	75 - 125	20	MS
Silver	107	0.0492	B	12.9880		13.6594		10.0000	9.9010	MG/KG	129	N	137	N	5		75 - 125	20	MS
Sodium		188.8618		1130.6120		1132.9554		1000.0000	990.0990	MG/KG	94		95		0		75 - 125	20	P
Strontium	88	29.5490		39.3800		40.0198		8.0000	7.9208	MG/KG	123		132	N	2		75 - 125	20	MS
Thallium	203	0.3551		0.9366		1.0297		0.4000	0.3960	MG/KG	145	N	170	N	9		75 - 125	20	MS
Tin		2.6853	B	352.8890		340.0842		400.0000	396.0396	MG/KG	88		85		4		75 - 125	20	P
Titanium		1066.2647		1567.8130		1544.6911		100.0000	99.0099	MG/KG	502		480		2				P
Vanadium		48.6137		112.4480		116.0386		50.0000	49.5050	MG/KG	128	N	136	N	3		75 - 125	20	P
Zinc		57.8284		112.6700		112.0010		50.0000	49.5050	MG/KG	110		109		1		75 - 125	20	P
Zirconium		6.4069		103.5870		99.7673		100.0000	99.0099	MG/KG	97		94		4		75 - 125	20	P

Note: Results shown are reported on an as-received basis.

<p>METHODS: P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>		<p>CONCENTRATION QUALIFIERS: U= Below MDL, B= Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS</p>	
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Background Lab Sample ID: 7126351BKG
 Batch ID(s): P19937A, P20338A
 Concentration Units: MG/KG

Duplicate Lab Sample ID: 7126354DUP

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			22338.0059		26275.4370		16		P
Antimony			1.3814	B	1.8020	B	-26		P
Arsenic		3.9	5.0451		5.6130		11		P
Barium			102.4892		124.9910		20		P
Beryllium			0.7333	B	0.8620	B	16		P
Boron			2.4324	B	2.6440	B	8		P
Cadmium			0.4716	B	0.4420	B	6		P
Calcium			4514.8951		4808.4500		6		P
Chromium			26.2716		30.9410		16		P
Cobalt			7.5324		9.0150		18		P
Copper			14.9814		15.9390		6		P
Iron			24293.8755		27804.0160		13		P
Lead		2.9	7.5814		7.8350		3		P
Lithium		3.9	18.1314		21.0910		15		P
Magnesium			4918.2892		5526.7920		12		P
Manganese			330.0039		496.5370		40	*	P
Mercury			0.0095	U	0.0096	U			CV
Molybdenum			0.3676	B	0.2680	B	31		P
Nickel			15.7833		18.6740		17		P
Phosphorus			252.6676		224.3600		12		P
Potassium			4596.2324		4491.3120		2		P
Selenium	78		0.2451	B	0.2450	B	0		MS
Silver	107		0.0492	B	0.0723	B	38		MS
Sodium		98.0	188.8618		196.3800		4		P
Strontium	88		29.5490		34.3400		15		MS
Thallium	203	0.2	0.3551		0.3626		2		MS
Tin			2.6853	B	2.8280	B	5		P
Titanium			1066.2647		1151.3980		8		P
Vanadium			48.6137		57.8000		17		P
Zinc			57.8284		62.5160		8		P
Zirconium		4.9	6.4069		7.4830		15		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).
 The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

OK based on control

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: = Duplicate Out of Spec
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Background Lab Sample ID: 7126351BKG Serial Dilution Lab Sample ID: 7126351L
 Batch ID(s): P19937A
 Concentration Units: UG/L

Analyte	Mass	Initial Sample Result (I)	C	Serial Dilution Result (S)	C	% Diff.	Q	M
Aluminum		227847.6600		240455.5000		6		P
Antimony		7.4000	U	37.0000	U			P
Arsenic		51.4600		35.0000	U	100		P
Barium		1045.3900		1156.6000		11		P
Beryllium		7.4800	B	8.1500	B	9		P
Boron		24.8100	B	66.8000	B	169		P
Cadmium		4.8100	B	5.4500	B	13		P
Calcium		46051.9300		48618.6000		6		P
Chromium		267.9700		295.5000		10		P
Cobalt		76.8300		82.3000		7		P
Copper		152.8100		189.6000		24		P
Iron		247797.5300		242794.2000		2		P
Lead		77.3300		83.9000	B	8		P
Lithium		184.9400		199.3000	B	8		P
Magnesium		50166.5500		52707.3000		5		P
Manganese		3366.0400		3736.4500		11		P
Molybdenum		3.7500	B	11.4500	B	205		P
Nickel		160.9900		174.9500		9		P
Phosphorus		2577.2100		2541.9500		1		P
Potassium		46881.5700		49311.8000		5		P
Selenium	78	1.2500	B	2.5000	U	100		MS
Silver	107	0.2511	B	0.6500	U	100		MS
Sodium		1926.3900		1946.4500	B	1		P
Strontium	88	150.7000		153.8000		2		MS
Thallium	203	1.8110		1.6135	B	11		MS
Tin		27.3900	B	32.6500	B	19		P
Titanium		10875.9000		11653.3500		7		P
Vanadium		495.8600		535.5500		8		P
Zinc		589.8500		605.1500		3		P
Zirconium		65.3500		94.7500	B	45		P

NOTE: An E in column Q indicates the presence of a chemical or physical interference in the matrix when the % difference is greater than 10%. This applies only when (I) is greater than or equal to 50x MDL for ICP, 100x MDL for ICP-MS (6020), 50x MDL for ICP-MS (200.8), or 25x MDL for GFAA.

5/5/11

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: E = Matrix Effects exist as proven by Serial Dilution or Spiked Dilution
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**Data Validation Report
Santa Susana Field Laboratory**

Subarea 8

SDG: PH067

Prepared for

CDM

555 17th Street, Suite 1100
Denver, CO 80202

Prepared by

Laboratory Data Consultants, Inc

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Carlsbad, California 92009

September 24, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on July 15th, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOAs) by EPA SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Pesticides by EPA SW 846 Method 8081B
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A, and 7471B
Herbicides by EPA SW 846 Method 846 Method 8151A
Total Petroleum Hydrocarbon as Gasoline (TPH-G) by EPA SW 846 Method 8015M
Total Petroleum Hydrocarbons as Extractable (TPH-E) by EPA SW 846 Method 8015M
Dioxins/Dibenzofurans by EPA Method 1613B

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks, and field duplicate samples. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for dioxins and metals. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for SVOAs, Herbicides, TPH-E, dioxins, and metals. The 2,4,5-T, 2,4,5-TP (Silvex), 2,4-D, and 2,4-DB results in sample SL-501-SA8-SB-0.0-0.5 were qualified as rejected (R) due to MS/MSD %Rs grossly outside QC limits (i.e., $\leq 0\%$). The remainder of the associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ), as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits with the exception of one DUP for metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one LCS/LCSD pair for herbicides. Since the percent recoveries were out high and the results in the associated samples were non-detect, no data were qualified due to high %Rs. The details regarding LCS/LCSD outliers are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met with the following exceptions:

Diluted Sample	Analyte	%D (Limits)	Associated Samples	Flag	A or P
SL-501-SA8-SB-0.0-0.5	Cobalt Lead Lithium	12 (≤ 10) 14 (≤ 10) 13 (≤ 10)	All soil samples in SDG PH067	J (all detects) UJ (all non-detects)	A

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH067	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pair was collected and analyzed for SVOAs, Pesticides, PCBs, metals, herbicides, TPH-E, and dioxins. All RPDs were within QC limits with the exception of several SVOAs, pesticides, dioxins, and metals. In these duplicate pairs, the associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for TPH-G. No contaminant concentrations were found in the trip blank.

Two equipment blanks (from SDG PH069) were collected and analyzed for SVOAs, Pesticides, PCBs, herbicides, metals, TPH-G, TPH-E, and dioxins. The equipment blanks had detections for several SVOAs, metals, and dioxins. All associated sample results were not detected or were significantly greater than the concentrations found in the equipment blanks and were not qualified.

One field blank (from SDG PH029) was collected and analyzed for SVOAs, Pesticides, PCBs, herbicides, metals, TPH-G, TPH-E, and dioxins. The field blank had detections for several semivolatiles, metals and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

Four herbicide results were rejected due to MS/MSD %Rs grossly outside of QC limits. These results are not useable for all purposes.

The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. The remainder of the data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
15-Jul-2013	TB-071513	7128155	TB	5030B	8015M	III
15-Jul-2013	SL-502-SA8-SB-0.0-0.5	7128152	N	3050B	6010C	III
15-Jul-2013	SL-502-SA8-SB-0.0-0.5	7128152	N	3050B	6020A	III
15-Jul-2013	SL-502-SA8-SB-0.0-0.5	7128152	N	3546	8015M	III
15-Jul-2013	SL-502-SA8-SB-0.0-0.5	7128152	N	3546	8082A	III
15-Jul-2013	SL-502-SA8-SB-0.0-0.5	7128152	N	3546	8270D SIM	III
15-Jul-2013	SL-502-SA8-SB-0.0-0.5	7128152	N	METHOD	1613B	III
15-Jul-2013	SL-502-SA8-SB-0.0-0.5	7128152	N	METHOD	7471B	III
15-Jul-2013	SL-502-SA8-SB-4.0-5.0	7128153	N	3050B	6010C	III
15-Jul-2013	SL-502-SA8-SB-4.0-5.0	7128153	N	3050B	6020A	III
15-Jul-2013	SL-502-SA8-SB-4.0-5.0	7128153	N	3546	8015M	III
15-Jul-2013	SL-502-SA8-SB-4.0-5.0	7128153	N	3546	8082A	III
15-Jul-2013	SL-502-SA8-SB-4.0-5.0	7128153	N	3546	8270D SIM	III
15-Jul-2013	SL-502-SA8-SB-4.0-5.0	7128153	N	5035A	8015M	III
15-Jul-2013	SL-502-SA8-SB-4.0-5.0	7128153	N	METHOD	1613B	III
15-Jul-2013	SL-502-SA8-SB-4.0-5.0	7128153	N	METHOD	7471B	III
15-Jul-2013	SL-502-SA8-SB-7.5-8.5	7128154	N	3050B	6010C	III
15-Jul-2013	SL-502-SA8-SB-7.5-8.5	7128154	N	3050B	6020A	III
15-Jul-2013	SL-502-SA8-SB-7.5-8.5	7128154	N	3546	8015M	III
15-Jul-2013	SL-502-SA8-SB-7.5-8.5	7128154	N	3546	8082A	III
15-Jul-2013	SL-502-SA8-SB-7.5-8.5	7128154	N	3546	8270D SIM	III
15-Jul-2013	SL-502-SA8-SB-7.5-8.5	7128154	N	5035A	8015M	III
15-Jul-2013	SL-502-SA8-SB-7.5-8.5	7128154	N	METHOD	1613B	III
15-Jul-2013	SL-502-SA8-SB-7.5-8.5	7128154	N	METHOD	7471B	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5	7128144	N	3050B	6010C	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5	7128144	N	3050B	6020A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
15-Jul-2013	SL-501-SA8-SB-0.0-0.5	7128144	N	3546	8015M	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5	7128144	N	3546	8081B	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5	7128144	N	3546	8082A	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5	7128144	N	3546	8270D SIM	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5	7128144	N	3550B	8151A	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5	7128144	N	METHOD	1613B	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5	7128144	N	METHOD	7471B	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MS	7128145	MS	3050B	6010C	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MS	7128145	MS	3050B	6020A	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MS	7128145	MS	3546	8015M	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MS	7128145	MS	3546	8081B	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MS	7128145	MS	3546	8082A	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MS	7128145	MS	3546	8270D SIM	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MS	7128145	MS	3550B	8151A	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MS	7128145	MS	METHOD	1613B	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MS	7128145	MS	METHOD	7471B	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MSD	7128146	MSD	3050B	6010C	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MSD	7128146	MSD	3050B	6020A	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MSD	7128146	MSD	3546	8015M	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MSD	7128146	MSD	3546	8081B	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MSD	7128146	MSD	3546	8082A	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MSD	7128146	MSD	3546	8270D SIM	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MSD	7128146	MSD	3550B	8151A	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MSD	7128146	MSD	METHOD	1613B	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5MSD	7128146	MSD	METHOD	7471B	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5DUP	7128147	DUP	3050B	6010C	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
15-Jul-2013	SL-501-SA8-SB-0.0-0.5DUP	7128147	DUP	3050B	6020A	III
15-Jul-2013	SL-501-SA8-SB-0.0-0.5DUP	7128147	DUP	METHOD	7471B	III
15-Jul-2013	SL-801-SA8-SB-0.0-0.5	7128148	FD	3050B	6010C	III
15-Jul-2013	SL-801-SA8-SB-0.0-0.5	7128148	FD	3050B	6020A	III
15-Jul-2013	SL-801-SA8-SB-0.0-0.5	7128148	FD	3546	8015M	III
15-Jul-2013	SL-801-SA8-SB-0.0-0.5	7128148	FD	3546	8081B	III
15-Jul-2013	SL-801-SA8-SB-0.0-0.5	7128148	FD	3546	8082A	III
15-Jul-2013	SL-801-SA8-SB-0.0-0.5	7128148	FD	3546	8270D SIM	III
15-Jul-2013	SL-801-SA8-SB-0.0-0.5	7128148	FD	3550B	8151A	III
15-Jul-2013	SL-801-SA8-SB-0.0-0.5	7128148	FD	METHOD	1613B	III
15-Jul-2013	SL-801-SA8-SB-0.0-0.5	7128148	FD	METHOD	7471B	III
15-Jul-2013	SL-501-SA8-SB-4.0-5.0	7128149	N	3050B	6010C	III
15-Jul-2013	SL-501-SA8-SB-4.0-5.0	7128149	N	3050B	6020A	III
15-Jul-2013	SL-501-SA8-SB-4.0-5.0	7128149	N	3546	8015M	III
15-Jul-2013	SL-501-SA8-SB-4.0-5.0	7128149	N	3546	8081B	III
15-Jul-2013	SL-501-SA8-SB-4.0-5.0	7128149	N	3546	8082A	III
15-Jul-2013	SL-501-SA8-SB-4.0-5.0	7128149	N	3546	8270D SIM	III
15-Jul-2013	SL-501-SA8-SB-4.0-5.0	7128149	N	3550B	8151A	III
15-Jul-2013	SL-501-SA8-SB-4.0-5.0	7128149	N	5035A	8015M	III
15-Jul-2013	SL-501-SA8-SB-4.0-5.0	7128149	N	METHOD	1613B	III
15-Jul-2013	SL-501-SA8-SB-4.0-5.0	7128149	N	METHOD	7471B	III
15-Jul-2013	SL-501-SA8-SB-9.0-10.0	7128150	N	3050B	6010C	III
15-Jul-2013	SL-501-SA8-SB-9.0-10.0	7128150	N	3050B	6020A	III
15-Jul-2013	SL-501-SA8-SB-9.0-10.0	7128150	N	3546	8015M	III
15-Jul-2013	SL-501-SA8-SB-9.0-10.0	7128150	N	3546	8081B	III
15-Jul-2013	SL-501-SA8-SB-9.0-10.0	7128150	N	3546	8082A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
15-Jul-2013	SL-501-SA8-SB-9.0-10.0	7128150	N	3546	8270D SIM	III
15-Jul-2013	SL-501-SA8-SB-9.0-10.0	7128150	N	3550B	8151A	III
15-Jul-2013	SL-501-SA8-SB-9.0-10.0	7128150	N	5035A	8015M	III
15-Jul-2013	SL-501-SA8-SB-9.0-10.0	7128150	N	METHOD	1613B	III
15-Jul-2013	SL-501-SA8-SB-9.0-10.0	7128150	N	METHOD	7471B	III
15-Jul-2013	SL-501-SA8-SB-12.0-13.0	7128151	N	3050B	6010C	III
15-Jul-2013	SL-501-SA8-SB-12.0-13.0	7128151	N	3050B	6020A	III
15-Jul-2013	SL-501-SA8-SB-12.0-13.0	7128151	N	3546	8015M	III
15-Jul-2013	SL-501-SA8-SB-12.0-13.0	7128151	N	3546	8081B	III
15-Jul-2013	SL-501-SA8-SB-12.0-13.0	7128151	N	3546	8082A	III
15-Jul-2013	SL-501-SA8-SB-12.0-13.0	7128151	N	3546	8270D SIM	III
15-Jul-2013	SL-501-SA8-SB-12.0-13.0	7128151	N	3550B	8151A	III
15-Jul-2013	SL-501-SA8-SB-12.0-13.0	7128151	N	5035A	8015M	III
15-Jul-2013	SL-501-SA8-SB-12.0-13.0	7128151	N	METHOD	1613B	III
15-Jul-2013	SL-501-SA8-SB-12.0-13.0	7128151	N	METHOD	7471B	III
15-Jul-2013	SL-511-SA8-SB-0.0-0.5	7128158	N	3050B	6010C	III
15-Jul-2013	SL-511-SA8-SB-0.0-0.5	7128158	N	3050B	6020A	III
15-Jul-2013	SL-511-SA8-SB-0.0-0.5	7128158	N	3546	8015M	III
15-Jul-2013	SL-511-SA8-SB-0.0-0.5	7128158	N	3546	8082A	III
15-Jul-2013	SL-511-SA8-SB-0.0-0.5	7128158	N	3546	8270D SIM	III
15-Jul-2013	SL-511-SA8-SB-0.0-0.5	7128158	N	METHOD	1613B	III
15-Jul-2013	SL-511-SA8-SB-0.0-0.5	7128158	N	METHOD	7471B	III
15-Jul-2013	SL-511-SA8-SB-6.0-7.0	7128159	N	3050B	6010C	III
15-Jul-2013	SL-511-SA8-SB-6.0-7.0	7128159	N	3050B	6020A	III
15-Jul-2013	SL-511-SA8-SB-6.0-7.0	7128159	N	3546	8015M	III
15-Jul-2013	SL-511-SA8-SB-6.0-7.0	7128159	N	3546	8082A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
15-Jul-2013	SL-511-SA8-SB-6.0-7.0	7128159	N	3546	8270D SIM	III
15-Jul-2013	SL-511-SA8-SB-6.0-7.0	7128159	N	5035A	8015M	III
15-Jul-2013	SL-511-SA8-SB-6.0-7.0	7128159	N	METHOD	1613B	III
15-Jul-2013	SL-511-SA8-SB-6.0-7.0	7128159	N	METHOD	7471B	III
15-Jul-2013	SL-503-SA8-SB-0.0-0.5	7128156	N	3050B	6010C	III
15-Jul-2013	SL-503-SA8-SB-0.0-0.5	7128156	N	3050B	6020A	III
15-Jul-2013	SL-503-SA8-SB-0.0-0.5	7128156	N	3546	8015M	III
15-Jul-2013	SL-503-SA8-SB-0.0-0.5	7128156	N	3546	8082A	III
15-Jul-2013	SL-503-SA8-SB-0.0-0.5	7128156	N	3546	8270D SIM	III
15-Jul-2013	SL-503-SA8-SB-0.0-0.5	7128156	N	METHOD	1613B	III
15-Jul-2013	SL-503-SA8-SB-0.0-0.5	7128156	N	METHOD	7471B	III
15-Jul-2013	SL-503-SA8-SB-3.0-4.0	7128157	N	3050B	6010C	III
15-Jul-2013	SL-503-SA8-SB-3.0-4.0	7128157	N	3050B	6020A	III
15-Jul-2013	SL-503-SA8-SB-3.0-4.0	7128157	N	3546	8015M	III
15-Jul-2013	SL-503-SA8-SB-3.0-4.0	7128157	N	3546	8082A	III
15-Jul-2013	SL-503-SA8-SB-3.0-4.0	7128157	N	3546	8270D SIM	III
15-Jul-2013	SL-503-SA8-SB-3.0-4.0	7128157	N	5035A	8015M	III
15-Jul-2013	SL-503-SA8-SB-3.0-4.0	7128157	N	METHOD	1613B	III
15-Jul-2013	SL-503-SA8-SB-3.0-4.0	7128157	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: METALS

Method: 6010C

Matrix: SO

Sample ID: SL-501-SA8-SB-0.0-0.5

Collected: 7/15/2013 11:10:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.11	U	0.760	MDL	4.11	PQL	mg/Kg	UJ	Q
ARSENIC	3.89	J	0.719	MDL	4.11	PQL	mg/Kg	J	Z
BERYLLIUM	0.605	J	0.0689	MDL	1.03	PQL	mg/Kg	J	Z
BORON	6.11	J	0.863	MDL	10.3	PQL	mg/Kg	UJ	B, FD
CADMIUM	0.352	J	0.0781	MDL	1.03	PQL	mg/Kg	J	Z
CALCIUM	3910		3.43	MDL	20.6	PQL	mg/Kg	J	E
COBALT	5.94		0.102	MDL	1.03	PQL	mg/Kg	J	A
LEAD	35.3		0.514	MDL	3.08	PQL	mg/Kg	J	E, Q, FD, A
LITHIUM	21.5		0.35	MDL	4.1	PQL	mg/Kg	J	A
MOLYBDENUM	0.441	J	0.175	MDL	2.06	PQL	mg/Kg	UJ	FD, F
SODIUM	77.0	J	17.2	MDL	103	PQL	mg/Kg	J	Z
TIN	2.77	J	0.226	MDL	10.3	PQL	mg/Kg	U	B
Zirconium	1.14	J	0.863	MDL	5.14	PQL	mg/Kg	J	Z

Sample ID: SL-501-SA8-SB-12.0-13.0

Collected: 7/15/2013 12:00:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.14	U	0.765	MDL	4.14	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.767	J	0.0693	MDL	1.03	PQL	mg/Kg	J	Z
BORON	5.26	J	0.869	MDL	10.3	PQL	mg/Kg	U	B
CADMIUM	0.0910	J	0.0786	MDL	1.03	PQL	mg/Kg	J	Z
CALCIUM	3410		3.45	MDL	20.7	PQL	mg/Kg	J	E
COBALT	5.85		0.102	MDL	1.03	PQL	mg/Kg	J	A
LEAD	5.75		0.517	MDL	3.10	PQL	mg/Kg	J	E, Q, A
LITHIUM	32.5		0.35	MDL	4.1	PQL	mg/Kg	J	A
TIN	2.89	J	0.227	MDL	10.3	PQL	mg/Kg	U	B
Zirconium	2.62	J	0.869	MDL	5.17	PQL	mg/Kg	J	Z

Sample ID: SL-501-SA8-SB-4.0-5.0

Collected: 7/15/2013 11:30:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.15	U	0.768	MDL	4.15	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.833	J	0.0695	MDL	1.04	PQL	mg/Kg	J	Z
BORON	5.00	J	0.871	MDL	10.4	PQL	mg/Kg	U	B
CADMIUM	0.111	J	0.0788	MDL	1.04	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: METALS

Method: 6010C

Matrix: SO

Sample ID: SL-501-SA8-SB-4.0-5.0

Collected: 7/15/2013 11:30:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	3080		3.46	MDL	20.7	PQL	mg/Kg	J	E
COBALT	6.02		0.103	MDL	1.04	PQL	mg/Kg	J	A
LEAD	6.77		0.519	MDL	3.11	PQL	mg/Kg	J	E, Q, A
LITHIUM	23.9		0.35	MDL	4.1	PQL	mg/Kg	J	A
MOLYBDENUM	0.417	J	0.176	MDL	2.07	PQL	mg/Kg	U	F
TIN	3.04	J	0.228	MDL	10.4	PQL	mg/Kg	U	B
Zirconium	1.89	J	0.871	MDL	5.19	PQL	mg/Kg	J	Z

Sample ID: SL-501-SA8-SB-9.0-10.0

Collected: 7/15/2013 11:45:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.33	U	0.801	MDL	4.33	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.863	J	0.0725	MDL	1.08	PQL	mg/Kg	J	Z
BORON	6.30	J	0.909	MDL	10.8	PQL	mg/Kg	U	B
CADMIUM	0.110	J	0.0823	MDL	1.08	PQL	mg/Kg	J	Z
CALCIUM	3440		3.61	MDL	21.6	PQL	mg/Kg	J	E
COBALT	6.98		0.107	MDL	1.08	PQL	mg/Kg	J	A
LEAD	6.16		0.541	MDL	3.25	PQL	mg/Kg	J	E, Q, A
LITHIUM	29.7		0.37	MDL	4.3	PQL	mg/Kg	J	A
TIN	3.12	J	0.238	MDL	10.8	PQL	mg/Kg	U	B
Zirconium	2.66	J	0.909	MDL	5.41	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-0.0-0.5

Collected: 7/15/2013 8:40:00

Analysis Type: RES

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.21	U	0.778	MDL	4.21	PQL	mg/Kg	UJ	Q
ARSENIC	4.11	J	0.736	MDL	4.21	PQL	mg/Kg	J	Z
BERYLLIUM	0.562	J	0.0705	MDL	1.05	PQL	mg/Kg	J	Z
BORON	7.28	J	0.883	MDL	10.5	PQL	mg/Kg	U	B
CADMIUM	0.787	J	0.0799	MDL	1.05	PQL	mg/Kg	J	Z
CALCIUM	5170		3.51	MDL	21.0	PQL	mg/Kg	J	E
COBALT	5.81		0.104	MDL	1.05	PQL	mg/Kg	J	A
LEAD	30.3		0.526	MDL	3.15	PQL	mg/Kg	J	E, Q, A
LITHIUM	20.3		0.36	MDL	4.2	PQL	mg/Kg	J	A
SODIUM	82.3	J	17.6	MDL	105	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-502-SA8-SB-0.0-0.5	Collected: 7/15/2013 8:40:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
TIN	2.89	J	0.231	MDL	10.5	PQL	mg/Kg	U	B
Zirconium	2.27	J	0.883	MDL	5.26	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-4.0-5.0	Collected: 7/15/2013 9:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.15	U	0.768	MDL	4.15	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.753	J	0.0696	MDL	1.04	PQL	mg/Kg	J	Z
BORON	4.78	J	0.872	MDL	10.4	PQL	mg/Kg	U	B
CADMIUM	0.221	J	0.0789	MDL	1.04	PQL	mg/Kg	J	Z
CALCIUM	4150		3.47	MDL	20.8	PQL	mg/Kg	J	E
COBALT	6.39		0.103	MDL	1.04	PQL	mg/Kg	J	A
LEAD	20.5		0.519	MDL	3.12	PQL	mg/Kg	J	E, Q, A
LITHIUM	26.8		0.35	MDL	4.2	PQL	mg/Kg	J	A
MOLYBDENUM	0.278	J	0.177	MDL	2.08	PQL	mg/Kg	U	F
TIN	3.08	J	0.228	MDL	10.4	PQL	mg/Kg	U	B
Zirconium	2.50	J	0.872	MDL	5.19	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-7.5-8.5	Collected: 7/15/2013 10:15:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.26	U	0.789	MDL	4.26	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.602	J	0.0714	MDL	1.07	PQL	mg/Kg	J	Z
BORON	1.86	J	0.895	MDL	10.7	PQL	mg/Kg	U	B
CADMIUM	0.220	J	0.0810	MDL	1.07	PQL	mg/Kg	J	Z
CALCIUM	3610		3.56	MDL	21.3	PQL	mg/Kg	J	E
COBALT	6.32		0.106	MDL	1.07	PQL	mg/Kg	J	A
LEAD	7.93		0.533	MDL	3.20	PQL	mg/Kg	J	E, Q, A
LITHIUM	29.9		0.36	MDL	4.3	PQL	mg/Kg	J	A
TIN	2.95	J	0.234	MDL	10.7	PQL	mg/Kg	U	B
Zirconium	2.80	J	0.895	MDL	5.33	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-503-SA8-SB-0.0-0.5 Collected: 7/15/2013 2:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.14	U	0.765	MDL	4.14	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.614	J	0.0693	MDL	1.03	PQL	mg/Kg	J	Z
BORON	5.29	J	0.868	MDL	10.3	PQL	mg/Kg	U	B
CADMIUM	0.375	J	0.0786	MDL	1.03	PQL	mg/Kg	J	Z
CALCIUM	11700		3.45	MDL	20.7	PQL	mg/Kg	J	E
COBALT	6.76		0.102	MDL	1.03	PQL	mg/Kg	J	A
LEAD	9.56		0.517	MDL	3.10	PQL	mg/Kg	J	E, Q, A
LITHIUM	21.9		0.35	MDL	4.1	PQL	mg/Kg	J	A
MOLYBDENUM	0.191	J	0.176	MDL	2.07	PQL	mg/Kg	U	F
TIN	2.88	J	0.227	MDL	10.3	PQL	mg/Kg	U	B
Zirconium	2.26	J	0.868	MDL	5.17	PQL	mg/Kg	J	Z

Sample ID: SL-503-SA8-SB-3.0-4.0 Collected: 7/15/2013 3:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.19	U	0.774	MDL	4.19	PQL	mg/Kg	UJ	Q
ARSENIC	2.66	J	0.732	MDL	4.19	PQL	mg/Kg	J	Z
BERYLLIUM	0.400	J	0.0701	MDL	1.05	PQL	mg/Kg	J	Z
BORON	2.99	J	0.879	MDL	10.5	PQL	mg/Kg	U	B
CADMIUM	0.243	J	0.0795	MDL	1.05	PQL	mg/Kg	J	Z
CALCIUM	5060		3.49	MDL	20.9	PQL	mg/Kg	J	E
COBALT	4.88		0.104	MDL	1.05	PQL	mg/Kg	J	A
LEAD	3.71		0.523	MDL	3.14	PQL	mg/Kg	J	E, Q, A
LITHIUM	23.2		0.36	MDL	4.2	PQL	mg/Kg	J	A
TIN	2.71	J	0.230	MDL	10.5	PQL	mg/Kg	U	B
Zirconium	1.71	J	0.879	MDL	5.23	PQL	mg/Kg	J	Z

Sample ID: SL-511-SA8-SB-0.0-0.5 Collected: 7/15/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.09	U	0.757	MDL	4.09	PQL	mg/Kg	UJ	Q
ARSENIC	3.19	J	0.716	MDL	4.09	PQL	mg/Kg	J	Z
BERYLLIUM	0.578	J	0.0685	MDL	1.02	PQL	mg/Kg	J	Z
BORON	3.23	J	0.859	MDL	10.2	PQL	mg/Kg	U	B
CADMIUM	0.269	J	0.0777	MDL	1.02	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-511-SA8-SB-0.0-0.5 Collected: 7/15/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	2630		3.42	MDL	20.5	PQL	mg/Kg	J	E
COBALT	6.06		0.101	MDL	1.02	PQL	mg/Kg	J	A
LEAD	4.63		0.511	MDL	3.07	PQL	mg/Kg	J	E, Q, A
LITHIUM	23.2		0.35	MDL	4.1	PQL	mg/Kg	J	A
MOLYBDENUM	0.217	J	0.174	MDL	2.05	PQL	mg/Kg	U	F
SODIUM	75.7	J	17.1	MDL	102	PQL	mg/Kg	J	Z
TIN	2.71	J	0.225	MDL	10.2	PQL	mg/Kg	U	B
Zirconium	1.47	J	0.859	MDL	5.11	PQL	mg/Kg	J	Z

Sample ID: SL-511-SA8-SB-6.0-7.0 Collected: 7/15/2013 2:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.54	U	0.839	MDL	4.54	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.595	J	0.0760	MDL	1.13	PQL	mg/Kg	J	Z
BORON	1.56	J	0.953	MDL	11.3	PQL	mg/Kg	U	B
CADMIUM	0.128	J	0.0862	MDL	1.13	PQL	mg/Kg	J	Z
CALCIUM	3090		3.79	MDL	22.7	PQL	mg/Kg	J	E
COBALT	5.33		0.112	MDL	1.13	PQL	mg/Kg	J	A
LEAD	4.95		0.567	MDL	3.40	PQL	mg/Kg	J	E, Q, A
LITHIUM	27.2		0.39	MDL	4.5	PQL	mg/Kg	J	A
TIN	3.54	J	0.250	MDL	11.3	PQL	mg/Kg	U	B
Zirconium	1.99	J	0.953	MDL	5.67	PQL	mg/Kg	J	Z

Sample ID: SL-801-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:15:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
COBALT	5.67		0.101	MDL	1.02	PQL	mg/Kg	J	A

Sample ID: SL-801-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.06	U	0.752	MDL	4.06	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.842	J	0.0680	MDL	1.02	PQL	mg/Kg	J	Z
BORON	16.1		0.853	MDL	10.2	PQL	mg/Kg	J	FD
CADMIUM	0.374	J	0.0772	MDL	1.02	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-801-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CALCIUM	5920		3.39	MDL	20.3	PQL	mg/Kg	J	E
LEAD	18.4		0.508	MDL	3.05	PQL	mg/Kg	J	E, Q, FD, A
LITHIUM	21.7		0.35	MDL	4.1	PQL	mg/Kg	J	A
MOLYBDENUM	2.03	U	0.173	MDL	2.03	PQL	mg/Kg	UJ	FD
SODIUM	85.6	J	17.0	MDL	102	PQL	mg/Kg	J	Z
TIN	2.68	J	0.223	MDL	10.2	PQL	mg/Kg	U	B
Zirconium	0.923	J	0.853	MDL	5.08	PQL	mg/Kg	J	Z

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-501-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:10:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.245	J	0.103	MDL	0.411	PQL	mg/Kg	J	Z

Sample ID: SL-501-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:10:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0421	J	0.0267	MDL	0.206	PQL	mg/Kg	J	Z

Sample ID: SL-501-SA8-SB-12.0-13.0 Collected: 7/15/2013 12:00:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0697	J	0.0269	MDL	0.207	PQL	mg/Kg	J	Z

Sample ID: SL-501-SA8-SB-4.0-5.0 Collected: 7/15/2013 11:30:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.283	J	0.104	MDL	0.415	PQL	mg/Kg	J	Z

Sample ID: SL-501-SA8-SB-4.0-5.0 Collected: 7/15/2013 11:30:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0344	J	0.0270	MDL	0.207	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-501-SA8-SB-9.0-10.0		Collected: 7/15/2013 11:45:00		Analysis Type: RES		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0487	J	0.0281	MDL	0.216	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-0.0-0.5		Collected: 7/15/2013 8:40:00		Analysis Type: REA2		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.255	J	0.105	MDL	0.421	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-0.0-0.5		Collected: 7/15/2013 8:40:00		Analysis Type: RES		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0425	J	0.0273	MDL	0.210	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-4.0-5.0		Collected: 7/15/2013 9:30:00		Analysis Type: REA2		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.189	J	0.104	MDL	0.415	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-4.0-5.0		Collected: 7/15/2013 9:30:00		Analysis Type: RES		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0428	J	0.0270	MDL	0.208	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-7.5-8.5		Collected: 7/15/2013 10:15:00		Analysis Type: REA2		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.158	J	0.107	MDL	0.426	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-7.5-8.5		Collected: 7/15/2013 10:15:00		Analysis Type: RES		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0505	J	0.0277	MDL	0.213	PQL	mg/Kg	J	Z

Sample ID: SL-503-SA8-SB-0.0-0.5		Collected: 7/15/2013 2:55:00		Analysis Type: REA2		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.225	J	0.103	MDL	0.414	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-503-SA8-SB-0.0-0.5	Collected: 7/15/2013 2:55:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0964	J	0.0269	MDL	0.207	PQL	mg/Kg	J	Z

Sample ID: SL-511-SA8-SB-0.0-0.5	Collected: 7/15/2013 2:15:00	Analysis Type: REA2	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.204	J	0.102	MDL	0.409	PQL	mg/Kg	J	Z

Sample ID: SL-511-SA8-SB-0.0-0.5	Collected: 7/15/2013 2:15:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0411	J	0.0266	MDL	0.205	PQL	mg/Kg	J	Z

Sample ID: SL-511-SA8-SB-6.0-7.0	Collected: 7/15/2013 2:30:00	Analysis Type: REA2	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.142	J	0.113	MDL	0.454	PQL	mg/Kg	J	Z

Sample ID: SL-511-SA8-SB-6.0-7.0	Collected: 7/15/2013 2:30:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.104	J	0.0295	MDL	0.227	PQL	mg/Kg	J	Z

Sample ID: SL-801-SA8-SB-0.0-0.5	Collected: 7/15/2013 11:15:00	Analysis Type: REA2	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.246	J	0.102	MDL	0.406	PQL	mg/Kg	J	Z

Sample ID: SL-801-SA8-SB-0.0-0.5	Collected: 7/15/2013 11:15:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0479	J	0.0264	MDL	0.203	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	7471B	Matrix: SO

Sample ID: SL-501-SA8-SB-0.0-0.5	Collected: 7/15/2013 11:10:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0276		0.0102	MDL	0.0171	PQL	mg/Kg	J	FD

Sample ID: SL-501-SA8-SB-4.0-5.0	Collected: 7/15/2013 11:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0107	J	0.0107	MDL	0.0178	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-4.0-5.0	Collected: 7/15/2013 9:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0164	J	0.0106	MDL	0.0176	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-7.5-8.5	Collected: 7/15/2013 10:15:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0118	J	0.0103	MDL	0.0172	PQL	mg/Kg	J	Z

Sample ID: SL-801-SA8-SB-0.0-0.5	Collected: 7/15/2013 11:15:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0493		0.010	MDL	0.0166	PQL	mg/Kg	J	FD

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-501-SA8-SB-0.0-0.5	Collected: 7/15/2013 11:10:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	3.22	JB	0.0327	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.317	JQ	0.0567	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.303	JQ	0.0729	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDF	0.312	JBQ	0.0405	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	1.14	J	0.0805	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.379	JBQ	0.0411	MDL	5.05	PQL	ng/Kg	J	Z, FD
1,2,3,7,8,9-HxCDD	0.996	JB	0.0819	MDL	5.05	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA		
Method:	1613B	Matrix:	SO

Sample ID: SL-501-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,7,8,9-HXCDF	0.415	J	0.0454	MDL	5.05	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.263	JQ	0.0829	MDL	5.05	PQL	ng/Kg	J	Z, FD
1,2,3,7,8-PECDF	1.09	JB	0.0476	MDL	5.05	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.262	JB	0.0407	MDL	5.05	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.653	JQ	0.0470	MDL	5.05	PQL	ng/Kg	J	Z, FD
2,3,7,8-TCDD	0.0816	JQ	0.0656	MDL	1.01	PQL	ng/Kg	J	Z, FD
2,3,7,8-TCDF	0.540	J	0.0809	MDL	1.01	PQL	ng/Kg	J	Z
OCDD	240	B	0.0610	MDL	10.1	PQL	ng/Kg	J	Q, Q, FD
OCDF	5.69	JB	0.0547	MDL	10.1	PQL	ng/Kg	J	Z

Sample ID: SL-501-SA8-SB-12.0-13.0 Collected: 7/15/2013 12:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.252	JBQ	0.0397	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0569	JBQ	0.0122	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0304	J	0.0292	MDL	5.26	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0331	JBQ	0.0153	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0301	JBQ	0.0157	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0562	JBQ	0.0315	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0198	JQ	0.0181	MDL	5.26	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.0530	JB	0.0271	MDL	5.26	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0238	JBQ	0.0155	MDL	5.26	PQL	ng/Kg	U	B
OCDD	2.26	JB	0.0289	MDL	10.5	PQL	ng/Kg	J	Z
OCDF	0.183	JB	0.0439	MDL	10.5	PQL	ng/Kg	U	B

Sample ID: SL-501-SA8-SB-4.0-5.0 Collected: 7/15/2013 11:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.33	JBQ	0.0818	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.243	JB	0.0184	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.0621	JQ	0.0560	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0457	JBQ	0.0272	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.301	JQ	0.0609	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0985	JB	0.0295	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.471	JBQ	0.0603	MDL	5.30	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-501-SA8-SB-4.0-5.0 Collected: 7/15/2013 11:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,7,8,9-HXCDF	0.123	JQ	0.0362	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0925	JQ	0.0694	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.132	JBQ	0.0454	MDL	5.30	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0570	JBQ	0.0274	MDL	5.30	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0585	JQ	0.0399	MDL	5.30	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.105	J	0.0638	MDL	1.06	PQL	ng/Kg	J	Z
OCDF	0.543	JB	0.0591	MDL	10.6	PQL	ng/Kg	J	Z

Sample ID: SL-501-SA8-SB-9.0-10.0 Collected: 7/15/2013 11:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.138	JB	0.0455	MDL	5.33	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0422	JBQ	0.0117	MDL	5.33	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0295	J	0.0162	MDL	5.33	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0259	JBQ	0.0189	MDL	5.33	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0503	JBQ	0.0472	MDL	5.33	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0522	JQ	0.0211	MDL	5.33	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0265	JBQ	0.0174	MDL	5.33	PQL	ng/Kg	U	B
OCDD	1.06	JBQ	0.0333	MDL	10.7	PQL	ng/Kg	U	B
OCDF	0.198	JB	0.0508	MDL	10.7	PQL	ng/Kg	U	B

Sample ID: SL-502-SA8-SB-0.0-0.5 Collected: 7/15/2013 8:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HXCDF	4.75	JB	0.107	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	3.19	JB	0.112	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	4.56	JB	0.104	MDL	5.10	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	2.77	JB	0.100	MDL	5.10	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	1.46	JQ	0.107	MDL	5.10	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.571	J	0.193	MDL	1.02	PQL	ng/Kg	J	Z

Sample ID: SL-502-SA8-SB-4.0-5.0 Collected: 7/15/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	4.77	JB	0.0373	MDL	5.38	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA		
Method:	1613B	Matrix:	SO

Sample ID: SL-502-SA8-SB-4.0-5.0 Collected: 7/15/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8,9-HPCDF	0.577	JQ	0.0554	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	1.75	J	0.111	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.715	JB	0.0533	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	2.30	J	0.116	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.566	JBQ	0.0511	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	2.14	JB	0.118	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.513	J	0.0599	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.906	JQ	0.145	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	1.75	JB	0.0532	MDL	5.38	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.396	JB	0.0474	MDL	5.38	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.440	J	0.0482	MDL	5.38	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.149	JQ	0.0830	MDL	1.08	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.205	J	0.0906	MDL	1.08	PQL	ng/Kg	J	Z
OCDF	8.63	JB	0.0499	MDL	10.8	PQL	ng/Kg	J	Z

Sample ID: SL-502-SA8-SB-7.5-8.5 Collected: 7/15/2013 10:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	1.83	JBQ	0.0257	MDL	5.24	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.230	JQ	0.0395	MDL	5.24	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.665	J	0.0848	MDL	5.24	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.309	JB	0.0313	MDL	5.24	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.871	JQ	0.0870	MDL	5.24	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.228	JBQ	0.0315	MDL	5.24	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.638	JB	0.0884	MDL	5.24	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.324	JQ	0.0394	MDL	5.24	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.268	JQ	0.0834	MDL	5.24	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.541	JB	0.0336	MDL	5.24	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.227	JBQ	0.0325	MDL	5.24	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.206	JQ	0.0311	MDL	5.24	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0961	JQ	0.0579	MDL	1.05	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0929	JQ	0.0541	MDL	1.05	PQL	ng/Kg	J	Z
OCDF	3.40	JB	0.0419	MDL	10.5	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-503-SA8-SB-0.0-0.5 Collected: 7/15/2013 2:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	2.16	JB	0.0445	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.209	JQ	0.0690	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.270	JQ	0.0794	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.303	JB	0.0389	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.868	J	0.0852	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.441	JB	0.0396	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.815	JB	0.0831	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.471	J	0.0453	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.148	JQ	0.0922	MDL	5.19	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	2.52	JB	0.0504	MDL	5.19	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.213	JBQ	0.0392	MDL	5.19	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.210	JQ	0.0522	MDL	5.19	PQL	ng/Kg	J	Z
OCDF	3.21	JB	0.0604	MDL	10.4	PQL	ng/Kg	J	Z

Sample ID: SL-503-SA8-SB-3.0-4.0 Collected: 7/15/2013 3:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.403	JBQ	0.0435	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.0653	JBQ	0.0131	MDL	5.17	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0431	J	0.0174	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0484	JBQ	0.0121	MDL	5.17	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.140	JQ	0.0375	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0283	JBQ	0.0150	MDL	5.17	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.222	JBQ	0.0372	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.436	J	0.0204	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0679	JQ	0.0479	MDL	5.17	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.118	JBQ	0.0253	MDL	5.17	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0300	JQ	0.0233	MDL	5.17	PQL	ng/Kg	J	Z
OCDD	2.77	JB	0.0266	MDL	10.3	PQL	ng/Kg	J	Z
OCDF	0.162	JB	0.0393	MDL	10.3	PQL	ng/Kg	U	B

Sample ID: SL-511-SA8-SB-0.0-0.5 Collected: 7/15/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0766	JBQ	0.0352	MDL	5.01	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA		
Method:	1613B	Matrix:	SO

Sample ID: SL-511-SA8-SB-0.0-0.5 Collected: 7/15/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	0.0266	JBQ	0.0121	MDL	5.01	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0230	JQ	0.0196	MDL	5.01	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.0339	JQ	0.0253	MDL	5.01	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0362	JBQ	0.0138	MDL	5.01	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0499	JBQ	0.0273	MDL	5.01	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0613	JQ	0.0158	MDL	5.01	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0157	JB	0.0135	MDL	5.01	PQL	ng/Kg	U	B
OCDD	0.345	JBQ	0.0267	MDL	10.0	PQL	ng/Kg	U	B
OCDF	0.0970	JBQ	0.0380	MDL	10.0	PQL	ng/Kg	U	B

Sample ID: SL-511-SA8-SB-6.0-7.0 Collected: 7/15/2013 2:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.112	JBQ	0.0530	MDL	5.77	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0397	JBQ	0.0176	MDL	5.77	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0478	JQ	0.0264	MDL	5.77	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0353	JBQ	0.0223	MDL	5.77	PQL	ng/Kg	U	B
OCDD	0.347	JBQ	0.0505	MDL	11.5	PQL	ng/Kg	U	B

Sample ID: SL-801-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	2.24	JB	0.0371	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.224	JQ	0.0616	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.265	J	0.0643	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.228	JBQ	0.0373	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.803	J	0.0691	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.132	JBQ	0.0376	MDL	5.10	PQL	ng/Kg	UJ	B, FD
1,2,3,7,8,9-HXCDD	0.808	JB	0.0701	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.431	J	0.0416	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.149	JQ	0.0809	MDL	5.10	PQL	ng/Kg	J	Z, FD
1,2,3,7,8-PECDF	1.18	JB	0.0427	MDL	5.10	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.208	JBQ	0.0358	MDL	5.10	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.208	JQ	0.0396	MDL	5.10	PQL	ng/Kg	J	Z, FD
2,3,7,8-TCDD	1.02	U	0.0473	MDL	1.02	PQL	ng/Kg	UJ	FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-801-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,3,7,8-TCDF	0.685	J	0.0794	MDL	1.02	PQL	ng/Kg	J	Z
OCDD	137	B	0.0573	MDL	10.2	PQL	ng/Kg	J	FD
OCDF	4.72	JB	0.0482	MDL	10.2	PQL	ng/Kg	J	Z

Method Category:	SVOA	
Method:	8015M	Matrix: SO

Sample ID: SL-501-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	2.6	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z

Sample ID: SL-502-SA8-SB-0.0-0.5 Collected: 7/15/2013 8:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	2.9	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z

Sample ID: SL-801-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	2.4	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z

Method Category:	SVOA	
Method:	8081B	Matrix: SO

Sample ID: SL-501-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:10:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
4,4'-DDE	2.7		0.34	MDL	1.8	PQL	ug/Kg	J	FD

Sample ID: SL-801-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
4,4'-DDE	1.3	J	0.34	MDL	1.8	PQL	ug/Kg	J	Z, FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	8082A	Matrix:	SO
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Sample ID: SL-501-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:10:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCOLOR 1254	8.1	J	4.6	MDL	18	PQL	ug/Kg	J	Z

Sample ID: SL-502-SA8-SB-4.0-5.0 Collected: 7/15/2013 9:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCOLOR 1254	6.8	J	4.8	MDL	18	PQL	ug/Kg	J	Z

Sample ID: SL-502-SA8-SB-7.5-8.5 Collected: 7/15/2013 10:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
Aroclor 5460	17	J	11	MDL	36	PQL	ug/Kg	J	Z

Sample ID: SL-801-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCOLOR 1254	13	J	4.6	MDL	18	PQL	ug/Kg	J	Z

Method Category:	SVOA	Method:	8151A	Matrix:	SO
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Sample ID: SL-501-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,4,5-T	1.8	U	0.85	MDL	1.8	PQL	ug/Kg	R	Q
2,4,5-TP (Silvex)	1.8	U	0.78	MDL	1.8	PQL	ug/Kg	R	Q
2,4-D	37	U	12	MDL	37	PQL	ug/Kg	R	Q
2,4-DB	18	U	6.4	MDL	18	PQL	ug/Kg	R	Q

Method Category:	SVOA	Method:	8270D SIM	Matrix:	SO
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Sample ID: SL-501-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:10:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	1.7	U	0.70	MDL	1.7	PQL	ug/Kg	UJ	FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	8270D SIM	Matrix:	SO
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Sample ID: SL-501-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:10:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	1.3	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z
ANTHRACENE	0.49	J	0.35	MDL	1.7	PQL	ug/Kg	J	Z, FD
BENZO(A)ANTHRACENE	3.3		0.70	MDL	1.7	PQL	ug/Kg	J	FD
BENZO(A)PYRENE	5.2		0.70	MDL	1.7	PQL	ug/Kg	J	FD
BENZO(B)FLUORANTHENE	6.9		0.70	MDL	1.7	PQL	ug/Kg	J	FD
BENZO(E)PYRENE	4.4	J	3.5	MDL	18	PQL	ug/Kg	J	Z, FD
BENZO(G,H,I)PERYLENE	3.7		0.70	MDL	1.7	PQL	ug/Kg	J	FD
BIS(2-ETHYLHEXYL)PHTHALATE	13	J	6.3	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	8.0		0.35	MDL	1.7	PQL	ug/Kg	J	FD
DIBENZO(A,H)ANTHRACENE	1.2	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z, FD
Di-n-octylphthalate	19	U	6.3	MDL	19	PQL	ug/Kg	UJ	FD
FLUORANTHENE	6.5		0.70	MDL	1.7	PQL	ug/Kg	J	FD
INDENO(1,2,3-CD)PYRENE	3.4		0.70	MDL	1.7	PQL	ug/Kg	J	FD
PHENANTHRENE	3.2		0.70	MDL	1.7	PQL	ug/Kg	J	FD
PYRENE	6.9		0.70	MDL	1.7	PQL	ug/Kg	J	FD

Sample ID: SL-501-SA8-SB-4.0-5.0 Collected: 7/15/2013 11:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(K)FLUORANTHENE	1.2	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	0.65	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	1.5	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-502-SA8-SB-0.0-0.5 Collected: 7/15/2013 8:40:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BIS(2-ETHYLHEXYL)PHTHALATE	13	J	6.3	MDL	19	PQL	ug/Kg	J	Z
Di-n-butylphthalate	8.2	J	6.3	MDL	19	PQL	ug/Kg	J	Z

Sample ID: SL-502-SA8-SB-4.0-5.0 Collected: 7/15/2013 9:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	1.2	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
ANTHRACENE	0.67	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(A)ANTHRACENE	0.82	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-502-SA8-SB-4.0-5.0 Collected: 7/15/2013 9:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BIS(2-ETHYLHEXYL)PHTHALATE	11	J	6.5	MDL	19	PQL	ug/Kg	J	Z

Sample ID: SL-502-SA8-SB-7.5-8.5 Collected: 7/15/2013 10:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTHRACENE	0.43	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(B)FLUORANTHENE	0.88	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	15	J	6.5	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	1.4	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
FLUORANTHENE	1.5	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	1.7	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
PHENANTHRENE	1.1	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
PYRENE	1.4	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-503-SA8-SB-0.0-0.5 Collected: 7/15/2013 2:55:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.84	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(B)FLUORANTHENE	1.4	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	11	J	6.5	MDL	19	PQL	ug/Kg	J	Z
FLUORANTHENE	1.4	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
PHENANTHRENE	1.0	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
PYRENE	1.2	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-511-SA8-SB-0.0-0.5 Collected: 7/15/2013 2:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(K)FLUORANTHENE	1.5	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
CHRYSENE	0.58	J	0.34	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-801-SA8-SB-0.0-0.5 Collected: 7/15/2013 11:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	0.72	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z, FD
2-METHYLNAPHTHALENE	1.0	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-801-SA8-SB-0.0-0.5

Collected: 7/15/2013 11:15:00

Analysis Type: RES-BASE/NEUTRAL

Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTHRACENE	1.7	U	0.35	MDL	1.7	PQL	ug/Kg	UJ	FD
BENZO(A)ANTHRACENE	0.77	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z, FD
BENZO(A)PYRENE	0.94	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z, FD
BENZO(B)FLUORANTHENE	2.2		0.70	MDL	1.7	PQL	ug/Kg	J	FD
BENZO(E)PYRENE	18	U	3.5	MDL	18	PQL	ug/Kg	UJ	FD
BENZO(G,H,I)PERYLENE	0.94	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z, FD
BIS(2-ETHYLHEXYL)PHTHALATE	10	J	6.3	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	3.9		0.35	MDL	1.7	PQL	ug/Kg	J	FD
DIBENZO(A,H)ANTHRACENE	1.7	U	0.70	MDL	1.7	PQL	ug/Kg	UJ	FD
Di-n-octylphthalate	10	J	6.3	MDL	19	PQL	ug/Kg	J	Z, FD
FLUORANTHENE	1.7	J	0.70	MDL	1.7	PQL	ug/Kg	J	FD
INDENO(1,2,3-CD)PYRENE	0.70	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z, FD
PHENANTHRENE	1.4	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z, FD
PYRENE	2.1		0.70	MDL	1.7	PQL	ug/Kg	J	FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
*#	Professional Judgment
A	ICP Serial Dilution
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Field Blank Contamination
FD	Field Duplicate Precision
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
S	Surrogate/Tracer Recovery Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH067

Method Blank Outlier Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B				
Matrix: SO				
Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2040B371956	7/24/2013 7:56:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF OCDD OCDF	0.0781 ng/Kg 0.0390 ng/Kg 0.0595 ng/Kg 0.0378 ng/Kg 0.0383 ng/Kg 0.0603 ng/Kg 0.0469 ng/Kg 0.316 ng/Kg 0.0824 ng/Kg	SL-501-SA8-SB-0.0-0.5 SL-501-SA8-SB-12.0-13.0 SL-501-SA8-SB-4.0-5.0 SL-501-SA8-SB-9.0-10.0 SL-502-SA8-SB-0.0-0.5 SL-502-SA8-SB-4.0-5.0 SL-502-SA8-SB-7.5-8.5 SL-503-SA8-SB-0.0-0.5 SL-503-SA8-SB-3.0-4.0 SL-511-SA8-SB-0.0-0.5 SL-511-SA8-SB-6.0-7.0 SL-801-SA8-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-501-SA8-SB-12.0-13.0(RES)	1,2,3,4,6,7,8-HPCDD	0.252 ng/Kg	0.252U ng/Kg
SL-501-SA8-SB-12.0-13.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0569 ng/Kg	0.0569U ng/Kg
SL-501-SA8-SB-12.0-13.0(RES)	1,2,3,4,7,8-HXCDF	0.0331 ng/Kg	0.0331U ng/Kg
SL-501-SA8-SB-12.0-13.0(RES)	1,2,3,6,7,8-HXCDF	0.0301 ng/Kg	0.0301U ng/Kg
SL-501-SA8-SB-12.0-13.0(RES)	1,2,3,7,8,9-HXCDD	0.0562 ng/Kg	0.0562U ng/Kg
SL-501-SA8-SB-12.0-13.0(RES)	1,2,3,7,8-PECDF	0.0530 ng/Kg	0.0530U ng/Kg
SL-501-SA8-SB-12.0-13.0(RES)	2,3,4,6,7,8-HXCDF	0.0238 ng/Kg	0.0238U ng/Kg
SL-501-SA8-SB-12.0-13.0(RES)	OCDF	0.183 ng/Kg	0.183U ng/Kg
SL-501-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HXCDF	0.0457 ng/Kg	0.0457U ng/Kg
SL-501-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDF	0.0985 ng/Kg	0.0985U ng/Kg
SL-501-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.132 ng/Kg	0.132U ng/Kg
SL-501-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0570 ng/Kg	0.0570U ng/Kg
SL-501-SA8-SB-9.0-10.0(RES)	1,2,3,4,6,7,8-HPCDD	0.138 ng/Kg	0.138U ng/Kg
SL-501-SA8-SB-9.0-10.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0422 ng/Kg	0.0422U ng/Kg
SL-501-SA8-SB-9.0-10.0(RES)	1,2,3,4,7,8-HXCDF	0.0259 ng/Kg	0.0259U ng/Kg
SL-501-SA8-SB-9.0-10.0(RES)	1,2,3,7,8,9-HXCDD	0.0503 ng/Kg	0.0503U ng/Kg
SL-501-SA8-SB-9.0-10.0(RES)	2,3,4,6,7,8-HXCDF	0.0265 ng/Kg	0.0265U ng/Kg
SL-501-SA8-SB-9.0-10.0(RES)	OCDD	1.06 ng/Kg	1.06U ng/Kg
SL-501-SA8-SB-9.0-10.0(RES)	OCDF	0.198 ng/Kg	0.198U ng/Kg
SL-502-SA8-SB-7.5-8.5(RES)	2,3,4,6,7,8-HXCDF	0.227 ng/Kg	0.227U ng/Kg
SL-503-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.213 ng/Kg	0.213U ng/Kg
SL-503-SA8-SB-3.0-4.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0653 ng/Kg	0.0653U ng/Kg
SL-503-SA8-SB-3.0-4.0(RES)	1,2,3,4,7,8-HXCDF	0.0484 ng/Kg	0.0484U ng/Kg
SL-503-SA8-SB-3.0-4.0(RES)	1,2,3,6,7,8-HXCDF	0.0283 ng/Kg	0.0283U ng/Kg
SL-503-SA8-SB-3.0-4.0(RES)	1,2,3,7,8-PECDF	0.118 ng/Kg	0.118U ng/Kg
SL-503-SA8-SB-3.0-4.0(RES)	OCDF	0.162 ng/Kg	0.162U ng/Kg
SL-511-SA8-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.0766 ng/Kg	0.0766U ng/Kg
SL-511-SA8-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0266 ng/Kg	0.0266U ng/Kg
SL-511-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0362 ng/Kg	0.0362U ng/Kg
SL-511-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.0499 ng/Kg	0.0499U ng/Kg
SL-511-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0157 ng/Kg	0.0157U ng/Kg
SL-511-SA8-SB-0.0-0.5(RES)	OCDD	0.345 ng/Kg	0.345U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-511-SA8-SB-0.0-0.5(RES)	OCDF	0.0970 ng/Kg	0.0970U ng/Kg
SL-511-SA8-SB-6.0-7.0(RES)	1,2,3,4,6,7,8-HPCDD	0.112 ng/Kg	0.112U ng/Kg
SL-511-SA8-SB-6.0-7.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0397 ng/Kg	0.0397U ng/Kg
SL-511-SA8-SB-6.0-7.0(RES)	1,2,3,4,7,8-HXCDF	0.0353 ng/Kg	0.0353U ng/Kg
SL-511-SA8-SB-6.0-7.0(RES)	OCDD	0.347 ng/Kg	0.347U ng/Kg
SL-801-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.228 ng/Kg	0.228U ng/Kg
SL-801-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.132 ng/Kg	0.132U ng/Kg
SL-801-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.208 ng/Kg	0.208U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19937BB221020	7/21/2013 10:20:00 AM	ALUMINUM BORON CALCIUM TIN ZINC	9.23 mg/Kg 2.24 mg/Kg 24.4 mg/Kg 1.60 mg/Kg 0.513 mg/Kg	SL-501-SA8-SB-0.0-0.5 SL-501-SA8-SB-12.0-13.0 SL-501-SA8-SB-4.0-5.0 SL-501-SA8-SB-9.0-10.0 SL-502-SA8-SB-0.0-0.5 SL-502-SA8-SB-4.0-5.0 SL-502-SA8-SB-7.5-8.5 SL-503-SA8-SB-0.0-0.5 SL-503-SA8-SB-3.0-4.0 SL-511-SA8-SB-0.0-0.5 SL-511-SA8-SB-6.0-7.0 SL-801-SA8-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-501-SA8-SB-0.0-0.5(RES)	BORON	6.11 mg/Kg	6.11U mg/Kg
SL-501-SA8-SB-0.0-0.5(RES)	TIN	2.77 mg/Kg	2.77U mg/Kg
SL-501-SA8-SB-12.0-13.0(RES)	BORON	5.26 mg/Kg	5.26U mg/Kg
SL-501-SA8-SB-12.0-13.0(RES)	TIN	2.89 mg/Kg	2.89U mg/Kg
SL-501-SA8-SB-4.0-5.0(RES)	BORON	5.00 mg/Kg	5.00U mg/Kg
SL-501-SA8-SB-4.0-5.0(RES)	TIN	3.04 mg/Kg	3.04U mg/Kg
SL-501-SA8-SB-9.0-10.0(RES)	BORON	6.30 mg/Kg	6.30U mg/Kg
SL-501-SA8-SB-9.0-10.0(RES)	TIN	3.12 mg/Kg	3.12U mg/Kg
SL-502-SA8-SB-0.0-0.5(RES)	BORON	7.28 mg/Kg	7.28U mg/Kg
SL-502-SA8-SB-0.0-0.5(RES)	TIN	2.89 mg/Kg	2.89U mg/Kg
SL-502-SA8-SB-4.0-5.0(RES)	BORON	4.78 mg/Kg	4.78U mg/Kg
SL-502-SA8-SB-4.0-5.0(RES)	TIN	3.08 mg/Kg	3.08U mg/Kg
SL-502-SA8-SB-7.5-8.5(RES)	BORON	1.86 mg/Kg	1.86U mg/Kg
SL-502-SA8-SB-7.5-8.5(RES)	TIN	2.95 mg/Kg	2.95U mg/Kg
SL-503-SA8-SB-0.0-0.5(RES)	BORON	5.29 mg/Kg	5.29U mg/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-503-SA8-SB-0.0-0.5(RES)	TIN	2.88 mg/Kg	2.88U mg/Kg
SL-503-SA8-SB-3.0-4.0(RES)	BORON	2.99 mg/Kg	2.99U mg/Kg
SL-503-SA8-SB-3.0-4.0(RES)	TIN	2.71 mg/Kg	2.71U mg/Kg
SL-511-SA8-SB-0.0-0.5(RES)	BORON	3.23 mg/Kg	3.23U mg/Kg
SL-511-SA8-SB-0.0-0.5(RES)	TIN	2.71 mg/Kg	2.71U mg/Kg
SL-511-SA8-SB-6.0-7.0(RES)	BORON	1.56 mg/Kg	1.56U mg/Kg
SL-511-SA8-SB-6.0-7.0(RES)	TIN	3.54 mg/Kg	3.54U mg/Kg
SL-801-SA8-SB-0.0-0.5(RES)	TIN	2.68 mg/Kg	2.68U mg/Kg

Method: 6020A
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19937BB222059A	7/25/2013 8:59:00 PM	STRONTIUM	0.161 mg/Kg	SL-501-SA8-SB-0.0-0.5 SL-501-SA8-SB-12.0-13.0 SL-501-SA8-SB-4.0-5.0 SL-501-SA8-SB-9.0-10.0 SL-502-SA8-SB-0.0-0.5 SL-502-SA8-SB-4.0-5.0 SL-502-SA8-SB-7.5-8.5 SL-503-SA8-SB-0.0-0.5 SL-503-SA8-SB-3.0-4.0 SL-511-SA8-SB-0.0-0.5 SL-511-SA8-SB-6.0-7.0 SL-801-SA8-SB-0.0-0.5

Field Blank Outlier Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(REA2)	4/11/2013 3:00:00 PM	COPPER MOLYBDENUM	0.0036 mg/L 0.0036 mg/L	SL-501-SA8-SB-0.0-0.5 SL-501-SA8-SB-12.0-13.0 SL-501-SA8-SB-4.0-5.0 SL-501-SA8-SB-9.0-10.0 SL-502-SA8-SB-0.0-0.5 SL-502-SA8-SB-4.0-5.0 SL-502-SA8-SB-7.5-8.5 SL-503-SA8-SB-0.0-0.5 SL-503-SA8-SB-3.0-4.0 SL-511-SA8-SB-0.0-0.5 SL-511-SA8-SB-6.0-7.0 SL-801-SA8-SB-0.0-0.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-501-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.441 mg/Kg	0.441U mg/Kg
SL-501-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.417 mg/Kg	0.417U mg/Kg
SL-502-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.278 mg/Kg	0.278U mg/Kg
SL-503-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.191 mg/Kg	0.191U mg/Kg
SL-511-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.217 mg/Kg	0.217U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8151A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-501-SA8-SB-0.0-0.5MSD (SL-501-SA8-SB-0.0-0.5)	DINOSEB	-	-	10.00-46.00	62 (35.00)	DINOSEB	J (all detects)
SL-501-SA8-SB-0.0-0.5MS SL-501-SA8-SB-0.0-0.5MSD (SL-501-SA8-SB-0.0-0.5)	2,4,5-T 2,4,5-TP (Silvex) 2,4-D 2,4-DB	0 0 0 0	0 0 0 0	10.00-156.00 24.00-141.00 17.00-180.00 10.00-201.00	- - - -	2,4,5-T 2,4,5-TP (Silvex) 2,4-D 2,4-DB	J(all detects) R(all non-detects)

Method: 8015M
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-501-SA8-SB-0.0-0.5MS SL-501-SA8-SB-0.0-0.5MSD (SL-501-SA8-SB-0.0-0.5)	EFH (C21-C30) EFH (C30-C40)	-182 -286	-241 -356	49.00-123.00 49.00-123.00	- -	EFH (C21-C30) EFH (C30-C40)	No Qual, >4X

Method: 1613B
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-501-SA8-SB-0.0-0.5MS SL-501-SA8-SB-0.0-0.5MSD (SL-501-SA8-SB-0.0-0.5)	OCDD	11	-3	40.00-135.00	-	OCDD	J(all detects) R(all non-detects)

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-501-SA8-SB-0.0-0.5MS (TOT) SL-501-SA8-SB-0.0-0.5MSD (TOT) (SL-501-SA8-SB-0.0-0.5 SL-501-SA8-SB-12.0-13.0 SL-501-SA8-SB-4.0-5.0 SL-501-SA8-SB-9.0-10.0 SL-502-SA8-SB-0.0-0.5 SL-502-SA8-SB-4.0-5.0 SL-502-SA8-SB-7.5-8.5 SL-503-SA8-SB-0.0-0.5 SL-503-SA8-SB-3.0-4.0 SL-511-SA8-SB-0.0-0.5 SL-511-SA8-SB-6.0-7.0 SL-801-SA8-SB-0.0-0.5)	ALUMINUM MAGNESIUM TITANIUM	1766 140 334	2388 269 450	75.00-125.00 75.00-125.00 75.00-125.00	- - -	ALUMINUM MAGNESIUM TITANIUM	No Qual, >4X

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-501-SA8-SB-0.0-0.5MS (TOT) SL-501-SA8-SB-0.0-0.5MSD (TOT) (SL-501-SA8-SB-0.0-0.5 SL-501-SA8-SB-12.0-13.0 SL-501-SA8-SB-4.0-5.0 SL-501-SA8-SB-9.0-10.0 SL-502-SA8-SB-0.0-0.5 SL-502-SA8-SB-4.0-5.0 SL-502-SA8-SB-7.5-8.5 SL-503-SA8-SB-0.0-0.5 SL-503-SA8-SB-3.0-4.0 SL-511-SA8-SB-0.0-0.5 SL-511-SA8-SB-6.0-7.0 SL-801-SA8-SB-0.0-0.5)	CALCIUM IRON LEAD MANGANESE PHOSPHORUS	-71 -25 -80 -17 24	-21 1180 -84 -35 33	75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00 75.00-125.00	- - - - -	CALCIUM IRON LEAD MANGANESE PHOSPHORUS	J(all detects) UJ(all non-detects) Pb, PS=124% Ca, Fe, Mn, P No Qual, >4X
SL-501-SA8-SB-0.0-0.5MS (TOT) SL-501-SA8-SB-0.0-0.5MSD (TOT) (SL-501-SA8-SB-0.0-0.5 SL-501-SA8-SB-12.0-13.0 SL-501-SA8-SB-4.0-5.0 SL-501-SA8-SB-9.0-10.0 SL-502-SA8-SB-0.0-0.5 SL-502-SA8-SB-4.0-5.0 SL-502-SA8-SB-7.5-8.5 SL-503-SA8-SB-0.0-0.5 SL-503-SA8-SB-3.0-4.0 SL-511-SA8-SB-0.0-0.5 SL-511-SA8-SB-6.0-7.0 SL-801-SA8-SB-0.0-0.5)	ANTIMONY	48	45	75.00-125.00	-	ANTIMONY	J(all detects) UJ(all non-detects)

Method: 8270D SIM
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-501-SA8-SB-0.0-0.5MS (SL-501-SA8-SB-0.0-0.5)	Diethylphthalate	128	-	76.00-127.00	-	Diethylphthalate	J(all detects)

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-501-SA8-SB-0.0-0.5DUP (TOT) (SL-501-SA8-SB-0.0-0.5 SL-501-SA8-SB-12.0-13.0 SL-501-SA8-SB-4.0-5.0 SL-501-SA8-SB-9.0-10.0 SL-502-SA8-SB-0.0-0.5 SL-502-SA8-SB-4.0-5.0 SL-502-SA8-SB-7.5-8.5 SL-503-SA8-SB-0.0-0.5 SL-503-SA8-SB-3.0-4.0 SL-511-SA8-SB-0.0-0.5 SL-511-SA8-SB-6.0-7.0 SL-801-SA8-SB-0.0-0.5)	CADMIUM CALCIUM LEAD	27 32 25.6042mg/kg	20.00 20.00 ≤6.16	J (all detects) UJ (all non-detects) Cd, No Qual, Ok by difference

Method: 6020A
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-501-SA8-SB-0.0-0.5DUP (TOT) (SL-501-SA8-SB-0.0-0.5 SL-501-SA8-SB-12.0-13.0 SL-501-SA8-SB-4.0-5.0 SL-501-SA8-SB-9.0-10.0 SL-502-SA8-SB-0.0-0.5 SL-502-SA8-SB-4.0-5.0 SL-502-SA8-SB-7.5-8.5 SL-503-SA8-SB-0.0-0.5 SL-503-SA8-SB-3.0-4.0 SL-511-SA8-SB-0.0-0.5 SL-511-SA8-SB-6.0-7.0 SL-801-SA8-SB-0.0-0.5)	SILVER	37	20.00	No Qual, Ok by difference

Method: 7471B
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-501-SA8-SB-0.0-0.5DUP (TOT) (SL-501-SA8-SB-0.0-0.5 SL-501-SA8-SB-12.0-13.0 SL-501-SA8-SB-4.0-5.0 SL-501-SA8-SB-9.0-10.0 SL-502-SA8-SB-0.0-0.5 SL-502-SA8-SB-4.0-5.0 SL-502-SA8-SB-7.5-8.5 SL-503-SA8-SB-0.0-0.5 SL-503-SA8-SB-3.0-4.0 SL-511-SA8-SB-0.0-0.5 SL-511-SA8-SB-6.0-7.0 SL-801-SA8-SB-0.0-0.5)	MERCURY	78	20.00	No Qual, Ok by difference

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PrepPH067_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8151A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P31988AQ241603A (SL-501-SA8-SB-0.0-0.5 SL -501-SA8-SB-12.0-13.0 SL -501-SA8-SB-4.0-5.0 SL -501-SA8-SB-9.0-10.0 SL -801-SA8-SB-0.0-0.5)	2,4-D DICHLOROPROP	153 172	- -	59.00-122.00 65.00-158.00	- -	2,4-D DICHLOROPROP	J (all detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 160.3M
Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-501-SA8-SB-0.0-0.5	SL-801-SA8-SB-0.0-0.5			
MOISTURE	4.6	4.4	4		No Qualifiers Applied

Method: 1613B
Matrix: SO

Analyte	Concentration (ng/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-501-SA8-SB-0.0-0.5	SL-801-SA8-SB-0.0-0.5			
1,2,3,4,6,7,8-HPCDD	19.1	11.7	48	50.00	No Qualifiers Applied
1,2,3,4,6,7,8-HPCDF	3.22	2.24	36	50.00	
1,2,3,4,7,8,9-HPCDF	0.317	0.224	34	50.00	
1,2,3,4,7,8-HxCDD	0.303	0.265	13	50.00	
1,2,3,4,7,8-HxCDF	0.312	0.228	31	50.00	
1,2,3,6,7,8-HxCDD	1.14	0.803	35	50.00	
1,2,3,7,8,9-HxCDD	0.996	0.808	21	50.00	
1,2,3,7,8,9-HxCDF	0.415	0.431	4	50.00	
1,2,3,7,8-PCDF	1.09	1.18	8	50.00	
2,3,4,6,7,8-HxCDF	0.262	0.208	23	50.00	
2,3,7,8-TCDF	0.540	0.685	24	50.00	
OCDF	5.69	4.72	19	50.00	
1,2,3,6,7,8-HxCDF	0.379	0.132	97	50.00	
1,2,3,7,8-PCDD	0.263	0.149	55	50.00	
2,3,4,7,8-PCDF	0.653	0.208	103	50.00	
2,3,7,8-TCDD	0.0816	1.02 U	200	50.00	
OCDD	240	137	55	50.00	

Method: 6010C
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-501-SA8-SB-0.0-0.5 (TOT)	SL-801-SA8-SB-0.0-0.5 (TOT)			
ALUMINUM	17500	18400	5	50.00	No Qualifiers Applied
ARSENIC	3.89	4.41	13	50.00	
BARIUM	103	104	1	50.00	
BERYLLIUM	0.605	0.842	33	50.00	
CADMIUM	0.352	0.374	6	50.00	
CALCIUM	3910	5920	41	50.00	
CHROMIUM	20.7	21.4	3	50.00	
COBALT	5.94	5.67	5	50.00	
COPPER	12.2	11.2	9	50.00	
IRON	20900	21900	5	50.00	
LITHIUM	21.5	21.7	1	50.00	
MAGNESIUM	4540	4900	8	50.00	
MANGANESE	393	410	4	50.00	
NICKEL	12.4	12.6	2	50.00	
PHOSPHORUS	434	466	7	50.00	
POTASSIUM	3510	3740	6	50.00	
SODIUM	77.0	85.6	11	50.00	
TIN	2.77	2.68	3	50.00	
TITANIUM	1050	1150	9	50.00	
VANADIUM	37.3	40.9	9	50.00	
ZINC	62.7	65.3	4	50.00	
Zirconium	1.14	0.923	21	50.00	
BORON	6.11	16.1	90	50.00	J(all detects) UJ(all non-detects)
LEAD	35.3	18.4	63	50.00	
MOLYBDENUM	0.441	2.03 U	200	50.00	

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Field Duplicate RPD Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-501-SA8-SB-0.0-0.5 (TOT)	SL-801-SA8-SB-0.0-0.5 (TOT)			
SELENIUM	0.245	0.246	0	50.00	No Qualifiers Applied
SILVER	0.0421	0.0479	13	50.00	
STRONTIUM	20.9	26.0	22	50.00	
THALLIUM	0.284	0.289	2	50.00	

Method: 7471B

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-501-SA8-SB-0.0-0.5 (TOT)	SL-801-SA8-SB-0.0-0.5 (TOT)			
MERCURY	0.0276	0.0493	56	50.00	J(all detects)

Method: 8015M

Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-501-SA8-SB-0.0-0.5	SL-801-SA8-SB-0.0-0.5			
EFH (C15-C20)	2.6	2.4	8	50.00	No Qualifiers Applied
EFH (C21-C30)	24	20	18	50.00	
EFH (C30-C40)	51	41	22	50.00	

Method: 8081B

Matrix: SO

Analyte	Concentration (ug/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-501-SA8-SB-0.0-0.5	SL-801-SA8-SB-0.0-0.5			
4,4'-DDT	2.0	1.8	11	50.00	No Qualifiers Applied
4,4'-DDE	2.7	1.3	70	50.00	J(all detects)

Method: 8082A

Matrix: SO

Analyte	Concentration (ug/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-501-SA8-SB-0.0-0.5	SL-801-SA8-SB-0.0-0.5			
AROCLOR 1254	8.1	13	46	50.00	No Qualifiers Applied

Method: 8270D SIM

Matrix: SO

Analyte	Concentration (ug/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-501-SA8-SB-0.0-0.5	SL-801-SA8-SB-0.0-0.5			
2-METHYLNAPHTHALENE	1.3	1.0	26	50.00	No Qualifiers Applied
BENZO(K)FLUORANTHENE	7.5	9.3	21	50.00	
BIS(2-ETHYLHEXYL)PHTHALATE	13	10	26	50.00	
NAPHTHALENE	2.3	2.4	4	50.00	

Field Duplicate RPD Report

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8270D SIM					
Matrix: SO					
1-METHYLNAPHTHALENE	1.7 U	0.72	200	50.00	J(all detects) UJ(all non-detects)
ANTHRACENE	0.49	1.7 U	200	50.00	
BENZO(A)ANTHRACENE	3.3	0.77	124	50.00	
BENZO(A)PYRENE	5.2	0.94	139	50.00	
BENZO(B)FLUORANTHENE	6.9	2.2	103	50.00	
BENZO(E)PYRENE	4.4	18 U	200	50.00	
BENZO(G,H,I)PERYLENE	3.7	0.94	119	50.00	
CHRYSENE	8.0	3.9	69	50.00	
DIBENZO(A,H)ANTHRACENE	1.2	1.7 U	200	50.00	
Di-n-octylphthalate	19 U	10	200	50.00	
FLUORANTHENE	6.5	1.7	117	50.00	
INDENO(1,2,3-CD)PYRENE	3.4	0.70	132	50.00	
PHENANTHRENE	3.2	1.4	78	50.00	
PYRENE	6.9	2.1	107	50.00	

Method: 9045M					
Matrix: SO					
Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-501-SA8-SB-0.0-0.5	SL-801-SA8-SB-0.0-0.5			
PH	6.26	7.77	22	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-501-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	3.22	5.05	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JQ	0.317	5.05	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.303	5.05	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.312	5.05	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	1.14	5.05	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.379	5.05	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.996	5.05	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.415	5.05	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.263	5.05	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	1.09	5.05	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.262	5.05	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.653	5.05	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0816	1.01	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.540	1.01	PQL	ng/Kg	
	OCDF	JB	5.69	10.1	PQL	ng/Kg	
SL-501-SA8-SB-12.0-13.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.252	5.26	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0569	5.26	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	J	0.0304	5.26	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0331	5.26	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0301	5.26	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.0562	5.26	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.0198	5.26	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.0530	5.26	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0238	5.26	PQL	ng/Kg	
	OCDD	JB	2.26	10.5	PQL	ng/Kg	
OCDF	JB	0.183	10.5	PQL	ng/Kg		
SL-501-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JBQ	1.33	5.30	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.243	5.30	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0621	5.30	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0457	5.30	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.301	5.30	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.0985	5.30	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.471	5.30	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.123	5.30	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0925	5.30	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.132	5.30	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0570	5.30	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0585	5.30	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.105	1.06	PQL	ng/Kg	
	OCDF	JB	0.543	10.6	PQL	ng/Kg	
SL-501-SA8-SB-9.0-10.0	1,2,3,4,6,7,8-HPCDD	JB	0.138	5.33	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0422	5.33	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	J	0.0295	5.33	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0259	5.33	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.0503	5.33	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.0522	5.33	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0265	5.33	PQL	ng/Kg	
	OCDD	JBQ	1.06	10.7	PQL	ng/Kg	
	OCDF	JB	0.198	10.7	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-502-SA8-SB-0.0-0.5	1,2,3,4,7,8-HXCDF	JB	4.75	5.10	PQL	ng/Kg	J (all detects)
	1,2,3,6,7,8-HXCDF	JB	3.19	5.10	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	4.56	5.10	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	2.77	5.10	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	1.46	5.10	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.571	1.02	PQL	ng/Kg	
SL-502-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDF	JB	4.77	5.38	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JQ	0.577	5.38	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	J	1.75	5.38	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.715	5.38	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	2.30	5.38	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.566	5.38	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	2.14	5.38	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.513	5.38	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.906	5.38	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	1.75	5.38	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.396	5.38	PQL	ng/Kg	
	2,3,4,7,8-PECDF	J	0.440	5.38	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.149	1.08	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.205	1.08	PQL	ng/Kg	
	OCDF	JB	8.63	10.8	PQL	ng/Kg	
SL-502-SA8-SB-7.5-8.5	1,2,3,4,6,7,8-HPCDF	JBQ	1.83	5.24	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JQ	0.230	5.24	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	J	0.665	5.24	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.309	5.24	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.871	5.24	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.228	5.24	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.638	5.24	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JQ	0.324	5.24	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.268	5.24	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.541	5.24	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.227	5.24	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.206	5.24	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0961	1.05	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0929	1.05	PQL	ng/Kg	
OCDF	JB	3.40	10.5	PQL	ng/Kg		
SL-503-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	2.16	5.19	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JQ	0.209	5.19	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.270	5.19	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.303	5.19	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	0.868	5.19	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.441	5.19	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.815	5.19	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.471	5.19	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.148	5.19	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	2.52	5.19	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.213	5.19	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.210	5.19	PQL	ng/Kg	
	OCDF	JB	3.21	10.4	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-503-SA8-SB-3.0-4.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.403	5.17	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0653	5.17	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	J	0.0431	5.17	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0484	5.17	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.140	5.17	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0283	5.17	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.222	5.17	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.436	5.17	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0679	5.17	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.118	5.17	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0300	5.17	PQL	ng/Kg	
	OCDD	JB	2.77	10.3	PQL	ng/Kg	
	OCDF	JB	0.162	10.3	PQL	ng/Kg	
	SL-511-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.0766	5.01	PQL	
1,2,3,4,6,7,8-HPCDF		JBQ	0.0266	5.01	PQL	ng/Kg	
1,2,3,4,7,8,9-HPCDF		JQ	0.0230	5.01	PQL	ng/Kg	
1,2,3,4,7,8-HxCDD		JQ	0.0339	5.01	PQL	ng/Kg	
1,2,3,4,7,8-HXCDF		JBQ	0.0362	5.01	PQL	ng/Kg	
1,2,3,7,8,9-HXCDD		JBQ	0.0499	5.01	PQL	ng/Kg	
1,2,3,7,8,9-HXCDF		JQ	0.0613	5.01	PQL	ng/Kg	
2,3,4,6,7,8-HXCDF		JB	0.0157	5.01	PQL	ng/Kg	
OCDD		JBQ	0.345	10.0	PQL	ng/Kg	
OCDF		JBQ	0.0970	10.0	PQL	ng/Kg	
SL-511-SA8-SB-6.0-7.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.112	5.77	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0397	5.77	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JQ	0.0478	5.77	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0353	5.77	PQL	ng/Kg	
	OCDD	JBQ	0.347	11.5	PQL	ng/Kg	
SL-801-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	2.24	5.10	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JQ	0.224	5.10	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	J	0.265	5.10	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.228	5.10	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	0.803	5.10	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.132	5.10	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.808	5.10	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	J	0.431	5.10	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.149	5.10	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	1.18	5.10	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.208	5.10	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.208	5.10	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.685	1.02	PQL	ng/Kg	
	OCDF	JB	4.72	10.2	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-501-SA8-SB-0.0-0.5	ARSENIC	J	3.89	4.11	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.605	1.03	PQL	mg/Kg	
	BORON	J	6.11	10.3	PQL	mg/Kg	
	CADMIUM	J	0.352	1.03	PQL	mg/Kg	
	MOLYBDENUM	J	0.441	2.06	PQL	mg/Kg	
	SODIUM	J	77.0	103	PQL	mg/Kg	
	TIN	J	2.77	10.3	PQL	mg/Kg	
Zirconium	J	1.14	5.14	PQL	mg/Kg		
SL-501-SA8-SB-12.0-13.0	BERYLLIUM	J	0.767	1.03	PQL	mg/Kg	J (all detects)
	BORON	J	5.26	10.3	PQL	mg/Kg	
	CADMIUM	J	0.0910	1.03	PQL	mg/Kg	
	TIN	J	2.89	10.3	PQL	mg/Kg	
	Zirconium	J	2.62	5.17	PQL	mg/Kg	
SL-501-SA8-SB-4.0-5.0	BERYLLIUM	J	0.833	1.04	PQL	mg/Kg	J (all detects)
	BORON	J	5.00	10.4	PQL	mg/Kg	
	CADMIUM	J	0.111	1.04	PQL	mg/Kg	
	MOLYBDENUM	J	0.417	2.07	PQL	mg/Kg	
	TIN	J	3.04	10.4	PQL	mg/Kg	
	Zirconium	J	1.89	5.19	PQL	mg/Kg	
SL-501-SA8-SB-9.0-10.0	BERYLLIUM	J	0.863	1.08	PQL	mg/Kg	J (all detects)
	BORON	J	6.30	10.8	PQL	mg/Kg	
	CADMIUM	J	0.110	1.08	PQL	mg/Kg	
	TIN	J	3.12	10.8	PQL	mg/Kg	
	Zirconium	J	2.66	5.41	PQL	mg/Kg	
SL-502-SA8-SB-0.0-0.5	ARSENIC	J	4.11	4.21	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.562	1.05	PQL	mg/Kg	
	BORON	J	7.28	10.5	PQL	mg/Kg	
	CADMIUM	J	0.787	1.05	PQL	mg/Kg	
	SODIUM	J	82.3	105	PQL	mg/Kg	
	TIN	J	2.89	10.5	PQL	mg/Kg	
	Zirconium	J	2.27	5.26	PQL	mg/Kg	
SL-502-SA8-SB-4.0-5.0	BERYLLIUM	J	0.753	1.04	PQL	mg/Kg	J (all detects)
	BORON	J	4.78	10.4	PQL	mg/Kg	
	CADMIUM	J	0.221	1.04	PQL	mg/Kg	
	MOLYBDENUM	J	0.278	2.08	PQL	mg/Kg	
	TIN	J	3.08	10.4	PQL	mg/Kg	
	Zirconium	J	2.50	5.19	PQL	mg/Kg	
SL-502-SA8-SB-7.5-8.5	BERYLLIUM	J	0.602	1.07	PQL	mg/Kg	J (all detects)
	BORON	J	1.86	10.7	PQL	mg/Kg	
	CADMIUM	J	0.220	1.07	PQL	mg/Kg	
	TIN	J	2.95	10.7	PQL	mg/Kg	
	Zirconium	J	2.80	5.33	PQL	mg/Kg	
SL-503-SA8-SB-0.0-0.5	BERYLLIUM	J	0.614	1.03	PQL	mg/Kg	J (all detects)
	BORON	J	5.29	10.3	PQL	mg/Kg	
	CADMIUM	J	0.375	1.03	PQL	mg/Kg	
	MOLYBDENUM	J	0.191	2.07	PQL	mg/Kg	
	TIN	J	2.88	10.3	PQL	mg/Kg	
	Zirconium	J	2.26	5.17	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-503-SA8-SB-3.0-4.0	ARSENIC	J	2.66	4.19	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.400	1.05	PQL	mg/Kg	
	BORON	J	2.99	10.5	PQL	mg/Kg	
	CADMIUM	J	0.243	1.05	PQL	mg/Kg	
	TIN	J	2.71	10.5	PQL	mg/Kg	
	Zirconium	J	1.71	5.23	PQL	mg/Kg	
SL-511-SA8-SB-0.0-0.5	ARSENIC	J	3.19	4.09	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.578	1.02	PQL	mg/Kg	
	BORON	J	3.23	10.2	PQL	mg/Kg	
	CADMIUM	J	0.269	1.02	PQL	mg/Kg	
	MOLYBDENUM	J	0.217	2.05	PQL	mg/Kg	
	SODIUM	J	75.7	102	PQL	mg/Kg	
	TIN	J	2.71	10.2	PQL	mg/Kg	
	Zirconium	J	1.47	5.11	PQL	mg/Kg	
SL-511-SA8-SB-6.0-7.0	BERYLLIUM	J	0.595	1.13	PQL	mg/Kg	J (all detects)
	BORON	J	1.56	11.3	PQL	mg/Kg	
	CADMIUM	J	0.128	1.13	PQL	mg/Kg	
	TIN	J	3.54	11.3	PQL	mg/Kg	
	Zirconium	J	1.99	5.67	PQL	mg/Kg	
SL-801-SA8-SB-0.0-0.5	BERYLLIUM	J	0.842	1.02	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.374	1.02	PQL	mg/Kg	
	SODIUM	J	85.6	102	PQL	mg/Kg	
	TIN	J	2.68	10.2	PQL	mg/Kg	
	Zirconium	J	0.923	5.08	PQL	mg/Kg	

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-501-SA8-SB-0.0-0.5	SELENIUM	J	0.245	0.411	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0421	0.206	PQL	mg/Kg	
SL-501-SA8-SB-12.0-13.0	SILVER	J	0.0697	0.207	PQL	mg/Kg	J (all detects)
SL-501-SA8-SB-4.0-5.0	SELENIUM	J	0.283	0.415	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0344	0.207	PQL	mg/Kg	
SL-501-SA8-SB-9.0-10.0	SILVER	J	0.0487	0.216	PQL	mg/Kg	J (all detects)
SL-502-SA8-SB-0.0-0.5	SELENIUM	J	0.255	0.421	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0425	0.210	PQL	mg/Kg	
SL-502-SA8-SB-4.0-5.0	SELENIUM	J	0.189	0.415	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0428	0.208	PQL	mg/Kg	
SL-502-SA8-SB-7.5-8.5	SELENIUM	J	0.158	0.426	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0505	0.213	PQL	mg/Kg	
SL-503-SA8-SB-0.0-0.5	SELENIUM	J	0.225	0.414	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0964	0.207	PQL	mg/Kg	
SL-511-SA8-SB-0.0-0.5	SELENIUM	J	0.204	0.409	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0411	0.205	PQL	mg/Kg	
SL-511-SA8-SB-6.0-7.0	SELENIUM	J	0.142	0.454	PQL	mg/Kg	J (all detects)
	SILVER	J	0.104	0.227	PQL	mg/Kg	

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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ADR version 1.7.0.207

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Reporting Limit Outliers

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-801-SA8-SB-0.0-0.5	SELENIUM SILVER	J J	0.246 0.0479	0.406 0.203	PQL PQL	mg/Kg mg/Kg	J (all detects)

Method: 7471B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-501-SA8-SB-4.0-5.0	MERCURY	J	0.0107	0.0178	PQL	mg/Kg	J (all detects)
SL-502-SA8-SB-4.0-5.0	MERCURY	J	0.0164	0.0176	PQL	mg/Kg	J (all detects)
SL-502-SA8-SB-7.5-8.5	MERCURY	J	0.0118	0.0172	PQL	mg/Kg	J (all detects)

Method: 8015M
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-501-SA8-SB-0.0-0.5	EFH (C15-C20)	J	2.6	5.2	PQL	mg/Kg	J (all detects)
SL-502-SA8-SB-0.0-0.5	EFH (C15-C20)	J	2.9	5.2	PQL	mg/Kg	J (all detects)
SL-801-SA8-SB-0.0-0.5	EFH (C15-C20)	J	2.4	5.2	PQL	mg/Kg	J (all detects)

Method: 8081B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-801-SA8-SB-0.0-0.5	4,4'-DDE	J	1.3	1.8	PQL	ug/Kg	J (all detects)

Method: 8082A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-501-SA8-SB-0.0-0.5	AROCLOR 1254	J	8.1	18	PQL	ug/Kg	J (all detects)
SL-502-SA8-SB-4.0-5.0	AROCLOR 1254	J	6.8	18	PQL	ug/Kg	J (all detects)
SL-502-SA8-SB-7.5-8.5	Aroclor 5460	J	17	36	PQL	ug/Kg	J (all detects)
SL-801-SA8-SB-0.0-0.5	AROCLOR 1254	J	13	18	PQL	ug/Kg	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH067

Laboratory: LL

EDD Filename: PH067_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8270D SIM
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-501-SA8-SB-0.0-0.5	2-METHYLNAPHTHALENE	J	1.3	1.7	PQL	ug/Kg	J (all detects)
	ANTHRACENE	J	0.49	1.7	PQL	ug/Kg	
	BENZO(E)PYRENE	J	4.4	18	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	13	19	PQL	ug/Kg	
	DIBENZO(A,H)ANTHRACENE	J	1.2	1.7	PQL	ug/Kg	
SL-501-SA8-SB-4.0-5.0	BENZO(K)FLUORANTHENE	J	1.2	1.8	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	0.65	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	1.5	1.8	PQL	ug/Kg	
SL-502-SA8-SB-0.0-0.5	BIS(2-ETHYLHEXYL)PHTHALATE	J	13	19	PQL	ug/Kg	J (all detects)
	Di-n-butylphthalate	J	8.2	19	PQL	ug/Kg	
SL-502-SA8-SB-4.0-5.0	2-METHYLNAPHTHALENE	J	1.2	1.8	PQL	ug/Kg	J (all detects)
	ANTHRACENE	J	0.67	1.8	PQL	ug/Kg	
	BENZO(A)ANTHRACENE	J	0.82	1.8	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	11	19	PQL	ug/Kg	
SL-502-SA8-SB-7.5-8.5	ANTHRACENE	J	0.43	1.8	PQL	ug/Kg	J (all detects)
	BENZO(B)FLUORANTHENE	J	0.88	1.8	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	15	19	PQL	ug/Kg	
	CHRYSENE	J	1.4	1.8	PQL	ug/Kg	
	FLUORANTHENE	J	1.5	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	1.7	1.8	PQL	ug/Kg	
	PHENANTHRENE	J	1.1	1.8	PQL	ug/Kg	
	PYRENE	J	1.4	1.8	PQL	ug/Kg	
SL-503-SA8-SB-0.0-0.5	2-METHYLNAPHTHALENE	J	0.84	1.8	PQL	ug/Kg	J (all detects)
	BENZO(B)FLUORANTHENE	J	1.4	1.8	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	11	19	PQL	ug/Kg	
	FLUORANTHENE	J	1.4	1.8	PQL	ug/Kg	
	PHENANTHRENE	J	1.0	1.8	PQL	ug/Kg	
	PYRENE	J	1.2	1.8	PQL	ug/Kg	
SL-511-SA8-SB-0.0-0.5	BENZO(K)FLUORANTHENE	J	1.5	1.7	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	0.58	1.7	PQL	ug/Kg	
SL-801-SA8-SB-0.0-0.5	1-METHYLNAPHTHALENE	J	0.72	1.7	PQL	ug/Kg	J (all detects)
	2-METHYLNAPHTHALENE	J	1.0	1.7	PQL	ug/Kg	
	BENZO(A)ANTHRACENE	J	0.77	1.7	PQL	ug/Kg	
	BENZO(A)PYRENE	J	0.94	1.7	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	0.94	1.7	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	10	19	PQL	ug/Kg	
	Di-n-octylphthalate	J	10	19	PQL	ug/Kg	
	INDENO(1,2,3-CD)PYRENE	J	0.70	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	1.4	1.7	PQL	ug/Kg	

LDC #: 30289B4

VALIDATION COMPLETENESS WORKSHEET

Date: 9/4/13

SDG #: PH067

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: [Signature]

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	—	Sampling dates: 7/15/13
II.	ICP/MS Tune	—	
III.	Calibration	—	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	—	
VI.	Matrix Spike Analysis	SW	MSD
VII.	Duplicate Sample Analysis	SW	DUP
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	SW	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	—	(1,2)
XV.	Field Blanks	SW	EB = EB1-071713 / PH069 FB = FB-041113 = EB2-071713 / (PH029)

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

D = Duplicate
TB = Trip blank
EB = Equipment blank

Validated Samples: soil

1	SL-501-SA8-SB-0.0-0.5	11	SL-511-SA8-SB-0.0-0.5	21		31	
2	SL-801-SA8-SB-0.0-0.5	12	SL-511-SA8-SB-6.0-7.0	22		32	
3	SL-501-SA8-SB-4.0-5.0	13	SL-501-SA8-SB-0.0-0.5MS	23		33	
4	SL-501-SA8-SB-9.0-10.0	14	SL-501-SA8-SB-0.0-0.5MSD	24		34	
5	SL-501-SA8-SB-12.0-13.0	15	SL-501-SA8-SB-0.0-0.5DUP	25		35	
6	SL-502-SA8-SB-0.0-0.5	16		26		36	
7	SL-502-SA8-SB-4.0-5.0	17		27		37	
8	SL-502-SA8-SB-7.5-8.5	18		28		38	
9	SL-503-SA8-SB-0.0-0.5	19		29		39	
10	SL-503-SA8-SB-3.0-4.0	20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F
Sampling date: 4/11/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: Associated Samples: All

Analyte	Blank ID	Sample Identification											
	FB-041113 (SDG: PH029)	Action Limit	1	3	7	9	11						
Cu	0.0036	1.8											
Mo	0.0036	1.8	0.44	0.42	0.28	0.19	0.22						

Sampling date: 7/17/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: Associated Samples: 1-5, 9-12

Analyte	Blank ID	Sample Identification											
	EB1-071713 (SDG: PH069)	Action Limit	No Qualifiers										
Sb	0.0069	3.45											

Sampling date: 7/17/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: Associated Samples: 6-8

Analyte	Blank ID	Sample Identification											
	EB2-071713 (SDG: PH069)	Action Limit	No Qualifiers										
Sb	0.0054	2.7											

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

QUALITY ASSURANCE SUMMARY

FORM 5A (MS/MSD)

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

SDG No.: PH067

Matrix: SOIL

Level

(low/med):

LOW

Background Lab Sample ID: 7128144BKG Matrix Spike Lab Sample ID: 7128145MS Matrix Spike Duplicate Lab Sample ID: 7128146MSD
Batch Id(s): P19937B, P19938B

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Control Limit		M
		Result	C	Result	C	Result	C				%R	Q	%R	Q		%R	RPD	
Aluminum		16668.3235		20097.2223		21350.3755		194.1748	196.0784	MG/KG	1766	2388			6			20 P
Antimony		0.7255	U	23.4816		22.2569		48.5437	49.0196	MG/KG	48	N	45	N	5	75 - 125		20 P
Arsenic		3.7137	B	17.0650		18.0255		14.5631	14.7059	MG/KG	92		97		5	75 - 125		20 P
Barium		98.1598		287.4534		296.9206		194.1748	196.0784	MG/KG	97		101		3	75 - 125		20 P
Beryllium		0.5775	B	5.5184		5.6098		4.8544	4.9020	MG/KG	102		103		2	75 - 125		20 P
Boron		5.8324	B	196.0505		198.0912		194.1748	196.0784	MG/KG	98		98		1	75 - 125		20 P
Cadmium		0.3363	B	4.9320		4.9608		4.8544	4.9020	MG/KG	95		94		1	75 - 125		20 P
Calcium		3729.5059		3453.8631		3648.5059		388.3495	392.1569	MG/KG	-71		-24		5			20 P
Chromium		19.7284		39.3893		40.7157		19.4175	19.6078	MG/KG	101		107		3	75 - 125		20 P
Cobalt		5.6647		51.7214		52.1118		48.5437	49.0196	MG/KG	95		95		1	75 - 125		20 P
Copper		11.6784		36.6650		37.4882		24.2718	24.5098	MG/KG	103		105		2	75 - 125		20 P
Iron		19928.7529		19904.9058		21085.4520		97.0874	98.0392	MG/KG	-25		1180		6			20 P
Lead		33.6775		22.0049		21.2569		14.5631	14.7059	MG/KG	-80	N	-84	N	3	75 - 125		20 P
Lithium		20.5216		116.5981		117.7167		97.0874	98.0392	MG/KG	99		99		1	75 - 125		20 P
Magnesium		4934.6922		4606.9660		4861.2500		194.1748	196.0784	MG/KG	140		269		5			20 P
Manganese		375.1471		366.8592		357.7461		48.5437	49.0196	MG/KG	-17		-35		3			20 P
Mercury		0.0263		0.1916		0.1849		0.1584	0.1542	MG/KG	104		103		4	65 - 135		20 CV
Molybdenum		0.4206	B	180.1320		180.7863		194.1748	196.0784	MG/KG	93		92		0	75 - 125		20 P
Nickel		11.8471		57.5641		58.3990		48.5437	49.0196	MG/KG	94		95		1	75 - 125		20 P
Phosphorus		414.4588		437.5893		447.2961		97.0874	98.0392	MG/KG	24		33		2			20 P
Potassium		3347.0755		4413.7845		4551.2029		970.8738	980.3922	MG/KG	110		123		3	75 - 125		20 P
Selenium	78	0.2337	B	2.1447		2.1180		1.9417	1.9608	MG/KG	98		96		1	75 - 125		20 MS
Silver	107	0.0402	B	10.4315		10.5716		9.7087	9.8039	MG/KG	107		107		1	75 - 125		20 MS
Sodium		73.4725	B	1041.5971		1043.9069		970.8738	980.3922	MG/KG	100		99		0	75 - 125		20 P
Strontium	88	19.9035		28.8581		27.5971		7.7670	7.8431	MG/KG	115		98		4	75 - 125		20 MS
Thallium	203	0.2710		0.6833		0.6675		0.3883	0.3922	MG/KG	106		101		2	75 - 125		20 MS

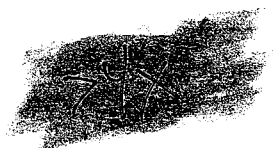
PS=124%
(S/U/A)

Note: Results shown are reported on an as-received basis.

<p>METHODS: P = ICP Atomic Emission Spectrometer CV = Cold Vapor MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS: U= Below MDL, B= Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS</p>
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Lancaster
Laboratories



QUALITY ASSURANCE SUMMARY

FORM 5A (MS/MSD)

MATRIX SPIKE/MATRIX SPIKE DUPLICATE

SDG No.: PH067

Matrix: SOIL

Level

(low/med):

LOW

Background Lab Sample ID: 7128144BKG Matrix Spike Lab Sample ID: 7128145MS Matrix Spike Duplicate Lab Sample ID: 7128146MSD
Batch Id(s): P19937B, P19938B

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		Control Limit		M		
		Result	C	Result	C	Result	C				%R	Q	%R	Q	RPD	Q		%R	RPD
Tin		2.6392	B	335.2583		335.0451		388.3495	392.1569	MG/KG	86		85		0	75 - 125	20	P	
Tin		2.6392	B	335.2583		335.0451		388.3495	392.1569	MG/KG	86		85		0				P
Vanadium		35.5667		86.1340		89.6196		48.5437	49.0196	MG/KG	104		110		4	75 - 125	20	P	
Zinc		59.8510		100.6777		103.1010		48.5437	49.0196	MG/KG	84		88		2	75 - 125	20	P	
Zirconium		1.0922	B	82.2029		81.4716		97.0874	98.0392	MG/KG	84		82		1	75 - 125	20	P	

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer CV = Cold Vapor</p> <p>MS = ICP Mass Spectrometry AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U= Below MDL, B= Below LOQ</p> <p>FLAGS:</p> <p>N = Matrix Spike OOS, * = Duplicate OOS</p>
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Background Lab Sample ID: 7128144BKG
 Batch ID(s): P19937B, P19938B
 Concentration Units: MG/KG

Duplicate Lab Sample ID: 7128147DUP

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			16668.3235		16490.7733		1		P
Antimony			1.2284	B	0.8911	B	32		P
Arsenic		3.9	3.7137	B	4.2119		13		P
Barium			98.1598		94.3050		4		P
Beryllium			0.5775	B	0.5931	B	3		P
Boron			5.8324	B	5.4723	B	6		P
Cadmium			0.3363	B	0.2554	B	27		P
Calcium			3729.5059		2698.5911		32	*	P
Chromium			19.7284		19.5139		1		P
Cobalt			5.6647		5.6158		1		P
Copper			11.6784		10.8525		7		P
Iron			19928.7529		19582.4980		2		P
Lead		2.9	33.6775		8.0733		123	*	P
Lithium		3.9	20.5216		19.1287		7		P
Magnesium			4334.6922		4227.9119		2		P
Manganese			375.1471		306.2782		20		P
Mercury		0.0	0.0263		0.0415	B	78		CV
Molybdenum			0.4206	B	0.4307	B	2		P
Nickel			11.8471		11.2366		5		P
Phosphorus			414.4588		368.2822		12		P
Potassium			3347.0755		2987.7941		11		P
Selenium	78		0.2337	B	0.2410	B	3		MS
Silver	107		0.0402	B	0.0277	B	37		MS
Sodium			73.4725	B	72.0960	B	2		P
Strontium	88		19.9035		20.5851		3		MS
Thallium	203	0.2	0.2710		0.2990		10		MS
Tin			2.6392	B	2.5465	B	4		P
Titanium			998.5137		1012.5406		1		P
Vanadium			35.5667		35.8624		1		P
Zinc			59.8510		51.8733		14		P
Zirconium			1.0922	B	0.9465	B	14		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).
 The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

*pb: out by difference
 25.6042 mg/kg (≤ 6.16)
 J/JJA*

25.6042 mg/kg difference

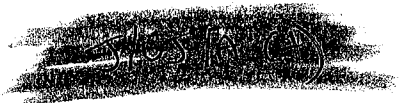
Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ FLAGS: Duplicate Out of Spec
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Background Lab Sample ID: 7128144BKG Serial Dilution Lab Sample ID: 7128144L
 Batch ID(s): P19937B
 Concentration Units: UG/L

Analyte	Mass	Initial Sample Result (I)	C	Serial Dilution Result (S)	C	% Diff.	Q	M
Aluminum		170016.9000		179629.4500		6		P
Antimony		7.4000	U	37.0000	U			P
Arsenic		37.8800	B	35.0000	U	100		P
Barium		1001.2300		1060.6500		6		P
Beryllium		5.8900	B	6.1500	B	4		P
Boron		59.4900	B	85.7000	B	44		P
Cadmium		3.4300	B	3.9500	B	15		P
Calcium		38040.9600		40342.2000		6		P
Chromium		201.2300		210.0000		4		P
Cobalt		57.7800		64.7000		12		P
Copper		119.1200		126.4000		6		P
Iron		203273.2800		209386.8000		3		P
Lead		343.5100		390.8000		14		P
Lithium		209.3200		235.9000		13		P
Magnesium		44213.8600		47144.4000		7		P
Manganese		3826.5000		4060.8000		6		P
Molybdenum		4.2900	B	15.4500	B	260		P
Nickel		120.8400		125.2500		4		P
Phosphorus		4227.4800		4347.9500		3		P
Potassium		34140.1700		35250.8000		3		P
Selenium	78	1.1920	B	2.5000	U	100		MS
Silver	107	0.2050	B	0.6500	U	100		MS
Sodium		749.4200	B	835.0000	U	100		P
Strontium	88	101.5080		99.7700		2		MS
Thallium	203	1.3820		1.5500	B	12		MS
Tin		26.9200	B	32.2000	B	20		P
Titanium		10184.8400		10504.0500		3		P
Vanadium		362.7800		373.2000		3		P
Zinc		610.4800		630.4000		3		P
Zirconium		11.1400	B	42.0000	U	100		P

NOTE: An E in column Q indicates the presence of a chemical or physical interference in the matrix when the % difference is greater than 10%. This applies only when (I) is greater than or equal to 50x MDL for ICP, 100x MDL for ICP-MS (6020), 50x MDL for ICP-MS (200.8), or 25x MDL for GFAA.



<p>METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry</p>	<p>CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ</p> <p>FLAGS: E = Matrix Effects exist as proven by Serial Dilution or Spiked Dilution</p>
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**Data Validation Report
Santa Susana Field Laboratory**

Subarea 8

SDG: PH068

Prepared for

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Prepared by

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October 3, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on July 16th, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Volatiles (VOAs) by EPA SW 846 Method 8260B
Semivolatiles (SVOAs) by EPA SW 846 Method 8270D
(SVOAs) by EPA SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A, and 7471B
Total Petroleum Hydrocarbon as Gasoline (TPH-G) by EPA SW 846 Method 8015M
Total Petroleum Hydrocarbons as Extractable (TPH-E) by EPA SW 846 Method 8015M
Dioxins/Dibenzofurans by EPA Method 1613B
Perchlorate by EPA method 6850
Hexavalent Chromium by EPA method 7199

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks, and field duplicate samples. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of two blanks for dioxins and metals. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for VOAs, SVOAs, and metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ), as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits with the exception of one DUP for metals and hexavalent chromium. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosure I.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of two LCS/LCSD pairs for VOAs. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding LCS/LCSD outliers are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met with the following exceptions:

Diluted Sample	Analyte	%D (Limits)	Associated Samples	Flag	A or P
SL-504-SA8-SB-2.5-3.5	Lithium	18 (≤10)	All soil samples in SDG PH068	J (all detects) UJ (all non-detects)	A

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH068	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

One field duplicate pair was collected and analyzed for VOAs, SVOAs, PCBs, metals, perchlorate, TPH-G, TPH-E, hexavalent chromium, and dioxins. All RPDs were within QC limits with the exception of several dioxins, metals, and hexavalent chromium. In these duplicate pairs, the associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The field duplicate result comparisons are provided in Enclosure I.

XIV. Field Blank Samples

One trip blank was collected and analyzed for VOAs and TPH-G. No contaminant concentrations were found in the trip blank.

Two equipment blanks (from SDG PH069) were collected and analyzed for VOAs, SVOAs, PCBs, metals, perchlorate, TPH-G, TPH-E, hexavalent chromium, and dioxins. The equipment

blanks had detections for several VOAs, SVOAs, metals, and dioxins. All associated sample results were not detected or were significantly greater than the concentrations found in the equipment blanks and were not qualified.

One field blank (from SDG PH029) was collected and analyzed for VOAs, SVOAs, PCBs, metals, perchlorate, TPH-G, TPH-E, hexavalent chromium, and dioxins. The field blank had detections for several VOAs, SVOAs, metals, and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. Overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
16-Jul-2013	TB-071613	7129592	TB	5030B	8015M	III
16-Jul-2013	TB-071613	7129592	TB	5030B	8260B	III
16-Jul-2013	SL-510-SA8-SB-0.0-0.5	7129591	N	3050B	6010C	III
16-Jul-2013	SL-510-SA8-SB-0.0-0.5	7129591	N	3050B	6020A	III
16-Jul-2013	SL-510-SA8-SB-0.0-0.5	7129591	N	3546	8015M	III
16-Jul-2013	SL-510-SA8-SB-0.0-0.5	7129591	N	3546	8082A	III
16-Jul-2013	SL-510-SA8-SB-0.0-0.5	7129591	N	3546	8270D SIM	III
16-Jul-2013	SL-510-SA8-SB-0.0-0.5	7129591	N	METHOD	1613B	III
16-Jul-2013	SL-510-SA8-SB-0.0-0.5	7129591	N	METHOD	7471B	III
16-Jul-2013	SL-505-SA8-SB-0.0-0.5	7129593	N	3050B	6010C	III
16-Jul-2013	SL-505-SA8-SB-0.0-0.5	7129593	N	3050B	6020A	III
16-Jul-2013	SL-505-SA8-SB-0.0-0.5	7129593	N	3546	8015M	III
16-Jul-2013	SL-505-SA8-SB-0.0-0.5	7129593	N	3546	8082A	III
16-Jul-2013	SL-505-SA8-SB-0.0-0.5	7129593	N	3546	8270D SIM	III
16-Jul-2013	SL-505-SA8-SB-0.0-0.5	7129593	N	METHOD	1613B	III
16-Jul-2013	SL-505-SA8-SB-0.0-0.5	7129593	N	METHOD	7471B	III
16-Jul-2013	SL-505-SA8-SB-0.0-0.5MSD	P129593M260649	MSD	3546	8270D SIM	III
16-Jul-2013	SL-505-SA8-SB-0.0-0.5MS	P129593R260618	MS	3546	8270D SIM	III
16-Jul-2013	SL-505-SA8-SB-4.0-5.0	7129594	N	3050B	6010C	III
16-Jul-2013	SL-505-SA8-SB-4.0-5.0	7129594	N	3050B	6020A	III
16-Jul-2013	SL-505-SA8-SB-4.0-5.0	7129594	N	3546	8015M	III
16-Jul-2013	SL-505-SA8-SB-4.0-5.0	7129594	N	3546	8082A	III
16-Jul-2013	SL-505-SA8-SB-4.0-5.0	7129594	N	3546	8270D SIM	III
16-Jul-2013	SL-505-SA8-SB-4.0-5.0	7129594	N	5035A	8015M	III
16-Jul-2013	SL-505-SA8-SB-4.0-5.0	7129594	N	METHOD	1613B	III
16-Jul-2013	SL-505-SA8-SB-4.0-5.0	7129594	N	METHOD	7471B	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
16-Jul-2013	SL-504-SA8-SB-0.0-0.5	7129595	N	3050B	6010C	III
16-Jul-2013	SL-504-SA8-SB-0.0-0.5	7129595	N	3050B	6020A	III
16-Jul-2013	SL-504-SA8-SB-0.0-0.5	7129595	N	3060A	7199	III
16-Jul-2013	SL-504-SA8-SB-0.0-0.5	7129595	N	3546	8015M	III
16-Jul-2013	SL-504-SA8-SB-0.0-0.5	7129595	N	3546	8082A	III
16-Jul-2013	SL-504-SA8-SB-0.0-0.5	7129595	N	3546	8270D	III
16-Jul-2013	SL-504-SA8-SB-0.0-0.5	7129595	N	3546	8270D SIM	III
16-Jul-2013	SL-504-SA8-SB-0.0-0.5	7129595	N	METHOD	1613B	III
16-Jul-2013	SL-504-SA8-SB-0.0-0.5	7129595	N	METHOD	6850	III
16-Jul-2013	SL-504-SA8-SB-0.0-0.5	7129595	N	METHOD	7471B	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	3050B	6010C	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	3050B	6020A	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	3060A	7199	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	3546	8015M	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	3546	8082A	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	3546	8270D	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	3546	8270D SIM	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	5035A	8015M	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	5035A	8260B	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	METHOD	1613B	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	METHOD	6850	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5	7129596	N	METHOD	7471B	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	3050B	6010C	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	3050B	6020A	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	3546	8015M	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	3546	8082A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	3546	8270D	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	3546	8270D SIM	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	5035A	8015M	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	5035A	8260B	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	METHOD	1613B	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	METHOD	6850	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129597	MS	METHOD	7471B	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	3050B	6010C	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	3050B	6020A	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	3546	8015M	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	3546	8082A	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	3546	8270D	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	3546	8270D SIM	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	5035A	8015M	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	5035A	8260B	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	METHOD	1613B	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	METHOD	6850	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MSD	7129598	MSD	METHOD	7471B	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129599	MS	3060A	7199	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5MS	7129600	MS	3060A	7199	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5DUP	7129602	DUP	3050B	6010C	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5DUP	7129602	DUP	3050B	6020A	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5DUP	7129602	DUP	3060A	7199	III
16-Jul-2013	SL-504-SA8-SB-2.5-3.5DUP	7129602	DUP	METHOD	7471B	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	3050B	6010C	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	3050B	6020A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	3060A	7199	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	3546	8015M	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	3546	8082A	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	3546	8270D	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	3546	8270D SIM	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	5035A	8015M	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	5035A	8260B	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	METHOD	1613B	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	METHOD	6850	III
16-Jul-2013	SL-804-SA8-SB-2.5-3.5	7129603	FD	METHOD	7471B	III
16-Jul-2013	SL-507-SA8-SB-0.0-0.5	7129604	N	3050B	6010C	III
16-Jul-2013	SL-507-SA8-SB-0.0-0.5	7129604	N	3050B	6020A	III
16-Jul-2013	SL-507-SA8-SB-0.0-0.5	7129604	N	3060A	7199	III
16-Jul-2013	SL-507-SA8-SB-0.0-0.5	7129604	N	3546	8015M	III
16-Jul-2013	SL-507-SA8-SB-0.0-0.5	7129604	N	3546	8082A	III
16-Jul-2013	SL-507-SA8-SB-0.0-0.5	7129604	N	3546	8270D	III
16-Jul-2013	SL-507-SA8-SB-0.0-0.5	7129604	N	3546	8270D SIM	III
16-Jul-2013	SL-507-SA8-SB-0.0-0.5	7129604	N	METHOD	1613B	III
16-Jul-2013	SL-507-SA8-SB-0.0-0.5	7129604	N	METHOD	6850	III
16-Jul-2013	SL-507-SA8-SB-0.0-0.5	7129604	N	METHOD	7471B	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	3050B	6010C	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	3050B	6020A	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	3060A	7199	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	3546	8015M	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	3546	8082A	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	3546	8270D	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	3546	8270D SIM	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	5035A	8015M	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	5035A	8260B	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	METHOD	1613B	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	METHOD	6850	III
16-Jul-2013	SL-507-SA8-SB-3.0-4.0	7129605	N	METHOD	7471B	III
16-Jul-2013	SL-508-SA8-SB-0.0-0.5	7129606	N	3050B	6010C	III
16-Jul-2013	SL-508-SA8-SB-0.0-0.5	7129606	N	3050B	6020A	III
16-Jul-2013	SL-508-SA8-SB-0.0-0.5	7129606	N	3060A	7199	III
16-Jul-2013	SL-508-SA8-SB-0.0-0.5	7129606	N	3546	8015M	III
16-Jul-2013	SL-508-SA8-SB-0.0-0.5	7129606	N	3546	8082A	III
16-Jul-2013	SL-508-SA8-SB-0.0-0.5	7129606	N	3546	8270D	III
16-Jul-2013	SL-508-SA8-SB-0.0-0.5	7129606	N	3546	8270D SIM	III
16-Jul-2013	SL-508-SA8-SB-0.0-0.5	7129606	N	METHOD	1613B	III
16-Jul-2013	SL-508-SA8-SB-0.0-0.5	7129606	N	METHOD	6850	III
16-Jul-2013	SL-508-SA8-SB-0.0-0.5	7129606	N	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-504-SA8-SB-0.0-0.5 Collected: 7/16/2013 11:45:00 Analysis Type: REA2 Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
LITHIUM	30.1		0.35	MDL	4.1	PQL	mg/Kg	J	A

Sample ID: SL-504-SA8-SB-0.0-0.5 Collected: 7/16/2013 11:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.06	U	0.752	MDL	4.06	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.641	J	0.0680	MDL	1.02	PQL	mg/Kg	J	Z
CADMIUM	0.513	J	0.0772	MDL	1.02	PQL	mg/Kg	J	Z
CHROMIUM	34.9		0.162	MDL	3.05	PQL	mg/Kg	J	Q
MOLYBDENUM	0.518	J	0.173	MDL	2.03	PQL	mg/Kg	U	F
POTASSIUM	6080		8.47	MDL	102	PQL	mg/Kg	J	Q
TIN	3.11	J	0.223	MDL	10.2	PQL	mg/Kg	U	B
Zirconium	2.16	J	0.853	MDL	5.08	PQL	mg/Kg	J	Z

Sample ID: SL-504-SA8-SB-2.5-3.5 Collected: 7/16/2013 12:00:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
LITHIUM	29.8		0.35	MDL	4.1	PQL	mg/Kg	J	A

Sample ID: SL-504-SA8-SB-2.5-3.5 Collected: 7/16/2013 12:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.08	U	0.755	MDL	4.08	PQL	mg/Kg	UJ	Q
ARSENIC	3.88	J	0.714	MDL	4.08	PQL	mg/Kg	J	Z
BERYLLIUM	0.394	J	0.0684	MDL	1.02	PQL	mg/Kg	J	Z
CADMIUM	0.370	J	0.0775	MDL	1.02	PQL	mg/Kg	J	Z
CHROMIUM	21.8		0.163	MDL	3.06	PQL	mg/Kg	J	Q
MOLYBDENUM	0.373	J	0.173	MDL	2.04	PQL	mg/Kg	UJ	FD, F
POTASSIUM	2220		8.51	MDL	102	PQL	mg/Kg	J	Q
TIN	2.86	J	0.224	MDL	10.2	PQL	mg/Kg	U	B
Zirconium	3.17	J	0.857	MDL	5.10	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-505-SA8-SB-0.0-0.5	Collected: 7/16/2013 8:55:00	Analysis Type: REA2	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
LITHIUM	33.0		0.35	MDL	4.2	PQL	mg/Kg	J	A

Sample ID: SL-505-SA8-SB-0.0-0.5	Collected: 7/16/2013 8:55:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.16	U	0.770	MDL	4.16	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.692	J	0.0697	MDL	1.04	PQL	mg/Kg	J	Z
CADMIUM	0.543	J	0.0791	MDL	1.04	PQL	mg/Kg	J	Z
CHROMIUM	34.5		0.167	MDL	3.12	PQL	mg/Kg	J	Q
MOLYBDENUM	0.627	J	0.177	MDL	2.08	PQL	mg/Kg	U	F
POTASSIUM	6120		8.68	MDL	104	PQL	mg/Kg	J	Q
TIN	3.19	J	0.229	MDL	10.4	PQL	mg/Kg	U	B
Zirconium	1.60	J	0.874	MDL	5.20	PQL	mg/Kg	J	Z

Sample ID: SL-505-SA8-SB-4.0-5.0	Collected: 7/16/2013 9:30:00	Analysis Type: REA	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
LITHIUM	29.0		0.37	MDL	4.4	PQL	mg/Kg	J	A

Sample ID: SL-505-SA8-SB-4.0-5.0	Collected: 7/16/2013 9:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.39	U	0.811	MDL	4.39	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.689	J	0.0735	MDL	1.10	PQL	mg/Kg	J	Z
CADMIUM	0.357	J	0.0833	MDL	1.10	PQL	mg/Kg	J	Z
CHROMIUM	26.9		0.175	MDL	3.29	PQL	mg/Kg	J	Q
POTASSIUM	2270		9.14	MDL	110	PQL	mg/Kg	J	Q
TIN	3.18	J	0.241	MDL	11.0	PQL	mg/Kg	U	B
Zirconium	3.50	J	0.921	MDL	5.48	PQL	mg/Kg	J	Z

Sample ID: SL-507-SA8-SB-0.0-0.5	Collected: 7/16/2013 12:55:00	Analysis Type: REA	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
LITHIUM	25.7		0.34	MDL	4.0	PQL	mg/Kg	J	A

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	Matrix:	SO
Method:	6010C		

Sample ID: SL-507-SA8-SB-0.0-0.5 Collected: 7/16/2013 12:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	3.97	U	0.734	MDL	3.97	PQL	mg/Kg	UJ	Q
ARSENIC	3.27	J	0.694	MDL	3.97	PQL	mg/Kg	J	Z
BERYLLIUM	0.403	J	0.0664	MDL	0.992	PQL	mg/Kg	J	Z
CADMIUM	0.322	J	0.0754	MDL	0.992	PQL	mg/Kg	J	Z
CHROMIUM	18.7		0.159	MDL	2.98	PQL	mg/Kg	J	Q
MOLYBDENUM	0.343	J	0.169	MDL	1.98	PQL	mg/Kg	U	F
POTASSIUM	3440		8.27	MDL	99.2	PQL	mg/Kg	J	Q
TIN	2.89	J	0.218	MDL	9.92	PQL	mg/Kg	U	B
Zirconium	2.37	J	0.833	MDL	4.96	PQL	mg/Kg	J	Z

Sample ID: SL-507-SA8-SB-3.0-4.0 Collected: 7/16/2013 1:10:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
LITHIUM	22.2		0.35	MDL	4.1	PQL	mg/Kg	J	A

Sample ID: SL-507-SA8-SB-3.0-4.0 Collected: 7/16/2013 1:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.10	U	0.759	MDL	4.10	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.381	J	0.0687	MDL	1.03	PQL	mg/Kg	J	Z
CADMIUM	0.232	J	0.0780	MDL	1.03	PQL	mg/Kg	J	Z
CHROMIUM	13.6		0.164	MDL	3.08	PQL	mg/Kg	J	Q
POTASSIUM	1840		8.56	MDL	103	PQL	mg/Kg	J	Q
TIN	2.96	J	0.226	MDL	10.3	PQL	mg/Kg	U	B
Zirconium	3.17	J	0.862	MDL	5.13	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA8-SB-0.0-0.5 Collected: 7/16/2013 1:30:00 Analysis Type: REA2 Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
LITHIUM	27.2		0.35	MDL	4.1	PQL	mg/Kg	J	A

Sample ID: SL-508-SA8-SB-0.0-0.5 Collected: 7/16/2013 1:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.11	U	0.759	MDL	4.11	PQL	mg/Kg	UJ	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-508-SA8-SB-0.0-0.5	Collected: 7/16/2013 1:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.582	J	0.0688	MDL	1.03	PQL	mg/Kg	J	Z
CADMIUM	0.251	J	0.0780	MDL	1.03	PQL	mg/Kg	J	Z
CHROMIUM	30.4		0.164	MDL	3.08	PQL	mg/Kg	J	Q
MOLYBDENUM	0.574	J	0.174	MDL	2.05	PQL	mg/Kg	U	F
POTASSIUM	5690		8.56	MDL	103	PQL	mg/Kg	J	Q
TIN	2.77	J	0.226	MDL	10.3	PQL	mg/Kg	U	B
Zirconium	2.38	J	0.862	MDL	5.13	PQL	mg/Kg	J	Z

Sample ID: SL-510-SA8-SB-0.0-0.5	Collected: 7/16/2013 8:30:00	Analysis Type: REA	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
LITHIUM	24.0		0.34	MDL	4.0	PQL	mg/Kg	J	A

Sample ID: SL-510-SA8-SB-0.0-0.5	Collected: 7/16/2013 8:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	3.97	U	0.735	MDL	3.97	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.462	J	0.0666	MDL	0.993	PQL	mg/Kg	J	Z
CADMIUM	0.393	J	0.0755	MDL	0.993	PQL	mg/Kg	J	Z
CHROMIUM	16.5		0.159	MDL	2.98	PQL	mg/Kg	J	Q
MOLYBDENUM	0.373	J	0.169	MDL	1.99	PQL	mg/Kg	U	F
POTASSIUM	3330		8.28	MDL	99.3	PQL	mg/Kg	J	Q
SODIUM	82.4	J	16.6	MDL	99.3	PQL	mg/Kg	J	Z
TIN	2.85	J	0.219	MDL	9.93	PQL	mg/Kg	U	B
Zirconium	1.57	J	0.834	MDL	4.97	PQL	mg/Kg	J	Z

Sample ID: SL-804-SA8-SB-2.5-3.5	Collected: 7/16/2013 12:10:00	Analysis Type: REA	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
LITHIUM	29.5		0.34	MDL	4.0	PQL	mg/Kg	J	A

Sample ID: SL-804-SA8-SB-2.5-3.5	Collected: 7/16/2013 12:10:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.05	U	0.748	MDL	4.05	PQL	mg/Kg	UJ	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-804-SA8-SB-2.5-3.5 Collected: 7/16/2013 12:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	3.68	J	0.708	MDL	4.05	PQL	mg/Kg	J	Z
BERYLLIUM	0.449	J	0.0678	MDL	1.01	PQL	mg/Kg	J	Z
CADMIUM	0.330	J	0.0769	MDL	1.01	PQL	mg/Kg	J	Z
CHROMIUM	22.0		0.162	MDL	3.03	PQL	mg/Kg	J	Q
MOLYBDENUM	2.02	U	0.172	MDL	2.02	PQL	mg/Kg	UJ	FD
POTASSIUM	2240		8.43	MDL	101	PQL	mg/Kg	J	Q
TIN	2.99	J	0.222	MDL	10.1	PQL	mg/Kg	U	B
Zirconium	2.85	J	0.850	MDL	5.06	PQL	mg/Kg	J	Z

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-504-SA8-SB-0.0-0.5 Collected: 7/16/2013 11:45:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.251	J	0.102	MDL	0.406	PQL	mg/Kg	J	Z

Sample ID: SL-504-SA8-SB-0.0-0.5 Collected: 7/16/2013 11:45:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0343	J	0.0264	MDL	0.203	PQL	mg/Kg	J	Z

Sample ID: SL-505-SA8-SB-0.0-0.5 Collected: 7/16/2013 8:55:00 Analysis Type: REA Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.329	J	0.104	MDL	0.416	PQL	mg/Kg	J	Z

Sample ID: SL-505-SA8-SB-0.0-0.5 Collected: 7/16/2013 8:55:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0558	J	0.0271	MDL	0.208	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-505-SA8-SB-4.0-5.0		Collected: 7/16/2013 9:30:00		Analysis Type: REA			Dilution: 2		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.132	J	0.110	MDL	0.439	PQL	mg/Kg	J	Z

Sample ID: SL-505-SA8-SB-4.0-5.0		Collected: 7/16/2013 9:30:00		Analysis Type: RES			Dilution: 2		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0328	J	0.0285	MDL	0.219	PQL	mg/Kg	J	Z

Sample ID: SL-507-SA8-SB-0.0-0.5		Collected: 7/16/2013 12:55:00		Analysis Type: REA			Dilution: 2		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.186	J	0.0992	MDL	0.397	PQL	mg/Kg	J	Z

Sample ID: SL-507-SA8-SB-3.0-4.0		Collected: 7/16/2013 1:10:00		Analysis Type: RES			Dilution: 2		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
THALLIUM	0.192	J	0.0308	MDL	0.205	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA8-SB-0.0-0.5		Collected: 7/16/2013 1:30:00		Analysis Type: REA			Dilution: 2		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.286	J	0.103	MDL	0.411	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA8-SB-0.0-0.5		Collected: 7/16/2013 1:30:00		Analysis Type: RES			Dilution: 2		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0409	J	0.0267	MDL	0.205	PQL	mg/Kg	J	Z

Sample ID: SL-510-SA8-SB-0.0-0.5		Collected: 7/16/2013 8:30:00		Analysis Type: REA			Dilution: 2		
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.133	J	0.0993	MDL	0.397	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	7199	Matrix: SO

Sample ID: SL-504-SA8-SB-0.0-0.5	Collected: 7/16/2013 11:45:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.52		0.15	MDL	0.43	PQL	mg/Kg	J	E

Sample ID: SL-504-SA8-SB-2.5-3.5	Collected: 7/16/2013 12:00:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	1.4		0.15	MDL	0.41	PQL	mg/Kg	J	E, FD

Sample ID: SL-507-SA8-SB-0.0-0.5	Collected: 7/16/2013 12:55:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.35	J	0.14	MDL	0.40	PQL	mg/Kg	J	Z, E

Sample ID: SL-507-SA8-SB-3.0-4.0	Collected: 7/16/2013 1:10:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.40	U	0.14	MDL	0.40	PQL	mg/Kg	UJ	E

Sample ID: SL-508-SA8-SB-0.0-0.5	Collected: 7/16/2013 1:30:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.30	J	0.15	MDL	0.44	PQL	mg/Kg	J	Z, E

Sample ID: SL-804-SA8-SB-2.5-3.5	Collected: 7/16/2013 12:10:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.43	U	0.15	MDL	0.43	PQL	mg/Kg	UJ	E, FD

Method Category:	METALS	
Method:	7471B	Matrix: SO

Sample ID: SL-505-SA8-SB-0.0-0.5	Collected: 7/16/2013 8:55:00	Analysis Type: RES	Dilution: 1						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0128	J	0.0105	MDL	0.0176	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: METALS
Method: 7471B **Matrix: SO**

Sample ID: SL-505-SA8-SB-4.0-5.0 Collected: 7/16/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0140	J	0.0105	MDL	0.0174	PQL	mg/Kg	J	Z

Sample ID: SL-510-SA8-SB-0.0-0.5 Collected: 7/16/2013 8:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0144	J	0.0099	MDL	0.0165	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 1613B **Matrix: SO**

Sample ID: SL-504-SA8-SB-0.0-0.5 Collected: 7/16/2013 11:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	2.93	JB	0.0253	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.408	JQ	0.0527	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.280	JBQ	0.0576	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.225	JB	0.0342	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.891	J	0.0624	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.102	JBQ	0.0319	MDL	5.18	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.497	JB	0.0649	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.106	JB	0.0432	MDL	5.18	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.188	JQ	0.0586	MDL	5.18	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.752	JB	0.0360	MDL	5.18	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.213	JB	0.0350	MDL	5.18	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.0795	JQ	0.0370	MDL	5.18	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0670	JQ	0.0616	MDL	1.04	PQL	ng/Kg	J	Z
OCDF	8.29	JB	0.0632	MDL	10.4	PQL	ng/Kg	J	Z

Sample ID: SL-504-SA8-SB-2.5-3.5 Collected: 7/16/2013 12:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.219	JB	0.0419	MDL	5.08	PQL	ng/Kg	UJ	B, FD
1,2,3,4,6,7,8-HPCDF	0.0823	JBQ	0.0111	MDL	5.08	PQL	ng/Kg	UJ	B, FD
1,2,3,4,7,8,9-HPCDF	0.0343	JQ	0.0229	MDL	5.08	PQL	ng/Kg	J	Z, FD
1,2,3,4,7,8-HxCDD	0.0400	JBQ	0.0313	MDL	5.08	PQL	ng/Kg	UJ	B, FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-504-SA8-SB-2.5-3.5 Collected: 7/16/2013 12:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HXCDF	0.0695	JBQ	0.0154	MDL	5.08	PQL	ng/Kg	UJ	B, FD
1,2,3,6,7,8-HXCDD	0.0391	JQ	0.0342	MDL	5.08	PQL	ng/Kg	J	Z, FD
1,2,3,6,7,8-HXCDF	0.0354	JB	0.0148	MDL	5.08	PQL	ng/Kg	UJ	B, FD
1,2,3,7,8,9-HXCDD	5.08	U	0.0324	MDL	5.08	PQL	ng/Kg	UJ	FD
1,2,3,7,8,9-HXCDF	5.08	U	0.0230	MDL	5.08	PQL	ng/Kg	UJ	FD
1,2,3,7,8-PECDD	5.08	U	0.0496	MDL	5.08	PQL	ng/Kg	UJ	FD
1,2,3,7,8-PECDF	0.203	JB	0.0235	MDL	5.08	PQL	ng/Kg	J	Z, FD
2,3,4,6,7,8-HXCDF	0.0365	JB	0.0170	MDL	5.08	PQL	ng/Kg	UJ	B, FD
2,3,4,7,8-PECDF	0.0347	JQ	0.0261	MDL	5.08	PQL	ng/Kg	J	Z, FD
OCDD	1.89	JB	0.0427	MDL	10.2	PQL	ng/Kg	J	Z, FD
OCDF	0.188	JBQ	0.0529	MDL	10.2	PQL	ng/Kg	UJ	B, FD

Sample ID: SL-505-SA8-SB-0.0-0.5 Collected: 7/16/2013 8:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	0.565	JB	0.0256	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.162	JQ	0.0465	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0906	JBQ	0.0307	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.195	JQ	0.0614	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0869	JB	0.0286	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.210	JB	0.0596	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.0867	JBQ	0.0358	MDL	5.22	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.141	JQ	0.0755	MDL	5.22	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	1.21	JB	0.0445	MDL	5.22	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0814	JBQ	0.0272	MDL	5.22	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.128	JQ	0.0464	MDL	5.22	PQL	ng/Kg	J	Z
OCDF	1.60	JB	0.0759	MDL	10.4	PQL	ng/Kg	J	Z

Sample ID: SL-505-SA8-SB-4.0-5.0 Collected: 7/16/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.392	JBQ	0.0579	MDL	5.41	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.0982	JBQ	0.0154	MDL	5.41	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0433	JBQ	0.0217	MDL	5.41	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0425	JQ	0.0412	MDL	5.41	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-505-SA8-SB-4.0-5.0 Collected: 7/16/2013 9:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,6,7,8-HXCDF	0.0333	JBQ	0.0212	MDL	5.41	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0400	JBQ	0.0394	MDL	5.41	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0357	JBQ	0.0296	MDL	5.41	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0674	JBQ	0.0220	MDL	5.41	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0459	JQ	0.0377	MDL	5.41	PQL	ng/Kg	J	Z
OCDD	3.36	JB	0.0466	MDL	10.8	PQL	ng/Kg	J	Z
OCDF	0.226	JBQ	0.0699	MDL	10.8	PQL	ng/Kg	U	B

Sample ID: SL-507-SA8-SB-0.0-0.5 Collected: 7/16/2013 12:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	2.30	JB	0.0250	MDL	4.93	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.427	J	0.0470	MDL	4.93	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.603	JB	0.0838	MDL	4.93	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.702	JB	0.0554	MDL	4.93	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	1.35	J	0.0902	MDL	4.93	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.698	JB	0.0466	MDL	4.93	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.960	JB	0.0883	MDL	4.93	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.408	JBQ	0.0802	MDL	4.93	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.493	J	0.0794	MDL	4.93	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.562	JB	0.0541	MDL	4.93	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	1.05	J	0.0694	MDL	4.93	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.132	J	0.0621	MDL	0.986	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.722	J	0.116	MDL	0.986	PQL	ng/Kg	J	Z
OCDF	3.70	JB	0.0720	MDL	9.86	PQL	ng/Kg	J	Z

Sample ID: SL-507-SA8-SB-3.0-4.0 Collected: 7/16/2013 1:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0934	JB	0.0325	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.109	JB	0.00953	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0662	JQ	0.0210	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.165	JB	0.0258	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.207	JB	0.0192	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.110	JQ	0.0275	MDL	5.10	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-507-SA8-SB-3.0-4.0 Collected: 7/16/2013 1:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,6,7,8-HXCDF	0.152	JBQ	0.0172	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.150	JBQ	0.0263	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.166	JBQ	0.0209	MDL	5.10	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.322	J	0.0413	MDL	5.10	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.402	JB	0.0216	MDL	5.10	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.111	JB	0.0175	MDL	5.10	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.324	J	0.0231	MDL	5.10	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0702	JQ	0.0363	MDL	1.02	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0858	J	0.0319	MDL	1.02	PQL	ng/Kg	J	Z
OCDD	0.390	JB	0.0296	MDL	10.2	PQL	ng/Kg	U	B
OCDF	0.144	JB	0.0454	MDL	10.2	PQL	ng/Kg	U	B

Sample ID: SL-508-SA8-SB-0.0-0.5 Collected: 7/16/2013 1:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.98	JB	0.0772	MDL	5.23	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	2.91	JB	0.0252	MDL	5.23	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.192	JQ	0.0563	MDL	5.23	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.0951	JBQ	0.0462	MDL	5.23	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.598	JB	0.0408	MDL	5.23	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.179	J	0.0494	MDL	5.23	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.746	JB	0.0352	MDL	5.23	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.231	JBQ	0.0505	MDL	5.23	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.263	JB	0.0550	MDL	5.23	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.216	JQ	0.0857	MDL	5.23	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.205	JB	0.0386	MDL	5.23	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.579	J	0.0636	MDL	5.23	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0654	JQ	0.0543	MDL	1.05	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.370	JQ	0.101	MDL	1.05	PQL	ng/Kg	J	Z
OCDF	7.54	JB	0.0869	MDL	10.5	PQL	ng/Kg	J	Z

Sample ID: SL-510-SA8-SB-0.0-0.5 Collected: 7/16/2013 8:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	1.98	JB	0.0286	MDL	4.95	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-510-SA8-SB-0.0-0.5 Collected: 7/16/2013 8:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8,9-HPCDF	0.226	JQ	0.0478	MDL	4.95	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.279	JB	0.0724	MDL	4.95	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.372	JBQ	0.0400	MDL	4.95	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.874	J	0.0789	MDL	4.95	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.339	JB	0.0397	MDL	4.95	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.790	JB	0.0779	MDL	4.95	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.187	JB	0.0463	MDL	4.95	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.142	JQ	0.0588	MDL	4.95	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.728	JB	0.0394	MDL	4.95	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.364	JBQ	0.0385	MDL	4.95	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.146	J	0.0424	MDL	4.95	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.151	JQ	0.0680	MDL	0.989	PQL	ng/Kg	J	Z
OCDF	2.98	JB	0.0522	MDL	9.89	PQL	ng/Kg	J	Z

Sample ID: SL-804-SA8-SB-2.5-3.5 Collected: 7/16/2013 12:10:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	2.15	JB	0.0848	MDL	5.19	PQL	ng/Kg	J	Z, FD
1,2,3,4,6,7,8-HPCDF	0.440	JB	0.0246	MDL	5.19	PQL	ng/Kg	J	Z, FD
1,2,3,4,7,8,9-HPCDF	0.130	JQ	0.0458	MDL	5.19	PQL	ng/Kg	J	Z, FD
1,2,3,4,7,8-HxCDD	0.169	JB	0.0503	MDL	5.19	PQL	ng/Kg	UJ	B, FD
1,2,3,4,7,8-HXCDF	0.158	JB	0.0329	MDL	5.19	PQL	ng/Kg	J	Z, FD
1,2,3,6,7,8-HXCDD	0.253	JQ	0.0566	MDL	5.19	PQL	ng/Kg	J	Z, FD
1,2,3,6,7,8-HXCDF	0.197	JB	0.0302	MDL	5.19	PQL	ng/Kg	J	Z, FD
1,2,3,7,8,9-HXCDD	0.181	JBQ	0.0575	MDL	5.19	PQL	ng/Kg	J	Z, FD
1,2,3,7,8,9-HXCDF	0.136	JBQ	0.0419	MDL	5.19	PQL	ng/Kg	UJ	B, FD
1,2,3,7,8-PECDD	0.168	JQ	0.0781	MDL	5.19	PQL	ng/Kg	J	Z, FD
1,2,3,7,8-PECDF	1.07	JB	0.0424	MDL	5.19	PQL	ng/Kg	J	Z, FD
2,3,4,6,7,8-HXCDF	0.0926	JBQ	0.0300	MDL	5.19	PQL	ng/Kg	UJ	B, FD
2,3,4,7,8-PECDF	0.203	JQ	0.0432	MDL	5.19	PQL	ng/Kg	J	Z, FD
OCDD	18.8	B	0.0517	MDL	10.4	PQL	ng/Kg	J	FD
OCDF	0.988	JBQ	0.0777	MDL	10.4	PQL	ng/Kg	J	Z, FD

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	8015M	Matrix:	SO
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Sample ID: SL-504-SA8-SB-0.0-0.5		Collected: 7/16/2013 11:45:00		Analysis Type: RES		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	4.9	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z

Sample ID: SL-507-SA8-SB-0.0-0.5		Collected: 7/16/2013 12:55:00		Analysis Type: RES		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	4.6	J	2.0	MDL	5.1	PQL	mg/Kg	J	Z

Sample ID: SL-508-SA8-SB-0.0-0.5		Collected: 7/16/2013 1:30:00		Analysis Type: RES		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	4.7	J	2.1	MDL	5.3	PQL	mg/Kg	J	Z

Method Category:	SVOA	Method:	8082A	Matrix:	SO
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Sample ID: SL-504-SA8-SB-0.0-0.5		Collected: 7/16/2013 11:45:00		Analysis Type: RES-BASE/NEUTRAL		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	9.6	J	4.5	MDL	18	PQL	ug/Kg	J	Z

Sample ID: SL-505-SA8-SB-0.0-0.5		Collected: 7/16/2013 8:55:00		Analysis Type: RES-BASE/NEUTRAL		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	16	J	4.7	MDL	18	PQL	ug/Kg	J	Z

Method Category:	SVOA	Method:	8270D	Matrix:	SO
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Sample ID: SL-504-SA8-SB-2.5-3.5		Collected: 7/16/2013 12:00:00		Analysis Type: RES-BASE/NEUTRAL		Dilution: 1			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZIDINE	1700	U	730	MDL	1700	PQL	ug/Kg	UJ	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-504-SA8-SB-0.0-0.5 Collected: 7/16/2013 11:45:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.87	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	1.1	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z
CHRYSENE	0.90	J	0.35	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-505-SA8-SB-0.0-0.5 Collected: 7/16/2013 8:55:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.73	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	1.0	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	1.1	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-505-SA8-SB-4.0-5.0 Collected: 7/16/2013 9:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ACENAPHTHYLENE	0.50	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-507-SA8-SB-0.0-0.5 Collected: 7/16/2013 12:55:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	1.1	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	7.2	J	6.1	MDL	18	PQL	ug/Kg	J	Z
PHENANTHRENE	1.1	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	1.3	J	0.68	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-508-SA8-SB-0.0-0.5 Collected: 7/16/2013 1:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BIS(2-ETHYLHEXYL)PHTHALATE	11	J	6.3	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	0.82	J	0.35	MDL	1.7	PQL	ug/Kg	J	Z
PHENANTHRENE	0.86	J	0.70	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-510-SA8-SB-0.0-0.5 Collected: 7/16/2013 8:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.71	J	0.66	MDL	1.7	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8270D SIM	Matrix: SO

Sample ID: SL-510-SA8-SB-0.0-0.5 Collected: 7/16/2013 8:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(K)FLUORANTHENE	1.2	J	0.66	MDL	1.7	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	14	J	6.0	MDL	18	PQL	ug/Kg	J	Z
CHRYSENE	1.1	J	0.33	MDL	1.7	PQL	ug/Kg	J	Z
FLUORANTHENE	0.93	J	0.66	MDL	1.7	PQL	ug/Kg	J	Z
PHENANTHRENE	0.70	J	0.66	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	0.84	J	0.66	MDL	1.7	PQL	ug/Kg	J	Z

Method Category:	VOA	
Method:	8260B	Matrix: AQ

Sample ID: TB-071613 Collected: 7/16/2013 8:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
Chlorotrifluoroethylene	5	U	2	MDL	5	PQL	ug/L	UJ	L

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
*#	Professional Judgment
A	ICP Serial Dilution
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Field Blank Contamination
FD	Field Duplicate Precision
L	Laboratory Control Spike Lower Estimation
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH068

Method Blank Outlier Report

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2060B372237	7/26/2013 10:37:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,7,8,9-HxCDD 1,2,3,7,8,9-HxCDF 1,2,3,7,8-PECDF 2,3,4,6,7,8-HxCDF OCDD OCDF	0.0714 ng/Kg 0.0256 ng/Kg 0.0432 ng/Kg 0.0229 ng/Kg 0.0231 ng/Kg 0.0358 ng/Kg 0.0413 ng/Kg 0.0307 ng/Kg 0.0401 ng/Kg 0.230 ng/Kg 0.159 ng/Kg	SL-504-SA8-SB-0.0-0.5 SL-504-SA8-SB-2.5-3.5 SL-505-SA8-SB-0.0-0.5 SL-505-SA8-SB-4.0-5.0 SL-507-SA8-SB-0.0-0.5 SL-507-SA8-SB-3.0-4.0 SL-508-SA8-SB-0.0-0.5 SL-510-SA8-SB-0.0-0.5 SL-804-SA8-SB-2.5-3.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-504-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HxCDF	0.102 ng/Kg	0.102U ng/Kg
SL-504-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HxCDF	0.106 ng/Kg	0.106U ng/Kg
SL-504-SA8-SB-2.5-3.5(RES)	1,2,3,4,6,7,8-HPCDD	0.219 ng/Kg	0.219U ng/Kg
SL-504-SA8-SB-2.5-3.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0823 ng/Kg	0.0823U ng/Kg
SL-504-SA8-SB-2.5-3.5(RES)	1,2,3,4,7,8-HxCDD	0.0400 ng/Kg	0.0400U ng/Kg
SL-504-SA8-SB-2.5-3.5(RES)	1,2,3,4,7,8-HxCDF	0.0695 ng/Kg	0.0695U ng/Kg
SL-504-SA8-SB-2.5-3.5(RES)	1,2,3,6,7,8-HxCDF	0.0354 ng/Kg	0.0354U ng/Kg
SL-504-SA8-SB-2.5-3.5(RES)	2,3,4,6,7,8-HxCDF	0.0365 ng/Kg	0.0365U ng/Kg
SL-504-SA8-SB-2.5-3.5(RES)	OCDF	0.188 ng/Kg	0.188U ng/Kg
SL-505-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.0906 ng/Kg	0.0906U ng/Kg
SL-505-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HxCDF	0.0869 ng/Kg	0.0869U ng/Kg
SL-505-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HxCDF	0.0867 ng/Kg	0.0867U ng/Kg
SL-505-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HxCDF	0.0814 ng/Kg	0.0814U ng/Kg
SL-505-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0982 ng/Kg	0.0982U ng/Kg
SL-505-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDF	0.0433 ng/Kg	0.0433U ng/Kg
SL-505-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HxCDF	0.0333 ng/Kg	0.0333U ng/Kg
SL-505-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDD	0.0400 ng/Kg	0.0400U ng/Kg
SL-505-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDF	0.0357 ng/Kg	0.0357U ng/Kg
SL-505-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HxCDF	0.0674 ng/Kg	0.0674U ng/Kg
SL-505-SA8-SB-4.0-5.0(RES)	OCDF	0.226 ng/Kg	0.226U ng/Kg
SL-507-SA8-SB-3.0-4.0(RES)	1,2,3,4,6,7,8-HPCDD	0.0934 ng/Kg	0.0934U ng/Kg
SL-507-SA8-SB-3.0-4.0(RES)	1,2,3,4,6,7,8-HPCDF	0.109 ng/Kg	0.109U ng/Kg
SL-507-SA8-SB-3.0-4.0(RES)	1,2,3,4,7,8-HxCDD	0.165 ng/Kg	0.165U ng/Kg
SL-507-SA8-SB-3.0-4.0(RES)	1,2,3,7,8,9-HxCDD	0.150 ng/Kg	0.150U ng/Kg
SL-507-SA8-SB-3.0-4.0(RES)	1,2,3,7,8,9-HxCDF	0.166 ng/Kg	0.166U ng/Kg
SL-507-SA8-SB-3.0-4.0(RES)	2,3,4,6,7,8-HxCDF	0.111 ng/Kg	0.111U ng/Kg
SL-507-SA8-SB-3.0-4.0(RES)	OCDD	0.390 ng/Kg	0.390U ng/Kg
SL-507-SA8-SB-3.0-4.0(RES)	OCDF	0.144 ng/Kg	0.144U ng/Kg
SL-508-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.0951 ng/Kg	0.0951U ng/Kg
SL-510-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HxCDF	0.187 ng/Kg	0.187U ng/Kg
SL-804-SA8-SB-2.5-3.5(RES)	1,2,3,4,7,8-HxCDD	0.169 ng/Kg	0.169U ng/Kg
SL-804-SA8-SB-2.5-3.5(RES)	1,2,3,7,8,9-HxCDF	0.136 ng/Kg	0.136U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-804-SA8-SB-2.5-3.5(RES)	2,3,4,6,7,8-HXCDF	0.0926 ng/Kg	0.0926U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P19837AB221510	7/24/2013 3:10:00 PM	TIN ZINC	1.34 mg/Kg 0.343 mg/Kg	SL-504-SA8-SB-0.0-0.5 SL-504-SA8-SB-2.5-3.5 SL-505-SA8-SB-0.0-0.5 SL-505-SA8-SB-4.0-5.0 SL-507-SA8-SB-0.0-0.5 SL-507-SA8-SB-3.0-4.0 SL-508-SA8-SB-0.0-0.5 SL-510-SA8-SB-0.0-0.5 SL-804-SA8-SB-2.5-3.5
P19837AB222016	7/25/2013 8:16:00 PM	COPPER	0.441 mg/Kg	SL-504-SA8-SB-0.0-0.5 SL-504-SA8-SB-2.5-3.5 SL-505-SA8-SB-0.0-0.5 SL-505-SA8-SB-4.0-5.0 SL-507-SA8-SB-0.0-0.5 SL-507-SA8-SB-3.0-4.0 SL-508-SA8-SB-0.0-0.5 SL-510-SA8-SB-0.0-0.5 SL-804-SA8-SB-2.5-3.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-504-SA8-SB-0.0-0.5(RES)	TIN	3.11 mg/Kg	3.11U mg/Kg
SL-504-SA8-SB-2.5-3.5(RES)	TIN	2.86 mg/Kg	2.86U mg/Kg
SL-505-SA8-SB-0.0-0.5(RES)	TIN	3.19 mg/Kg	3.19U mg/Kg
SL-505-SA8-SB-4.0-5.0(RES)	TIN	3.18 mg/Kg	3.18U mg/Kg
SL-507-SA8-SB-0.0-0.5(RES)	TIN	2.89 mg/Kg	2.89U mg/Kg
SL-507-SA8-SB-3.0-4.0(RES)	TIN	2.96 mg/Kg	2.96U mg/Kg
SL-508-SA8-SB-0.0-0.5(RES)	TIN	2.77 mg/Kg	2.77U mg/Kg
SL-510-SA8-SB-0.0-0.5(RES)	TIN	2.85 mg/Kg	2.85U mg/Kg
SL-804-SA8-SB-2.5-3.5(RES)	TIN	2.99 mg/Kg	2.99U mg/Kg

Field Blank Outlier Report

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(REA2)	4/11/2013 3:00:00 PM	COPPER MOLYBDENUM	0.0036 mg/L 0.0036 mg/L	SL-504-SA8-SB-0.0-0.5 SL-504-SA8-SB-2.5-3.5 SL-505-SA8-SB-0.0-0.5 SL-505-SA8-SB-4.0-5.0 SL-507-SA8-SB-0.0-0.5 SL-507-SA8-SB-3.0-4.0 SL-508-SA8-SB-0.0-0.5 SL-510-SA8-SB-0.0-0.5 SL-804-SA8-SB-2.5-3.5

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-504-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.518 mg/Kg	0.518U mg/Kg
SL-504-SA8-SB-2.5-3.5(RES)	MOLYBDENUM	0.373 mg/Kg	0.373U mg/Kg
SL-505-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.627 mg/Kg	0.627U mg/Kg
SL-507-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.343 mg/Kg	0.343U mg/Kg
SL-508-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.574 mg/Kg	0.574U mg/Kg
SL-510-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.373 mg/Kg	0.373U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-504-SA8-SB-2.5-3.5MS (TOT)	ALUMINUM	1060	1082	75.00-125.00	-	ALUMINUM	J (all detects) Al, Ca, Fe, Mg, Mn, P, Ti, No Qual, >4x
SL-504-SA8-SB-2.5-3.5MSD (TOT)	CALCIUM	202	167	75.00-125.00	-	CALCIUM	
SL-504-SA8-SB-2.5-3.5MSD (TOT)	CHROMIUM	134	-	75.00-125.00	-	CHROMIUM	
(SL-504-SA8-SB-0.0-0.5)	IRON	358	211	75.00-125.00	-	IRON	
SL-504-SA8-SB-2.5-3.5	MAGNESIUM	562	223	75.00-125.00	-	MAGNESIUM	
SL-505-SA8-SB-0.0-0.5	MANGANESE	133	-	75.00-125.00	-	MANGANESE	
SL-505-SA8-SB-4.0-5.0	PHOSPHORUS	133	-	75.00-125.00	-	PHOSPHORUS	
SL-507-SA8-SB-0.0-0.5	POTASSIUM	128	135	75.00-125.00	-	POTASSIUM	
SL-507-SA8-SB-3.0-4.0	TITANIUM	342	330	75.00-125.00	-	TITANIUM	
SL-508-SA8-SB-0.0-0.5							
SL-510-SA8-SB-0.0-0.5							
SL-804-SA8-SB-2.5-3.5)							
SL-504-SA8-SB-2.5-3.5MS (TOT)	ANTIMONY	72	70	75.00-125.00	-	ANTIMONY	J(all detects) UJ(all non-detects)
SL-504-SA8-SB-2.5-3.5MSD (TOT)							
(SL-504-SA8-SB-0.0-0.5)							
SL-504-SA8-SB-2.5-3.5							
SL-505-SA8-SB-0.0-0.5							
SL-505-SA8-SB-4.0-5.0							
SL-507-SA8-SB-0.0-0.5							
SL-507-SA8-SB-3.0-4.0							
SL-508-SA8-SB-0.0-0.5							
SL-510-SA8-SB-0.0-0.5							
SL-804-SA8-SB-2.5-3.5)							

Method: 8270D
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-504-SA8-SB-2.5-3.5MSD (SL-504-SA8-SB-2.5-3.5)	2,4-DIMETHYLPHENOL 4-CHLOROANILINE	-	-	39.00-147.00 11.00-114.00	31 (30.00) 47 (30.00)	2,4-DIMETHYLPHENOL 4-CHLOROANILINE	J(all detects)
SL-504-SA8-SB-2.5-3.5MS (SL-504-SA8-SB-2.5-3.5MSD)	BENZIDINE	13	10	35.00-141.00	-	BENZIDINE	J(all detects) UJ(all non-detects)

Method: 8260B
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-504-SA8-SB-2.5-3.5MS (SL-504-SA8-SB-2.5-3.5MSD)	ACROLEIN	169	239	10.00-165.00	36 (30.00)	ACROLEIN	J(all detects)
SL-504-SA8-SB-2.5-3.5MSD (SL-504-SA8-SB-2.5-3.5)	Tertiary butyl alcohol	218	294	47.00-153.00	32 (30.00)	Tertiary butyl alcohol	

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-504-SA8-SB-2.5-3.5DUP (TOT) (SL-504-SA8-SB-0.0-0.5 SL-504-SA8-SB-2.5-3.5 SL-505-SA8-SB-0.0-0.5 SL-505-SA8-SB-4.0-5.0 SL-507-SA8-SB-0.0-0.5 SL-507-SA8-SB-3.0-4.0 SL-508-SA8-SB-0.0-0.5 SL-510-SA8-SB-0.0-0.5 SL-804-SA8-SB-2.5-3.5)	MOLYBDENUM Zirconium	42 24	20.00 20.00	J (all detects) UJ (all non-detects)

Method: 7199
Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-504-SA8-SB-2.5-3.5DUP (TOT) (SL-504-SA8-SB-0.0-0.5 SL-504-SA8-SB-2.5-3.5 SL-507-SA8-SB-0.0-0.5 SL-507-SA8-SB-3.0-4.0 SL-508-SA8-SB-0.0-0.5 SL-804-SA8-SB-2.5-3.5)	HEXAVALENT CHROMIUM	1.24	≤ 0.82	J(all detects) UJ(all non-detects)

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8260B
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
LCSY77Q211110A LCSY77Y211131A (TB-071613)	Chlorotrifluoroethylene	44	43	47.00-120.00	-	Chlorotrifluoroethylene	J(all detects) UJ(all non-detects)

Method: 8260B
Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P9LELCSQ261256 (SL-504-SA8-SB-2.5-3.5 SL-507-SA8-SB-3.0-4.0 SL-804-SA8-SB-2.5-3.5)	VINYL ACETATE	135	125	29.00-111.00	-	VINYL ACETATE	J (all detects)

Field Duplicate RPD Report

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 160.3M
Matrix: SO

Analyte	Concentration (%)		Sample RPD	eQAPP RPD	Flag
	SL-504-SA8-SB-2.5-3.5	SL-804-SA8-SB-2.5-3.5			
MOISTURE	3.9	4.0	3		No Qualifiers Applied

Method: 1613B
Matrix: SO

Analyte	Concentration (ng/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-504-SA8-SB-2.5-3.5	SL-804-SA8-SB-2.5-3.5			
TEQ WHO 2005 - EDLx0.0	0.0160	0.116	152		No Qualifiers Applied
1,2,3,4,6,7,8-HPCDD	0.219	2.15	163	50.00	J(all detects) UJ(all non-detects)
1,2,3,4,6,7,8-HPCDF	0.0823	0.440	137	50.00	
1,2,3,4,7,8,9-HPCDF	0.0343	0.130	116	50.00	
1,2,3,4,7,8-HxCDD	0.0400	0.169	123	50.00	
1,2,3,4,7,8-HxCDF	0.0695	0.158	78	50.00	
1,2,3,6,7,8-HxCDD	0.0391	0.253	146	50.00	
1,2,3,6,7,8-HxCDF	0.0354	0.197	139	50.00	
1,2,3,7,8,9-HxCDD	5.08 U	0.181	200	50.00	
1,2,3,7,8,9-HxCDF	5.08 U	0.136	200	50.00	
1,2,3,7,8-PECDD	5.08 U	0.168	200	50.00	
1,2,3,7,8-PECDF	0.203	1.07	136	50.00	
2,3,4,6,7,8-HxCDF	0.0365	0.0926	87	50.00	
2,3,4,7,8-PECDF	0.0347	0.203	142	50.00	
OCDD	1.89	18.8	163	50.00	
OCDF	0.188	0.988	136	50.00	

Method: 6010C
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-504-SA8-SB-2.5-3.5 (TOT)	SL-804-SA8-SB-2.5-3.5 (TOT)			
ALUMINUM	11800	12100	3	50.00	No Qualifiers Applied
ARSENIC	3.88	3.68	5	50.00	
BARIUM	84.4	88.4	5	50.00	
BERYLLIUM	0.394	0.449	13	50.00	
BORON	18.1	16.0	12	50.00	
CADMIUM	0.370	0.330	11	50.00	
CALCIUM	3670	3610	2	50.00	
CHROMIUM	21.8	22.0	1	50.00	
COBALT	4.81	5.16	7	50.00	
COPPER	10.6	10.9	3	50.00	
IRON	22900	23200	1	50.00	
LEAD	4.43	4.49	1	50.00	
LITHIUM	29.8	29.5	1	50.00	
MAGNESIUM	6080	6410	5	50.00	
MANGANESE	422	313	30	50.00	
NICKEL	12.6	13.0	3	50.00	
PHOSPHORUS	423	424	0	50.00	
POTASSIUM	2220	2240	1	50.00	
SODIUM	177	197	11	50.00	
TIN	2.86	2.99	4	50.00	
TITANIUM	1460	1480	1	50.00	
VANADIUM	43.8	43.2	1	50.00	
ZINC	53.6	58.6	9	50.00	
Zirconium	3.17	2.85	11	50.00	
MOLYBDENUM	0.373	2.02 U	200	50.00	J(all detects) UJ(all non-detects)

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Field Duplicate RPD Report

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-504-SA8-SB-2.5-3.5 (TOT)	SL-804-SA8-SB-2.5-3.5 (TOT)			
STRONTIUM	14.9	14.9	0	50.00	No Qualifiers Applied
THALLIUM	0.238	0.212	12	50.00	

Method: 7199
Matrix: SO

Analyte	Concentration (mg/Kg)		Sample RPD	eQAPP RPD	Flag
	SL-504-SA8-SB-2.5-3.5 (TOT)	SL-804-SA8-SB-2.5-3.5 (TOT)			
HEXAVALENT CHROMIUM	1.4	0.43 U	200	50.00	J(all detects) UJ(all non-detects)

Method: 9045M
Matrix: SO

Analyte	Concentration (pH unit)		Sample RPD	eQAPP RPD	Flag
	SL-504-SA8-SB-2.5-3.5	SL-804-SA8-SB-2.5-3.5			
PH	8.56	8.55	0	50.00	No Qualifiers Applied

Reporting Limit Outliers

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-504-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	2.93	5.18	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JQ	0.408	5.18	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.280	5.18	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.225	5.18	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	0.891	5.18	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.102	5.18	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.497	5.18	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.106	5.18	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.188	5.18	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.752	5.18	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.213	5.18	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0795	5.18	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0670	1.04	PQL	ng/Kg	
	OCDF	JB	8.29	10.4	PQL	ng/Kg	
SL-504-SA8-SB-2.5-3.5	1,2,3,4,6,7,8-HPCDD	JB	0.219	5.08	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0823	5.08	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JQ	0.0343	5.08	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0400	5.08	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0695	5.08	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.0391	5.08	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.0354	5.08	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.203	5.08	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.0365	5.08	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0347	5.08	PQL	ng/Kg	
	OCDD	JB	1.89	10.2	PQL	ng/Kg	
OCDF	JBQ	0.188	10.2	PQL	ng/Kg		
SL-505-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	0.565	5.22	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JQ	0.162	5.22	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0906	5.22	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.195	5.22	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.0869	5.22	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.210	5.22	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.0867	5.22	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.141	5.22	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	1.21	5.22	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0814	5.22	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.128	5.22	PQL	ng/Kg	
OCDF	JB	1.60	10.4	PQL	ng/Kg		
SL-505-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.392	5.41	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0982	5.41	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0433	5.41	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.0425	5.41	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0333	5.41	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.0400	5.41	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.0357	5.41	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0674	5.41	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0459	5.41	PQL	ng/Kg	
	OCDD	JB	3.36	10.8	PQL	ng/Kg	
	OCDF	JBQ	0.226	10.8	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-507-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	2.30	4.93	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	J	0.427	4.93	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.603	4.93	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.702	4.93	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	1.35	4.93	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.698	4.93	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.960	4.93	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.408	4.93	PQL	ng/Kg	
	1,2,3,7,8-PECDD	J	0.493	4.93	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.562	4.93	PQL	ng/Kg	
	2,3,4,7,8-PECDF	J	1.05	4.93	PQL	ng/Kg	
	2,3,7,8-TCDD	J	0.132	0.986	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.722	0.986	PQL	ng/Kg	
	OCDF	JB	3.70	9.86	PQL	ng/Kg	
SL-507-SA8-SB-3.0-4.0	1,2,3,4,6,7,8-HPCDD	JB	0.0934	5.10	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.109	5.10	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JQ	0.0662	5.10	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.165	5.10	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.207	5.10	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.110	5.10	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.152	5.10	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.150	5.10	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.166	5.10	PQL	ng/Kg	
	1,2,3,7,8-PECDD	J	0.322	5.10	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.402	5.10	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.111	5.10	PQL	ng/Kg	
	2,3,4,7,8-PECDF	J	0.324	5.10	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0702	1.02	PQL	ng/Kg	
2,3,7,8-TCDF	J	0.0858	1.02	PQL	ng/Kg		
OCDD	JB	0.390	10.2	PQL	ng/Kg		
OCDF	JB	0.144	10.2	PQL	ng/Kg		
SL-508-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	1.98	5.23	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	2.91	5.23	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JQ	0.192	5.23	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0951	5.23	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.598	5.23	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	0.179	5.23	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.746	5.23	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.231	5.23	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.263	5.23	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.216	5.23	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.205	5.23	PQL	ng/Kg	
	2,3,4,7,8-PECDF	J	0.579	5.23	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0654	1.05	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.370	1.05	PQL	ng/Kg	
OCDF	JB	7.54	10.5	PQL	ng/Kg		

Reporting Limit Outliers

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-510-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	1.98	4.95	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JQ	0.226	4.95	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.279	4.95	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.372	4.95	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	0.874	4.95	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.339	4.95	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.790	4.95	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.187	4.95	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.142	4.95	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.728	4.95	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.364	4.95	PQL	ng/Kg	
	2,3,4,7,8-PECDF	J	0.146	4.95	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.151	0.989	PQL	ng/Kg	
	OCDF	JB	2.98	9.89	PQL	ng/Kg	
SL-804-SA8-SB-2.5-3.5	1,2,3,4,6,7,8-HPCDD	JB	2.15	5.19	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.440	5.19	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JQ	0.130	5.19	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.169	5.19	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.158	5.19	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.253	5.19	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.197	5.19	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.181	5.19	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.136	5.19	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.168	5.19	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	1.07	5.19	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0926	5.19	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.203	5.19	PQL	ng/Kg	
	OCDF	JBQ	0.988	10.4	PQL	ng/Kg	

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-504-SA8-SB-0.0-0.5	BERYLLIUM	J	0.641	1.02	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.513	1.02	PQL	mg/Kg	
	MOLYBDENUM	J	0.518	2.03	PQL	mg/Kg	
	TIN	J	3.11	10.2	PQL	mg/Kg	
	Zirconium	J	2.16	5.08	PQL	mg/Kg	
SL-504-SA8-SB-2.5-3.5	ARSENIC	J	3.88	4.08	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.394	1.02	PQL	mg/Kg	
	CADMIUM	J	0.370	1.02	PQL	mg/Kg	
	MOLYBDENUM	J	0.373	2.04	PQL	mg/Kg	
	TIN	J	2.86	10.2	PQL	mg/Kg	
	Zirconium	J	3.17	5.10	PQL	mg/Kg	
SL-505-SA8-SB-0.0-0.5	BERYLLIUM	J	0.692	1.04	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.543	1.04	PQL	mg/Kg	
	MOLYBDENUM	J	0.627	2.08	PQL	mg/Kg	
	TIN	J	3.19	10.4	PQL	mg/Kg	
	Zirconium	J	1.60	5.20	PQL	mg/Kg	
SL-505-SA8-SB-4.0-5.0	BERYLLIUM	J	0.689	1.10	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.357	1.10	PQL	mg/Kg	
	TIN	J	3.18	11.0	PQL	mg/Kg	
	Zirconium	J	3.50	5.48	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-507-SA8-SB-0.0-0.5	ARSENIC	J	3.27	3.97	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.403	0.992	PQL	mg/Kg	
	CADMIUM	J	0.322	0.992	PQL	mg/Kg	
	MOLYBDENUM	J	0.343	1.98	PQL	mg/Kg	
	TIN	J	2.89	9.92	PQL	mg/Kg	
	Zirconium	J	2.37	4.96	PQL	mg/Kg	
SL-507-SA8-SB-3.0-4.0	BERYLLIUM	J	0.381	1.03	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.232	1.03	PQL	mg/Kg	
	TIN	J	2.96	10.3	PQL	mg/Kg	
	Zirconium	J	3.17	5.13	PQL	mg/Kg	
SL-508-SA8-SB-0.0-0.5	BERYLLIUM	J	0.582	1.03	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.251	1.03	PQL	mg/Kg	
	MOLYBDENUM	J	0.574	2.05	PQL	mg/Kg	
	TIN	J	2.77	10.3	PQL	mg/Kg	
	Zirconium	J	2.38	5.13	PQL	mg/Kg	
SL-510-SA8-SB-0.0-0.5	BERYLLIUM	J	0.462	0.993	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.393	0.993	PQL	mg/Kg	
	MOLYBDENUM	J	0.373	1.99	PQL	mg/Kg	
	SODIUM	J	82.4	99.3	PQL	mg/Kg	
	TIN	J	2.85	9.93	PQL	mg/Kg	
	Zirconium	J	1.57	4.97	PQL	mg/Kg	
SL-804-SA8-SB-2.5-3.5	ARSENIC	J	3.68	4.05	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.449	1.01	PQL	mg/Kg	
	CADMIUM	J	0.330	1.01	PQL	mg/Kg	
	TIN	J	2.99	10.1	PQL	mg/Kg	
	Zirconium	J	2.85	5.06	PQL	mg/Kg	

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-504-SA8-SB-0.0-0.5	SELENIUM	J	0.251	0.406	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0343	0.203	PQL	mg/Kg	
SL-505-SA8-SB-0.0-0.5	SELENIUM	J	0.329	0.416	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0558	0.208	PQL	mg/Kg	
SL-505-SA8-SB-4.0-5.0	SELENIUM	J	0.132	0.439	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0328	0.219	PQL	mg/Kg	
SL-507-SA8-SB-0.0-0.5	SELENIUM	J	0.186	0.397	PQL	mg/Kg	J (all detects)
SL-507-SA8-SB-3.0-4.0	THALLIUM	J	0.192	0.205	PQL	mg/Kg	J (all detects)
SL-508-SA8-SB-0.0-0.5	SELENIUM	J	0.286	0.411	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0409	0.205	PQL	mg/Kg	
SL-510-SA8-SB-0.0-0.5	SELENIUM	J	0.133	0.397	PQL	mg/Kg	J (all detects)

Method: 7199
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-507-SA8-SB-0.0-0.5	HEXAVALENT CHROMIUM	J	0.35	0.40	PQL	mg/Kg	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 7199

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-508-SA8-SB-0.0-0.5	HEXAVALENT CHROMIUM	J	0.30	0.44	PQL	mg/Kg	J (all detects)

Method: 7471B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-505-SA8-SB-0.0-0.5	MERCURY	J	0.0128	0.0176	PQL	mg/Kg	J (all detects)
SL-505-SA8-SB-4.0-5.0	MERCURY	J	0.0140	0.0174	PQL	mg/Kg	J (all detects)
SL-510-SA8-SB-0.0-0.5	MERCURY	J	0.0144	0.0165	PQL	mg/Kg	J (all detects)

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-504-SA8-SB-0.0-0.5	EFH (C21-C30)	J	4.9	5.2	PQL	mg/Kg	J (all detects)
SL-507-SA8-SB-0.0-0.5	EFH (C21-C30)	J	4.6	5.1	PQL	mg/Kg	J (all detects)
SL-508-SA8-SB-0.0-0.5	EFH (C21-C30)	J	4.7	5.3	PQL	mg/Kg	J (all detects)

Method: 8082A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-504-SA8-SB-0.0-0.5	AROCLOR 1254	J	9.6	18	PQL	ug/Kg	J (all detects)
SL-505-SA8-SB-0.0-0.5	AROCLOR 1254	J	16	18	PQL	ug/Kg	J (all detects)

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-504-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	0.87	1.7	PQL	ug/Kg	J (all detects)
	BENZO(K)FLUORANTHENE	J	1.1	1.7	PQL	ug/Kg	
	CHRYSENE	J	0.90	1.7	PQL	ug/Kg	
SL-505-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	0.73	1.8	PQL	ug/Kg	J (all detects)
	BENZO(K)FLUORANTHENE	J	1.0	1.8	PQL	ug/Kg	
	CHRYSENE	J	1.1	1.8	PQL	ug/Kg	
SL-505-SA8-SB-4.0-5.0	ACENAPHTHYLENE	J	0.50	1.8	PQL	ug/Kg	J (all detects)
SL-507-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	1.1	1.7	PQL	ug/Kg	J (all detects)
	BIS(2-ETHYLHEXYL)PHTHALATE	J	7.2	18	PQL	ug/Kg	
	PHENANTHRENE	J	1.1	1.7	PQL	ug/Kg	
	PYRENE	J	1.3	1.7	PQL	ug/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH068

Laboratory: LL

EDD Filename: PH068_v2

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-508-SA8-SB-0.0-0.5	BIS(2-ETHYLHEXYL)PHTHALATE	J	11	19	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	0.82	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	0.86	1.7	PQL	ug/Kg	
SL-510-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	0.71	1.7	PQL	ug/Kg	J (all detects)
	BENZO(K)FLUORANTHENE	J	1.2	1.7	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	14	18	PQL	ug/Kg	
	CHRYSENE	J	1.1	1.7	PQL	ug/Kg	
	FLUORANTHENE	J	0.93	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	0.70	1.7	PQL	ug/Kg	
	PYRENE	J	0.84	1.7	PQL	ug/Kg	

LDC #: 30289C4

VALIDATION COMPLETENESS WORKSHEET

Date: 9/4/13

SDG #: PH068

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: [Signature]

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	-	Sampling dates: 7/16/13
II.	ICP/MS Tune	-	
III.	Calibration	-	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	SW	MS/D
VII.	Duplicate Sample Analysis	SW	DUP
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	SW	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	-	(5,6)
XV.	Field Blanks	SW	EB = EB1-071713 } PH069 FB = FB-041113 = EB2-071713 (PH029)

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: soil

1	SL-510-SA8-SB-0.0-0.5	11	SL-504-SA8-SB-2.5-3.5MSD	21		31	
2	SL-505-SA8-SB-0.0-0.5	12	SL-504-SA8-SB-2.5-3.5DUP	22		32	
3	SL-505-SA8-SB-4.0-5.0	13		23		33	
4	SL-504-SA8-SB-0.0-0.5	14		24		34	
5	SL-504-SA8-SB-2.5-3.5	15		25		35	
6	SL-804-SA8-SB-2.5-3.5	16		26		36	
7	SL-507-SA8-SB-0.0-0.5	17		27		37	
8	SL-507-SA8-SB-3.0-4.0	18		28		38	
9	SL-508-SA8-SB-0.0-0.5	19		29		39	
10	SL-504-SA8-SB-2.5-3.5MS	20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F
Sampling date: 4/11/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification										
	FB-041113 (SDG: PH029)	Action Limit	1	2	4	5	7	9				
Cu	0.0036	1.8										
Mo	0.0036	1.8	0.37	0.63	0.52	0.37	0.34	0.57				

Sampling date: 7/17/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: 2-9

Analyte	Blank ID	Sample Identification										
	EB1-071713 (SDG: PH069)	Action Limit	No Qualifiers									
Sb	0.0069	3.45										

Sampling date: 7/17/13 Soil factor applied 100x
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: 1

Analyte	Blank ID	Sample Identification										
	EB2-071713 (SDG: PH069)	Action Limit	No Qualifiers									
Sb	0.0054	2.7										

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

Background Lab Sample ID: 7129596BKG Matrix Spike Lab Sample ID: 7129597MS Matrix Spike Duplicate Lab Sample ID: 7129598MSD
Batch Id(s): P19837A, P19838A

Analyte	Mass	BKG Sample		MS Sample		MSD Sample		MS Spike Added	MSD Spike Added	Units	MS		MSD		RPD	Control Limit		
		Result	C	Result	C	Result	C				%R	Q	%R	Q		%R	RPD	M
Aluminum		1327.6627		1385.3184		13428.4718		194.1748	194.1748	MG/KG	1060		1082		0			20 P
Antimony		0.7255	U	34.9291		33.8476		48.5437	48.5437	MG/KG	72	N	70	N	3		75 - 125	20 P
Arsenic		3.7275	B	18.0049		18.1631		14.5631	14.5631	MG/KG	98		99		1		75 - 125	20 P
Barium		81.0618		307.7767		273.9204		194.1748	194.1748	MG/KG	117		99		12		75 - 125	20 P
Beryllium		0.3784	B	5.2631		5.2864		4.8544	4.8544	MG/KG	101		101		0		75 - 125	20 P
Boron		17.3794		199.4437		198.6350		194.1748	194.1748	MG/KG	94		93		0		75 - 125	20 P
Cadmium		0.3559	B	5.0592		5.1650		4.8544	4.8544	MG/KG	97		99		2		75 - 125	20 P
Carbon		3523.0082		4307.0825		4170.4408		388.3495	388.3495	MG/KG	202		167		3			20 P
Chromium		20.9186		46.9544		41.4087		19.4175	19.4175	MG/KG	134	N	106		13		75 - 125	20 P
Cobalt		4.6186		50.3835		50.6864		48.5437	48.5437	MG/KG	94		95		1		75 - 125	20 P
Copper		10.2118		37.2107		37.1495		24.2718	24.2718	MG/KG	111		111		0		75 - 125	20 P
Iron		21975.0941		22322.6282		22179.5563		97.0874	97.0874	MG/KG	358		211		1			20 P
Lead		4.2559		18.0942		18.4777		14.5631	14.5631	MG/KG	95		98		2		75 - 125	20 P
Lithium		28.6725		119.7583		120.4107		97.0874	97.0874	MG/KG	94		94		1		75 - 125	20 P
Magnesium		5846.4451		6937.3311		6279.2155		194.1748	194.1748	MG/KG	562		223		10			20 P
Manganese		405.2775		469.9534		449.7078		48.5437	48.5437	MG/KG	133		92		4			20 P
Mercury		0.0097	U	0.1890		0.1826		0.1599	0.1603	MG/KG	118		114		3		65 - 135	20 CV
Molybdenum		0.3588	B	186.4670		187.1864		194.1748	194.1748	MG/KG	96		96		0		75 - 125	20 P
Nickel		12.1176		64.9097		59.6282		48.5437	48.5437	MG/KG	109		98		8		75 - 125	20 P
Phosphorus		406.5922		535.7379		494.3117		97.0874	97.0874	MG/KG	133		90		8			20 P
Potassium		2129.3402		3367.6641		3436.6699		970.8738	970.8738	MG/KG	128	N	135	N	2		75 - 125	20 P
Selenium	78	0.0980	U	2.0058		2.2485		1.9417	1.9417	MG/KG	103		116		11		75 - 125	20 MS
Silver	107	0.0255	U	10.0893		9.7282		9.7087	9.7087	MG/KG	104		100		4		75 - 125	20 MS
Sodium		170.4441		1156.7019		1154.8262		970.8738	970.8738	MG/KG	102		101		0		75 - 125	20 P
Strontium	88	14.3431		22.0388		22.2330		7.7670	7.7670	MG/KG	99		102		1		75 - 125	20 MS
Thallium	203	0.2286		0.6208		0.6159		0.3883	0.3883	MG/KG	101		100		1		75 - 125	20 MS
Tin		2.7510	B	355.3340		353.8466		388.3495	388.3495	MG/KG	91		90		0		75 - 125	20 P
Titanium		1403.0598		1735.5447		1723.4524		97.0874	97.0874	MG/KG	342		330		1			20 P
Vanadium		42.0873		91.8524		90.2825		48.5437	48.5437	MG/KG	103		99		2		75 - 125	20 P
Zinc		51.5088		101.0981		101.8146		48.5437	48.5437	MG/KG	102		104		1		75 - 125	20 P
Zirconium		3.0451	B	99.2233		99.3951		97.0874	97.0874	MG/KG	99		99		0		75 - 125	20 P

Note: Results shown are reported on an as-received basis.

METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry	CV = Cold Vapor AF = Cold Vapor Atomic Fluorescence	CONCENTRATION QUALIFIERS: U= Below MDL, B= Below LOQ FLAGS: N = Matrix Spike OOS, * = Duplicate OOS
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QUALITY ASSURANCE SUMMARY

FORM 6

DUPLICATES

SDG No.: PH068

Matrix: SOIL Level (low/med): LOW

Background Lab Sample ID: 7129596BKG

Duplicate Lab Sample ID: 7129602DUP

Batch ID(s): P19837A, P19838A

Concentration Units: MG/KG

5x LOQ

Analyte	Mass	Control Limit	Samples (S)	C	Duplicate (D)	C	RPD	Q	M
Aluminum			11327.6627		11207.6235		1		P
Antimony			-0.7412	B	0.7255	U	200		P
Arsenic			3.7275	B	3.2853	B	13		P
Barium			81.0618		80.5412		1		P
Beryllium			0.3784	B	0.3824	B	1		P
Boron		9.8	17.3794		16.3549		6		P
Cadmium			0.3559	B	0.3363	B	6		P
Calcium			3523.0088		3411.9392		3		P
Chromium			20.9186		20.5784		2		P
Cobalt		1.0	4.6186		4.4471		4		P
Copper			10.2118		10.4049		2		P
Iron			21975.0941		22008.4010		0		P
Lead		2.9	4.2559		3.8814		9		P
Lithium			28.6725		27.3255		5		P
Magnesium			5846.4451		5923.3647		1		P
Manganese			405.2775		406.3461		0		P
Mercury			0.0097	U	0.0097	U			CV
Molybdenum			0.3588	B	0.2383	B	42		P
Nickel			12.1176		12.1069		0		P
Phosphorus			406.5922		403.1422		1		P
Potassium			2129.3402		2166.8676		2		P
Selenium	78		0.0980	U	0.0980	U			MS
Silver	107		0.0255	U	0.0255	U			MS
Sodium		98.0	170.4441		168.4804		1		P
Strontium	88		14.3431		13.7255		4		MS
Thallium	203	0.2	0.2286		0.2006		13		MS
Tin			2.7510	B	2.5853	B	6		P
Titanium			1403.0598		1357.2049		3		P
Vanadium			42.0873		40.8549		3		P
Zinc			51.5088		51.3980		0		P
Zirconium			3.0451	B	3.8922	B	24		P

NOTE: An asterisk (*) in column "Q" indicates poor duplicate precision (RPD > 20% OR |(S) - (D)| > LOQ for values < 5x LOQ).

The data are considered to be valid because the laboratory control sample is within the control limits. See the Laboratory Control Sample.

ok by reference

Note: Results shown are reported on an as-received basis.

<p>METHODS:</p> <p>P = ICP Atomic Emission Spectrometer</p> <p>MS = ICP Mass Spectrometry</p> <p>CV = Cold Vapor</p> <p>AF = Cold Vapor Atomic Fluorescence</p>	<p>CONCENTRATION QUALIFIERS:</p> <p>U= Below MDL</p> <p>B= Below LOQ</p> <p>FLAGS:</p> <p>3433 of 3852</p> <p>Duplicate Out of Spec</p>
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Background Lab Sample ID: 7129596BKG Serial Dilution Lab Sample ID: 7129596L
 Batch ID(s): P19837A
 Concentration Units: UG/L

Analyte	Mass	Initial Sample Result (I)	C	Serial Dilution Result (S)	C	% Diff.	Q	M
Aluminum		115542.1600		108978.8500		6		P
Antimony		7.4000	U	37.0000	U			P
Arsenic		38.0200	B	37.1500	B	2		P
Barium		826.8300		785.5500		5		P
Beryllium		3.8600	B	3.3500	U	100		P
Boron		177.2700		191.3000	B	8		P
Cadmium		3.6300	B	3.8000	U	100		P
Calcium		35934.6900		34214.4000		5		P
Chromium		213.3700		192.8500		10		P
Cobalt		47.1100		48.1000	B	2		P
Copper		104.1600		99.4500	B	5		P
Iron		224145.9600		210061.5000		6		P
Lead		43.4100		29.7000	B	32		P
Lead		22.4600		240.4000		18		E P
Magnesium		59633.7400		56247.8500		6		P
Manganese		4133.8300		3956.6000		4		P
Molybdenum		3.6600	B	11.2000	B	206		P
Nickel		123.6000		118.9000		4		P
Phosphorus		4147.2400		3893.5500		6		P
Potassium		21719.2700		20985.6000		3		P
Selenium	78	0.5000	U	2.5000	U			MS
Silver	107	0.1300	U	0.6500	U			MS
Sodium		1738.5300		1473.9500	B	15		P
Strontium	88	73.1500		75.1500		3		MS
Thallium	203	1.1660		0.7500	U	100		MS
Tin		28.0600	B	25.3500	B	10		P
Titanium		14311.2100		13194.8500		8		P
Vanadium		429.2900		390.7000		9		P
Zinc		525.3900		486.6000		7		P
Zirconium		31.0600	B	42.0000	U	100		P

NOTE: An E in column Q indicates the presence of a chemical or physical interference in the matrix when the % difference is greater than 10%. This applies only when (I) is greater than or equal to 50x MDL for ICP, 100x MDL for ICP-MS (6020), 50x MDL for ICP-MS (200.8), or 25x MDL for GFAA.



<p>METHODS: P = ICP Atomic Emission Spectrometer MS = ICP Mass Spectrometry</p>	<p>CONCENTRATION QUALIFIERS: U= Below MDL B= Below LOQ</p> <p>FLAGS: E = Matrix Effects exist as proven by Serial Dilution or Spiked Dilution</p>
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**Data Validation Report
Santa Susana Field Laboratory**

Subarea 8

SDG: PH069

Prepared for

CDM
555 17th Street, Suite 1100
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Prepared by

Laboratory Data Consultants, Inc
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September 24, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on July 17th, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Volatiles (VOAs) by EPA SW 846 Method 8260B
Semivolatiles (SVOAs) by EPA 846 Method 8270D
SVOAs by EPA SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Pesticides by EPA SW 846 Method 8081B
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A, 7470A and 7471B
Herbicides by EPA SW 846 Method 8151A
Total Petroleum Hydrocarbons as Gasoline (TPH-G) by EPA SW 846 Method 8015M
Total Petroleum Hydrocarbons as Extractables (TPH-E) by EPA SW 846 Method 8015M
Dioxins/Dibenzofurans by EPA Method 1613B
Perchlorate by EPA Method 6850
Hexavalent Chromium by EPA Method 7199

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I. DVRs for samples on which EPA Level IV validation was performed are presented in Enclosure II.

All sample results were subjected to Level IV data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibrations (ICB/CCBs), surrogates, internal standards, interference check (ICSA and ICSAB) samples, matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), method blanks, trip blanks, equipment blanks, field blanks, and review of the QC summary forms as well as the raw data, to confirm sample quantitation and identification.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the calibrations, internal standard, and interference check samples, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

II. Initial Calibration

All criteria for the initial calibration of each method were met.

III. Continuing Calibration

All criteria for the initial calibration verifications and continuing calibration of each method were met with the exception of several VOAs, SVOAs, and Herbicides. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosures I and II.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of four blanks for SVOAs, dioxins, and metals. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosures I and II.

No contaminant concentrations were detected in the initial or continuing calibration blanks with the exception of several metals. The associated sample results were qualified as non-detected (U) due to initial or continuing calibration blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure II.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

The frequency of analysis was met.

The criteria for analysis were met.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for herbicides, TPH-E, and metals. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosures I and II.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of several LCS/LCSD pairs for VOAs, SVOAs, pesticides, and herbicides. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding the qualification of data are provided in Enclosures I and II.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met with the exception of Lithium. The details regarding the qualification of data are provided in Enclosure II.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

The 2nd column 2,3,7,8-TCDF confirmation for dioxin analysis was not performed for several samples. No data were qualified based on lack of 2,3,7,8-TCDF confirmation analysis. Details are provided in Enclosure II.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

No field duplicates were identified in this SDG.

XIV. Field Blank Samples

One trip blank was collected and analyzed for VOAs and TPH-G. No contaminant concentrations were found in the trip blank.

Two equipment blanks were collected and analyzed for VOAs, SVOAs, pesticides, PCBs, metals, herbicides, perchlorate, TPH-G, TPH-E, hexavalent chromium, and dioxins. The associated sample results were qualified as non-detected (U) due to equipment blank

contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the equipment blank were not qualified. The equipment blank outlier reports are presented in Enclosures I and II.

One field blank (from SDG PH029) was collected and analyzed for VOAs, SVOAs, pesticides, PCBs, metals, herbicides, perchlorate, TPH-G, TPH-E, hexavalent chromium, and dioxins. The field blank had detections for several VOAs, SVOAs, metals, and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosures I and II.

XV. Overall Assessment of Data

No data associated with this sampling event were rejected. The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. All data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
17-Jul-2013	SL-512-SA8-SB-0.0-0.5	7130897	N	3050B	6010C	IV
17-Jul-2013	SL-512-SA8-SB-0.0-0.5	7130897	N	3050B	6020A	IV
17-Jul-2013	SL-512-SA8-SB-0.0-0.5	7130897	N	3546	8015M	IV
17-Jul-2013	SL-512-SA8-SB-0.0-0.5	7130897	N	3546	8082A	IV
17-Jul-2013	SL-512-SA8-SB-0.0-0.5	7130897	N	3546	8270D SIM	IV
17-Jul-2013	SL-512-SA8-SB-0.0-0.5	7130897	N	METHOD	1613B	IV
17-Jul-2013	SL-512-SA8-SB-0.0-0.5	7130897	N	METHOD	7471B	IV
17-Jul-2013	TB-071713	7130905	TB	5030B	8015M	IV
17-Jul-2013	TB-071713	7130905	TB	5030B	8260B	IV
17-Jul-2013	SL-512-SA8-SB-0.0-0.5DUP	P130897D220947	DUP	METHOD	7471B	IV
17-Jul-2013	SL-512-SA8-SB-0.0-0.5MSD	P130897M220951	MSD	METHOD	7471B	IV
17-Jul-2013	SL-512-SA8-SB-0.0-0.5MS	P130897R220949	MS	METHOD	7471B	IV
17-Jul-2013	SL-514-SA8-SB-0.0-0.5	7130898	N	3050B	6010C	IV
17-Jul-2013	SL-514-SA8-SB-0.0-0.5	7130898	N	3050B	6020A	IV
17-Jul-2013	SL-514-SA8-SB-0.0-0.5	7130898	N	3546	8015M	IV
17-Jul-2013	SL-514-SA8-SB-0.0-0.5	7130898	N	3546	8081B	IV
17-Jul-2013	SL-514-SA8-SB-0.0-0.5	7130898	N	3546	8270D SIM	IV
17-Jul-2013	SL-514-SA8-SB-0.0-0.5	7130898	N	3550B	8151A	IV
17-Jul-2013	SL-514-SA8-SB-0.0-0.5	7130898	N	METHOD	1613B	IV
17-Jul-2013	SL-514-SA8-SB-0.0-0.5	7130898	N	METHOD	7471B	IV
17-Jul-2013	SL-514-SA8-SB-0.0-0.5MSD	P130898M240418A	MSD	3550B	8151A	IV
17-Jul-2013	SL-514-SA8-SB-0.0-0.5MS	P130898R240351A	MS	3550B	8151A	IV
17-Jul-2013	SL-514-SA8-SB-4.0-5.0	7130899	N	3050B	6010C	IV
17-Jul-2013	SL-514-SA8-SB-4.0-5.0	7130899	N	3050B	6020A	IV
17-Jul-2013	SL-514-SA8-SB-4.0-5.0	7130899	N	3546	8015M	IV
17-Jul-2013	SL-514-SA8-SB-4.0-5.0	7130899	N	3546	8081B	IV

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
17-Jul-2013	SL-514-SA8-SB-4.0-5.0	7130899	N	3546	8270D SIM	IV
17-Jul-2013	SL-514-SA8-SB-4.0-5.0	7130899	N	3550B	8151A	IV
17-Jul-2013	SL-514-SA8-SB-4.0-5.0	7130899	N	5035A	8015M	IV
17-Jul-2013	SL-514-SA8-SB-4.0-5.0	7130899	N	METHOD	1613B	IV
17-Jul-2013	SL-514-SA8-SB-4.0-5.0	7130899	N	METHOD	7471B	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5	7130900	N	3050B	6010C	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5	7130900	N	3050B	6020A	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5	7130900	N	3546	8015M	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5	7130900	N	3546	8081B	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5	7130900	N	3546	8270D SIM	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5	7130900	N	3550B	8151A	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5	7130900	N	5035A	8015M	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5	7130900	N	METHOD	7471B	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5DUP	P130900D220901	DUP	3050B	6010C	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5DUP	P130900D221804A	DUP	3050B	6020A	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5DUP	P130900D221912B	DUP	3050B	6020A	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5MSD	P130900M220909	MSD	3050B	6010C	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5MSD	P130900M221808A	MSD	3050B	6020A	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5MSD	P130900M221917B	MSD	3050B	6020A	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5MSD	P130900M321931A	MSD	3546	8015M	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5MS	P130900R220905	MS	3050B	6010C	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5MS	P130900R221806A	MS	3050B	6020A	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5MS	P130900R221914B	MS	3050B	6020A	IV
17-Jul-2013	SL-514-SA8-SB-7.5-8.5MS	P130900R321909A	MS	3546	8015M	IV
17-Jul-2013	SL-515-SA8-SB-0.0-0.5	7130901	N	3050B	6010C	IV
17-Jul-2013	SL-515-SA8-SB-0.0-0.5	7130901	N	3050B	6020A	IV

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
17-Jul-2013	SL-515-SA8-SB-0.0-0.5	7130901	N	3546	8015M	IV
17-Jul-2013	SL-515-SA8-SB-0.0-0.5	7130901	N	3546	8082A	IV
17-Jul-2013	SL-515-SA8-SB-0.0-0.5	7130901	N	3546	8270D SIM	IV
17-Jul-2013	SL-515-SA8-SB-0.0-0.5	7130901	N	METHOD	1613B	IV
17-Jul-2013	SL-515-SA8-SB-0.0-0.5	7130901	N	METHOD	6850	IV
17-Jul-2013	SL-515-SA8-SB-0.0-0.5	7130901	N	METHOD	7471B	IV
17-Jul-2013	SL-515-SA8-SB-4.0-5.0	7130902	N	3050B	6010C	IV
17-Jul-2013	SL-515-SA8-SB-4.0-5.0	7130902	N	3050B	6020A	IV
17-Jul-2013	SL-515-SA8-SB-4.0-5.0	7130902	N	3546	8015M	IV
17-Jul-2013	SL-515-SA8-SB-4.0-5.0	7130902	N	3546	8082A	IV
17-Jul-2013	SL-515-SA8-SB-4.0-5.0	7130902	N	3546	8270D SIM	IV
17-Jul-2013	SL-515-SA8-SB-4.0-5.0	7130902	N	5035A	8015M	IV
17-Jul-2013	SL-515-SA8-SB-4.0-5.0	7130902	N	METHOD	1613B	IV
17-Jul-2013	SL-515-SA8-SB-4.0-5.0	7130902	N	METHOD	6850	IV
17-Jul-2013	SL-515-SA8-SB-4.0-5.0	7130902	N	METHOD	7471B	IV
17-Jul-2013	SL-515-SA8-SB-9.0-10.0	7130903	N	3050B	6010C	IV
17-Jul-2013	SL-515-SA8-SB-9.0-10.0	7130903	N	3050B	6020A	IV
17-Jul-2013	SL-515-SA8-SB-9.0-10.0	7130903	N	3546	8015M	IV
17-Jul-2013	SL-515-SA8-SB-9.0-10.0	7130903	N	3546	8082A	IV
17-Jul-2013	SL-515-SA8-SB-9.0-10.0	7130903	N	3546	8270D SIM	IV
17-Jul-2013	SL-515-SA8-SB-9.0-10.0	7130903	N	5035A	8015M	IV
17-Jul-2013	SL-515-SA8-SB-9.0-10.0	7130903	N	METHOD	1613B	IV
17-Jul-2013	SL-515-SA8-SB-9.0-10.0	7130903	N	METHOD	6850	IV
17-Jul-2013	SL-515-SA8-SB-9.0-10.0	7130903	N	METHOD	7471B	IV
17-Jul-2013	SL-515-SA8-SB-14.0-15.0	7130904	N	3050B	6010C	IV
17-Jul-2013	SL-515-SA8-SB-14.0-15.0	7130904	N	3050B	6020A	IV

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
17-Jul-2013	SL-515-SA8-SB-14.0-15.0	7130904	N	3546	8015M	IV
17-Jul-2013	SL-515-SA8-SB-14.0-15.0	7130904	N	3546	8082A	IV
17-Jul-2013	SL-515-SA8-SB-14.0-15.0	7130904	N	3546	8270D SIM	IV
17-Jul-2013	SL-515-SA8-SB-14.0-15.0	7130904	N	5035A	8015M	IV
17-Jul-2013	SL-515-SA8-SB-14.0-15.0	7130904	N	METHOD	1613B	IV
17-Jul-2013	SL-515-SA8-SB-14.0-15.0	7130904	N	METHOD	6850	IV
17-Jul-2013	SL-515-SA8-SB-14.0-15.0	7130904	N	METHOD	7471B	IV
17-Jul-2013	SL-513-SA8-SB-0.0-0.5	7130906	N	3050B	6010C	IV
17-Jul-2013	SL-513-SA8-SB-0.0-0.5	7130906	N	3050B	6020A	IV
17-Jul-2013	SL-513-SA8-SB-0.0-0.5	7130906	N	3060A	7199	IV
17-Jul-2013	SL-513-SA8-SB-0.0-0.5	7130906	N	3546	8015M	IV
17-Jul-2013	SL-513-SA8-SB-0.0-0.5	7130906	N	3546	8082A	IV
17-Jul-2013	SL-513-SA8-SB-0.0-0.5	7130906	N	3546	8270D	IV
17-Jul-2013	SL-513-SA8-SB-0.0-0.5	7130906	N	3546	8270D SIM	IV
17-Jul-2013	SL-513-SA8-SB-0.0-0.5	7130906	N	METHOD	1613B	IV
17-Jul-2013	SL-513-SA8-SB-0.0-0.5	7130906	N	METHOD	6850	IV
17-Jul-2013	SL-513-SA8-SB-0.0-0.5	7130906	N	METHOD	7471B	IV
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	3050B	6010C	IV
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	3050B	6020A	IV
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	3060A	7199	IV
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	3546	8015M	IV
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	3546	8082A	IV
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	3546	8270D	IV
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	3546	8270D SIM	IV
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	5035A	8015M	IV
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	METHOD	1613B	IV

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	METHOD	6850	IV
17-Jul-2013	SL-513-SA8-SB-4.0-5.0	7130907	N	METHOD	7471B	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	3050B	6010C	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	3050B	6020A	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	3060A	7199	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	3546	8015M	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	3546	8082A	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	3546	8270D	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	3546	8270D SIM	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	5035A	8015M	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	5035A	8260B	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	METHOD	1613B	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	METHOD	6850	IV
17-Jul-2013	SL-513-SA8-SB-8.0-9.0	7130908	N	METHOD	7471B	IV
17-Jul-2013	EB1-071713	7130895	EB	3005A	6010C	IV
17-Jul-2013	EB1-071713	7130895	EB	3510C	8015M	IV
17-Jul-2013	EB1-071713	7130895	EB	3510C	8081B	IV
17-Jul-2013	EB1-071713	7130895	EB	3510C	8082A	IV
17-Jul-2013	EB1-071713	7130895	EB	3510C	8270D	IV
17-Jul-2013	EB1-071713	7130895	EB	3510C	8270D SIM	IV
17-Jul-2013	EB1-071713	7130895	EB	5030B	8015M	IV
17-Jul-2013	EB1-071713	7130895	EB	5030B	8260B	IV
17-Jul-2013	EB1-071713	7130895	EB	Gen Prep	6850	IV
17-Jul-2013	EB1-071713	7130895	EB	Gen Prep	7199	IV
17-Jul-2013	EB1-071713	7130895	EB	M3010A	6020A	IV
17-Jul-2013	EB1-071713	7130895	EB	METHOD	1613B	IV

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
17-Jul-2013	EB1-071713	7130895	EB	METHOD	7470A	IV
17-Jul-2013	EB1-071713	7130895	EB	METHOD	8151A	IV
17-Jul-2013	EB1-071713MSD	P130895M241736A	MSD	Gen Prep	6850	IV
17-Jul-2013	EB1-071713MS	P130895R241723A	MS	Gen Prep	6850	IV
17-Jul-2013	EB2-071713	7130896	EB	3005A	6010C	IV
17-Jul-2013	EB2-071713	7130896	EB	3510C	8015M	IV
17-Jul-2013	EB2-071713	7130896	EB	3510C	8081B	IV
17-Jul-2013	EB2-071713	7130896	EB	3510C	8082A	IV
17-Jul-2013	EB2-071713	7130896	EB	3510C	8270D SIM	IV
17-Jul-2013	EB2-071713	7130896	EB	5030B	8015M	IV
17-Jul-2013	EB2-071713	7130896	EB	Gen Prep	7199	IV
17-Jul-2013	EB2-071713	7130896	EB	M3010A	6020A	IV
17-Jul-2013	EB2-071713	7130896	EB	METHOD	1613B	IV
17-Jul-2013	EB2-071713	7130896	EB	METHOD	7470A	IV
17-Jul-2013	EB2-071713	7130896	EB	METHOD	8151A	IV

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: AQ

Sample ID: EB1-071713 Collected: 7/17/2013 3:00:00 Analysis Type: REA3 Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	0.0069	J	0.0053	MDL	0.0400	PQL	mg/L	U	B

Sample ID: EB2-071713 Collected: 7/17/2013 3:30:00 Analysis Type: REA3 Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	0.0054	J	0.0053	MDL	0.0400	PQL	mg/L	U	B

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-512-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	3.90	U	0.722	MDL	3.90	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.593	J	0.0654	MDL	0.976	PQL	mg/Kg	J	Z
CADMIUM	0.235	J	0.0742	MDL	0.976	PQL	mg/Kg	J	Z
LITHIUM	22.8		0.33	MDL	3.9	PQL	mg/Kg	J	A
MOLYBDENUM	0.340	J	0.166	MDL	1.95	PQL	mg/Kg	U	F
POTASSIUM	3090		8.14	MDL	97.6	PQL	mg/Kg	J	Q
SODIUM	77.2	J	16.3	MDL	97.6	PQL	mg/Kg	J	Z
TIN	2.98	J	0.215	MDL	9.76	PQL	mg/Kg	U	B

Sample ID: SL-513-SA8-SB-0.0-0.5 Collected: 7/17/2013 1:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	1.20	J	0.792	MDL	4.28	PQL	mg/Kg	UJ	Q, B, F
BERYLLIUM	0.659	J	0.0717	MDL	1.07	PQL	mg/Kg	J	Z
BORON	2.19	J	0.899	MDL	10.7	PQL	mg/Kg	J	Z
CADMIUM	0.104	J	0.0813	MDL	1.07	PQL	mg/Kg	J	Z
LITHIUM	25.9		0.36	MDL	4.3	PQL	mg/Kg	J	A
MOLYBDENUM	0.861	J	0.182	MDL	2.14	PQL	mg/Kg	U	F
POTASSIUM	5480		8.93	MDL	107	PQL	mg/Kg	J	Q
TIN	3.28	J	0.235	MDL	10.7	PQL	mg/Kg	U	B
Zirconium	2.54	J	0.899	MDL	5.35	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-513-SA8-SB-4.0-5.0 Collected: 7/17/2013 2:00:00 Analysis Type: REA2 Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.40	U	0.814	MDL	4.40	PQL	mg/Kg	UJ	Q
BORON	10.2	J	0.924	MDL	11.0	PQL	mg/Kg	J	Z
CADMIUM	0.178	J	0.0836	MDL	1.10	PQL	mg/Kg	J	Z
LITHIUM	22.3		0.37	MDL	4.4	PQL	mg/Kg	J	A

Sample ID: SL-513-SA8-SB-4.0-5.0 Collected: 7/17/2013 2:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.202	J	0.0737	MDL	1.10	PQL	mg/Kg	J	Z
POTASSIUM	2320		9.17	MDL	110	PQL	mg/Kg	J	Q
TIN	2.78	J	0.242	MDL	11.0	PQL	mg/Kg	U	B
Zirconium	2.87	J	0.924	MDL	5.50	PQL	mg/Kg	J	Z

Sample ID: SL-513-SA8-SB-8.0-9.0 Collected: 7/17/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	3.94	U	0.730	MDL	3.94	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.408	J	0.0661	MDL	0.986	PQL	mg/Kg	J	Z
CADMIUM	0.218	J	0.0750	MDL	0.986	PQL	mg/Kg	J	Z
LITHIUM	21.9		0.34	MDL	3.9	PQL	mg/Kg	J	A
POTASSIUM	1830		8.22	MDL	98.6	PQL	mg/Kg	J	Q
TIN	3.13	J	0.217	MDL	9.86	PQL	mg/Kg	U	B
Zirconium	1.91	J	0.828	MDL	4.93	PQL	mg/Kg	J	Z

Sample ID: SL-514-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.22	U	0.781	MDL	4.22	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.845	J	0.0707	MDL	1.06	PQL	mg/Kg	J	Z
LITHIUM	23.2		0.36	MDL	4.2	PQL	mg/Kg	J	A
POTASSIUM	5440		8.81	MDL	106	PQL	mg/Kg	J	Q
TIN	3.33	J	0.232	MDL	10.6	PQL	mg/Kg	U	B
Zirconium	3.38	J	0.887	MDL	5.28	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-514-SA8-SB-4.0-5.0 Collected: 7/17/2013 9:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	7.33		0.868	MDL	4.69	PQL	mg/Kg	J	Q
BERYLLIUM	0.378	J	0.0786	MDL	1.17	PQL	mg/Kg	J	Z
BORON	2.51	J	0.985	MDL	11.7	PQL	mg/Kg	J	Z
CADMIUM	0.223	J	0.0892	MDL	1.17	PQL	mg/Kg	J	Z
LITHIUM	21.4		0.40	MDL	4.7	PQL	mg/Kg	J	A
POTASSIUM	2590		9.78	MDL	117	PQL	mg/Kg	J	Q
TIN	3.05	J	0.258	MDL	11.7	PQL	mg/Kg	U	B
Zirconium	3.34	J	0.985	MDL	5.87	PQL	mg/Kg	J	Z

Sample ID: SL-514-SA8-SB-7.5-8.5 Collected: 7/17/2013 10:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	5.55		0.863	MDL	4.66	PQL	mg/Kg	J	Q
BERYLLIUM	0.409	J	0.0781	MDL	1.17	PQL	mg/Kg	J	Z
BORON	1.24	J	0.979	MDL	11.7	PQL	mg/Kg	J	Z
CADMIUM	0.117	J	0.0886	MDL	1.17	PQL	mg/Kg	J	Z
LITHIUM	21.0		0.40	MDL	4.7	PQL	mg/Kg	J	A
MOLYBDENUM	2.30	J	0.198	MDL	2.33	PQL	mg/Kg	J	Z
POTASSIUM	2690		9.72	MDL	117	PQL	mg/Kg	J	Q
TIN	3.00	J	0.256	MDL	11.7	PQL	mg/Kg	U	B
Zirconium	3.38	J	0.979	MDL	5.83	PQL	mg/Kg	J	Z

Sample ID: SL-515-SA8-SB-0.0-0.5 Collected: 7/17/2013 10:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.42	U	0.817	MDL	4.42	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.890	J	0.0740	MDL	1.10	PQL	mg/Kg	J	Z
CADMIUM	0.116	J	0.0839	MDL	1.10	PQL	mg/Kg	J	Z
LITHIUM	23.0		0.38	MDL	4.4	PQL	mg/Kg	J	A
POTASSIUM	6010		9.21	MDL	110	PQL	mg/Kg	J	Q
TIN	3.43	J	0.243	MDL	11.0	PQL	mg/Kg	U	B
Zirconium	3.76	J	0.928	MDL	5.52	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-515-SA8-SB-14.0-15.0 Collected: 7/17/2013 11:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.48	U	0.830	MDL	4.48	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.869	J	0.0751	MDL	1.12	PQL	mg/Kg	J	Z
LITHIUM	19.0		0.38	MDL	4.5	PQL	mg/Kg	J	A
POTASSIUM	2480		9.35	MDL	112	PQL	mg/Kg	J	Q
TIN	3.34	J	0.247	MDL	11.2	PQL	mg/Kg	U	B
Zirconium	2.60	J	0.942	MDL	5.61	PQL	mg/Kg	J	Z

Sample ID: SL-515-SA8-SB-4.0-5.0 Collected: 7/17/2013 11:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.42	U	0.817	MDL	4.42	PQL	mg/Kg	UJ	Q
BERYLLIUM	0.984	J	0.0740	MDL	1.10	PQL	mg/Kg	J	Z
LITHIUM	26.5		0.38	MDL	4.4	PQL	mg/Kg	J	A
POTASSIUM	3110		9.21	MDL	110	PQL	mg/Kg	J	Q
TIN	3.38	J	0.243	MDL	11.0	PQL	mg/Kg	U	B
Zirconium	3.04	J	0.928	MDL	5.52	PQL	mg/Kg	J	Z

Sample ID: SL-515-SA8-SB-9.0-10.0 Collected: 7/17/2013 11:20:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ANTIMONY	4.71	U	0.872	MDL	4.71	PQL	mg/Kg	UJ	Q
BERYLLIUM	1.13	J	0.0789	MDL	1.18	PQL	mg/Kg	J	Z
LITHIUM	27.4		0.40	MDL	4.7	PQL	mg/Kg	J	A
POTASSIUM	2930		9.83	MDL	118	PQL	mg/Kg	J	Q
TIN	3.76	J	0.259	MDL	11.8	PQL	mg/Kg	U	B
Zirconium	4.44	J	0.990	MDL	5.89	PQL	mg/Kg	J	Z

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-512-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:00:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.204	J	0.0976	MDL	0.390	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-512-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:00:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0426	J	0.0254	MDL	0.195	PQL	mg/Kg	J	Z

Sample ID: SL-513-SA8-SB-0.0-0.5 Collected: 7/17/2013 1:45:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.270	J	0.107	MDL	0.428	PQL	mg/Kg	J	Z

Sample ID: SL-513-SA8-SB-0.0-0.5 Collected: 7/17/2013 1:45:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0396	J	0.0278	MDL	0.214	PQL	mg/Kg	J	Z

Sample ID: SL-513-SA8-SB-4.0-5.0 Collected: 7/17/2013 2:00:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.119	J	0.110	MDL	0.440	PQL	mg/Kg	J	Z

Sample ID: SL-513-SA8-SB-4.0-5.0 Collected: 7/17/2013 2:00:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0473	J	0.0286	MDL	0.220	PQL	mg/Kg	J	Z
THALLIUM	0.215	J	0.0330	MDL	0.220	PQL	mg/Kg	J	Z

Sample ID: SL-514-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:55:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.256	J	0.106	MDL	0.422	PQL	mg/Kg	J	Z

Sample ID: SL-514-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:55:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0469	J	0.0275	MDL	0.211	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-514-SA8-SB-4.0-5.0		Collected: 7/17/2013 9:55:00		Analysis Type: RES		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0467	J	0.0305	MDL	0.235	PQL	mg/Kg	J	Z

Sample ID: SL-514-SA8-SB-7.5-8.5		Collected: 7/17/2013 10:15:00		Analysis Type: REA2		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.171	J	0.117	MDL	0.466	PQL	mg/Kg	J	Z

Sample ID: SL-515-SA8-SB-0.0-0.5		Collected: 7/17/2013 10:50:00		Analysis Type: REA2		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.258	J	0.110	MDL	0.442	PQL	mg/Kg	J	Z

Sample ID: SL-515-SA8-SB-0.0-0.5		Collected: 7/17/2013 10:50:00		Analysis Type: RES		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0440	J	0.0287	MDL	0.221	PQL	mg/Kg	J	Z

Sample ID: SL-515-SA8-SB-4.0-5.0		Collected: 7/17/2013 11:05:00		Analysis Type: RES		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0353	J	0.0287	MDL	0.221	PQL	mg/Kg	J	Z

Sample ID: SL-515-SA8-SB-9.0-10.0		Collected: 7/17/2013 11:20:00		Analysis Type: REA2		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.137	J	0.118	MDL	0.471	PQL	mg/Kg	J	Z

Sample ID: SL-515-SA8-SB-9.0-10.0		Collected: 7/17/2013 11:20:00		Analysis Type: RES		Dilution: 2			
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0431	J	0.0306	MDL	0.236	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	7199	Matrix: SO

Sample ID: SL-513-SA8-SB-4.0-5.0 Collected: 7/17/2013 2:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.18	J	0.16	MDL	0.44	PQL	mg/Kg	J	Z

Method Category:	METALS	
Method:	7471B	Matrix: SO

Sample ID: SL-515-SA8-SB-0.0-0.5 Collected: 7/17/2013 10:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0136	J	0.0110	MDL	0.0184	PQL	mg/Kg	J	Z

Method Category:	SVOA	
Method:	1613B	Matrix: AQ

Sample ID: EB1-071713 Collected: 7/17/2013 3:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.413	JBQ	0.170	MDL	9.98	PQL	pg/L	U	B
1,2,3,4,6,7,8-HPCDF	0.238	JB	0.0620	MDL	9.98	PQL	pg/L	U	B
1,2,3,4,7,8,9-HPCDF	0.191	JBQ	0.0696	MDL	9.98	PQL	pg/L	U	B
1,2,3,4,7,8-HxCDD	0.161	JBQ	0.148	MDL	9.98	PQL	pg/L	U	B
1,2,3,4,7,8-HxCDF	0.222	JBQ	0.0723	MDL	9.98	PQL	pg/L	U	B
1,2,3,6,7,8-HxCDF	0.191	JBQ	0.0753	MDL	9.98	PQL	pg/L	U	B
1,2,3,7,8,9-HxCDF	0.186	JB	0.0815	MDL	9.98	PQL	pg/L	U	B
1,2,3,7,8-PECDF	0.406	JB	0.142	MDL	9.98	PQL	pg/L	U	B
2,3,4,6,7,8-HxCDF	0.224	JB	0.0716	MDL	9.98	PQL	pg/L	U	B
2,3,4,7,8-PECDF	0.132	JBQ	0.131	MDL	9.98	PQL	pg/L	U	B
OCDD	0.814	JBQ	0.169	MDL	20.0	PQL	pg/L	U	B
OCDF	0.512	JBQ	0.212	MDL	20.0	PQL	pg/L	U	B

Sample ID: EB2-071713 Collected: 7/17/2013 3:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.352	JBQ	0.277	MDL	9.97	PQL	pg/L	U	B
1,2,3,4,6,7,8-HPCDF	0.174	JBQ	0.110	MDL	9.97	PQL	pg/L	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	1613B	Matrix:	AQ
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Sample ID: EB2-071713 Collected: 7/17/2013 3:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8,9-HPCDF	0.519	JBQ	0.127	MDL	9.97	PQL	pg/L	U	B
1,2,3,4,7,8-HxCDD	0.373	JBQ	0.221	MDL	9.97	PQL	pg/L	U	B
1,2,3,6,7,8-HXCDF	0.396	JBQ	0.146	MDL	9.97	PQL	pg/L	U	B
1,2,3,7,8,9-HXCDF	0.283	JBQ	0.170	MDL	9.97	PQL	pg/L	U	B
1,2,3,7,8-PECDD	0.542	JQ	0.387	MDL	9.97	PQL	pg/L	J	Z
1,2,3,7,8-PECDF	0.532	JB	0.249	MDL	9.97	PQL	pg/L	U	B
2,3,4,6,7,8-HXCDF	0.359	JBQ	0.140	MDL	9.97	PQL	pg/L	U	B
2,3,4,7,8-PECDF	0.245	JB	0.217	MDL	9.97	PQL	pg/L	U	B
OCDD	0.797	JBQ	0.229	MDL	19.9	PQL	pg/L	U	B
OCDF	0.733	JBQ	0.346	MDL	19.9	PQL	pg/L	U	B

Method Category:	SVOA	Method:	1613B	Matrix:	SO
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Sample ID: SL-512-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	4.39	JB	0.0564	MDL	4.90	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.827	JBQ	0.0208	MDL	4.90	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.111	J	0.0447	MDL	4.90	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.156	JBQ	0.0592	MDL	4.90	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.193	JBQ	0.0364	MDL	4.90	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.473	J	0.0591	MDL	4.90	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.235	JBQ	0.0305	MDL	4.90	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.587	JB	0.0595	MDL	4.90	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.328	JBQ	0.0408	MDL	4.90	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.243	JQ	0.0480	MDL	4.90	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.930	JB	0.0348	MDL	4.90	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.184	JBQ	0.0327	MDL	4.90	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.384	J	0.0392	MDL	4.90	PQL	ng/Kg	J	Z
OCDF	1.57	JB	0.0641	MDL	9.80	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA		
Method:	1613B	Matrix:	SO

Sample ID: SL-513-SA8-SB-0.0-0.5 Collected: 7/17/2013 1:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	3.96	JB	0.0318	MDL	5.36	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.472	J	0.0367	MDL	5.36	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.714	JBQ	0.0920	MDL	5.36	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDF	0.658	JBQ	0.0386	MDL	5.36	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	1.57	J	0.104	MDL	5.36	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.443	JBQ	0.0442	MDL	5.36	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	1.06	JBQ	0.0986	MDL	5.36	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDF	0.423	JB	0.0459	MDL	5.36	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.296	J	0.0762	MDL	5.36	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	3.88	JB	0.0559	MDL	5.36	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HxCDF	0.318	JB	0.0439	MDL	5.36	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.599	J	0.0488	MDL	5.36	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.365	J	0.0936	MDL	1.07	PQL	ng/Kg	J	Z
OCDF	6.36	JB	0.0322	MDL	10.7	PQL	ng/Kg	J	Z

Sample ID: SL-513-SA8-SB-4.0-5.0 Collected: 7/17/2013 2:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.995	JB	0.0432	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.539	JB	0.0197	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0465	J	0.0201	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.0567	JBQ	0.0289	MDL	5.38	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDF	0.0433	JBQ	0.0139	MDL	5.38	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.0702	JQ	0.0311	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.0453	JBQ	0.0150	MDL	5.38	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDD	0.0790	JB	0.0304	MDL	5.38	PQL	ng/Kg	U	B
1,2,3,7,8,9-HxCDF	0.0828	JB	0.0163	MDL	5.38	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0586	JQ	0.0383	MDL	5.38	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.151	JB	0.0273	MDL	5.38	PQL	ng/Kg	U	B
2,3,4,6,7,8-HxCDF	0.0507	JBQ	0.0164	MDL	5.38	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0431	JQ	0.0388	MDL	1.08	PQL	ng/Kg	J	Z
OCDF	3.42	JB	0.0364	MDL	10.8	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	1613B
		Matrix:	SO

Sample ID: SL-513-SA8-SB-8.0-9.0 Collected: 7/17/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0671	JBQ	0.0363	MDL	4.94	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0293	JBQ	0.0122	MDL	4.94	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0564	J	0.0201	MDL	4.94	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0311	JBQ	0.0124	MDL	4.94	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0177	JBQ	0.0130	MDL	4.94	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0460	JB	0.0256	MDL	4.94	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0393	JBQ	0.0207	MDL	4.94	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0189	JBQ	0.0149	MDL	4.94	PQL	ng/Kg	U	B
OCDD	0.192	JBQ	0.0331	MDL	9.87	PQL	ng/Kg	U	B
OCDF	0.126	JB	0.0419	MDL	9.87	PQL	ng/Kg	U	B

Sample ID: SL-514-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	2.65	JB	0.0224	MDL	5.21	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.196	JQ	0.0437	MDL	5.21	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDD	0.164	JB	0.0494	MDL	5.21	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.198	JB	0.0286	MDL	5.21	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.515	JQ	0.0523	MDL	5.21	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.159	JB	0.0273	MDL	5.21	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.557	JB	0.0505	MDL	5.21	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.272	JB	0.0362	MDL	5.21	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.0902	JQ	0.0547	MDL	5.21	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	1.28	JB	0.0415	MDL	5.21	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.141	JB	0.0291	MDL	5.21	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.171	JQ	0.0450	MDL	5.21	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0488	JQ	0.0420	MDL	1.04	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.678	J	0.0684	MDL	1.04	PQL	ng/Kg	J	Z
OCDF	9.21	JB	0.0668	MDL	10.4	PQL	ng/Kg	J	Z

Sample ID: SL-514-SA8-SB-4.0-5.0 Collected: 7/17/2013 9:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.133	JBQ	0.0391	MDL	5.67	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0499	JBQ	0.0147	MDL	5.67	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-514-SA8-SB-4.0-5.0 Collected: 7/17/2013 9:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HXCDF	0.0270	JBQ	0.0204	MDL	5.67	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0534	JQ	0.0368	MDL	5.67	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.0729	JB	0.0353	MDL	5.67	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0427	JBQ	0.0282	MDL	5.67	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0833	JQ	0.0572	MDL	5.67	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0234	JB	0.0200	MDL	5.67	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0591	JQ	0.0313	MDL	5.67	PQL	ng/Kg	J	Z
OCDD	0.864	JB	0.0364	MDL	11.3	PQL	ng/Kg	U	B
OCDF	0.0864	JBQ	0.0566	MDL	11.3	PQL	ng/Kg	U	B

Sample ID: SL-515-SA8-SB-0.0-5.0 Collected: 7/17/2013 10:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	1.64	JB	0.0271	MDL	5.54	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.185	J	0.0307	MDL	5.54	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.142	JB	0.0620	MDL	5.54	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0707	JB	0.0314	MDL	5.54	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.698	J	0.0651	MDL	5.54	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.143	JBQ	0.0303	MDL	5.54	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.637	JB	0.0606	MDL	5.54	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.724	JB	0.0322	MDL	5.54	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDD	0.191	JQ	0.0702	MDL	5.54	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.476	JB	0.0464	MDL	5.54	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.169	JBQ	0.0324	MDL	5.54	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0615	JQ	0.0512	MDL	1.11	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0827	JQ	0.0711	MDL	1.11	PQL	ng/Kg	J	Z
OCDF	3.24	JB	0.0346	MDL	11.1	PQL	ng/Kg	J	Z

Sample ID: SL-515-SA8-SB-14.0-15.0 Collected: 7/17/2013 11:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.106	JBQ	0.0470	MDL	5.71	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0865	JB	0.0177	MDL	5.71	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.136	JQ	0.0286	MDL	5.71	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.194	JBQ	0.0409	MDL	5.71	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	1613B
		Matrix:	SO

Sample ID: SL-515-SA8-SB-14.0-15.0 Collected: 7/17/2013 11:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8-HXCDF	0.165	JBQ	0.0236	MDL	5.71	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.121	JQ	0.0448	MDL	5.71	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.164	JB	0.0245	MDL	5.71	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.121	JB	0.0442	MDL	5.71	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.134	JBQ	0.0353	MDL	5.71	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.333	JBQ	0.0375	MDL	5.71	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	0.151	JQ	0.0353	MDL	5.71	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0811	JQ	0.0546	MDL	1.14	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0687	JQ	0.0505	MDL	1.14	PQL	ng/Kg	J	Z
OCDD	0.675	JB	0.0417	MDL	11.4	PQL	ng/Kg	U	B
OCDF	0.116	JBQ	0.0535	MDL	11.4	PQL	ng/Kg	U	B

Sample ID: SL-515-SA8-SB-4.0-5.0 Collected: 7/17/2013 11:05:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0770	JBQ	0.0402	MDL	5.59	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0819	JBQ	0.0117	MDL	5.59	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0563	JQ	0.0225	MDL	5.59	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.0544	JBQ	0.0280	MDL	5.59	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0599	JBQ	0.0167	MDL	5.59	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0382	J	0.0279	MDL	5.59	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.0461	JB	0.0151	MDL	5.59	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.115	JBQ	0.0279	MDL	5.59	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0755	JBQ	0.0207	MDL	5.59	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.125	JBQ	0.0225	MDL	5.59	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0345	JBQ	0.0164	MDL	5.59	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0722	JQ	0.0246	MDL	5.59	PQL	ng/Kg	J	Z
OCDD	0.658	JB	0.0392	MDL	11.2	PQL	ng/Kg	U	B
OCDF	0.135	JBQ	0.0563	MDL	11.2	PQL	ng/Kg	U	B

Sample ID: SL-515-SA8-SB-9.0-10.0 Collected: 7/17/2013 11:20:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.121	JBQ	0.0522	MDL	5.82	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0679	JB	0.0183	MDL	5.82	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-515-SA8-SB-9.0-10.0 Collected: 7/17/2013 11:20:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,7,8,9-HPCDF	0.0377	JQ	0.0306	MDL	5.82	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.0603	JBQ	0.0369	MDL	5.82	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.118	JB	0.0224	MDL	5.82	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDD	0.105	JQ	0.0373	MDL	5.82	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.118	JBQ	0.0198	MDL	5.82	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.120	JB	0.0400	MDL	5.82	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.145	JBQ	0.0308	MDL	5.82	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.101	JQ	0.0674	MDL	5.82	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.152	JB	0.0418	MDL	5.82	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0653	JBQ	0.0204	MDL	5.82	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.146	J	0.0389	MDL	5.82	PQL	ng/Kg	J	Z
2,3,7,8-TCDD	0.0765	JQ	0.0536	MDL	1.16	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0766	JQ	0.0567	MDL	1.16	PQL	ng/Kg	J	Z
OCDD	0.562	JBQ	0.0464	MDL	11.6	PQL	ng/Kg	U	B
OCDF	0.129	JBQ	0.0603	MDL	11.6	PQL	ng/Kg	U	B

Method Category:	SVOA	
Method:	6850	Matrix: SO

Sample ID: SL-513-SA8-SB-0.0-0.5 Collected: 7/17/2013 1:45:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
PERCHLORATE	3.2	J	2.3	MDL	5.5	PQL	ug/Kg	J	Z

Sample ID: SL-513-SA8-SB-4.0-5.0 Collected: 7/17/2013 2:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
PERCHLORATE	3.6	J	2.3	MDL	5.6	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	8015M	Matrix:	SO
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Sample ID: SL-512-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	3.0	J	2.0	MDL	5.1	PQL	mg/Kg	J	Z

Sample ID: SL-514-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	2.9	J	2.1	MDL	5.3	PQL	mg/Kg	J	Z

Sample ID: SL-514-SA8-SB-4.0-5.0 Collected: 7/17/2013 9:55:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	5.0	J	2.4	MDL	5.9	PQL	mg/Kg	J	Z

Sample ID: SL-514-SA8-SB-7.5-8.5 Collected: 7/17/2013 10:15:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	3.6	J	2.4	MDL	5.9	PQL	mg/Kg	J	Z, Q, Q
EFH (C30-C40)	12		4.8	MDL	12	PQL	mg/Kg	J	Q

Sample ID: SL-515-SA8-SB-14.0-15.0 Collected: 7/17/2013 11:35:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	2.7	J	2.3	MDL	5.8	PQL	mg/Kg	J	Z
EFH (C30-C40)	10	J	4.6	MDL	12	PQL	mg/Kg	J	Z

Method Category:	SVOA	Method:	8081B	Matrix:	AQ
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Sample ID: EB1-071713 Collected: 7/17/2013 3:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ENDRIN	0.017	U	0.0068	MDL	0.017	PQL	ug/L	UJ	E

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 8081B **Matrix:** AQ

Sample ID: EB2-071713 **Collected:** 7/17/2013 3:30:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ENDRIN	0.017	U	0.0067	MDL	0.017	PQL	ug/L	UJ	E

Method Category: SVOA
Method: 8081B **Matrix:** SO

Sample ID: SL-514-SA8-SB-0.0-0.5 **Collected:** 7/17/2013 8:55:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
4,4'-DDT	1.6	J	0.37	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-514-SA8-SB-4.0-5.0 **Collected:** 7/17/2013 9:55:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
4,4'-DDT	0.76	J	0.41	MDL	2.0	PQL	ug/Kg	J	Z

Method Category: SVOA
Method: 8082A **Matrix:** SO

Sample ID: SL-512-SA8-SB-0.0-0.5 **Collected:** 7/17/2013 8:00:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	8.2	J	4.5	MDL	17	PQL	ug/Kg	J	Z, *XIII

Method Category: SVOA
Method: 8151A **Matrix:** AQ

Sample ID: EB1-071713 **Collected:** 7/17/2013 3:00:00 **Analysis Type:** RES **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DALAPON	1.2	U	0.24	MDL	1.2	PQL	ug/L	UJ	C
DINOSEB	0.48	U	0.12	MDL	0.48	PQL	ug/L	UJ	C

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	8151A	Matrix:	AQ
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Sample ID: EB2-071713 Collected: 7/17/2013 3:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
DALAPON	1.2	U	0.24	MDL	1.2	PQL	ug/L	UJ	C
DINOSEB	0.48	U	0.12	MDL	0.48	PQL	ug/L	UJ	C

Method Category:	SVOA	Method:	8151A	Matrix:	SO
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Sample ID: SL-514-SA8-SB-0.0-0.5 Collected: 7/17/2013 8:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,4,5-T	1.8	U	0.86	MDL	1.8	PQL	ug/Kg	UJ	C
2,4-D	38	U	13	MDL	38	PQL	ug/Kg	UJ	C
2,4-DB	18	U	6.5	MDL	18	PQL	ug/Kg	UJ	C
DALAPON	94	U	46	MDL	94	PQL	ug/Kg	UJ	C
MCPA	2600	U	800	MDL	2600	PQL	ug/Kg	UJ	C
MCPD	2600	U	790	MDL	2600	PQL	ug/Kg	UJ	C

Sample ID: SL-514-SA8-SB-4.0-5.0 Collected: 7/17/2013 9:55:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,4,5-T	2.0	U	0.97	MDL	2.0	PQL	ug/Kg	UJ	C
2,4-D	43	U	14	MDL	43	PQL	ug/Kg	UJ	C
2,4-DB	20	U	7.3	MDL	20	PQL	ug/Kg	UJ	C
DALAPON	110	U	52	MDL	110	PQL	ug/Kg	UJ	C
MCPA	3000	U	900	MDL	3000	PQL	ug/Kg	UJ	C
MCPD	3000	U	890	MDL	3000	PQL	ug/Kg	UJ	C

Sample ID: SL-514-SA8-SB-7.5-8.5 Collected: 7/17/2013 10:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,4,5-T	2.0	U	0.97	MDL	2.0	PQL	ug/Kg	UJ	C
2,4-D	43	U	14	MDL	43	PQL	ug/Kg	UJ	C
2,4-DB	20	U	7.3	MDL	20	PQL	ug/Kg	UJ	C
DALAPON	110	U	52	MDL	110	PQL	ug/Kg	UJ	C
MCPA	3000	U	900	MDL	3000	PQL	ug/Kg	UJ	C
MCPD	3000	U	890	MDL	3000	PQL	ug/Kg	UJ	C

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	8151A	Matrix: SO

Method Category:	SVOA	
Method:	8270D	Matrix: AQ

Sample ID: EB1-071713 **Collected:** 7/17/2013 3:00:00 **Analysis Type:** RES-ACID **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZOIC ACID	17	U	7	MDL	17	PQL	ug/L	UJ	E
HEXACHLOROCYCLOPENTADIENE	17	U	6	MDL	17	PQL	ug/L	UJ	C

Method Category:	SVOA	
Method:	8270D	Matrix: SO

Sample ID: SL-513-SA8-SB-0.0-0.5 **Collected:** 7/17/2013 1:45:00 **Analysis Type:** RES-ACID **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,4-DINITROPHENOL	1100	U	320	MDL	1100	PQL	ug/Kg	UJ	C
BENZO(A)ANTHRACENE	5	J	4	MDL	18	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	9	J	4	MDL	18	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	8	J	4	MDL	18	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	5	J	4	MDL	18	PQL	ug/Kg	J	Z
BIPHENYL	25	J	18	MDL	36	PQL	ug/Kg	J	Z
HEXACHLOROCYCLOPENTADIENE	540	U	180	MDL	540	PQL	ug/Kg	UJ	C
NAPHTHALENE	4	J	4	MDL	18	PQL	ug/Kg	J	Z
PHENANTHRENE	15	J	4	MDL	18	PQL	ug/Kg	J	Z

Sample ID: SL-513-SA8-SB-4.0-5.0 **Collected:** 7/17/2013 2:00:00 **Analysis Type:** RES-ACID **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,4-DICHLOROPHENOL	37	U	18	MDL	37	PQL	ug/Kg	UJ	C
BENZO(B)FLUORANTHENE	7	J	4	MDL	19	PQL	ug/Kg	J	Z
HEXACHLOROCYCLOPENTADIENE	550	U	180	MDL	550	PQL	ug/Kg	UJ	C
PYRENE	4	J	4	MDL	19	PQL	ug/Kg	J	Z

Sample ID: SL-513-SA8-SB-8.0-9.0 **Collected:** 7/17/2013 2:15:00 **Analysis Type:** RES-ACID **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,4-DICHLOROPHENOL	34	U	17	MDL	34	PQL	ug/Kg	UJ	C

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069
 EDD Filename: PrepPH069_v1

Laboratory: LL
 eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 8270D **Matrix:** SO

Sample ID: SL-513-SA8-SB-8.0-9.0 **Collected:** 7/17/2013 2:15:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXACHLOROCYCLOPENTADIENE	510	U	170	MDL	510	PQL	ug/Kg	UJ	C

Method Category: SVOA
Method: 8270D SIM **Matrix:** AQ

Sample ID: EB1-071713 **Collected:** 7/17/2013 3:00:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.014	J	0.011	MDL	0.055	PQL	ug/L	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	0.19	J	0.055	MDL	1.1	PQL	ug/L	U	B
Diethylphthalate	0.24	J	0.055	MDL	1.1	PQL	ug/L	J	Z
Di-n-butylphthalate	0.24	J	0.055	MDL	1.1	PQL	ug/L	J	Z
NAPHTHALENE	0.046	J	0.033	MDL	0.055	PQL	ug/L	J	Z

Sample ID: EB2-071713 **Collected:** 7/17/2013 3:30:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2-METHYLNAPHTHALENE	0.014	J	0.010	MDL	0.051	PQL	ug/L	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	0.14	J	0.051	MDL	1.0	PQL	ug/L	U	B
Diethylphthalate	0.22	J	0.051	MDL	1.0	PQL	ug/L	J	Z
Di-n-butylphthalate	0.14	J	0.051	MDL	1.0	PQL	ug/L	J	Z
NAPHTHALENE	0.045	J	0.031	MDL	0.051	PQL	ug/L	J	Z

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-512-SA8-SB-0.0-0.5 **Collected:** 7/17/2013 8:00:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	1.1	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	11	J	6.0	MDL	18	PQL	ug/Kg	J	Z
Di-n-butylphthalate	6.9	J	6.0	MDL	18	PQL	ug/Kg	J	Z
FLUORANTHENE	1.4	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z
NAPHTHALENE	0.78	J	0.67	MDL	1.7	PQL	ug/Kg	U	F

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	Method:	8270D SIM	Matrix:	SO
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Sample ID: SL-512-SA8-SB-0.0-0.5 **Collected:** 7/17/2013 8:00:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
PHENANTHRENE	0.82	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z
PYRENE	1.1	J	0.67	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-513-SA8-SB-0.0-0.5 **Collected:** 7/17/2013 1:45:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	13	J	7.2	MDL	18	PQL	ug/Kg	J	Z

Sample ID: SL-513-SA8-SB-4.0-5.0 **Collected:** 7/17/2013 2:00:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	11	J	3.7	MDL	18	PQL	ug/Kg	J	Z

Sample ID: SL-514-SA8-SB-0.0-0.5 **Collected:** 7/17/2013 8:55:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(A)ANTHRACENE	0.90	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	0.92	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BENZO(G,H,I)PERYLENE	0.92	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	11	J	6.2	MDL	19	PQL	ug/Kg	J	Z
NAPHTHALENE	0.72	J	0.69	MDL	1.7	PQL	ug/Kg	U	F
PHENANTHRENE	1.5	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-514-SA8-SB-7.5-8.5 **Collected:** 7/17/2013 10:15:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	0.91	J	0.78	MDL	1.9	PQL	ug/Kg	J	Z

Sample ID: SL-515-SA8-SB-0.0-0.5 **Collected:** 7/17/2013 10:50:00 **Analysis Type:** RES-BASE/NEUTRAL **Dilution:** 10

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	5.1	J	3.7	MDL	19	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	VOA	
Method:	8260B	Matrix: AQ

Sample ID: EB1-071713 Collected: 7/17/2013 3:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ACETONE	7	J	6	MDL	20	PQL	ug/L	J	Z
Chlorotrifluoroethylene	5	U	2	MDL	5	PQL	ug/L	UJ	L

Sample ID: TB-071713 Collected: 7/17/2013 8:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
Chlorotrifluoroethylene	5	U	2	MDL	5	PQL	ug/L	UJ	L

Method Category:	VOA	
Method:	8260B	Matrix: SO

Sample ID: SL-513-SA8-SB-8.0-9.0 Collected: 7/17/2013 2:15:00 Analysis Type: RES Dilution: 0.97

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
VINYL ACETATE	10	U	2	MDL	10	PQL	ug/Kg	UJ	C

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
*XII	RPD Between Columns
A	ICP Serial Dilution
B	Calibration Blank Contamination
B	Method Blank Contamination
C	Continuing Calibration Verification Percent Difference Lower Estimation
C	Initial Calibration Verification Percent Difference Lower Estimation
E	Laboratory Control Precision
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Equipment Blank Contamination
F	Field Blank Contamination
L	Laboratory Control Precision
L	Laboratory Control Spike Lower Estimation
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Estimation
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

9/21/2013 12:18:47 PM

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH069

Method Blank Outlier Report

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2000B372114	7/19/2013 9:14:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF OCDD OCDF	0.373 pg/L 0.413 pg/L 0.325 pg/L 0.368 pg/L 0.527 pg/L 0.364 pg/L 0.476 pg/L 0.327 pg/L 0.586 pg/L 0.465 pg/L 0.238 pg/L 0.511 pg/L 0.838 pg/L	EB1-071713 EB2-071713

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB1-071713(RES)	1,2,3,4,6,7,8-HPCDD	0.413 pg/L	0.413U pg/L
EB1-071713(RES)	1,2,3,4,6,7,8-HPCDF	0.238 pg/L	0.238U pg/L
EB1-071713(RES)	1,2,3,4,7,8,9-HPCDF	0.191 pg/L	0.191U pg/L
EB1-071713(RES)	1,2,3,4,7,8-HxCDD	0.161 pg/L	0.161U pg/L
EB1-071713(RES)	1,2,3,4,7,8-HXCDF	0.222 pg/L	0.222U pg/L
EB1-071713(RES)	1,2,3,6,7,8-HXCDF	0.191 pg/L	0.191U pg/L
EB1-071713(RES)	1,2,3,7,8,9-HXCDF	0.186 pg/L	0.186U pg/L
EB1-071713(RES)	1,2,3,7,8-PECDF	0.406 pg/L	0.406U pg/L
EB1-071713(RES)	2,3,4,6,7,8-HXCDF	0.224 pg/L	0.224U pg/L
EB1-071713(RES)	2,3,4,7,8-PECDF	0.132 pg/L	0.132U pg/L
EB1-071713(RES)	OCDD	0.814 pg/L	0.814U pg/L
EB1-071713(RES)	OCDF	0.512 pg/L	0.512U pg/L
EB2-071713(RES)	1,2,3,4,6,7,8-HPCDD	0.352 pg/L	0.352U pg/L
EB2-071713(RES)	1,2,3,4,6,7,8-HPCDF	0.174 pg/L	0.174U pg/L
EB2-071713(RES)	1,2,3,4,7,8,9-HPCDF	0.519 pg/L	0.519U pg/L
EB2-071713(RES)	1,2,3,4,7,8-HxCDD	0.373 pg/L	0.373U pg/L
EB2-071713(RES)	1,2,3,6,7,8-HXCDF	0.396 pg/L	0.396U pg/L
EB2-071713(RES)	1,2,3,7,8,9-HXCDF	0.283 pg/L	0.283U pg/L
EB2-071713(RES)	1,2,3,7,8-PECDF	0.532 pg/L	0.532U pg/L
EB2-071713(RES)	2,3,4,6,7,8-HXCDF	0.359 pg/L	0.359U pg/L
EB2-071713(RES)	2,3,4,7,8-PECDF	0.245 pg/L	0.245U pg/L
EB2-071713(RES)	OCDD	0.797 pg/L	0.797U pg/L
EB2-071713(RES)	OCDF	0.733 pg/L	0.733U pg/L

Method Blank Outlier Report

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2060B372237	7/26/2013 10:37:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDD 1,2,3,7,8,9-HxCDF 1,2,3,7,8-PECDF 2,3,4,6,7,8-HxCDF OCDD OCDF	0.0714 ng/Kg 0.0256 ng/Kg 0.0432 ng/Kg 0.0229 ng/Kg 0.0231 ng/Kg 0.0358 ng/Kg 0.0413 ng/Kg 0.0307 ng/Kg 0.0401 ng/Kg 0.230 ng/Kg 0.159 ng/Kg	SL-512-SA8-SB-0.0-0.5 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-14.0-15.0 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-512-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.156 ng/Kg	0.156U ng/Kg
SL-512-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HxCDF	0.184 ng/Kg	0.184U ng/Kg
SL-513-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDD	0.0567 ng/Kg	0.0567U ng/Kg
SL-513-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDF	0.0433 ng/Kg	0.0433U ng/Kg
SL-513-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HxCDF	0.0453 ng/Kg	0.0453U ng/Kg
SL-513-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDD	0.0790 ng/Kg	0.0790U ng/Kg
SL-513-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDF	0.0828 ng/Kg	0.0828U ng/Kg
SL-513-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.151 ng/Kg	0.151U ng/Kg
SL-513-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HxCDF	0.0507 ng/Kg	0.0507U ng/Kg
SL-513-SA8-SB-8.0-9.0(RES)	1,2,3,4,6,7,8-HPCDD	0.0671 ng/Kg	0.0671U ng/Kg
SL-513-SA8-SB-8.0-9.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0293 ng/Kg	0.0293U ng/Kg
SL-513-SA8-SB-8.0-9.0(RES)	1,2,3,4,7,8-HxCDF	0.0311 ng/Kg	0.0311U ng/Kg
SL-513-SA8-SB-8.0-9.0(RES)	1,2,3,6,7,8-HxCDF	0.0177 ng/Kg	0.0177U ng/Kg
SL-513-SA8-SB-8.0-9.0(RES)	1,2,3,7,8,9-HxCDD	0.0460 ng/Kg	0.0460U ng/Kg
SL-513-SA8-SB-8.0-9.0(RES)	1,2,3,7,8,9-HxCDF	0.0393 ng/Kg	0.0393U ng/Kg
SL-513-SA8-SB-8.0-9.0(RES)	2,3,4,6,7,8-HxCDF	0.0189 ng/Kg	0.0189U ng/Kg
SL-513-SA8-SB-8.0-9.0(RES)	OCDD	0.192 ng/Kg	0.192U ng/Kg
SL-513-SA8-SB-8.0-9.0(RES)	OCDF	0.126 ng/Kg	0.126U ng/Kg
SL-514-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.164 ng/Kg	0.164U ng/Kg
SL-514-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HxCDF	0.141 ng/Kg	0.141U ng/Kg
SL-514-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDD	0.133 ng/Kg	0.133U ng/Kg
SL-514-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0499 ng/Kg	0.0499U ng/Kg
SL-514-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDF	0.0270 ng/Kg	0.0270U ng/Kg
SL-514-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDD	0.0729 ng/Kg	0.0729U ng/Kg
SL-514-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDF	0.0427 ng/Kg	0.0427U ng/Kg
SL-514-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HxCDF	0.0234 ng/Kg	0.0234U ng/Kg
SL-514-SA8-SB-4.0-5.0(RES)	OCDD	0.864 ng/Kg	0.864U ng/Kg
SL-514-SA8-SB-4.0-5.0(RES)	OCDF	0.0864 ng/Kg	0.0864U ng/Kg
SL-515-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDD	0.142 ng/Kg	0.142U ng/Kg
SL-515-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HxCDF	0.0707 ng/Kg	0.0707U ng/Kg
SL-515-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HxCDF	0.169 ng/Kg	0.169U ng/Kg
SL-515-SA8-SB-14.0-15.0(RES)	1,2,3,4,6,7,8-HPCDD	0.106 ng/Kg	0.106U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-515-SA8-SB-14.0-15.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0865 ng/Kg	0.0865U ng/Kg
SL-515-SA8-SB-14.0-15.0(RES)	1,2,3,4,7,8-HxCDD	0.194 ng/Kg	0.194U ng/Kg
SL-515-SA8-SB-14.0-15.0(RES)	1,2,3,7,8,9-HxCDD	0.121 ng/Kg	0.121U ng/Kg
SL-515-SA8-SB-14.0-15.0(RES)	1,2,3,7,8,9-HxCDF	0.134 ng/Kg	0.134U ng/Kg
SL-515-SA8-SB-14.0-15.0(RES)	OCDD	0.675 ng/Kg	0.675U ng/Kg
SL-515-SA8-SB-14.0-15.0(RES)	OCDF	0.116 ng/Kg	0.116U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDD	0.0770 ng/Kg	0.0770U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0819 ng/Kg	0.0819U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDD	0.0544 ng/Kg	0.0544U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HxCDF	0.0599 ng/Kg	0.0599U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HxCDF	0.0461 ng/Kg	0.0461U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDD	0.115 ng/Kg	0.115U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HxCDF	0.0755 ng/Kg	0.0755U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.125 ng/Kg	0.125U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HxCDF	0.0345 ng/Kg	0.0345U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	OCDD	0.658 ng/Kg	0.658U ng/Kg
SL-515-SA8-SB-4.0-5.0(RES)	OCDF	0.135 ng/Kg	0.135U ng/Kg
SL-515-SA8-SB-9.0-10.0(RES)	1,2,3,4,6,7,8-HPCDD	0.121 ng/Kg	0.121U ng/Kg
SL-515-SA8-SB-9.0-10.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0679 ng/Kg	0.0679U ng/Kg
SL-515-SA8-SB-9.0-10.0(RES)	1,2,3,4,7,8-HxCDD	0.0603 ng/Kg	0.0603U ng/Kg
SL-515-SA8-SB-9.0-10.0(RES)	1,2,3,7,8,9-HxCDD	0.120 ng/Kg	0.120U ng/Kg
SL-515-SA8-SB-9.0-10.0(RES)	1,2,3,7,8,9-HxCDF	0.145 ng/Kg	0.145U ng/Kg
SL-515-SA8-SB-9.0-10.0(RES)	1,2,3,7,8-PECDF	0.152 ng/Kg	0.152U ng/Kg
SL-515-SA8-SB-9.0-10.0(RES)	2,3,4,6,7,8-HxCDF	0.0653 ng/Kg	0.0653U ng/Kg
SL-515-SA8-SB-9.0-10.0(RES)	OCDD	0.562 ng/Kg	0.562U ng/Kg
SL-515-SA8-SB-9.0-10.0(RES)	OCDF	0.129 ng/Kg	0.129U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P20337AB220845	7/26/2013 8:45:00 AM	TIN	1.73 mg/Kg	SL-512-SA8-SB-0.0-0.5 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-14.0-15.0 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0

Method Blank Outlier Report

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-512-SA8-SB-0.0-0.5(RES)	TIN	2.98 mg/Kg	2.98U mg/Kg
SL-513-SA8-SB-0.0-0.5(RES)	TIN	3.28 mg/Kg	3.28U mg/Kg
SL-513-SA8-SB-4.0-5.0(RES)	TIN	2.78 mg/Kg	2.78U mg/Kg
SL-513-SA8-SB-8.0-9.0(RES)	TIN	3.13 mg/Kg	3.13U mg/Kg
SL-514-SA8-SB-0.0-0.5(RES)	TIN	3.33 mg/Kg	3.33U mg/Kg
SL-514-SA8-SB-4.0-5.0(RES)	TIN	3.05 mg/Kg	3.05U mg/Kg
SL-514-SA8-SB-7.5-8.5(RES)	TIN	3.00 mg/Kg	3.00U mg/Kg
SL-515-SA8-SB-0.0-0.5(RES)	TIN	3.43 mg/Kg	3.43U mg/Kg
SL-515-SA8-SB-14.0-15.0(RES)	TIN	3.34 mg/Kg	3.34U mg/Kg
SL-515-SA8-SB-4.0-5.0(RES)	TIN	3.38 mg/Kg	3.38U mg/Kg
SL-515-SA8-SB-9.0-10.0(RES)	TIN	3.76 mg/Kg	3.76U mg/Kg

Method: 8270D SIM
Matrix: AQ

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
PLKWB20B262058	7/29/2013 8:58:00 PM	BIS(2-ETHYLHEXYL)PHTHALATE	0.051 ug/L	EB1-071713 EB2-071713

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
EB1-071713(RES)	BIS(2-ETHYLHEXYL)PHTHALATE	0.19 ug/L	1.1U ug/L
EB2-071713(RES)	BIS(2-ETHYLHEXYL)PHTHALATE	0.14 ug/L	1.0U ug/L

Field Blank Outlier Report

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(REA2)	4/11/2013 3:00:00 PM	COPPER MOLYBDENUM	0.0036 mg/L 0.0036 mg/L	SL-512-SA8-SB-0.0-0.5 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-14.0-15.0 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-512-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.340 mg/Kg	0.340U mg/Kg
SL-513-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.861 mg/Kg	0.861U mg/Kg

Method: 8270D SIM
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(RES)	4/11/2013 3:00:00 PM	1-METHYLNAPHTHALENE 2-METHYLNAPHTHALENE BIS(2-ETHYLHEXYL)PHTHALATE Diethylphthalate Di-n-butylphthalate NAPHTHALENE	0.019 ug/L 0.024 ug/L 0.082 ug/L 0.18 ug/L 0.17 ug/L 0.17 ug/L	SL-512-SA8-SB-0.0-0.5 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-14.0-15.0 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-512-SA8-SB-0.0-0.5(RES)	NAPHTHALENE	0.78 ug/Kg	1.7U ug/Kg
SL-514-SA8-SB-0.0-0.5(RES)	NAPHTHALENE	0.72 ug/Kg	1.7U ug/Kg

Equipment Rinsate Blank Outlier Report

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Equipment Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
EB1-071713(REA3)	7/17/2013 3:00:00 PM	ANTIMONY	0.0069 mg/L	SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-14.0-15.0 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-513-SA8-SB-0.0-0.5(RES)	ANTIMONY	1.20 mg/Kg	1.20U mg/Kg

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8015M
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-514-SA8-SB-7.5-8.5MS	EFH (C15-C20)	162	138	49.00-123.00	-	EFH (C15-C20)	J (all detects)
SL-514-SA8-SB-7.5-8.5MSD	EFH (C21-C30)	172	-	49.00-123.00	28 (20.00)	EFH (C21-C30)	
(SL-514-SA8-SB-7.5-8.5)	EFH (C30-C40)	131	-	49.00-123.00	-	EFH (C30-C40)	

Method: 8151A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-514-SA8-SB-0.0-0.5MSD (SL-514-SA8-SB-0.0-0.5)	DALAPON	-	-	10.00-125.00	65 (50.00)	DALAPON	J(all detects)

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-514-SA8-SB-7.5-8.5MS (TOT)	ALUMINUM	3157	2639	75.00-125.00	-	ALUMINUM	J(all detects) Al, Fe, Mn, Mg, P, and Ti No Qual, >4X
SL-514-SA8-SB-7.5-8.5MSD (TOT)	IRON	1337	667	75.00-125.00	-	IRON	
(SL-512-SA8-SB-0.0-0.5)	MAGNESIUM	341	251	75.00-125.00	-	MAGNESIUM	
SL-513-SA8-SB-0.0-0.5	MANGANESE	266	360	75.00-125.00	-	MANGANESE	
SL-513-SA8-SB-4.0-5.0	PHOSPHORUS	161	-	75.00-125.00	-	PHOSPHORUS	
SL-513-SA8-SB-8.0-9.0	POTASSIUM	177	162	75.00-125.00	-	POTASSIUM	
SL-514-SA8-SB-0.0-0.5	TITANIUM	405	367	75.00-125.00	-	TITANIUM	
SL-514-SA8-SB-4.0-5.0							
SL-514-SA8-SB-7.5-8.5							
SL-515-SA8-SB-0.0-0.5							
SL-515-SA8-SB-14.0-15.0							
SL-515-SA8-SB-4.0-5.0							
SL-515-SA8-SB-9.0-10.0)							
SL-514-SA8-SB-7.5-8.5MS (TOT)	CALCIUM	54	-325	75.00-125.00	-	CALCIUM	
SL-514-SA8-SB-7.5-8.5MSD (TOT)							
(SL-512-SA8-SB-0.0-0.5)							
SL-513-SA8-SB-0.0-0.5							
SL-513-SA8-SB-4.0-5.0							
SL-513-SA8-SB-8.0-9.0							
SL-514-SA8-SB-0.0-0.5							
SL-514-SA8-SB-4.0-5.0							
SL-514-SA8-SB-7.5-8.5							
SL-515-SA8-SB-0.0-0.5							
SL-515-SA8-SB-14.0-15.0							
SL-515-SA8-SB-4.0-5.0							
SL-515-SA8-SB-9.0-10.0)							

Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-514-SA8-SB-7.5-8.5MS (TOT) SL-514-SA8-SB-7.5-8.5MSD (TOT) (SL-512-SA8-SB-0.0-0.5 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-14.0-15.0 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0)	ANTIMONY	74	73	75.00-125.00	-	ANTIMONY	J(all detects) UJ(all non-detects)

Method: 6020A
Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-514-SA8-SB-7.5-8.5MS (TOT) SL-514-SA8-SB-7.5-8.5MSD (TOT) (SL-512-SA8-SB-0.0-0.5 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-14.0-15.0 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0)	STRONTIUM	19	-174	75.00-125.00	-	STRONTIUM	No Qual, >4X

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8081B
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P32004AY241931A (EB1-071713 EB2-071713)	ENDRIN	-	-	43.00-139.00	35 (30.00)	ENDRIN	J (all detects) UJ (all non-detects)

Method: 8151A
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P32039AY241147A (EB1-071713 EB2-071713)	DINOSEB	-	184	16.00-163.00	-	DINOSEB	J(all detects)

Method: 8270D
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P5WCLCSQ260439 P5WCLCSY260506 (EB1-071713)	4-NITROPHENOL	75	80	13.00-71.00	-	4-NITROPHENOL	J(all detects)
P5WCLCSY260506 (EB1-071713)	BENZOIC ACID	-	-	10.00-69.00	54 (30.00)	BENZOIC ACID	J(all detects) UJ(all non-detects)

Method: 8260B
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
LCSY77Q211110A LCSY77Y211131A (EB1-071713 TB-071713)	Chlorotrifluoroethylene	44	43	47.00-120.00	-	Chlorotrifluoroethylene	J(all detects) UJ(all non-detects)

Method: 8260B
Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
LCSB68Q212141A LCSB68Y212203A (SL-513-SA8-SB-8.0-9.0)	VINYL ACETATE	135	125	29.00-111.00	-	VINYL ACETATE	J(all detects)

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PrepPH069_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C

Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-514-SA8-SB-7.5-8.5DUP (TOT) (SL-512-SA8-SB-0.0-0.5 SL -513-SA8-SB-0.0-0.5 SL -513-SA8-SB-4.0-5.0 SL -513-SA8-SB-8.0-9.0 SL -514-SA8-SB-0.0-0.5 SL -514-SA8-SB-4.0-5.0 SL -514-SA8-SB-7.5-8.5 SL -515-SA8-SB-0.0-0.5 SL -515-SA8-SB-14.0-15.0 SL -515-SA8-SB-4.0-5.0 SL -515-SA8-SB-9.0-10.0)	MOLYBDENUM	28	20.00	No Qual, OK by Difference

Reporting Limit Outliers

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-071713	1,2,3,4,6,7,8-HPCDD	JBQ	0.413	9.98	PQL	pg/L	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.238	9.98	PQL	pg/L	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.191	9.98	PQL	pg/L	
	1,2,3,4,7,8-HxCDD	JBQ	0.161	9.98	PQL	pg/L	
	1,2,3,4,7,8-HXCDF	JBQ	0.222	9.98	PQL	pg/L	
	1,2,3,6,7,8-HXCDF	JBQ	0.191	9.98	PQL	pg/L	
	1,2,3,7,8,9-HXCDF	JB	0.186	9.98	PQL	pg/L	
	1,2,3,7,8-PECDF	JB	0.406	9.98	PQL	pg/L	
	2,3,4,6,7,8-HXCDF	JB	0.224	9.98	PQL	pg/L	
	2,3,4,7,8-PECDF	JBQ	0.132	9.98	PQL	pg/L	
	OCDD	JBQ	0.814	20.0	PQL	pg/L	
	OCDF	JBQ	0.512	20.0	PQL	pg/L	
EB2-071713	1,2,3,4,6,7,8-HPCDD	JBQ	0.352	9.97	PQL	pg/L	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.174	9.97	PQL	pg/L	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.519	9.97	PQL	pg/L	
	1,2,3,4,7,8-HxCDD	JBQ	0.373	9.97	PQL	pg/L	
	1,2,3,6,7,8-HXCDF	JBQ	0.396	9.97	PQL	pg/L	
	1,2,3,7,8,9-HXCDF	JBQ	0.283	9.97	PQL	pg/L	
	1,2,3,7,8-PECDD	JQ	0.542	9.97	PQL	pg/L	
	1,2,3,7,8-PECDF	JB	0.532	9.97	PQL	pg/L	
	2,3,4,6,7,8-HXCDF	JBQ	0.359	9.97	PQL	pg/L	
	2,3,4,7,8-PECDF	JB	0.245	9.97	PQL	pg/L	
	OCDD	JBQ	0.797	19.9	PQL	pg/L	
	OCDF	JBQ	0.733	19.9	PQL	pg/L	

Method: 6010C

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-071713	ANTIMONY	J	0.0069	0.0400	PQL	mg/L	J (all detects)
EB2-071713	ANTIMONY	J	0.0054	0.0400	PQL	mg/L	J (all detects)

Method: 8260B

Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-071713	ACETONE	J	7	20	PQL	ug/L	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8270D SIM
Matrix: AQ

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
EB1-071713	2-METHYLNAPHTHALENE	J	0.014	0.055	PQL	ug/L	J (all detects)
	BIS(2-ETHYLHEXYL)PHTHALATE	J	0.19	1.1	PQL	ug/L	
	Diethylphthalate	J	0.24	1.1	PQL	ug/L	
	Di-n-butylphthalate	J	0.24	1.1	PQL	ug/L	
	NAPHTHALENE	J	0.046	0.055	PQL	ug/L	
EB2-071713	2-METHYLNAPHTHALENE	J	0.014	0.051	PQL	ug/L	J (all detects)
	BIS(2-ETHYLHEXYL)PHTHALATE	J	0.14	1.0	PQL	ug/L	
	Diethylphthalate	J	0.22	1.0	PQL	ug/L	
	Di-n-butylphthalate	J	0.14	1.0	PQL	ug/L	
	NAPHTHALENE	J	0.045	0.051	PQL	ug/L	

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-512-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	4.39	4.90	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.827	4.90	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	J	0.111	4.90	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.156	4.90	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.193	4.90	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	J	0.473	4.90	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.235	4.90	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.587	4.90	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JBQ	0.328	4.90	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.243	4.90	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.930	4.90	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.184	4.90	PQL	ng/Kg	
	2,3,4,7,8-PECDF	J	0.384	4.90	PQL	ng/Kg	
	OCDF	JB	1.57	9.80	PQL	ng/Kg	
	SL-513-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	3.96	5.36	PQL	
1,2,3,4,7,8,9-HPCDF		J	0.472	5.36	PQL	ng/Kg	
1,2,3,4,7,8-HxCDD		JBQ	0.714	5.36	PQL	ng/Kg	
1,2,3,4,7,8-HxCDF		JBQ	0.658	5.36	PQL	ng/Kg	
1,2,3,6,7,8-HxCDD		J	1.57	5.36	PQL	ng/Kg	
1,2,3,6,7,8-HxCDF		JBQ	0.443	5.36	PQL	ng/Kg	
1,2,3,7,8,9-HxCDD		JBQ	1.06	5.36	PQL	ng/Kg	
1,2,3,7,8,9-HxCDF		JB	0.423	5.36	PQL	ng/Kg	
1,2,3,7,8-PECDD		J	0.296	5.36	PQL	ng/Kg	
1,2,3,7,8-PECDF		JB	3.88	5.36	PQL	ng/Kg	
2,3,4,6,7,8-HxCDF		JB	0.318	5.36	PQL	ng/Kg	
2,3,4,7,8-PECDF		J	0.599	5.36	PQL	ng/Kg	
2,3,7,8-TCDF		J	0.365	1.07	PQL	ng/Kg	
OCDF		JB	6.36	10.7	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-513-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	0.995	5.38	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.539	5.38	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	J	0.0465	5.38	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0567	5.38	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.0433	5.38	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JQ	0.0702	5.38	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JBQ	0.0453	5.38	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.0790	5.38	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.0828	5.38	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0586	5.38	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.151	5.38	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JBQ	0.0507	5.38	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0431	1.08	PQL	ng/Kg	
	OCDF	JB	3.42	10.8	PQL	ng/Kg	
	SL-513-SA8-SB-8.0-9.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.0671	4.94	PQL	
1,2,3,4,6,7,8-HPCDF		JBQ	0.0293	4.94	PQL	ng/Kg	
1,2,3,4,7,8,9-HPCDF		J	0.0564	4.94	PQL	ng/Kg	
1,2,3,4,7,8-HxCDF		JBQ	0.0311	4.94	PQL	ng/Kg	
1,2,3,6,7,8-HxCDF		JBQ	0.0177	4.94	PQL	ng/Kg	
1,2,3,7,8,9-HxCDD		JB	0.0460	4.94	PQL	ng/Kg	
1,2,3,7,8,9-HxCDF		JBQ	0.0393	4.94	PQL	ng/Kg	
2,3,4,6,7,8-HxCDF		JBQ	0.0189	4.94	PQL	ng/Kg	
OCDD		JBQ	0.192	9.87	PQL	ng/Kg	
OCDF		JB	0.126	9.87	PQL	ng/Kg	
SL-514-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	2.65	5.21	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JQ	0.196	5.21	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.164	5.21	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.198	5.21	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JQ	0.515	5.21	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDF	JB	0.159	5.21	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.557	5.21	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JB	0.272	5.21	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0902	5.21	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	1.28	5.21	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.141	5.21	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.171	5.21	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0488	1.04	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.678	1.04	PQL	ng/Kg	
	OCDF	JB	9.21	10.4	PQL	ng/Kg	
SL-514-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.133	5.67	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0499	5.67	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JBQ	0.0270	5.67	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JQ	0.0534	5.67	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.0729	5.67	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDF	JBQ	0.0427	5.67	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0833	5.67	PQL	ng/Kg	
	2,3,4,6,7,8-HxCDF	JB	0.0234	5.67	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0591	5.67	PQL	ng/Kg	
	OCDD	JB	0.864	11.3	PQL	ng/Kg	
	OCDF	JBQ	0.0864	11.3	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-515-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	1.64	5.54	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	J	0.185	5.54	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JB	0.142	5.54	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.0707	5.54	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	0.698	5.54	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.143	5.54	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.637	5.54	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.724	5.54	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.191	5.54	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.476	5.54	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.169	5.54	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0615	1.11	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0827	1.11	PQL	ng/Kg	
	OCDF	JB	3.24	11.1	PQL	ng/Kg	
SL-515-SA8-SB-14.0-15.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.106	5.71	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.0865	5.71	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JQ	0.136	5.71	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.194	5.71	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.165	5.71	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.121	5.71	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.164	5.71	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.121	5.71	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.134	5.71	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.333	5.71	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.151	5.71	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0811	1.14	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0687	1.14	PQL	ng/Kg	
	OCDD	JB	0.675	11.4	PQL	ng/Kg	
OCDF	JBQ	0.116	11.4	PQL	ng/Kg		
SL-515-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.0770	5.59	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0819	5.59	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JQ	0.0563	5.59	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0544	5.59	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0599	5.59	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	J	0.0382	5.59	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.0461	5.59	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.115	5.59	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.0755	5.59	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.125	5.59	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0345	5.59	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JQ	0.0722	5.59	PQL	ng/Kg	
	OCDD	JB	0.658	11.2	PQL	ng/Kg	
	OCDF	JBQ	0.135	11.2	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-515-SA8-SB-9.0-10.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.121	5.82	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.0679	5.82	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JQ	0.0377	5.82	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JBQ	0.0603	5.82	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDF	JB	0.118	5.82	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JQ	0.105	5.82	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.118	5.82	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.120	5.82	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.145	5.82	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.101	5.82	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.152	5.82	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0653	5.82	PQL	ng/Kg	
	2,3,4,7,8-PECDF	J	0.146	5.82	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0765	1.16	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0766	1.16	PQL	ng/Kg	
	OCDD	JBQ	0.562	11.6	PQL	ng/Kg	
OCDF	JBQ	0.129	11.6	PQL	ng/Kg		

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-512-SA8-SB-0.0-0.5	BERYLLIUM	J	0.593	0.976	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.235	0.976	PQL	mg/Kg	
	MOLYBDENUM	J	0.340	1.95	PQL	mg/Kg	
	SODIUM	J	77.2	97.6	PQL	mg/Kg	
SL-513-SA8-SB-0.0-0.5	TIN	J	2.98	9.76	PQL	mg/Kg	J (all detects)
	ANTIMONY	J	1.20	4.28	PQL	mg/Kg	
	BERYLLIUM	J	0.659	1.07	PQL	mg/Kg	
	BORON	J	2.19	10.7	PQL	mg/Kg	
	CADMIUM	J	0.104	1.07	PQL	mg/Kg	
	MOLYBDENUM	J	0.861	2.14	PQL	mg/Kg	
SL-513-SA8-SB-4.0-5.0	TIN	J	3.28	10.7	PQL	mg/Kg	J (all detects)
	Zirconium	J	2.54	5.35	PQL	mg/Kg	
	BERYLLIUM	J	0.202	1.10	PQL	mg/Kg	
	BORON	J	10.2	11.0	PQL	mg/Kg	
	CADMIUM	J	0.178	1.10	PQL	mg/Kg	
SL-513-SA8-SB-8.0-9.0	TIN	J	2.78	11.0	PQL	mg/Kg	J (all detects)
	Zirconium	J	2.87	5.50	PQL	mg/Kg	
	BERYLLIUM	J	0.408	0.986	PQL	mg/Kg	
	CADMIUM	J	0.218	0.986	PQL	mg/Kg	
SL-514-SA8-SB-0.0-0.5	TIN	J	3.13	9.86	PQL	mg/Kg	J (all detects)
	Zirconium	J	1.91	4.93	PQL	mg/Kg	
	BERYLLIUM	J	0.845	1.06	PQL	mg/Kg	
SL-514-SA8-SB-0.0-0.5	TIN	J	3.33	10.6	PQL	mg/Kg	J (all detects)
	Zirconium	J	3.38	5.28	PQL	mg/Kg	
	BERYLLIUM	J	0.845	1.06	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-514-SA8-SB-4.0-5.0	BERYLLIUM	J	0.378	1.17	PQL	mg/Kg	J (all detects)
	BORON	J	2.51	11.7	PQL	mg/Kg	
	CADMIUM	J	0.223	1.17	PQL	mg/Kg	
	TIN	J	3.05	11.7	PQL	mg/Kg	
	Zirconium	J	3.34	5.87	PQL	mg/Kg	
SL-514-SA8-SB-7.5-8.5	BERYLLIUM	J	0.409	1.17	PQL	mg/Kg	J (all detects)
	BORON	J	1.24	11.7	PQL	mg/Kg	
	CADMIUM	J	0.117	1.17	PQL	mg/Kg	
	MOLYBDENUM	J	2.30	2.33	PQL	mg/Kg	
	TIN	J	3.00	11.7	PQL	mg/Kg	
SL-515-SA8-SB-0.0-0.5	Zirconium	J	3.38	5.83	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.890	1.10	PQL	mg/Kg	
	CADMIUM	J	0.116	1.10	PQL	mg/Kg	
	TIN	J	3.43	11.0	PQL	mg/Kg	
SL-515-SA8-SB-14.0-15.0	Zirconium	J	3.76	5.52	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.869	1.12	PQL	mg/Kg	
	TIN	J	3.34	11.2	PQL	mg/Kg	
SL-515-SA8-SB-4.0-5.0	Zirconium	J	2.60	5.61	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.984	1.10	PQL	mg/Kg	
	TIN	J	3.38	11.0	PQL	mg/Kg	
SL-515-SA8-SB-9.0-10.0	Zirconium	J	3.04	5.52	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	1.13	1.18	PQL	mg/Kg	
	TIN	J	3.76	11.8	PQL	mg/Kg	
SL-515-SA8-SB-9.0-10.0	Zirconium	J	4.44	5.89	PQL	mg/Kg	J (all detects)
	TIN	J	3.76	11.8	PQL	mg/Kg	

Method: 6020A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-512-SA8-SB-0.0-0.5	SELENIUM	J	0.204	0.390	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0426	0.195	PQL	mg/Kg	
SL-513-SA8-SB-0.0-0.5	SELENIUM	J	0.270	0.428	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0396	0.214	PQL	mg/Kg	
SL-513-SA8-SB-4.0-5.0	SELENIUM	J	0.119	0.440	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0473	0.220	PQL	mg/Kg	
	THALLIUM	J	0.215	0.220	PQL	mg/Kg	
SL-514-SA8-SB-0.0-0.5	SELENIUM	J	0.256	0.422	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0469	0.211	PQL	mg/Kg	
SL-514-SA8-SB-4.0-5.0	SILVER	J	0.0467	0.235	PQL	mg/Kg	J (all detects)
SL-514-SA8-SB-7.5-8.5	SELENIUM	J	0.171	0.466	PQL	mg/Kg	J (all detects)
SL-515-SA8-SB-0.0-0.5	SELENIUM	J	0.258	0.442	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0440	0.221	PQL	mg/Kg	
SL-515-SA8-SB-4.0-5.0	SILVER	J	0.0353	0.221	PQL	mg/Kg	J (all detects)
SL-515-SA8-SB-9.0-10.0	SELENIUM	J	0.137	0.471	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0431	0.236	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6850

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-513-SA8-SB-0.0-0.5	PERCHLORATE	J	3.2	5.5	PQL	ug/Kg	J (all detects)
SL-513-SA8-SB-4.0-5.0	PERCHLORATE	J	3.6	5.6	PQL	ug/Kg	J (all detects)

Method: 7199

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-513-SA8-SB-4.0-5.0	HEXAVALENT CHROMIUM	J	0.18	0.44	PQL	mg/Kg	J (all detects)

Method: 7471B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-515-SA8-SB-0.0-0.5	MERCURY	J	0.0136	0.0184	PQL	mg/Kg	J (all detects)

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-512-SA8-SB-0.0-0.5	EFH (C15-C20)	J	3.0	5.1	PQL	mg/Kg	J (all detects)
SL-514-SA8-SB-0.0-0.5	EFH (C15-C20)	J	2.9	5.3	PQL	mg/Kg	J (all detects)
SL-514-SA8-SB-4.0-5.0	EFH (C21-C30)	J	5.0	5.9	PQL	mg/Kg	J (all detects)
SL-514-SA8-SB-7.5-8.5	EFH (C21-C30)	J	3.6	5.9	PQL	mg/Kg	J (all detects)
SL-515-SA8-SB-14.0-15.0	EFH (C21-C30)	J	2.7	5.8	PQL	mg/Kg	J (all detects)
	EFH (C30-C40)	J	10	12	PQL	mg/Kg	

Method: 8081B

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-514-SA8-SB-0.0-0.5	4,4'-DDT	J	1.6	1.8	PQL	ug/Kg	J (all detects)
SL-514-SA8-SB-4.0-5.0	4,4'-DDT	J	0.76	2.0	PQL	ug/Kg	J (all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH069

Laboratory: LL

EDD Filename: PH069_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8082A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-512-SA8-SB-0.0-0.5	AROCLOR 1254	J	8.2	17	PQL	ug/Kg	J (all detects)

Method: 8270D

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-513-SA8-SB-0.0-0.5	BENZO(A)ANTHRACENE	J	5	18	PQL	ug/Kg	J (all detects)
	BENZO(A)PYRENE	J	9	18	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	8	18	PQL	ug/Kg	
	BENZO(K)FLUORANTHENE	J	5	18	PQL	ug/Kg	
	BIPHENYL	J	25	36	PQL	ug/Kg	
	NAPHTHALENE	J	4	18	PQL	ug/Kg	
PHENANTHRENE	J	15	18	PQL	ug/Kg		
SL-513-SA8-SB-4.0-5.0	BENZO(B)FLUORANTHENE	J	7	19	PQL	ug/Kg	J (all detects)
	PYRENE	J	4	19	PQL	ug/Kg	

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-512-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	1.1	1.7	PQL	ug/Kg	J (all detects)
	BIS(2-ETHYLHEXYL)PHTHALATE	J	11	18	PQL	ug/Kg	
	Di-n-butylphthalate	J	6.9	18	PQL	ug/Kg	
	FLUORANTHENE	J	1.4	1.7	PQL	ug/Kg	
	NAPHTHALENE	J	0.78	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	0.82	1.7	PQL	ug/Kg	
	PYRENE	J	1.1	1.7	PQL	ug/Kg	
SL-513-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	13	18	PQL	ug/Kg	J (all detects)
SL-513-SA8-SB-4.0-5.0	CHRYSENE	J	11	18	PQL	ug/Kg	J (all detects)
SL-514-SA8-SB-0.0-0.5	BENZO(A)ANTHRACENE	J	0.90	1.7	PQL	ug/Kg	J (all detects)
	BENZO(A)PYRENE	J	0.92	1.7	PQL	ug/Kg	
	BENZO(G,H,I)PERYLENE	J	0.92	1.7	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	11	19	PQL	ug/Kg	
	NAPHTHALENE	J	0.72	1.7	PQL	ug/Kg	
	PHENANTHRENE	J	1.5	1.7	PQL	ug/Kg	
SL-514-SA8-SB-7.5-8.5	NAPHTHALENE	J	0.91	1.9	PQL	ug/Kg	J (all detects)
SL-515-SA8-SB-0.0-0.5	CHRYSENE	J	5.1	19	PQL	ug/Kg	J (all detects)

Enclosure II
EPA Level IV Data Validation Reports

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name: Santa Susana Field Laboratory

Collection Date: July 17, 2013

LDC Report Date: September 23, 2013

Matrix: Soil/Water

Parameters: Volatiles

Validation Level: Level IV

Laboratory: Eurofins Lancaster Laboratories

Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713

TB-071713

SL-513-SA8-SB-8.0-9.0

Introduction

This data review covers one soil sample and 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8260B for Volatiles.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a Laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance was checked at 12 hour intervals.

All ion abundance requirements were met.

III. Initial Calibration

Initial calibration was performed using required standard concentrations.

Percent relative standard deviations (%RSD) were less than or equal to 30.0% for all compounds.

In the case where the laboratory used a calibration curve to evaluate the compounds, all coefficients of determination (r^2) were greater than or equal to 0.990 .

Average relative response factors (RRF) for all compounds were within method and validation criteria.

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

Percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within the method criteria of less than or equal to 20.0% for calibration check compounds (CCCs) and 25.0% for all other compounds.

The percent differences (%D) of the second source calibration standard were less than or equal to 25.0% for all compounds with the following exceptions:

Date	Compound	%D	Associated Samples	Flag	A or P
7/11/13	Vinyl acetate	44	All soil samples in SDG PH069	J (all detects) UJ (all non-detects)	A

All of the continuing calibration relative response factors (RRF) were within method and validation criteria.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No volatile contaminants were found in the method blanks.

Sample TB-071713 was identified as a trip blank. No volatile contaminants were found.

Sample EB1-071713 was identified as an equipment blank. No volatile contaminants were found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
EB1-071713	7/17/13	Acetone	7 ug/L	All soil samples in SDG PH069

Sample FB-041113 (from SDG PH029) was identified as a field blank. No volatile contaminants were found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB-041113	4/11/13	Methylene chloride	2 ug/L	All soil samples in SDG PH069

Sample concentrations were compared to concentrations detected in the field blanks. The sample concentrations were either not detected or were significantly greater (>10X for common contaminants, >5X for other contaminants) than the concentrations found in the associated field blanks.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

LCS ID (Associated Samples)	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Flag	A or P
LCSB68 (All soil samples in SDG PH069)	Vinyl acetate	135 (29-111)	125 (29-111)	-	J (all detects)	P
LCS/D Y77 (All water samples in SDG PH069)	Chlorotrifluoroethene	44 (47-120)	43 (47-120)	-	J (all detects) UJ (all non-detects)	P

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. Target Compound Identifications

All target compound identifications were within validation criteria.

XII. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported below the RL.	J (all detects)	A

XIII. Tentatively Identified Compounds (TICs)

Tentatively identified compounds were not reported by the laboratory.

XIV. System Performance

The system performance was acceptable.

XV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XVI. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Volatiles - Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH069	SL-513-SA8-SB-8.0-9.0	Vinyl acetate	J (all detects) UJ (all non-detects)	A	Continuing calibration (ICV %D) (C)
PH069	SL-513-SA8-SB-8.0-9.0	Vinyl acetate	J (all detects)	P	Laboratory control samples (%R) (L)
PH069	EB1-071713 TB-071713	Chlorotrifluoroethene	J (all detects) UJ (all non-detects)	P	Laboratory control samples (%R) (L)
PH069	EB1-071713 TB-071713 SL-513-SA8-SB-8.0-9.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Volatiles - Laboratory Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Volatiles - Field Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

METHOD: GC/MS Volatiles (EPA SW 846 Method 8260B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	Δ	Sampling dates: 7/17/13
II.	GC/MS Instrument performance check	Δ	
III.	Initial calibration	Δ	% RSD ≤ 30, r ²
IV.	Continuing calibration/ICV	SW	100/100 ≤ 25
V.	Blanks	Δ	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	N	client specified
VIII.	Laboratory control samples	SW	less 10
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	Δ	
XI.	Target compound identification	Δ	
XII.	Compound quantitation/RL/LOQ/LODs	Δ	
XIII.	Tentatively identified compounds (TICs)	N	
XIV.	System performance	Δ	
XV.	Overall assessment of data	Δ	
XVI.	Field duplicates	N	
XVII.	Field blanks	SW	EB = 1 * TB = 2 FB = FB-041113 (PH029)

Note: A = Acceptable * ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples:

water 2012

1	EB1-071713	w	11	VBLK476	21	31
2	TB-071713	w	12	VBLK467	22	32
3	SL-513-SA8-SB-8.0-9.0	✓	13		23	33
4		✓	14		24	34
5			15		25	35
6			16		26	36
7			17		27	37
8			18		28	38
9			19		29	39
10			20		30	40

Method: Volatiles (EPA SW 846 Method 8260C)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. GC/MS Instrument performance check				
Were the BFB performance results reviewed and found to be within the specified criteria?	/			
Were all samples analyzed within the 12 hour clock criteria?	/			
III. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	/			
Were all percent relative standard deviations (%RSD) \leq 20% ¹⁷ and relative response factors (RRF) within the method criteria? ₃₀	/			
Was a curve fit used for evaluation?	/			
Did the initial calibration meet the curve fit acceptance criteria of > 0.990 ?	/			
IV. Continuing calibration				
Was a continuing calibration standard analyzed at least once every 12 hours for each instrument?	/			
Were all percent differences (%D) \leq 20% ₂₅ and relative response factors (RRF) within the method criteria?		/		
V. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was a method blank analyzed at least once every 12 hours for each matrix and concentration?	/			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.		/		
VI. Surrogate spikes				
Were all surrogate %R within QC limits?	/			
If the percent recovery (%R) for one or more surrogates was out of QC limits, was a reanalysis performed to confirm samples with %R outside of criteria?			/	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.			/	
Was a MS/MSD analyzed every 20 samples of each matrix?			/	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?			/	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per analytical batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?		/		

VALIDATION FINDINGS CHECKLIST

Validation Area	Yes	No	NA	Findings/Comments
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?			<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?			<input checked="" type="checkbox"/>	
X. Internal standards				
Were internal standard area counts within -50% or +100% of the associated calibration standard?	<input checked="" type="checkbox"/>			
Were retention times within + 30 seconds of the associated calibration standard?	<input checked="" type="checkbox"/>			
XI. Target compound identification				
Were relative retention times (RRT's) within + 0.06 RRT units of the standard?	<input checked="" type="checkbox"/>			
Did compound spectra meet specified EPA "Functional Guidelines" criteria?	<input checked="" type="checkbox"/>			
Were chromatogram peaks verified and accounted for?	<input checked="" type="checkbox"/>			
XII. Compound quantitation/RLs				
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	<input checked="" type="checkbox"/>			
Were compound quantitation and RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	<input checked="" type="checkbox"/>			
XIII. Tentatively identified compounds (TICs)				
Were the major ions (> 10 percent relative intensity) in the reference spectrum evaluated in sample spectrum?			<input checked="" type="checkbox"/>	
Were relative intensities of the major ions within + 20% between the sample and the reference spectra?			<input checked="" type="checkbox"/>	
Did the raw data indicate that the laboratory performed a library search for all required peaks in the chromatograms (samples and blanks)?			<input checked="" type="checkbox"/>	
XIV. System performance				
System performance was found to be acceptable.	<input checked="" type="checkbox"/>			
XV. Overall assessment of data				
Overall assessment of data was found to be acceptable.	<input checked="" type="checkbox"/>			
XVI. Field duplicates				
Field duplicate pairs were identified in this SDG.		<input checked="" type="checkbox"/>		
Target compounds were detected in the field duplicates.			<input checked="" type="checkbox"/>	
XVII. Field blanks				
Field blanks were identified in this SDG.	<input checked="" type="checkbox"/>			
Target compounds were detected in the field blanks.	<input checked="" type="checkbox"/>			

TARGET COMPOUND WORKSHEET

METHOD: VOA

A. Chloromethane	U. 1,1,2-Trichloroethane	OO. 2,2-Dichloropropane	III. n-Butylbenzene	CCCC. 1-Chlorohexane
B. Bromomethane	V. Benzene	PP. Bromochloromethane	JJJ. 1,2-Dichlorobenzene	DDDD. Isopropyl alcohol
C. Vinyl chloride	W. trans-1,3-Dichloropropene	QQ. 1,1-Dichloropropene	KKK. 1,2,4-Trichlorobenzene	EEEE. Acetonitrile
D. Chloroethane	X. Bromoform	RR. Dibromomethane	LLL. Hexachlorobutadiene	FFFF. Acrolein
E. Methylene chloride	Y. 4-Methyl-2-pentanone	SS. 1,3-Dichloropropane	MMM. Naphthalene	GGGG. Acrylonitrile
F. Acetone	Z. 2-Hexanone	TT. 1,2-Dibromoethane	NNN. 1,2,3-Trichlorobenzene	HHHH. 1,4-Dioxane
G. Carbon disulfide	AA. Tetrachloroethene	UU. 1,1,1,2-Tetrachloroethane	OOO. 1,3,5-Trichlorobenzene	IIII. Isobutyl alcohol
H. 1,1-Dichloroethene	BB. 1,1,2,2-Tetrachloroethane	VV. Isopropylbenzene	PPP. trans-1,2-Dichloroethene	JJJJ. Methacrylonitrile
I. 1,1-Dichloroethane	CC. Toluene	WW. Bromobenzene	QQQ. cis-1,2-Dichloroethene	KKKK. Propionitrile
J. 1,2-Dichloroethene, total	DD. Chlorobenzene	XX. 1,2,3-Trichloropropane	RRR. m,p-Xylenes	LLLL. Ethyl ether
K. Chloroform	EE. Ethylbenzene	YY. n-Propylbenzene	SSS. o-Xylene	MMMM. Benzyl chloride
L. 1,2-Dichloroethane	FF. Styrene	ZZ. 2-Chlorotoluene	TTT. 1,1,2-Trichloro-1,2,2-trifluoroethane	NNNN. Iodomethane
M. 2-Butanone	GG. Xylenes, total	AAA. 1,3,5-Trimethylbenzene	UUU. 1,2-Dichlorotetrafluoroethane	OOOO. 1,1-Difluoroethane
N. 1,1,1-Trichloroethane	HH. Vinyl acetate	BBB. 4-Chlorotoluene	VVV. 4-Ethyltoluene	PPPP. <i>Chlorotrifluoroethene</i>
O. Carbon tetrachloride	II. 2-Chloroethylvinyl ether	CCC. tert-Butylbenzene	WWW. Ethanol	QQQQ.
P. Bromodichloromethane	JJ. Dichlorodifluoromethane	DDD. 1,2,4-Trimethylbenzene	XXX. Di-isopropyl ether	RRRR.
Q. 1,2-Dichloropropane	KK. Trichlorofluoromethane	EEE. sec-Butylbenzene	YYY. tert-Butanol	SSSS.
R. cis-1,3-Dichloropropene	LL. Methyl-tert-butyl ether	FFF. 1,3-Dichlorobenzene	ZZZ. tert-Butyl alcohol	TTTT.
S. Trichloroethene	MM. 1,2-Dibromo-3-chloropropane	GGG. p-Isopropyltoluene	AAAA. Ethyl tert-butyl ether	UUUU.
T. Dibromochloromethane	NN. Methyl ethyl ketone	HHH. 1,4-Dichlorobenzene	BBBB. tert-Amyl methyl ether	VVVV.

LDC #: 3028901a

VALIDATION FINDINGS WORKSHEET
Continuing Calibration

Page: 1 of 1
Reviewer: FT
2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260C)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

N N/A Was a continuing calibration standard analyzed at least once every 12 hours for each instrument?

N N/A Were percent differences (%D) $\leq 20\%$ and relative response factors (RRF) within the method criteria?

code = C

#	Date	Standard ID	Compound	Finding %D ^{F1} (Limit: $\leq 20.0\%$)	Finding RRF (Limit)	Associated Samples	Qualifications
	7/11/13	ICV	HH	44		All 8012	J/UJ/A

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: GC/MS VOA (EPA SW 846 Method 8260)

Y N /N/A Were field blanks identified in this SDG?
Y N /N/A Were target compounds detected in the field blanks?

Blank units: ug/L Associated sample units: ug/kg
Field blank type: (circle one) Field Blank / Rinsate / Trip Blank / Other: EB Associated Samples: All soil (ND)

Compound	Blank ID <u>1</u>	Blank ID	Sample Identification						
Sampling Date	<u>7/17/13</u>								
<u>F</u>	<u>7</u>								

Blank units: ug/L Associated sample units: ug/kg
Field blank type: (circle one) Field Blank / Rinsate / Trip Blank / Other: _____ Associated Samples: All soil (ND)

Compound	Blank ID	Blank ID	Sample Identification						
Sampling Date	<u>FB-041113</u>								
<u>E</u>	<u>2</u>								

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Common contaminants such as Methylene chloride, Acetone, 2-Butanone and Carbon disulfide that were detected in samples within ten times the associated field blank concentration were qualified as not detected, "U". Other contaminants within five times the field blank concentration were also qualified as not detected, "U".

LDC #: 3028901a

VALIDATION FINDINGS WORKSHEET
Laboratory Control Samples (LCS)

Page: 1 of 1
 Reviewer: FT
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260B)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Y N N/A Was a LCS required?
Y N N/A Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?

#	Date	LCS/LCSD ID	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Associated Samples	Qualifications
		<u>WS068</u>	<u>HH</u>	<u>135 (29-111)</u>	<u>125 (29-111)</u>	<u>()</u>	<u>All soils</u>	<u>J/Pdt (L)</u>
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
		<u>77</u> <u>Les/O 476_m</u>	<u>PPPP</u>	<u>44 (47-120)</u>	<u>43 (47-120)</u>	<u>()</u>	<u>All water</u>	<u>J/US/P</u> <u>↓ (L)</u>
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		
				<u>()</u>	<u>()</u>	<u>()</u>		

VALIDATION FINDINGS WORKSHEET

Initial Calibration Calculation Verification

METHOD: GC/MS VOA (EPA SW 846 Method 8260C)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$$RRF = (A_x)(C_{is}) / (A_{is})(C_x)$$

average RRF = sum of the RRFs/number of standards

$$\%RSD = 100 * (S/X)$$

A_x = Area of compound,

C_x = Concentration of compound,

S = Standard deviation of the RRFs

X = Mean of the RRFs

A_{is} = Area of associated internal standard

C_{is} = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				RRF (30 std)	RRF (50 std)	Average RRF (initial)	Average RRF (initial)	%RSD	%RSD
1	ICAL b	7/16/13	z z z (1st internal standard)	1.3094	1.3094	1.2195	1.2195	9	9
			C (2nd internal standard)	0.3413	0.3413	0.3579	0.3579	3	3
			CC (3rd internal standard)	0.9940	0.9940	0.9976	0.9976	1	1
			BB (4th internal standard)	0.7132	0.7132	0.7169	0.7169	5	5
2	ICAL	7/11/13	(1st internal standard)	(20)	(20)				
			Freon 133a (2nd internal standard)	0.6560	0.6560	0.6807	0.6807	4	4
			(3rd internal standard)						
			(4th internal standard)	(50)	(50)				
3	ICAL y	7/12/13	z z z (1st internal standard)	1.2475	1.2475	1.3105	1.3105	4	4
			C (2nd internal standard)	0.3813	0.3813	0.3694	0.3694	8	8
			CC (3rd internal standard)	1.0359	1.0359	0.9983	0.9983	5	5
			BB (4th internal standard)	1.1915	1.1915	1.1616	1.1616	4	4
4	ICAL	7/02/13	(1st internal standard)	(50)	(50)				
			Freon 133a (2nd internal standard)	0.7640	0.7640	0.7987	0.7987	5	5
			(3rd internal standard)						
			(4th internal standard)						

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results

LDC #: 80289D/2

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
 Reviewer: FT
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260C)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

$$\% \text{ Difference} = 100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$$

$$\text{RRF} = (A_x)(C_{is}) / (A_{is})(C_x)$$

Where: ave. RRF = initial calibration average RRF

RRF = continuing calibration RRF

A_x = Area of compound,A_{is} = Area of associated internal standardC_x = Concentration of compound,C_{is} = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference internal Standard)	Average RRF (initial)	Reported RRF (CC)	Recalculated RRF (CC)	Reported %D	Recalculated %D
1	cen 20.03	7/24/13	ZZZ (1st internal standard)	1.2195	1.2373	1.2373	1	1
			C (2nd internal standard)	0.3579	0.3917	0.3917	9	9
			CC (3rd internal standard)	0.9976	0.9965	0.9965	0	0.11
			BB (4th internal standard)	0.7169	0.7215	0.7215	1	1
2	cen 20.20	7/24/13	(1st internal standard)					
			Frem 133a (2nd internal standard)	0.6807	0.7448	0.7448	9	9
			(3rd internal standard)					
			(4th internal standard)					
3	cen 09.23	7/23/13	ZZZ (1st internal standard)	1.3105	1.4363	1.4363	10	10
			C (2nd internal standard)	0.3694	0.3466	0.3466	6	6
			CC (3rd internal standard)	0.9983	0.9752	0.9752	2	2
			BB (4th internal standard)	1.1616	1.1116	1.1116	4	4
4	cen 09.44	7/23/13	(1st internal standard)					
			Frem 133a (2nd internal standard)	0.7987	0.7813	0.7813	2	2
			(3rd internal standard)					
			(4th internal standard)					

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

METHOD: GC/MS VOA (EPA SW 846 Method 8260C)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
 SS = Surrogate Spiked

Sample ID: 3

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Dibromofluoromethane	50.0	50.334	101	101	0
1,2-Dichloroethane-d4		53.589	107	107	
Toluene-d8	↓	48.515	97	97	↓
Bromofluorobenzene	↓	47.821	96	96	↓

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Dibromofluoromethane					
1,2-Dichloroethane-d4					
Toluene-d8					
Bromofluorobenzene					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Dibromofluoromethane					
1,2-Dichloroethane-d4					
Toluene-d8					
Bromofluorobenzene					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Dibromofluoromethane					
1,2-Dichloroethane-d4					
Toluene-d8					
Bromofluorobenzene					

Sample ID: _____

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Dibromofluoromethane					
1,2-Dichloroethane-d4					
Toluene-d8					
Bromofluorobenzene					

LDC #: 30289D/a

VALIDATION FINDINGS WORKSHEET
Laboratory Control Sample Results Verification

Page: 1 of 1
 Reviewer: FT
 2nd Reviewer: [Signature]

METHOD: GC/MS VOA (EPA SW 846 Method 8260C)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the laboratory control sample and laboratory control sample duplicate (if applicable) were recalculated for the compounds identified below using the following calculation:

% Recovery = 100 * SSC/SA

Where: SSC = Spiked sample concentration
 SA = Spike added

RPD = | LCSC - LCSDC | * 2 / (LCSC + LCSDC)

LCSC = Laboratory control sample concentration LCSDC = Laboratory control sample duplicate concentration

LCS ID: LCS B67

Compound	Spike Added (ug/kg)		Spiked Sample Concentration (ug/kg)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD	LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
					Reported	Recalc.	Reported	Recalc.	Reported	Recalculated
1,1-Dichloroethene	20	NA	19.87	NA	99	99				
Trichloroethene	↓		19.81		99	99				
Benzene	↓	↓	19.68	↓	98	98				
Toluene	↓		19.33	↓	97	97				
Chlorobenzene	↓	↓	19.31	↓	97	97	NA			

Comments: Refer to Laboratory Control Sample findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 17, 2013
Matrix: Soil/Water
Parameters: Semivolatiles
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713
SL-513-SA8-SB-0.0-0.5
SL-513-SA8-SB-4.0-5.0
SL-513-SA8-SB-8.0-9.0

Introduction

This data review covers 3 soil samples and one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8270D for Semivolatiles.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance was checked at 12 hour intervals. All ion abundance requirements were met.

III. Initial Calibration

Initial calibration was performed using required standard concentrations.

Percent relative standard deviations (%RSD) were less than or equal to 30.0% for all compounds.

In the case where the laboratory used a calibration curve to evaluate the compounds all coefficients of determination (r^2) were greater than or equal to 0.990 .

Average relative response factors (RRF) for all compounds were within method and validation criteria.

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

Percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within the method criteria of less than or equal to 20.0% for calibration check compounds (CCCs) and 25.0% for all other compounds.

The percent differences (%D) of the second source calibration standard were less than or equal to 25.0% for all compounds with the following exceptions:

Date	Compound	%D	Associated Samples	Flag	A or P
7/25/13	Hexachlorocyclopentadiene	31	All water samples in SDG PH069	J (all detects) UJ (all non-detects)	A
7/22/13	Hexachlorocyclopentadiene	29	All soil samples in SDG PH069	J (all detects) UJ (all non-detects)	A
	2,4-Dinitrophenol	32		J (all detects) UJ (all non-detects)	

All of the continuing calibration relative response factors (RRF) were within method and validation criteria.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No semivolatile contaminants were found in the method blanks.

Sample EB1-071713 was identified as an equipment blank. No semivolatile contaminants were found.

Sample FB-041113 was identified as a field blank. No semivolatile contaminants were found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB-041113	4/11/13	Naphthalene	0.2 ug/L	SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0

Sample concentrations were compared to concentrations detected in the field blanks. The sample concentrations were either not detected or were significantly greater (>10X for common contaminants, >5X for other contaminants) than the concentrations found in the associated field blanks.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

LCS ID (Associated Samples)	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Flag	A or P
205WCLCS/D (All water samples in SDG PH069)	Benzoic acid	-	-	54 (≤30)	J (all detects) UJ (all non-detects)	P

LCS ID (Associated Samples)	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Flag	A or P
205WCLCS/D (All water samples in SDG PH069)	4-Nitrophenol	75 (13-71)	80 (13-71)	-	J (all detects)	P

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. Target Compound Identifications

All target compound identifications were within validation criteria.

XII. Compound Quantitation

All compound quantitation were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported below the RL.	J (all detects)	A

XIII. Tentatively Identified Compounds (TICs)

Tentatively identified compounds were not reported by the laboratory.

XIV. System Performance

The system performance was acceptable.

XV. Overall Assessment

Data flags are summarized at the end of this report if data has been qualified.

XVI. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Semivolatiles - Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH069	EB1-071713	Hexachlorocyclopentadiene	J (all detects) UJ (all non-detects)	A	Continuing calibration (%D) (C)
PH069	SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	Hexachlorocyclopentadiene 2,4-Dinitrophenol	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	A	Continuing calibration (%D) (C)
PH069	EB1-071713	Benzoic acid	J (all detects) UJ (all non-detects)	P	Laboratory control samples (RPD) (E)
PH069	EB1-071713	4-Nitrophenol	J (all detects)	P	Laboratory control samples (%R) (L)
PH069	EB1-071713 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Semivolatiles - Laboratory Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Semivolatiles - Field Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

METHOD: GC/MS Semivolatiles (EPA SW 846 Method 8270^D)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 7/17/13
II.	GC/MS Instrument performance check	A	
III.	Initial calibration	A	RSD ≤ 30%, P ²
IV.	Continuing calibration/ICV	SW	CCV/ICV ≤ 25%
V.	Blanks	A	
VI.	Surrogate spikes	A	
VII.	Matrix spike/Matrix spike duplicates	N	
VIII.	Laboratory control samples	SW	LCS/D
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	A	
XI.	Target compound identification	A	
XII.	Compound quantitation/RL/LOQ/LODs	A	
XIII.	Tentatively identified compounds (TICs)	N	
XIV.	System performance	A	
XV.	Overall assessment of data	SW	
XVI.	Field duplicates	N	
XVII.	Field blanks	SW	EB = 1 FB = 1804 1113 (SDG PH029)

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples:

1	EB1-071713	W	11	SBLKWC225	21	31
2	SL-513-SA8-SB-0.0-0.5	S	12	SBLKLE199	22	32
3	SL-513-SA8-SB-4.0-5.0		13		23	33
4	SL-513-SA8-SB-8.0-9.0	✓	14		24	34
5			15		25	35
6			16		26	36
7			17		27	37
8			18		28	38
9			19		29	39
10			20		30	40

Method: Semivolatiles (EPA SW 846 Method 8270)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. GC/MS Instrument performance check				
Were the DFTPP performance results reviewed and found to be within the specified criteria?	/			
Were all samples analyzed within the 12 hour clock criteria?	/			
III. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	/			
Were all percent relative standard deviations (%RSD) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	/			
Was a curve fit used for evaluation?	/			
Did the initial calibration meet the curve fit acceptance criteria of ≥ 0.990 ?	/			
Were all percent relative standard deviations (%RSD) $\leq 30\%/15\%$ and relative response factors (RRF) ≥ 0.05 ?	/			
IV. Continuing calibration				
Was a continuing calibration standard analyzed at least once every 12 hours for each instrument?	/			
Were all percent differences (%D) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	/			
Were all percent differences (%D) $\leq 5\%$ and relative response factors (RRF) ≥ 0.05 ?		/		
V. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was a method blank analyzed for each matrix and concentration?	/			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.		/		
VI. Surrogate spikes				
Were all surrogate %R within QC limits?	/			
If 2 or more base neutral or acid surrogates were outside QC limits, was a reanalysis performed to confirm %R?			/	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?			/	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.		/		
Was a MS/MSD analyzed every 20 samples of each matrix?		/		
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?			/	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			


Validation Area	Yes	No	NA	Findings/Comments
Was an LCS analyzed per extraction batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?		/		
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?			/	
Were the performance evaluation (PE) samples within the acceptance limits?			/	
X. Internal standards				
Were internal standard area counts within -50% or +100% of the associated calibration standard?	/			
Were retention times within + 30 seconds from the associated calibration standard?	/			
XI. Target compound identification				
Were relative retention times (RRT's) within + 0.06 RRT units of the standard?	/			
Did compound spectra meet specified EPA "Functional Guidelines" criteria?	/			
Were chromatogram peaks verified and accounted for?	/			
XII. Compound quantitation/CRQLs				
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	/			
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XIII. Tentatively identified compounds (TICs)				
Were the major ions (> 10 percent relative intensity) in the reference spectrum evaluated in sample spectrum?			/	
Were relative intensities of the major ions within ± 20% between the sample and the reference spectra?			/	
Did the raw data indicate that the laboratory performed a library search for all required peaks in the chromatograms (samples and blanks)?			/	
XIV. System performance				
System performance was found to be acceptable.	/			
XV. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XVI. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
XVII. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.		/		

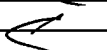
VALIDATION FINDINGS WORKSHEET

METHOD: GC/MS SVOA

A. Phenol	T. 4-Chloroaniline	MM. 4-Chlorophenyl-phenyl ether	FFF. Di-n-octylphthalate	YYY. 2,3,5-Trimethylnaphthalene
B. Bis (2-chloroethyl) ether	U. Hexachlorobutadiene	NN. Fluorene	GGG. Benzo(b)fluoranthene	ZZZ. Perylene
C. 2-Chlorophenol	V. 4-Chloro-3-methylphenol	OO. 4-Nitroaniline	HHH. Benzo(k)fluoranthene	AAAA. Dibenzothiophene
D. 1,3-Dichlorobenzene	W. 2-Methylnaphthalene	PP. 4,6-Dinitro-2-methylphenol	III. Benzo(a)pyrene	BBBB. Benzo(a)fluoranthene
E. 1,4-Dichlorobenzene	X. Hexachlorocyclopentadiene	QQ. N-Nitrosodiphenylamine	JJJ. Indeno(1,2,3-cd)pyrene	CCCC. Benzo(b)fluorene
F. 1,2-Dichlorobenzene	Y. 2,4,6-Trichlorophenol	RR. 4-Bromophenyl-phenylether	KKK. Dibenz(a,h)anthracene	DDDD. cis/trans-Decalin
G. 2-Methylphenol	Z. 2,4,5-Trichlorophenol	SS. Hexachlorobenzene	LLL. Benzo(g,h,i)perylene	EEEE. Biphenyl
H. 2,2'-Oxybis(1-chloropropane)	AA. 2-Chloronaphthalene	TT. Pentachlorophenol	MMM. Bis(2-Chloroisopropyl)ether	FFFF. Retene
I. 4-Methylphenol	BB. 2-Nitroaniline	UU. Phenanthrene	NNN. Aniline	GGGG. C30-Hopane
J. N-Nitroso-di-n-propylamine	CC. Dimethylphthalate	VV. Anthracene	OOO. N-Nitrosodimethylamine	HHHH. 1-Methylphenanthrene
K. Hexachloroethane	DD. Acenaphthylene	WW. Carbazole	PPP. Benzoic Acid	IIII. 1,4-Dioxane
L. Nitrobenzene	EE. 2,6-Dinitrotoluene	XX. Di-n-butylphthalate	QQQ. Benzyl alcohol	JJJJ. Acetophenone
M. Isophorone	FF. 3-Nitroaniline	YY. Fluoranthene	RRR. Pyridine	KKKK. Atrazine
N. 2-Nitrophenol	GG. Acenaphthene	ZZ. Pyrene	SSS. Benzidine	LLLL. Benzaldehyde
O. 2,4-Dimethylphenol	HH. 2,4-Dinitrophenol	AAA. Butylbenzylphthalate	TTT. 1-Methylnaphthalene	MMMM. Caprolactam
P. Bis(2-chloroethoxy)methane	II. 4-Nitrophenol	BBB. 3,3'-Dichlorobenzidine	UUU. Benzo(b)thiophene	NNNN.
Q. 2,4-Dichlorophenol	JJ. Dibenzofuran	CCC. Benzo(a)anthracene	VVV. Benzonaphthothiophene	OOOO.
R. 1,2,4-Trichlorobenzene	KK. 2,4-Dinitrotoluene	DDD. Chrysene	WWW. Benzo(e)pyrene	PPPP.
S. Naphthalene	LL. Diethylphthalate	EEE. Bis(2-ethylhexyl)phthalate	XXX. 2,6-Dimethylnaphthalene	QQQQ.

VALIDATION FINDINGS WORKSHEET
Continuing Calibration

Reviewer: 

2nd Reviewer: 

METHOD: GC/MS BNA (EPA SW 846 Method 8270D)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

N N/A Was a continuing calibration standard analyzed at least once every 12 hours for each instrument?

N N/A Were percent differences (%D) $\leq 20\%$ and relative response factors (RRF) within the method criteria?

#	Date	Standard ID	Compound	Finding %D (Limit: $\leq 20.0\%$)	Finding RRF (Limit)	Associated Samples	Qualifications
	7/25/13	d90521 (10V)	X	31		tel W	J/H/A
							↓
	7/22/13	eg0711 (10V)	X HH	29 32		tel soil	↓

LDC #: 30289D2a

VALIDATION FINDINGS WORKSHEET Field Blanks

Page: 1 of 1
Reviewer: SM
2nd Reviewer: A

METHOD: GC/MS BNA (EPA SW 846 Method 8270^B)
(N) N/A Were field blanks identified in this SDG?
(N) N/A Were target compounds detected in the field blanks?
Blank units: ug/L Associated sample units: ug/kg
Sampling date: 04/11/13
Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: 2-4

Compound	Blank ID	Sample Identification							
	FB-041113	action limit							
Naphthalene	0.2	1.0	No	Qual					

Blank units: _____ Associated sample units: _____
Sampling date: _____
Field blank type: (circle one) _____ / Rinsate / Other: _____ Associated Samples: _____

Compound	Blank ID	Sample Identification							

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Common contaminants such as the phthalates and TICs noted above that were detected in samples within ten times the associated field blank concentration were qualified as not detected, "U". Other contaminants within five times the field blank concentration were also qualified as not detected, "U".

VALIDATION FINDINGS WORKSHEET
Laboratory Control Samples (LCS)

METHOD: GC/MS BNA (EPA SW 846 Method 8270D)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Y N N/A

Was a LCS required?

Y N N/A

Were the LCS/LCSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?

#	Date	LCS/LCSD ID	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Associated Samples	Qualifications
	205W	CLCS/D	PPP	()	()	54 (=30)	See W	JMJ/P
				75 (13-71)	80 (13-71)	()		JMJ/P
				()	()	()		
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LDC #: 30287029

VALIDATION FINDINGS WORKSHEET
Overall Assessment of Data

Page: 1 of 1

Reviewer: [Signature]

2nd Reviewer: [Signature]

METHOD: GC/MS BNA (EPA SW 846 Method 8270D)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

All available information pertaining to the data were reviewed using professional judgement to compliment the determination of the overall quality of the data.

N N/A Was the overall quality and usability of the data acceptable?

#	Date	Compound	Finding	Associated Samples	Qualifications
			Lab indicated that extract	4	
			spilled after bottling, since		
			80% PSD w/in limits. No qual		

Comments: _____

VALIDATION FINDINGS WORKSHEET Initial Calibration Calculation Verification

METHOD: GC/MS BNA (EPA SW 846 Method 8270)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$$RRF = (A_x)(C_{is}) / (A_{is})(C_x)$$

average RRF = sum of the RRFs/number of standards

$$\%RSD = 100 * (S/X)$$

A_x = Area of compound,

C_x = Concentration of compound,

S = Standard deviation of the RRFs,

A_{is} = Area of associated internal standard

C_{is} = Concentration of internal standard

X = Mean of the RRFs

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				RRF (7.5 std)	RRF (7.5 std)	Average RRF (initial)	Average RRF (initial)	%RSD	%RSD
1	ICAL	7/24/13	Phenol (1st internal standard)	2.907	2.907	2.811	2.811	4	4
			Naphthalene (2nd internal standard)	1.090	1.090	1.054	1.054	3	3
			Fluorene (3rd internal standard)	1.290	1.290	1.257	1.257	5	5
			Pentachlorophenol (4th internal standard)	0.143	0.143	0.132	0.132	16	16
			Bis(2-ethylhexyl)phthalate (5th internal standard)	0.685	0.685	0.639	0.639	10	10
			Benzo(a)pyrene (6th internal standard)	1.130	1.130	0.991	0.991	17	17
2	LOK	7/22/13	Phenol (1st internal standard) (30 std)	2.749	2.749	2.696	2.696	9	9
			Naphthalene (2nd internal standard)	1.003	1.003	1.003	1.003	5	5
			Fluorene (3rd internal standard)	1.326	1.326	1.332	1.332	4	4
			Pentachlorophenol (4th internal standard)	0.098	0.098	0.094	0.094	19	19
			Bis(2-ethylhexyl)phthalate (5th internal standard)	0.660	0.660	0.660	0.660	5	5
			Benzo(a)pyrene (6th internal standard)	1.210	1.210	1.123	1.123	10	10
3			Phenol (1st internal standard)						
			Naphthalene (2nd internal standard)						
			Fluorene (3rd internal standard)						
			Pentachlorophenol (4th internal standard)						
			Bis(2-ethylhexyl)phthalate (5th internal standard)						
			Benzo(a)pyrene (6th internal standard)						

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET

Continuing Calibration Results Verification

METHOD: GC/MS BNA (EPA SW 846 Method 8270)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

$$\% \text{ Difference} = 100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$$

$$\text{RRF} = (A_x)(C_{is}) / (A_{is})(C_x)$$

Where: ave. RRF = initial calibration average RRF

RRF = continuing calibration RRF

 A_x = Area of compound, C_x = Concentration of compound, A_{is} = Area of associated internal standard C_{is} = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Average RRF (initial)	Reported	Recalculated	Reported	Recalculated
					RRF (CC)	RRF (CC)	%D	%D
1	dg0581	7/25/13	Phenol (1st internal standard)	2.811	2.733	2.733	3	3
			Naphthalene (2nd internal standard)	1.054	1.027	1.027	3	3
			Fluorene (3rd internal standard)	1.257	1.260	1.260	0	0
			Pentachlorophenol (4th internal standard)	0.132	0.140	0.140	6	6
			Bis(2-ethylhexyl)phthalate (5th internal standard)	0.639	0.658	0.658	3	3
			Benzo(a)pyrene (6th internal standard)	0.991	1.099	1.099	11	11
2	eg0901	7/28/13	Phenol (1st internal standard)	2.696	2.440	2.440	10	10
			Naphthalene (2nd internal standard)	1.003	1.000	1.0006 ^{cc}	0	0
			Fluorene (3rd internal standard)	1.332	1.355	1.355	2	2
			Pentachlorophenol (4th internal standard)	0.094	0.099	0.099	5	5
			Bis(2-ethylhexyl)phthalate (5th internal standard)	0.660	0.654	0.654	1	1
			Benzo(a)pyrene (6th internal standard)	1.123	1.141	1.141	2	2
3	eg0941	7/29/13	Phenol (1st internal standard)	2.696	2.533	2.533	6	6
			Naphthalene (2nd internal standard)	1.003	1.001	1.001	0	0
			Fluorene (3rd internal standard)	1.332	1.333	1.333	0	0
			Pentachlorophenol (4th internal standard)	0.094	0.099	0.099	6	6
			Bis(2-ethylhexyl)phthalate (5th internal standard)	0.660	0.679	0.679	3	3
			Benzo(a)pyrene (6th internal standard)	1.123	1.183	1.183	5	5

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET

Surrogate Results Verification

METHOD: GC/MS Semivolatiles (EPA SW 846 Method 8270)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: $SF/SS * 100$

Where: SF = Surrogate Found
 SS = Surrogate Spiked

Sample ID: 2

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5	100	83.689	84	84	0
2-Fluorobiphenyl	↓	90.234	90	90	↓
Terphenyl-d14	↓	90.577	91	91	↓
Phenol-d6	200	154.817	77	77	↓
2-Fluorophenol	↓	170.592	85	85	↓
2,4,6-Tribromophenol	↓	162.438	81	81	↓
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

Sample ID:

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5					
2-Fluorobiphenyl					
Terphenyl-d14					
Phenol-d5					
2-Fluorophenol					
2,4,6-Tribromophenol					
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

Sample ID:

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5					
2-Fluorobiphenyl					
Terphenyl-d14					
Phenol-d5					
2-Fluorophenol					
2,4,6-Tribromophenol					
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

VALIDATION FINDINGS WORKSHEET

Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

METHOD: GC/MS BNA (EPA SW 846 Method 8270)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

% Recovery = 100 * (SC/SA)

Where: SSC = Spike concentration
SA = Spike added

RPD = | LCSC - LCSDC | * 2 / (LCSC + LCSDC)

LCSC = Laboratory control sample concentration LCSDC = Laboratory control sample duplicate concentration

LCS/LCSD samples: 199LELCS

Compound	Spike Added (ug/kg)		Spike Concentration (ug/kg)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD	LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
					Reported	Recalc	Reported	Recalc	Reported	Recalculated
Phenol	1666.67		1610.49		97	97				
N-Nitroso-di-n-propylamine	↓		1512.06		91	91				
4-Chloro-3-methylphenol	↓		1728.87		104	104				
Acenaphthene	-									
Pentachlorophenol	1666.67		1755.68		105	105				
Pyrene	↓		1620.79		97	97				

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

**VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification**

METHOD: GC/MS BNA (EPA SW 846 Method 8270)

(Y) N N/A
(V) N N/A

Were all reported results recalculated and verified for all level IV samples?

Were all recalculated results for detected target compounds agree within 10.0% of the reported results?

$$\text{Concentration} = \frac{(A_x)(I_s)(V_i)(DF)(2.0)}{(A_{is})(RRF)(V_o)(V_i)(\%S)}$$

- A_x = Area of the characteristic ion (EICP) for the compound to be measured
- A_{is} = Area of the characteristic ion (EICP) for the specific internal standard
- I_s = Amount of internal standard added in nanograms (ng)
- V_o = Volume or weight of sample extract in milliliters (ml) or grams (g).
- V_i = Volume of extract injected in microliters (ul)
- V_c = Volume of the concentrated extract in microliters (ul)
- Df = Dilution Factor.
- %S = Percent solids, applicable to soil and solid matrices only.
- 2.0 = Factor of 2 to account for GPC cleanup

Example:

Sample I.D. 2, S:

$$\text{Conc.} = \frac{2373 \times 20 \times 1000 \times () \times ()}{(594265)(1.002)(30.3)(2.92) \times ()}$$

= 4 ug/kg

#	Sample ID	Compound	Reported Concentration ()	Calculated Concentration ()	Qualification

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 23, 2013
Matrix: Soil/Water
Parameters: Semivolatiles
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713
EB2-071713
SL-512-SA8-SB-0.0-0.5
SL-514-SA8-SB-0.0-0.5
SL-514-SA8-SB-4.0-5.0
SL-514-SA8-SB-7.5-8.5
SL-515-SA8-SB-0.0-0.5
SL-515-SA8-SB-4.0-5.0
SL-515-SA8-SB-9.0-10.0
SL-515-SA8-SB-14.0-15.0
SL-513-SA8-SB-0.0-0.5
SL-513-SA8-SB-4.0-5.0
SL-513-SA8-SB-8.0-9.0

Introduction

This data review covers 11 soil samples and 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8270C using Selected Ion Monitoring (SIM) for Semivolatiles.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/MS Instrument Performance Check

Instrument performance was checked at 12 hour intervals. All ion abundance requirements were met.

III. Initial Calibration

Initial calibration was performed using required standard.

Percent relative standard deviations (%RSD) were less than or equal to 30.0% for all compounds.

Average relative response factors (RRF) for all compounds were within method and validation criteria.

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

Percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within the method criteria of less than or equal to 20.0% for calibration check compounds (CCCs) and 25.0% for all other compounds.

The percent differences (%D) of the second source calibration standard were less than or equal to 25.0% for all compounds.

All of the continuing calibration relative response factors (RRF) were within method and validation criteria.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No semivolatile contaminants were found in the method with the following exceptions:

Method Blank ID	Analysis Date	Compound	Concentration	Associated Samples
SBLKWB205	7/29/13	Bis(2-ethylhexyl)phthalate	0.051 ug/L	All water samples in SDG PH069

Sample concentrations were compared to concentrations detected in the method blanks. The sample concentrations were either not detected or were significantly greater (>10X for common contaminants, >5X for other contaminants) than the concentrations found in the associated method blanks with the following exceptions:

Sample	Compound	Reported Concentration	Modified Final Concentration
EB1-071713	Bis(2-ethylhexyl)phthalate	0.19 ug/L	1.1U ug/L
EB2-071713	Bis(2-ethylhexyl)phthalate	0.14 ug/L	1.0U ug/L

Samples EB1-071713 and EB2-071713 were identified as equipment blanks. No semivolatle contaminants were found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
EB1-071713	7/17/13	Di-n-butylphthalate Diethylphthalate Bis(2-ethylhexyl)phthalate 2-Methylnaphthalene Naphthalene	0.24 ug/L 0.24 ug/L 0.19 ug/L 0.014 ug/L 0.046 ug/L	SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0
EB2-071713	7/17/13	Di-n-butylphthalate Diethylphthalate Bis(2-ethylhexyl)phthalate 2-Methylnaphthalene Naphthalene	0.14 ug/L 0.22 ug/L 0.14 ug/L 0.014 ug/L 0.045 ug/L	SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5

Sample FB-041113 (from SDG PH029) was identified as a field blank. No semivolatle contaminants were found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB-041113	4/11/13	Di-n-butylphthalate Diethylphthalate Bis(2-ethylhexyl)phthalate 1-Methylnaphthalene 2-Methylnaphthalene Naphthalene	0.17 ug/L 0.18 ug/L 0.082 ug/L 0.019 ug/L 0.024 ug/L 0.17 ug/L	SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0

Sample concentrations were compared to concentrations detected in the field blanks. The sample concentrations were either not detected or were significantly greater (>5X blank contaminants) than the concentrations found in the associated field blanks with the following exceptions:

Sample	Compound	Reported Concentration	Modified Final Concentration
SL-512-SA8-SB-0.0-0.5	Naphthalene	0.78 ug/Kg	1.7U ug/Kg
SL-514-SA8-SB-0.0-0.5	Naphthalene	0.72 ug/Kg	1.7U ug/Kg

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Internal Standards

All internal standard areas and retention times were within QC limits.

XI. Target Compound Identifications

All target compound identifications were within validation criteria.

XII. Compound Quantitation

All compound quantitation were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported below the RL.	J (all detects)	A

XIII. Tentatively Identified Compounds (TICs)

Tentatively identified compounds were not reported by the laboratory.

XIV. System Performance

The system performance was acceptable.

XV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XVI. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Semivolatiles - Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH069	EB1-071713 EB2-071713 SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Semivolatiles - Laboratory Blank Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
PH069	EB1-071713	Bis(2-ethylhexyl)phthalate	1.1U ug/L	A	B
PH069	EB2-071713	Bis(2-ethylhexyl)phthalate	1.0U ug/L	A	B

**Santa Susana Field Laboratory
Semivolatiles - Field Blank Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
PH069	SL-512-SA8-SB-0.0-0.5	Naphthalene	1.7U ug/Kg	A	F
PH069	SL-514-SA8-SB-0.0-0.5	Naphthalene	1.7U ug/Kg	A	F

METHOD: GC/MS Polynuclear Aromatic Hydrocarbons (EPA SW 846 Method 8270C-SIM)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	Δ	Sampling dates: 7/17/13
II.	GC/MS Instrument performance check	Δ	
III.	Initial calibration	Δ	% RSD \leq 30
IV.	Continuing calibration/ICV	Δ	ICV/CCV \leq 25
V.	Blanks	SW	
VI.	Surrogate spikes	Δ	
VII.	Matrix spike/Matrix spike duplicates	N	client specified
VIII.	Laboratory control samples	Δ	LCS10
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	Δ	
XI.	Target compound identification	Δ	
XII.	Compound quantitation/RL/LOQ/LODs	Δ	
XIII.	Tentatively identified compounds (TICs)	N	
XIV.	System performance	Δ	
XV.	Overall assessment of data	Δ	
XVI.	Field duplicates	N	
XVII.	Field blanks	SW	EB = 1, 2 *FB = FB-041113 (SDG# PH029)

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

*ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples:
 soil + water

13	EB1-071713	w	11	SL-513-SA8-SB-0.0-0.5	21	SBLKLF199	31
23	EB2-071713	↓	12	SL-513-SA8-SB-4.0-5.0	22	SBLKLA201 -	32
32	SL-512-SA8-SB-0.0-0.5		13	SL-513-SA8-SB-8.0-9.0	23	SBLKW/B205 -	33
42	SL-514-SA8-SB-0.0-0.5		14		24	SBLKLA207 -	34
52	SL-514-SA8-SB-4.0-5.0		15		25		35
62	SL-514-SA8-SB-7.5-8.5		16		26		36
72	SL-515-SA8-SB-0.0-0.5		17		27		37
82	SL-515-SA8-SB-4.0-5.0		18		28		38
92	SL-515-SA8-SB-9.0-10.0		19		29		39
102	SL-515-SA8-SB-14.0-15.0		20		30		40

Method: Semivolatiles (EPA SW 846 Method 8270C)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
II. GC/MS Instrument performance check				
Were the DFTPP performance results reviewed and found to be within the specified criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all samples analyzed within the 12 hour clock criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
III. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a curve fit used for evaluation?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Did the initial calibration meet the curve fit acceptance criteria of ≥ 0.990 ?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were all percent relative standard deviations (%RSD) $\leq 30\%/15\%$ and relative response factors (RRF) ≥ 0.05 ?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IV. Continuing calibration				
Was a continuing calibration standard analyzed at least once every 12 hours for each instrument?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) and relative response factors (RRF) within method criteria for all CCCs and SPCCs?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) $\leq 20\%$ and relative response factors (RRF) ≥ 0.05 ?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
V. Blanks				
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank analyzed for each matrix and concentration?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
VI. Surrogate spikes				
Were all surrogate %R within QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
If 2 or more base neutral or acid surrogates were outside QC limits, was a reanalysis performed to confirm %R?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	

Validation Area	Yes	No	NA	Findings/Comments
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
X. Internal standards				
Were internal standard area counts within -50% or +100% of the associated calibration standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were retention times within + 30 seconds from the associated calibration standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XI. Target compound identification				
Were relative retention times (RRT's) within + 0.06 RRT units of the standard?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Did compound spectra meet specified EPA "Functional Guidelines" criteria?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were chromatogram peaks verified and accounted for?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XII. Compound quantitation/CRQLs				
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XIII. Tentatively identified compounds (TICs)				
Were the major ions (> 10 percent relative intensity) in the reference spectrum evaluated in sample spectrum?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were relative intensities of the major ions within ± 20% between the sample and the reference spectra?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Did the raw data indicate that the laboratory performed a library search for all required peaks in the chromatograms (samples and blanks)?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
XIV. System performance				
System performance was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XV. Overall assessment of data				
Overall assessment of data was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XVI. Field duplicates				
Field duplicate pairs were identified in this SDG.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field duplicates.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
XVII. Field blanks				
Field blanks were identified in this SDG.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field blanks.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	

VALIDATION FINDINGS WORKSHEET

METHOD: GC/MS SVOA

A. Phenol	T. 4-Chloroaniline	MM. 4-Chlorophenyl-phenyl ether	FFF. Di-n-octylphthalate	YYY. 2,3,5-Trimethylnaphthalene
B. Bis (2-chloroethyl) ether	U. Hexachlorobutadiene	NN. Fluorene	GGG. Benzo(b)fluoranthene	ZZZ. Perylene
C. 2-Chlorophenol	V. 4-Chloro-3-methylphenol	OO. 4-Nitroaniline	HHH. Benzo(k)fluoranthene	AAAA. Dibenzothiophene
D. 1,3-Dichlorobenzene	W. 2-Methylnaphthalene	PP. 4,6-Dinitro-2-methylphenol	III. Benzo(a)pyrene	BBBB. Benzo(a)fluoranthene
E. 1,4-Dichlorobenzene	X. Hexachlorocyclopentadiene	QQ. N-Nitrosodiphenylamine	JJJ. Indeno(1,2,3-cd)pyrene	CCCC. Benzo(b)fluorene
F. 1,2-Dichlorobenzene	Y. 2,4,6-Trichlorophenol	RR. 4-Bromophenyl-phenylether	KKK. Dibenz(a,h)anthracene	DDDD. cis/trans-Decalin
G. 2-Methylphenol	Z. 2,4,5-Trichlorophenol	SS. Hexachlorobenzene	LLL. Benzo(g,h,i)perylene	EEEE. Biphenyl
H. 2,2'-Oxybis(1-chloropropane)	AA. 2-Chloronaphthalene	TT. Pentachlorophenol	MMM. Bis(2-Chloroisopropyl)ether	FFFF. Retene
I. 4-Methylphenol	BB. 2-Nitroaniline	UU. Phenanthrene	NNN. Aniline	GGGG. C30-Hopane
J. N-Nitroso-di-n-propylamine	CC. Dimethylphthalate	VV. Anthracene	OOO. N-Nitrosodimethylamine	HHHH. 1-Methylphenanthrene
K. Hexachloroethane	DD. Acenaphthylene	WW. Carbazole	PPP. Benzoic Acid	IIII. 1,4-Dioxane
L. Nitrobenzene	EE. 2,6-Dinitrotoluene	XX. Di-n-butylphthalate	QQQ. Benzyl alcohol	JJJJ. Acetophenone
M. Isophorone	FF. 3-Nitroaniline	YY. Fluoranthene	RRR. Pyridine	KKKK. Atrazine
N. 2-Nitrophenol	GG. Acenaphthene	ZZ. Pyrene	SSS. Benzidine	LLLL. Benzaldehyde
O. 2,4-Dimethylphenol	HH. 2,4-Dinitrophenol	AAA. Butylbenzylphthalate	TTT. 1-Methylnaphthalene	MMMM. Caprolactam
P. Bis(2-chloroethoxy)methane	II. 4-Nitrophenol	BBB. 3,3'-Dichlorobenzidine	UUU. Benzo(b)thiophene	NNNN.
Q. 2,4-Dichlorophenol	JJ. Dibenzofuran	CCC. Benzo(a)anthracene	VVV. Benzonaphthothiophene	OOOO.
R. 1,2,4-Trichlorobenzene	KK. 2,4-Dinitrotoluene	DDD. Chrysene	WWW. Benzo(e)pyrene	PPPP.
S. Naphthalene	LL. Diethylphthalate	EEE. Bis(2-ethylhexyl)phthalate	XXX. 2,6-Dimethylnaphthalene	QQQQ.

VALIDATION FINDINGS WORKSHEET

Blanks

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- Y ~~N~~ N/A Was a method blank analyzed for each matrix?
- Y ~~N~~ N/A Was a method blank analyzed for each concentration preparation level?
- Y ~~N~~ N/A Was a method blank associated with every sample?
- Y ~~N~~ N/A Was the blank contaminated? If yes, please see qualification below.

(B)

Blank extraction date: _____ Blank analysis date: 7/29/13

Conc. units: ug/l Associated Samples: All water

Compound	Blank ID	Sample Identification							
	SBLKWB205	1	2						
EEE	0.051	0.19/1.10	0.14/1.04						

Blank extraction date: _____ Blank analysis date: _____

Conc. units: _____ Associated Samples: _____

Compound	Blank ID	Sample Identification							

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 Common contaminants such as the phthalates and TICs noted above that were detected in samples within ten times the associated method blank concentration were qualified as not detected, "U". Other contaminants within five times the method blank concentration were also qualified as not detected, "U".

LDC #: 3028702b

VALIDATION FINDINGS WORKSHEET

Page: 1 of 7

Field Blanks

Reviewer: FT

2nd Reviewer: [Signature]

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

Y N N/A Were field blanks identified in this SDG?

Y N N/A Were target compounds detected in the field blanks?

Blank units: ug/L Associated sample units: ug/L

Sampling date: 7/17/13

Field blank type: (circle one) Field Blank / Rinsate / Other: EB Associated Samples: None 7 → 13 (ND)

Compound	Blank ID	Sample Identification							
	<u>1</u>	<u>5x/10x</u>							
<u>XX</u>	<u>0.24</u>	<u>2.4</u>							
<u>LL</u>	<u>0.24</u>	<u>2.4</u>							
<u>EEE</u>	<u>0.19</u>	<u>1.9</u>							
<u>W</u>	<u>0.014</u>	<u>0.070</u>							
<u>S</u>	<u>0.046</u>	<u>0.230</u>							

Blank units: ug/L Associated sample units: ug/L

Sampling date: 7/17/13

Field blank type: (circle one) Field Blank / Rinsate / Other: EB Associated Samples: 3 → 6 (ND + 75x, 10x)

Compound	Blank ID	Sample Identification							
	<u>2</u>	<u>5x/10x</u>							
<u>XX</u>	<u>0.14</u>	<u>1.4</u>							
<u>LL</u>	<u>0.22</u>	<u>2.2</u>							
<u>EEE</u>	<u>0.14</u>	<u>1.4</u>							
<u>W</u>	<u>0.014</u>	<u>0.070</u>							
<u>S</u>	<u>0.045</u>	<u>0.225</u>							

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:

Common contaminants such as the phthalates and TICs noted above that were detected in samples within ten times the associated field blank concentration were qualified as not detected, "U". Other contaminants within five times the field blank concentration were also qualified as not detected, "U".

LDC #: 30289D26

VALIDATION FINDINGS WORKSHEET

Field Blanks

Reviewer: FT
2nd Reviewer: [Signature]

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

Y N N/A Were field blanks identified in this SDG?
Y N N/A Were target compounds detected in the field blanks?

FB = FB = 041113
SDG # PH029

Blank units: ug/L Associated sample units: ug/kg

Sampling date: 4/11/13

Field blank type: (circle one) Field Blank / Rinsate / Other: FB Associated Samples: 3 → 13 Code = F

Compound	Blank ID	Sample Identification							
	<u>FB</u>	<u>5X/10X</u>	<u>3</u>	<u>4</u>					
<u>XX</u>	<u>0.17</u>	<u>1.7</u>							
<u>LL</u>	<u>0.18</u>	<u>1.8</u>							
<u>EEE</u>	<u>0.082</u>	<u>0.82</u>							
<u>TTT</u>	<u>0.019</u>	<u>0.095</u>							
<u>W</u>	<u>0.024</u>	<u>0.12</u>							
<u>S</u>	<u>0.17</u>	<u>0.85</u>	<u>0.78/1.7u</u>	<u>0.72/1.7u</u>					

Blank units: _____ Associated sample units: _____

Sampling date: _____

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: _____

Compound	Blank ID	Sample Identification							

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Common contaminants such as the phthalates and TICs noted above that were detected in samples within ten times the associated field blank concentration were qualified as not detected, "U". Other contaminants within five times the field blank concentration were also qualified as not detected, "U".

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$RRF = (A_x)(C_{is}) / (A_{is})(C_x)$
 average RRF = sum of the RRFs/number of standards
 %RSD = 100 * (S/X)

A_x = Area of compound,
 C_x = Concentration of compound,
 S = Standard deviation of the RRFs,
 A_{is} = Area of associated internal standard
 C_{is} = Concentration of internal standard
 X = Mean of the RRFs

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				RRF (0.5 std)	RRF (0.5 std)	Average RRF (initial)	Average RRF (initial)	%RSD	%RSD
1	ICAL	7/29/13	NDMA (1st internal standard)	0.918	0.918	0.877	0.877	8	8
			Naphthalene (2nd internal standard)	1.059	1.059	1.078	1.078	4	4
			Fluorene (3rd internal standard)	1.349	1.349	1.356	1.356	3	3
			Anthracene (4th internal standard)	1.180	1.180	1.186	1.186	4	4
			Pyrene (5th internal standard)	1.459	1.459	1.478	1.478	2	2
			Benzo(a)pyrene (6th internal standard)	1.215	1.215	1.178	1.178	2	2
2	ICAL	6/24/13		0.829	0.829	0.839	0.839	3	3
				1.050	1.050	1.046	1.046	3	3
				1.419	1.419	1.396	1.396	3	3
				1.255	1.255	1.239	1.239	4	4
				1.695	1.695	1.665	1.665	3	3
				1.192	1.192	1.167	1.167	3	3
3									

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET

Continuing Calibration Results Verification

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

$$\% \text{ Difference} = 100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$$

$$\text{RRF} = (A_x)(C_{is}) / (A_{is})(C_x)$$

Where: ave. RRF = initial calibration average RRF
 RRF = continuing calibration RRF
 A_x = Area of compound,
 C_x = Concentration of compound,

A_{is} = Area of associated internal standard
 C_{is} = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Average RRF (initial)	Reported	Recalculated	Reported	Recalculated
					RRF (CC)	RRF (CC)	%D	%D
1	ccv 3:48	7/23/13	NDMA (1st internal standard)	0.839	0.722	0.722	14	14
			Naphthalene (2nd internal standard)	1.046	1.051	1.051	0	0
			Fluorene (3rd internal standard)	1.396	1.394	1.394	0	0
			Anthracene (4th internal standard)	1.239	1.252	1.252	1	1
			Pyrene (5th internal standard)	1.665	1.812	1.812	9	9
			Benzo(a)pyrene (6th internal standard)	1.167	1.234	1.234	6	6
2	ccv 04:09	7/24/13	(1st internal standard)		0.684	0.684	18	18
			(2nd internal standard)		1.051	1.051	1	1
			(3rd internal standard)		1.383	1.383	1	1
			(4th internal standard)		1.253	1.253	1	1
			(5th internal standard)		1.796	1.796	8	8
			(6th internal standard)		1.251	1.251	7	7
3	ccv 3:14	7/25/13	157 (1st internal standard)		0.705	0.705	16	16
			(2nd internal standard)		1.021	1.021	2	2
			(3rd internal standard)		1.393	1.393	0	0
			(4th internal standard)		1.226	1.226	1	1
			(5th internal standard)		1.799	1.799	8	8
			(6th internal standard)		1.218	1.218	4	4

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET Continuing Calibration Results Verification

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

% Difference = $100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$
 $\text{RRF} = (A_x)(C_{is}) / (A_{is})(C_x)$

Where: ave. RRF = initial calibration average RRF
 RRF = continuing calibration RRF
 A_x = Area of compound, A_{is} = Area of associated internal standard
 C_x = Concentration of compound, C_{is} = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Average RRF (initial)	Reported	Recalculated	Reported	Recalculated
					RRF (CC)	RRF (CC)	%D	%D
1	001041X	7/27/13	NDMA (1st internal standard)	0.839	0.696	0.696	17	17
			Naphthalene (2nd internal standard)	1.046	1.040	1.040	1	1
			Fluorene (3rd internal standard)	1.396	1.388	1.388	1	1
			Anthracene (4th internal standard)	1.239	1.247	1.247	1	1
			Pyrene (5th internal standard)	1.665	1.758	1.758	6	6
			Benz(a)pyrene (6th internal standard)	1.167	1.221	1.221	5	5
2			(1st internal standard)					
			(2nd internal standard)					
			(3rd internal standard)					
			(4th internal standard)					
			(5th internal standard)					
			(6th internal standard)					
3			(2nd internal standard)					
			(3rd internal standard)					
			(4th internal standard)					
			(5th internal standard)					
			(6th internal standard)					

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

METHOD: GC/MS Semivolatiles (EPA SW 846 Method 8270C)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
 SS = Surrogate Spiked

Sample ID: # 3

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5 <i>Fluoranthene d10</i>	1.0	0.703	70	70	0
2-Fluorobiphenyl <i>Benzo(a) pyrene -d12</i>	↓	0.718	72	72	↓
Terphenyl-d14 <i>1-methyl naphthalene</i>	↓	0.858	86	86	↓
Phenol-d5					
2-Fluorophenol					
2,4,6-Tribromophenol					
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

Sample ID:

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5					
2-Fluorobiphenyl					
Terphenyl-d14					
Phenol-d5					
2-Fluorophenol					
2,4,6-Tribromophenol					
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

Sample ID:

	Surrogate Spiked	Surrogate Found	Percent Recovery Reported	Percent Recovery Recalculated	Percent Difference
Nitrobenzene-d5					
2-Fluorobiphenyl					
Terphenyl-d14					
Phenol-d5					
2-Fluorophenol					
2,4,6-Tribromophenol					
2-Chlorophenol-d4					
1,2-Dichlorobenzene-d4					

VALIDATION FINDINGS WORKSHEET
Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

% Recovery = 100 * (SC/SA)

Where: SSC = Spike concentration
 SA = Spike added

RPD = | LCSC - LCSDC | * 2 / (LCSC + LCSDC)

LCSC = Laboratory control sample concentration LCSDC = Laboratory control sample duplicate concentration

LCS/LCSD samples: 205 WBLCS / LCSD

Compound	Spike Added (ug/L)		Spike Concentration (ug/L)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD	LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
					Reported	Recalc	Reported	Recalc	Reported	Recalculated
Phenol										
N-Nitroso-di-n-propylamine										
4-Chloro-3-methylphenol										
Acenaphthene	1	1	1.0	1.01	100	100	101	101	0	0
Pentachlorophenol										
Pyrene	1	1	1.01	1.01	101	100	101	101	1	1

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET

Sample Calculation Verification

METHOD: GC/MS BNA (EPA SW 846 Method 8270C)

Y N N/A
Y N N/A

Were all reported results recalculated and verified for all level IV samples?

Were all recalculated results for detected target compounds agree within 10.0% of the reported results?

$$\text{Concentration} = \frac{(A_x)(I_s)(V_i)(DF)(2.0)}{(A_{is})(RRF)(V_o)(V_f)(\%S)}$$

- A_x = Area of the characteristic ion (EICP) for the compound to be measured
- A_{is} = Area of the characteristic ion (EICP) for the specific internal standard
- I_s = Amount of internal standard added in nanograms (ng)
- V_o = Volume or weight of sample extract in milliliters (ml) or grams (g).
- V_i = Volume of extract injected in microliters (ul)
- V_f = Volume of the concentrated extract in microliters (ul)
- Df = Dilution Factor.
- %S = Percent solids, applicable to soil and solid matrices only.
- 2.0 = Factor of 2 to account for GPC cleanup

Example:

Sample I.D. #1, 9:

$$\text{Conc.} = \frac{(11603)(0.25)(1000)}{29024(1.059)() () ()}$$

= 0.046585 ug/L

#	Sample ID	Compound	Reported Concentration ()	Calculated Concentration ()	Qualification

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 23, 2013
Matrix: Soil/Water
Parameters: Chlorinated Pesticides
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713
EB2-071713
SL-514-SA8-SB-0.0-0.5
SL-514-SA8-SB-4.0-5.0
SL-514-SA8-SB-7.5-8.5

Introduction

This data review covers 3 soil samples and 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8081A for Chlorinated Pesticides.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/ECD Instrument Performance Check

Instrument performance was acceptable unless noted otherwise under initial calibration and continuing calibration sections.

III. Initial Calibration

Initial calibration of single compounds was performed for the primary (quantitation) column and confirmation column as required by this method.

Percent relative standard deviations (%RSD) were less than or equal to 20.0% for all compounds.

Retention time windows were evaluated and considered technically acceptable.

IV. Continuing Calibration

Continuing calibration was performed at required frequencies.

The percent differences (%D) of calibration factors in continuing standard mixtures were within the 20.0% QC limits.

The percent differences (%D) of the second source calibration standard were less than or equal to 20.0% for all compounds.

Retention times (RT) of all compounds in the calibration standards were within QC limits.

The individual 4,4'-DDT and Endrin breakdowns (%BD) were less than or equal to 15.0%.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No chlorinated pesticide contaminants were found in the method blanks.

Samples EB1-071713 and EB2-071713 were identified as equipment blanks. No chlorinated pesticide contaminants were found.

Sample FB-041113 (from SDG PH029) were identified a field blank. No chlorinated pesticide contaminants were found.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

LCS ID (Associated Samples)	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Flag	A or P
LCS14200 (All water samples in SDG PH069)	Endrin	-	-	35 (≤30)	J (all detects) UJ (all non-detects)	P

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Florisil Cartridge Check

Florisil cleanup was not required and therefore not performed in this SDG.

XI. GPC Calibration

GPC cleanup was not required and therefore not performed in this SDG.

XII. Target Compound Identification

All target compound identifications were within validation criteria.

XIII. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported below the RL.	J (all detects)	A

XIV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XV. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Chlorinated Pesticides - Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH069	EB1-071713 EB2-071713	Endrin	J (all detects) UJ (all non-detects)	P	Laboratory control sample (RPD) (E)
PH069	EB1-071713 EB2-071713 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Chlorinated Pesticides - Laboratory Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Chlorinated Pesticides - Field Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

METHOD: GC Chlorinated Pesticides (EPA SW846 Method 8081A)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	Δ	Sampling dates: 7/17/13
II.	GC/ECD Instrument Performance Check	Δ	
III.	Initial calibration	Δ	% RSD ≤ 20
IV.	Continuing calibration/ICV	A	ICV/CCV ≤ 20
V.	Blanks	Δ	
VI.	Surrogate spikes	Δ	
VII.	Matrix spike/Matrix spike duplicates	N	client specified
VIII.	Laboratory control samples	SW	res ID
IX.	Regional quality assurance and quality control	N	
X.	Florisil cartridge check	N	
XI.	GPC Calibration	N	
XII.	Target compound identification	A	
XIII.	Compound quantitation/RL/LOQ/LODs	Δ	
XIV.	Overall assessment of data	Δ	
XV.	Field duplicates	Δ	
XVI.	Field blanks	ND	EB = 1, 2 FB = FB-04113

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

D = Duplicate
TB = Trip blank
EB = Equipment blank

SDG # PH029

Validated Samples:

soil + water

12	EB1-071713	w	111	PBLK11201	21		31
22	EB2-071713	w	122	PBLK14200	22		32
31	SL-514-SA8-SB-0.0-0.5		13		23		33
41	SL-514-SA8-SB-4.0-5.0		14		24		34
51	SL-514-SA8-SB-7.5-8.5		15		25		35
6			16		26		36
7			17		27		37
8			18		28		38
9			19		29		39
10			20		30		40

Notes: _____

Method: Pesticides/PCBs (EPA SW 846 Method 8081A/8082)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. GC/ECD Instrument performance check				
Was the instrument performance found to be acceptable?	/			
III. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	/			
Was a linear fit used for evaluation? If yes, were all percent relative standard deviations (%RSD) \leq 20%?	/			
Was a curve fit used for evaluation? If Yes, what was the acceptance criteria used?		/		
Did the initial calibration meet the curve fit acceptance criteria?			/	
Were the RT windows properly established?	/			
Were the required standard concentrations analyzed in the initial calibration?	/			
IV. Continuing calibration				
What type of continuing calibration calculation was performed? ____%D or ____%R	/			
Were Evaluation mix standards analyzed prior to the initial calibration and sample analysis?	/			
Were endrin and 4,4'-DDT breakdowns \leq 15% for individual breakdown in the Evaluation mix standards?	/			
Was a continuing calibration analyzed daily?	/			
Were all percent differences (%D) \leq 20%?	/			
Were all the retention times within the acceptance windows?	/			
V. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was a method blank analyzed for each matrix and concentration?	/			
Were extract cleanup blanks analyzed with every batch requiring clean-up?		/		
Was there contamination in the method blanks or clean-up blanks? If yes, please see the Blanks validation completeness worksheet.	/			
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?	/			
If the percent recovery (%R) of one or more surrogates was outside QC limits, was a reanalysis performed to confirm %R?			/	
If any %R was less than 10 percent, was a reanalysis performed to confirm %R?			/	

Validation Area	Yes	No	NA	Findings/Comments
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.			/	
Was a MS/MSD analyzed every 20 samples of each matrix?			/	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	/			
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per extraction batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	/			
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?			/	
Were the performance evaluation (PE) samples within the acceptance limits?			/	
X. Target compound identification				
Were the retention times of reported detects within the RT windows?	/			
XI. Compound quantitation/CRQLs				
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions, dry weight factors, and clean-up activities applicable to level IV validation?	/			
XII. System performance				
System performance was found to be acceptable.	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.		/		

VALIDATION FINDINGS WORKSHEET

METHOD: Pesticide/PCBs (EPASW 846 Method 8081/8082)

A. alpha-BHC	I. Dieldrin	Q. Endrin ketone	Y. Aroclor-1242	GG.
B. beta-BHC	J. 4,4'-DDE	R. Endrin aldehyde	Z. Aroclor-1248	HH.
C. delta-BHC	K. Endrin	S. alpha-Chlordane	AA. Aroclor-1254	II.
D. gamma-BHC	L. Endosulfan II	T. gamma-Chlordane	BB. Aroclor-1260	JJ.
E. Heptachlor	M. 4,4'-DDD	U. Toxaphene	CC. DB 608	KK.
F. Aldrin	N. Endosulfan sulfate	V. Aroclor-1016	DD. DB 1701	LL.
G. Heptachlor epoxide	O. 4,4'-DDT	W. Aroclor-1221	EE.	MM.
H. Endosulfan I	P. Methoxychlor	X. Aroclor-1232	FF.	NN.

Notes: _____

VALIDATION FINDINGS WORKSHEET
Laboratory Control Samples (LCS)

METHOD: GC HPLC

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".
 N N/A Were a laboratory control samples (LCS) and laboratory control sample duplicate (LCSD) analyzed for each matrix in this SDG?
 N N/A Were the LCS percent recoveries (%R) and relative percent differences (RPD) within the QC limits?

Level IV/D Only
 N N/A Was an LCS analyzed every 20 samples for each matrix or whenever a sample extraction was performed?

#	LCS/LCSD ID	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Associated Samples	Qualifications
	LC514200	K	()	()	35 (30)	All water	J/W/P (E)
			()	()	()		
			()	()	()		
			()	()	()		
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			()	()	()		

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

METHOD: GC HPLC

The calibration factors (CF) and relative standard deviation (%RSD) were recalculated using the following calculations:

CF = A/C
 Average CF = sum of the CF/number of standards
 %RSD = 100 * (S/X)

Where: A = Area of compound
 C = Concentration of compound
 S = Standard deviation of calibration factors
 X = Mean of calibration factors

#	Standard ID	Calibration Date	Compound	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				CF/100 (10/std)	CF/100 (10/std)	CF (initial)	CF (initial)	%RSD	%RSD
1	ICAL	7/15/13	endosulfan 1	2.65×10^5	2.65×10^5	2.87×10^5	2.87×10^5	18	18
	RTXCLP		methoxychlor	1.04×10^5	1.04×10^5	1.05×10^5	1.05×10^5	9	9
2	RTXCLP II		↓	6.63×10^4	6.63×10^4	6.73×10^4	6.73×10^4	5	5
				3.51×10^4	3.51×10^4	3.50×10^4	3.50×10^4	5	5
3	ICAL	7/15/13	↓	2.71×10^5	2.71×10^5	2.84×10^5	2.84×10^5	14	14
	RTXCLP			1.03×10^5	1.03×10^5	1.06×10^5	1.06×10^5	11	11
4	RTXCLP II		↓	7.18×10^4	7.18×10^4	7.33×10^4	7.33×10^4	4	4
				3.49×10^4	3.49×10^4	3.50×10^4	3.50×10^4	5	5

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30289D39

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
 Reviewer: F7
 2nd Reviewer: ←

METHOD: GC HPLC

The percent difference (%D) of the initial calibration average Calibration Factors (CF) and the continuing calibration CF were recalculated for the compounds identified below using the following calculation:

% Difference = 100 * (ave. CF - CF)/ave.CF

Where: ave. CF = initial calibration average CF
 CF = continuing calibration CF
 A = Area of compound
 C = Concentration of compound

#	Standard ID	Calibration Date	Compound	Average CF(Ical)/ CCV Conc.	Reported	Recalculated	Reported	Recalculated
					CF/ Conc. CCV	CF/ Conc. CCV	%D	%D
1	cen 13:16	7/23/13	endosulfan /	10.00	9.96	9.96	0	0
	RTX cup		methoxychlor	100.00	104.41	104.41	4	4
2	RTX cup 2		↓	10.00	10.54	10.54	5	5
				100.00	104.83	104.83	5	5
3	cen 18:14	7/23/13	↓	↓	9.52	9.52	5	5
	RTX cup				98.33	98.33	2	2
4			↓	↓	9.74	9.74	3	3
					100.50	100.50	1	1

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30289D3g

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

Page: 1 of 1
Reviewer: F7
2nd reviewer: [Signature]

METHOD: GC Pesticides (EPA SW 846 Method 8081)

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
SS = Surrogate Spiked

Sample ID: 3

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
TCMX	RTX CLP2	10	8.717306	87	87	0
PCB	↓	↓	10.136194	100	100	0

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

LDC #: 30289D35

VALIDATION FINDINGS WORKSHEET
Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Page: 1 of 1
 Reviewer: F7
 2nd Reviewer: [Signature]

METHOD: GC Pesticides (EPA SW 846 method 8081)

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

$\% \text{Recovery} = 100 * (\text{SSC} - \text{SC}) / \text{SA}$

Where SSC = Spiked sample concentration
 SA = Spike added
 LCS = Laboratory Control Sample

SC = Sample concentration

$\text{RPD} = (((\text{SSCLCS} - \text{SSCLCSD}) * 2) / (\text{SSCLCS} + \text{SSCLCSD})) * 100$

LCSD = Laboratory Control Sample duplicate

LCS/LCSD samples: LCS 14200

Compound	Spike Added (ug/L)		Sample Conc. (ug/L)	Spike Sample Concentration (ug/L)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD		LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
						Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gamma-BHC	0.1	0.1	ND	0.096656	0.097779	95	95	96	96	1	1
4,4'-DDT											

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

METHOD: GC Pesticides (EPA SW 846 Method 8081)

(Y) N N/A
(Y) N N/A

Were all reported results recalculated and verified for all level IV samples?
 Were all recalculated results for detected target compounds within 10% of the reported results?

Concentration = $\frac{(A)(Fv)(Df)}{(RF)(Vs \text{ or } Ws)(\%S/100)}$

Example:

Sample ID. 3 Compound Name Beta BHC

- A= Area or height of the compound to be measured
- Fv= Final Volume of extract
- Df= Dilution Factor
- RF= Average response factor of the compound
In the initial calibration
- Vs= Initial volume of the sample
- Ws= Initial weight of the sample
- %S= Percent Solid

Concentration = $\frac{(1022988)(10)}{(1.77 \times 10^5)(30.1)(0.947)}$

2.027 ug/kg

#	Sample ID	Compound	Reported Concentrations ()	Recalculated Results Concentrations ()	Qualifications

Comments: _____

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 23, 2013
Matrix: Soil/Water
Parameters: Polychlorinated Biphenyls
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713
EB2-071713
SL-512-SA8-SB-0.0-0.5
SL-515-SA8-SB-0.0-0.5
SL-515-SA8-SB-4.0-5.0
SL-515-SA8-SB-9.0-10.0
SL-515-SA8-SB-14.0-15.0
SL-513-SA8-SB-0.0-0.5
SL-513-SA8-SB-4.0-5.0
SL-513-SA8-SB-8.0-9.0

Introduction

This data review covers 8 soil samples and 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8082A for Polychlorinated Biphenyls.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. GC/ECD Instrument Performance Check

Instrument performance was acceptable unless noted otherwise under initial calibration and continuing calibration sections.

III. Initial Calibration

Initial calibration of multicomponent compounds was performed for the primary (quantitation) column as required by the method.

The percent relative standard deviations (%RSD) were less than or equal to 20.0% for all compounds.

Retention time windows were evaluated and considered technically acceptable.

IV. Continuing Calibration

Continuing calibration was performed at required frequencies.

The percent differences (%D) of calibration factors in continuing standard mixtures were within the 20.0% QC limits.

The percent difference (%D) of the second source calibration standard were less than or equal to 20.0% for all compounds.

Retention times (RT) of all compounds in the calibration standards were within QC limits.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No polychlorinated biphenyl contaminants were found in the method blanks.

Samples EB1-071713 and EB2-071713 were identified as equipment blanks. No polychlorinated biphenyl contaminants were found.

Sample FB-041113 (from SDG PH029) was identified as a field blank. No polychlorinated biphenyl contaminants were found.

VI. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VII. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

IX. Regional Quality Assurance and Quality Control

Not applicable.

X. Florisil Cartridge Check

Florisil cleanup was not required and therefore not performed in this SDG.

XI. GPC Calibration

GPC cleanup was not required and therefore not performed in this SDG.

XII. Target Compound Identification

All target compound identifications were within validation criteria.

XIII. Compound Quantitation

All compound quantitations were within validation criteria.

The sample results for detected compounds from the two columns were within 40% relative percent difference (RPD) with the following exceptions:

Sample	Compound	RPD	Flag	A or P
SL-512-SA8-SB-0.0-0.5	Aroclor-1254	51	J (all detects)	A

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported below the RL.	J (all detects)	A

XIV. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XV. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Polychlorinated Biphenyls - Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH069	SL-512-SA8-SB-0.0-0.5	Aroclor-1254	J (all detects)	A	Compound quantitation (RPD between columns) (*XIII)
PH069	EB1-071713 EB2-071713 SL-512-SA8-SB-0.0-0.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Polychlorinated Biphenyls - Laboratory Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Polychlorinated Biphenyls - Field Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

METHOD: GC Polychlorinated Biphenyls (EPA SW 846 Method 8082A)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	Δ	Sampling dates: 7/17/13
II.	GC/ECD Instrument Performance Check	NA	
III.	Initial calibration	Δ	% PSD ≤ 20
IV.	Continuing calibration/ICV	Δ	ICV/CCV ≤ 20
V.	Blanks	Δ	
VI.	Surrogate spikes	Δ	
VII.	Matrix spike/Matrix spike duplicates	N	client specified
VIII.	Laboratory control samples	Δ	LCS 10
IX.	Regional quality assurance and quality control	N	
X.	Florisil cartridge check	N	
XI.	GPC Calibration	N	
XII.	Target compound identification	Δ	
XIII.	Compound quantitation/RL/LOQ/LODs	sw	
XIV.	Overall assessment of data	Δ	
XV.	Field duplicates	N	
XVI.	Field blanks	ND	EB = 1, 2 FB = FB-041113 (SDG # PH029)

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples: soil + water

1	EB1-071713	w	11	PBLK12201	21	31
2	EB2-071713	w	12	PBLK13200	22	32
3	SL-512-SA8-SB-0.0-0.5		13		23	33
4	SL-515-SA8-SB-0.0-0.5		14		24	34
5	SL-515-SA8-SB-4.0-5.0		15		25	35
6	SL-515-SA8-SB-9.0-10.0		16		26	36
7	SL-515-SA8-SB-14.0-15.0		17		27	37
8	SL-513-SA8-SB-0.0-0.5		18		28	38
9	SL-513-SA8-SB-4.0-5.0		19		29	39
10	SL-513-SA8-SB-8.0-9.0		20		30	40

Notes:

Method: GC HPLC

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
II. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) < 20%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a curve fit used for evaluation?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Did the initial calibration meet the curve fit acceptance criteria of > 0.990?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the RT windows properly established?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IV. Continuing calibration				
Was a continuing calibration analyzed daily?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) < 20%.0 or percent recoveries 80-120%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all the retention times within the acceptance windows?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
V. Blanks				
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank analyzed for each matrix and concentration?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
If the percent recovery (%R) for one or more surrogates was out of QC limits, was a reanalysis performed to confirm samples with %R outside of criteria?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	

LDC #: 30289P3b

VALIDATION FINDINGS CHECKLIST

Page: 2 of 2
 Reviewer: FT
 2nd Reviewer: [Signature]

Validation Area	Yes	No	NA	Findings/Comments
X. Target compound identification				
Were the retention times of reported detects within the RT windows?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XI. Compound quantitation/CRQLs				
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XII. System performance				
System performance was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field duplicates.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
XV. Field blanks				
Field blanks were identified in this SDG.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Target compounds were detected in the field blanks.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	

VALIDATION FINDINGS WORKSHEET

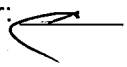
METHOD: Pesticide/PCBs (EPASW 846 Method 8081/8082)

A. alpha-BHC	I. Dieldrin	Q. Endrin ketone	Y. Aroclor-1242	GG.
B. beta-BHC	J. 4,4'-DDE	R. Endrin aldehyde	Z. Aroclor-1248	HH.
C. delta-BHC	K. Endrin	S. alpha-Chlordane	AA. Aroclor-1254	II.
D. gamma-BHC	L. Endosulfan II	T. gamma-Chlordane	BB. Aroclor-1260	JJ.
E. Heptachlor	M. 4,4'-DDD	U. Toxaphene	CC. DB 608	KK.
F. Aldrin	N. Endosulfan sulfate	V. Aroclor-1016	DD. DB 1701	LL.
G. Heptachlor epoxide	O. 4,4'-DDT	W. Aroclor-1221	EE.	MM.
H. Endosulfan I	P. Methoxychlor	X. Aroclor-1232	FF.	NN.

Notes: _____

LDC #: 30289 D36

VALIDATION FINDINGS WORKSHEET
Compound Quantitation and Reported CRQLs

Page: 1 of 1
 Reviewer: FT
 2nd Reviewer: 

METHOD: GC HPLC

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Level IV/D Only

- Y N N/A Were CRQLs adjusted for sample dilutions, dry weight factors, etc.?
- Y N N/A Did the reported results for detected target compounds agree within 10.0% of the recalculated results?

#	Compound Name	% RPD Finding	Bot 2 col ≤ 40	Associated Samples	Qualifications
	AA	51		3	J/Adt * X111

Comments: See sample calculation verification worksheet for recalculations

LDC #: 30289D3b

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: F7
 2nd Reviewer: [Signature]

METHOD: GC ✓ HPLC

The calibration factors (CF) and relative standard deviation (%RSD) were recalculated using the following calculations:

CF = A/C
 Average CF = sum of the CF/number of standards
 %RSD = 100 * (S/X)

Where: A = Area of compound
 C = Concentration of compound
 S = Standard deviation of calibration factors
 X = Mean of calibration factors

#	Standard ID	Calibration Date	Compound	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				CF (200 std)	CF (200 std)	Ave CF (initial)	Ave CF (initial)	%RSD	%RSD
1	KAL	7/9/13 (1T20190)	Aroclor 1260-1	18234	18234	18917	18917	11	11
	ZB Multi R1								
	ZB Multi R2	7/9/13	↓	41696	41696	41997	41997	7	7
2									
3	KAL	7/24/13 1T20205	↓	19814	19814	20206	20206	9	9
	- R1								
	R2		↓	60265	60265	62522	62522	7	7
4									

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30289036

VALIDATION FINDINGS WORKSHEET Continuing Calibration Results Verification

Page: 1 of 1
Reviewer: FJ
2nd Reviewer: A

METHOD: GC HPLC

The percent difference (%D) of the initial calibration average Calibration Factors (CF) and the continuing calibration CF were recalculated for the compounds identified below using the following calculation:

% Difference = 100 * (ave. CF - CF)/ave.CF

Where: ave. CF = initial calibration average CF
CF = continuing calibration CF
A = Area of compound
C = Concentration of compound

#	Standard ID	Calibration Date	Compound	Average CF(Ical)/ CCV Conc.	Reported	Recalculated	Reported	Recalculated
					CF/ Conc. CCV	CF/ Conc. CCV	%D	%D
1	ceV 23:5D	7/21/13	Aroclor 1260	200.0	189.70	189.70	5	5
	ZB multi R1							
	R2		↓	200.0	188.38	188.38	6	6
2	ceV 0:16	7/25/13	↓	200.0	189.55	189.55	5	5
	ZB multi R1							
	R2		↓	200.0	193.55	193.55	3	3
3	ceV 7:39	7/25/13	↓	200.0	189.24	189.24	5	5
	R1							
	R2		↓	200.0	201.83	201.83	1	1
4								

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

METHOD: GC HPLC

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: $SF/SS * 100$

Where: SF = Surrogate Found
 SS = Surrogate Spiked

Sample ID: 3

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
TCMX	ZB-R2	10	11.209	112	112	0
DCB	ZB-R1	10	9.785	97	98	1

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

LDC #: 30289036

VALIDATION FINDINGS WORKSHEET

Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Reviewer: FJ

2nd Reviewer: [Signature]

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

%Recovery = 100 * (SSC - SC)/SA

Where SSC = Spiked sample concentration
SA = Spike added
LCS = Laboratory Control Sample

SC = Sample concentration

RPD = ((SSCLCS - SSCLCSD) * 2) / (SSCLCS + SSCLCSD) * 100

LCSD = Laboratory Control Sample duplicate

LCS/LCSD samples: LC513200

Compound	Spike Added (ug/L)		Sample Conc. (ug/L)	Spike Sample Concentration (ug/L)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD		LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
						Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)											
Diesel (8015)											
Benzene (8021B)											
Methane (RSK-175)											
2,4-D (8151)											
Dinoseb (8151)											
Naphthalene (8310)											
Anthracene (8310)											
HMX (8330)											
2,4,6-Trinitrotoluene (8330)											
PcB 1260	5	5	ND	4.11	5.09	82	82	102	102	21	21

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

METHOD: GC HPLC

Y N/N/A
Y N/N/A

Were all reported results recalculated and verified for all level IV samples?
 Were all recalculated results for detected target compounds within 10% of the reported results?

Concentration = $\frac{(A)(Fv)(Df)}{(RF)(Vs \text{ or } Ws)(\%S/100)}$

Example:

Sample ID: # 3 Compound Name 1254

- A= Area or height of the compound to be measured
- Fv= Final Volume of extract
- Df= Dilution Factor
- RF= Average response factor of the compound in the initial calibration
- Vs= Initial volume of the sample
- Ws= Initial weight of the sample
- %S= Percent Solid

Concentration = $\frac{8.06144}{0.985}$

= 8.2 ug/kg

#	Sample ID	Compound	Reported Concentrations ()	Recalculated Results Concentrations ()	Qualifications
	1254-2 =	47625	$\frac{1.1562 (10)}{44398 (30.1)} = 3.5637$	1254-2 = 3.5637	
				3 = 4.215194	
				4 = 11.947963	
				5 = 4.629203	
				6 = 15.951103	
				Ave = 8.06143	

Comments: _____

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 23, 2013
Matrix: Soil/Water
Parameters: Metals
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PG069

Sample Identification

EB1-071713
EB2-071713
SL-512-SA8-SB-0.0-0.5
SL-514-SA8-SB-0.0-0.5
SL-514-SA8-SB-4.0-5.0
SL-514-SA8-SB-7.5-8.5
SL-515-SA8-SB-0.0-0.5
SL-515-SA8-SB-4.0-5.0
SL-515-SA8-SB-9.0-10.0
SL-515-SA8-SB-14.0-15.0
SL-513-SA8-SB-0.0-0.5
SL-513-SA8-SB-4.0-5.0
SL-513-SA8-SB-8.0-9.0
SL-512-SA8-SB-0.0-0.5MS
SL-512-SA8-SB-0.0-0.5MSD
SL-512-SA8-SB-0.0-0.5DUP
SL-514-SA8-SB-7.5-8.5MS
SL-514-SA8-SB-7.5-8.5MSD
SL-514-SA8-SB-7.5-8.5DUP

Introduction

This data review covers 17 soil samples and 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Methods 6010C, 6020A, 7470A, and 7471B for Metals. The metals analyzed were Aluminum, Antimony, Arsenic, Barium, Beryllium, Boron, Cadmium, Calcium, Chromium, Cobalt, Copper, Iron, Lead, Lithium, Magnesium, Manganese, Mercury, Molybdenum, Nickel, Phosphorus, Potassium, Selenium, Silver, Sodium, Strontium, Thallium, Tin, Titanium, Vanadium, Zinc, and Zirconium.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review (January 2010).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. ICPMS Tune

The mass calibration was within 0.1 AMU and the percent relative standard deviation (%RSD) was less than or equal to 5%.

III. Calibration

The initial and continuing calibrations were performed at the required frequency.

The calibration standards criteria were met.

IV. Blanks

Method blanks were reviewed for each matrix as applicable. No metal contaminants were found in the initial, continuing and preparation blanks with the following exceptions:

Method Blank ID	Analyte	Maximum Concentration	Associated Samples
ICB/CCB	Antimony Lithium Magnesium Phosphorus	5.6 ug/L 2.7 ug/L 38.1 ug/L 2.3 ug/L	All water samples in SDG PH069
PB (prep blank)	Tin	1.732 mg/Kg	All soil samples in SDG PH069
ICB/CCB	Copper Phosphorus	3.2 ug/L 4.0 ug/L	All soil samples in SDG PH069
ICB/CCB	Antimony	4.5 ug/L	SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-8.0-9.0
ICB/CCB	Antimony	4.2 ug/L	SL-513-SA8-SB-4.0-5.0
ICB/CCB	Calcium	33.5 ug/L	SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5

Method Blank ID	Analyte	Maximum Concentration	Associated Samples
ICB/CCB	Calcium	39.2 ug/L	SL-513-SA8-SB-0.0-0.5
ICB/CCB	Iron	41.8 ug/L	SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5
ICB/CCB	Iron	43.0 ug/L	SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5

Data qualification by the initial, continuing and preparation blanks (ICB/CCB/PBs) was based on the maximum contaminant concentration in the ICB/CCB/PBs in the analysis of each analyte. The sample concentrations were either not detected or were significantly greater (>5X blank contaminants) than the concentrations found in the associated method blanks with the following exceptions:

Sample	Analyte	Reported Concentration	Modified Final Concentration
EB1-071713	Antimony	6.9 ug/L	6.9U ug/L
EB2-071713	Antimony	5.4 ug/L	5.4U ug/L
SL-512-SA8-SB-0.0-0.5	Tin	3.0 mg/Kg	3.0U mg/Kg
SL-514-SA8-SB-0.0-0.5	Tin	3.3 mg/Kg	3.3U mg/Kg
SL-514-SA8-SB-4.0-5.0	Tin	3.1 mg/Kg	3.1U mg/Kg
SL-514-SA8-SB-7.5-8.5	Tin	3.0 mg/Kg	3.0U mg/Kg
SL-515-SA8-SB-0.0-0.5	Tin	3.4 mg/Kg	3.4U mg/Kg
SL-515-SA8-SB-4.0-5.0	Tin	3.4 mg/Kg	3.4U mg/Kg
SL-515-SA8-SB-9.0-10.0	Tin	3.8 mg/Kg	3.8U mg/Kg
SL-515-SA8-SB-14.0-15.0	Tin	3.3 mg/Kg	3.3U mg/Kg
SL-513-SA8-SB-0.0-0.5	Antimony Tin	1.2 mg/Kg 3.3 mg/Kg	1.2U mg/Kg 3.3U mg/Kg
SL-513-SA8-SB-4.0-5.0	Tin	2.8 mg/Kg	2.8U mg/Kg
SL-513-SA8-SB-8.0-9.0	Tin	3.1 mg/Kg	3.1U mg/Kg

Samples EB1-071713 and EB2-071713 were identified as equipment blanks. No metal contaminants were found with the following exceptions:

Blank ID	Sampling Date	Analyte	Concentration	Associated Samples
EB1-071713	7/17/13	Antimony	0.0069 mg/L	SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0
EB2-071713	7/17/13	Antimony	0.0054 mg/L	SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5

Sample FB-041113 (from SDG PH029) was identified as a field blank. No metal contaminants were found with the following exceptions:

Blank ID	Sampling Date	Analyte	Concentration	Associated Samples
FB-041113	4/11/13	Copper Molybdenum	0.0036 mg/L 0.0036 mg/L	All soil samples in SDG PG069

Sample concentrations were compared to concentrations detected in the field blanks. The sample concentrations were either not detected or were significantly greater (>5X blank contaminants) than the concentrations found in the associated field blanks with the following exceptions:

Sample	Analyte	Reported Concentration	Modified Final Concentration
SL-512-SA8-SB-0.0-0.5	Molybdenum	0.34 mg/Kg	0.34U mg/Kg
SL-513-SA8-SB-0.0-0.5	Antimony Molybdenum	1.2 mg/Kg 0.86 mg/Kg	1.2U mg/Kg 0.86U mg/Kg

V. ICP Interference Check Sample (ICS) Analysis

The frequency of analysis was met.

The criteria for analysis were met.

VI. Matrix Spike Analysis

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

Spike ID (Associated Samples)	Analyte	MS (%R) (Limits)	MSD (%R) (Limits)	RPD (Limits)	Flag	A or P
SL-514-SA8-SB-7.5-8.5MS/MSD (All soil samples in SDG PH069)	Antimony	74 (75-125)	73 (75-125)	-	J (all detects) UJ (all non-detects)	A
SL-514-SA8-SB-7.5-8.5MS/MSD (All soil samples in SDG PH069)	Potassium	177 (75-125)	162 (75-125)	-	J (all detects)	A

VII. Duplicate Sample Analysis

Duplicate (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

VIII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

IX. Internal Standards

All internal standard percent recoveries (%R) were within QC limits.

X. Furnace Atomic Absorption QC

Graphite furnace atomic absorption was not utilized in this SDG.

XI. ICP Serial Dilution

ICP serial dilution analysis was performed by the laboratory. The analysis criteria were met with the following exceptions:

Diluted Sample	Analyte	%D (Limits)	Associated Samples	Flag	A or P
SL-514-SA8-SB-7.5-8.5	Lithium	14 (≤10)	All soil samples in SDG PH069	J (all detects) UJ (all non-detects)	A

XII. Sample Result Verification

All sample result verifications were acceptable.

XIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Metals - Data Qualification Summary - SDG PG069**

SDG	Sample	Analyte	Flag	A or P	Reason (Code)
PH069	SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	Antimony	J (all detects) UJ (all non-detects)	A	Matrix spike/Matrix spike duplicate (%R) (Q)
PH069	SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	Potassium	J (all detects)	A	Matrix spike/Matrix spike duplicate (%R) (Q)
PH069	SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	Lithium	J (all detects) UJ (all non-detects)	A	ICP serial dilution (%D) (A)

**Santa Susana Field Laboratory
Metals - Laboratory Blank Data Qualification Summary - SDG PG069**

SDG	Sample	Analyte	Modified Final Concentration	A or P	Code
PH069	EB1-071713	Antimony	6.9U ug/L	A	B
PH069	EB2-071713	Antimony	5.4U ug/L	A	B
PH069	SL-512-SA8-SB-0.0-0.5	Tin	3.0U mg/Kg	A	B
PH069	SL-514-SA8-SB-0.0-0.5	Tin	3.3U mg/Kg	A	B
PH069	SL-514-SA8-SB-4.0-5.0	Tin	3.1U mg/Kg	A	B

SDG	Sample	Analyte	Modified Final Concentration	A or P	Code
PH069	SL-514-SA8-SB-7.5-8.5	Tin	3.0U mg/Kg	A	B
PH069	SL-515-SA8-SB-0.0-0.5	Tin	3.4U mg/Kg	A	B
PH069	SL-515-SA8-SB-4.0-5.0	Tin	3.4U mg/Kg	A	B
PH069	SL-515-SA8-SB-9.0-10.0	Tin	3.8U mg/Kg	A	B
PH069	SL-515-SA8-SB-14.0-15.0	Tin	3.3U mg/Kg	A	B
PH069	SL-513-SA8-SB-0.0-0.5	Antimony Tin	1.2U mg/Kg 3.3U mg/Kg	A	B
PH069	SL-513-SA8-SB-4.0-5.0	Tin	2.8U mg/Kg	A	B
PH069	SL-513-SA8-SB-8.0-9.0	Tin	3.1U mg/Kg	A	B

**Santa Susana Field Laboratory
Metals - Field Blank Data Qualification Summary - SDG PG069**

SDG	Sample	Analyte	Modified Final Concentration	A or P	Code
PH069	SL-512-SA8-SB-0.0-0.5	Molybdenum	0.34U mg/Kg	A	F
PH069	SL-513-SA8-SB-0.0-0.5	Antimony Molybdenum	1.2U mg/Kg 0.86U mg/Kg	A	F

LDC #: 30289D4

VALIDATION COMPLETENESS WORKSHEET

Date: 9/14/13

SDG #: PH069

Level IV

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

7470A / 7471B

Reviewer: [Signature]

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 7/17/13
II.	ICP/MS Tune	A	
III.	Calibration	A	
IV.	Blanks	SW	
V.	ICP Interference Check Sample (ICS) Analysis	A	
VI.	Matrix Spike Analysis	SW	ms/D
VII.	Duplicate Sample Analysis	A	Dup
VIII.	Laboratory Control Samples (LCS)	A	LCS/D
IX.	Internal Standard (ICP-MS)	A	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	SW	
XII.	Sample Result Verification	A	
XIII.	Overall Assessment of Data	A	
XIV.	Field Duplicates	N	
XV.	Field Blanks	SW	EB = 1, 2 FB = FB-041113 (PH029)

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

D = Duplicate
TB = Trip blank
EB = Equipment blank

Validated Samples:

soil/water

1	EB1-071713	w	11	SL-513-SA8-SB-0.0-0.5	21	31
2	EB2-071713	↓	12	SL-513-SA8-SB-4.0-5.0	22	32
3	SL-512-SA8-SB-0.0-0.5		13	SL-513-SA8-SB-8.0-9.0	23	33
4	SL-514-SA8-SB-0.0-0.5		14	(3) ms	24	34
5	SL-514-SA8-SB-4.0-5.0	↓	15	msD	25	35
6	SL-514-SA8-SB-7.5-8.5	↓	16	Dup	26	36
7	SL-515-SA8-SB-0.0-0.5		17	(6) ms	27	37
8	SL-515-SA8-SB-4.0-5.0	↓	18	msD	28	38
9	SL-515-SA8-SB-9.0-10.0	↓	19	Dup	29	39
10	SL-515-SA8-SB-14.0-15.0		20		30	40

Notes: _____

Method:Metals (EPA SW 846 Method 6010B/7000/6020)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. ICP/MS Tune				
Were all isotopes in the tuning solution mass resolution within 0.1 amu?	/			
Were %RSD of isotopes in the tuning solution $\leq 5\%$?	/			
III. Calibration				
Were all instruments calibrated daily, each set-up time?	/			
Were the proper number of standards used?	/			
Were all initial and continuing calibration verification %Rs within the 90-110% (80-120% for mercury) QC limits?	/			
Were all initial calibration correlation coefficients ≥ 0.995 ?	/			
IV. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	/			
V. ICP Interference Check Sample				
Were ICP interference check samples performed daily?	/			
Were the AB solution percent recoveries (%R) with the 80-120% QC limits?	/			
VI. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and duplicate (DUP) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD or MS/DUP. Soil / Water.	/			
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the 75-125 QC limits? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.		/		
Were the MS/MSD or duplicate relative percent differences (RPD) $\leq 20\%$ for waters and $\leq 35\%$ for soil samples? A control limit of $\pm RL$ ($\pm 2X RL$ for soil) was used for samples that were $\leq 5X$ the RL, including when only one of the duplicate sample values were $< 5X$ the RL.	/			
VII. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per extraction batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the 80-120% QC limits for water samples and laboratory established QC limits for soils?	/			

Validation Area	Yes	No	NA	Findings/Comments
VIII. Furnace Atomic Absorption QC				
If MSA was performed, was the correlation coefficients > 0.995?			/	
Do all applicable analyses have duplicate injections? (Level IV only)			/	
For sample concentrations > RL, are applicable duplicate injection RSD values < 20%? (Level IV only)			/	
Were analytical spike recoveries within the 85-115% QC limits?			/	
IX. ICP Serial Dilution				
Was an ICP serial dilution analyzed if analyte concentrations were > 50X the MDL (ICP)/>100X the MDL(ICP/MS)?	/			
Were all percent differences (%Ds) < 10%?	/			
Was there evidence of negative interference? If yes, professional judgement will be used to qualify the data.		/		
X. Internal Standards (EPA SW 846 Method 6020/EPA 200.8)				
Were all the percent recoveries (%R) within the 30-120% (6020)/60-125% (200.8) of the intensity of the internal standard in the associated initial calibration?	/			
If the %Rs were outside the criteria, was a reanalysis performed?			/	
XI. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?		/		
Were the performance evaluation (PE) samples within the acceptance limits?			/	
XII. Sample Result Verification				
Were RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target analytes were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target analytes were detected in the field blanks.	/			

VALIDATION FINDINGS WORKSHEET
PB/ICB/CCB QUALIFIED SAMPLES

METHOD: Trace metals (EPA SW 864 Method 6010B/6020/7000)

Soil preparation factor applied: 100x Reason: B

Sample Concentration units, unless otherwise noted: ug/L

Associated Samples: All Water

					Sample Identification										
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/L)	Maximum ICB/CCB ^a (ug/L)	Action Level	1	2									
Sb			5.6	28	6.9	5.4									
Li			2.7	13.5											
Mg			38.1	190.5											
P			2.3	11.5											

Sample Concentration units, unless otherwise noted: mg/Kg

Associated Samples: All Soil

					Sample Identification										
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/L)	Maximum ICB/CCB ^a (ug/L)	Action Level	3	4	5	6	7	8	9	10	11	12	13
Cu			3.2	1.6											
P			4.0	2											
Sn	1.732			8.66	3.0	3.3	3.1	3.0	3.4	3.4	3.8	3.3	3.3	2.8	3.1

Sample Concentration units, unless otherwise noted: mg/Kg

Associated Samples: 3-11, 13

					Sample Identification										
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/L)	Maximum ICB/CCB ^a (ug/L)	Action Level	11										
Sb			4.5	2.25	1.2										

Sample Concentration units, unless otherwise noted: mg/Kg

Associated Samples: 12

					Sample Identification										
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/L)	Maximum ICB/CCB ^a (ug/L)	Action Level	No Qualifiers										
Sb			4.2	2.1											

Sample Concentration units, unless otherwise noted: mg/Kg

Associated Samples: 5, 6

					Sample Identification										
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/L)	Maximum ICB/CCB ^a (ug/L)	Action Level	No Qualifiers										
Ca			33.5	16.75											

LDC #: 30289D4

**VALIDATION FINDINGS WORKSHEET
PB/ICB/CCB QUALIFIED SAMPLES**

METHOD: Trace metals (EPA SW 864 Method 6010B/6020/7000)

Soil preparation factor applied: 100x Reason: B

Sample Concentration units, unless otherwise noted: mg/Kg Associated Samples: 11

					Sample Identification											
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/l)	Maximum ICB/CCB ^a (ug/l)	Action Level	No Qualifiers											
Ca			39.2	19.6												

Sample Concentration units, unless otherwise noted: mg/Kg Associated Samples: 4, 6, 7

					Sample Identification											
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/l)	Maximum ICB/CCB ^a (ug/l)	Action Level	No Qualifiers											
Fe			41.8	20.9												

Sample Concentration units, unless otherwise noted: mg/Kg Associated Samples: 8-11

					Sample Identification											
Analyte	Maximum PB ^a (mg/Kg)	Maximum PB ^a (ug/l)	Maximum ICB/CCB ^a (ug/l)	Action Level	No Qualifiers											
Fe			43.0	21.5												

Samples with analyte concentrations within five times the associated ICB, CCB or PB concentration are listed above with the identifications from the Validation Completeness Worksheet. These sample results were qualified as not detected, "U".

Note : a - The listed analyte concentration is the highest ICB, CCB, or PB detected in the analysis of each element.

VALIDATION FINDINGS WORKSHEET Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All Soil

Analyte	Blank ID	Sample Identification									
	FB-041113 (SDG: PH029)	Action Limit	3	11							
Cu	0.0036	1.8									
Mo	0.0036	1.8	0.34	0.86							

Sampling date: 7/17/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: 7-13

Analyte	Blank ID	Sample Identification									
	1	Action Limit	11								
Sb	0.0069	3.45	1.2								

Sampling date: 7/17/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: 3-6

Analyte	Blank ID	Sample Identification									
	2	Action Limit	No Qualifiers								
Sb	0.0054	2.7									

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

LDC #: 30289D4

VALIDATION FINDINGS WORKSHEET
Matrix Spike/Matrix Spike Duplicates

Page: 1 of 1
 Reviewer: SR
 2nd Reviewer: [Signature]

METHOD: Trace metals (EPA SW 846 Method 6010B/6020A/7000)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- N N/A Was a matrix spike analyzed for each matrix in this SDG?
 N N/A Were matrix spike percent recoveries (%R) within the control limits of 75-125%? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.
 N N/A Were all duplicate sample relative percent differences (RPD) $\leq 20\%$ for water samples and $\leq 35\%$ for soil samples?
LEVEL IV ONLY:
 N N/A Were recalculated results acceptable? See Level IV Recalculation Worksheet for recalculations.

#	MS/MSD ID	Matrix	Analyte	MS %Recovery	MSD %Recovery	RPD (Limits)	Associated Samples	Qualifications
	17/18	S	Sb	74	73		All Soil	J/USIA (Q)
			K	127	162		↓	Jdet/A ↓

Comments: 17/18: Al, Ca, Fe, Mg, Mn, P, Sr, Ti 74x

LDC #: 3028904

VALIDATION FINDINGS WORKSHEET
Initial and Continuing Calibration Calculation Verification

Page: 1 of 1
 Reviewer: GR
 2nd Reviewer: LA

METHOD: Trace Metals (EPA SW 846 Method 6010/6020/7000)

An initial and continuing calibration verification percent recovery (%R) was recalculated for each type of analysis using the following formula:

$$\%R = \frac{\text{Found}}{\text{True}} \times 100$$

Where, Found = concentration (in ug/L) of each analyte measured in the analysis of the ICV or CCV solution
 True = concentration (in ug/L) of each analyte in the ICV or CCV source

Standard ID	Type of Analysis	Element	Found (ug/L)	True (ug/L)	Recalculated	Reported	Acceptable (Y/N)
					%R	%R	
ICV	ICP (Initial calibration)	Cu	628.56	600	104.8	104.8	Y
ICV	ICP/MS (Initial calibration)	Se	52.06	50	104.1	104.1	Y
ICV	CVAA (Initial calibration)	Hg	2.46	2.5	98.4	98.4	Y
CCV	ICP (Continuing calibration)	V	504.94	500	101.0	101.0	Y
CCV	ICP/MS (Continuing calibration)	Tl	26.19	25	104.8	104.8	Y
CCV	CVAA (Continuing calibration)	Hg	1.06	1.0	106.0	106.0	Y
	GFAA (Initial calibration)						
	GFAA (Continuing calibration)						

Comments: Refer to Calibration Verification findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3028904

VALIDATION FINDINGS WORKSHEET
Level IV Recalculation Worksheet

Page: 1 of 1
Reviewer: QR
2nd Reviewer: h

METHOD: Trace Metals (EPA SW 846 Method 6010/6020/7000)

Percent recoveries (%R) for an ICP interference check sample, a laboratory control sample and a matrix spike sample were recalculated using the following formula:

$$\%R = \frac{\text{Found}}{\text{True}} \times 100$$
 Where, Found = Concentration of each analyte measured in the analysis of the sample. For the matrix spike calculation, Found = SSR (spiked sample result) - SR (sample result).
 True = Concentration of each analyte in the source.

A sample and duplicate relative percent difference (RPD) was recalculated using the following formula:

$$RPD = \frac{|S-D|}{(S+D)/2} \times 100$$
 Where, S = Original sample concentration
 D = Duplicate sample concentration

An ICP serial dilution percent difference (%D) was recalculated using the following formula:

$$\%D = \frac{|I-SDR|}{I} \times 100$$
 Where, I = Initial Sample Result (mg/L)
 SDR = Serial Dilution Result (mg/L) (Instrument Reading x 5)

Sample ID	Type of Analysis	Element	Found / S / I (units)	True / D / SDR (units)	Recalculated	Reported	Acceptable (Y/N)
					%R / RPD / %D	%R / RPD / %D	
ICSA3	ICP interference check	Cu	547.9	500	109.6	109.6	Y
LCS	Laboratory control sample	Fe	101749	100008	102	102	↓
17	Matrix spike	Ag ^(SSR-SR)	9.9627	9.8039	102	102	
19	Duplicate	Mn	382.333	319.2794	18	18	
6	ICP serial dilution	P	4140.59	4017.3	3	3	↓

Comments: Refer to appropriate worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

METHOD: Trace Metals (EPA SW 846 Method 6010/6020/7000)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- Y N N/A Have results been reported and calculated correctly?
- Y N N/A Are results within the calibrated range of the instruments and within the linear range of the ICP?
- Y N N/A Are all detection limits below the CRDL?

Detected analyte results for Be were recalculated and verified using the following equation:

Concentration = $\frac{(RD)(FV)(Dil)}{(In. Vol.)}$

Recalculation:

$\frac{100\text{mL} (0.00607\text{mg/L})}{0.985(1.04\text{g})} = 0.593\text{mg/kg}$

- RD = Raw data concentration
- FV = Final volume (ml)
- In. Vol. = Initial volume (ml) or weight (G)
- Dil = Dilution factor

#	Sample ID	Analyte	Reported Concentration (mg/Kg)	Calculated Concentration (mg/Kg)	Acceptable (Y/N)
	3	Aluminum	16900	16900	Y
		Arsenic	4.6	4.6	
		Barium	86.5	86.5	
		Beryllium	0.59	0.59	
		Cadmium	0.24	0.24	
		Calcium	2690	2690	
		Chromium	19.5	19.5	
		Cobalt	5.3	5.3	
		Copper	14.6	14.6	
		Iron	22400	22400	
		Lead	11.1	11.1	
		Lithium	22.8	22.8	
		Magnesium	4590	4590	
		Manganese	323	323	
		Mercury	0.021	0.021	
		Molybdenum	0.34	0.34	
		Nickel	11.1	11.1	
		Phosphorus	493	493	
		Potassium	3090	3090	
		Selenium	0.20	0.20	
		Silver	0.043	0.043	

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

METHOD: Trace Metals (EPA SW 846 Method 6010/6020/7000)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- Y N N/A Have results been reported and calculated correctly?
- Y N N/A Are results within the calibrated range of the instruments and within the linear range of the ICP?
- Y N N/A Are all detection limits below the CRDL?

Detected analyte results for Li were recalculated and verified using the following equation:

Concentration = $\frac{(RD)(FV)(Dil)}{(In. Vol.)}$

Recalculation:

$\frac{100 mL (0.21995 mg/L)}{0.947 (1g)} = 23.23 mg/kg$

- RD = Raw data concentration
- FV = Final volume (ml)
- In. Vol. = Initial volume (ml) or weight (G)
- Dil = Dilution factor

#	Sample ID	Analyte	Reported Concentration (mg/Kg)	Calculated Concentration (mg/Kg)	Acceptable (Y/N)
3		Sodium	77.2	77.2	Y
		Strontium	18.4	18.4	
		Thallium	0.25	0.25	
		Tin	3.0	3.0	
		Titanium	1230	1230	
		Vanadium	37.8	37.8	
		Zinc	57.9	57.9	
4		Aluminum	29400	29400	Y
		Arsenic	5.8	5.8	
		Barium	167	167	
		Beryllium	0.84	0.84	
		Calcium	9780	9780	
		Chromium	33.2	33.2	
		Cobalt	9.1	9.1	
		Copper	24.1	24.1	
		Iron	31900	31900	
		Lead	10.5	10.5	
		Lithium	23.2	23.2	
		Magnesium	6730	6730	
		Manganese	419	419	
	Mercury	0.030	0.030		

VALIDATION FINDINGS WORKSHEET

Sample Calculation Verification

METHOD: Trace Metals (EPA SW 846 Method 6010/6020/7000)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- Y N N/A Have results been reported and calculated correctly?
- Y N N/A Are results within the calibrated range of the instruments and within the linear range of the ICP?
- Y N N/A Are all detection limits below the CRDL?

Detected analyte results for _____ were recalculated and verified using the following equation:

Concentration = $\frac{(RD)(FV)(Dil)}{(In. Vol.)}$

Recalculation:

- RD = Raw data concentration
- FV = Final volume (ml)
- In. Vol. = Initial volume (ml) or weight (G)
- Dil = Dilution factor

See previous page

#	Sample ID	Analyte	Reported Concentration (mg/Kg)	Calculated Concentration (mg/Kg)	Acceptable (Y/N)
	4	Nickel	18.1	18.1	Y
		Phosphorus	1350	1350	
		Potassium	5440	5440	
		Selenium	0.26	0.26	
		Silver	0.047	0.047	
		Sodium	196	196	
		Strontium	41.3	41.3	
		Thallium	0.33	0.33	
		Tin	3.3	3.3	
		Titanium	1570	1570	
		Vanadium	63.2	63.2	
		Zinc	88.4	88.4	
		Zirconium	3.4	3.4	

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 23, 2013
Matrix: Soil/Water
Parameters: Herbicides
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713
EB2-071713
SL-514-SA8-SB-0.0-0.5
SL-514-SA8-SB-4.0-5.0
SL-514-SA8-SB-7.5-8.5
SL-514-SA8-SB-0.0-0.5MS
SL-514-SA8-SB-0.0-0.5MSD

Introduction

This data review covers 5 soil samples and 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8151A for Herbicides.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Initial Calibration

Initial calibration of compounds was performed for the primary (quantitation) column and confirmation column as required by this method.

The percent relative standard deviations (%RSD) of calibration factors for compounds were less than or equal to 20.0%.

Retention time windows were evaluated and considered technically acceptable.

III. Continuing Calibration

Continuing calibration was performed at the required frequencies.

The percent differences (%D) were less than or equal to 20.0% for all compounds with the following exceptions:

Date	Standard	Column	Compound	%D	Associated Samples	Flag	A or P
7/24/13	CCV-056	ZBXLB	Dalapon Dinoseb	40 31	All water samples in SDG PH069	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	A
7/24/13	CCV-056	ZB35	Dalapon Dinoseb	40 60	All water samples in SDG PH069	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	A
7/26/13	CCV-037	ZBXLB	Dalapon 2,4-D 2,4,5-T 2,4-DB	28 28 48 56	All soil samples in SDG PH069	J (all detects) UJ (all non-detects)	A
7/26/13	CCV-037	ZB35	MCPPP MCPA	61 29	All soil samples in SDG PH069	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	A

The percent differences (%D) of the second source calibration standard were less than or equal to 20.0% for all compounds.

Retention times (RT) of all compounds in the calibration standards were within QC limits.

IV. Blanks

Method blanks were reviewed for each matrix as applicable. No herbicide contaminants were found in the method blanks.

Samples EB1-071713 and EB2-071713 were identified as equipment blanks. No herbicide contaminants were found.

Sample FB-041113 (from SDG PH029) was identified as a field blank. No herbicide contaminants were found.

V. Surrogate Recovery

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. Matrix Spike/Matrix Spike Duplicate

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

Spike ID (Associated Samples)	Compound	MS (%R) (Limits)	MSD (%R) (Limits)	RPD (Limits)	Flag	A or P
SL-514-SA8-SB-0.0-0.5MS/MSD (SL-514-SA8-SB-0.0-0.5)	Dalapon	-	-	65 (≤50)	J (all detects)	A

VII. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

LCS ID (Associated Samples)	Compound	LCS %R (Limits)	LCS %R (Limits)	RPD (Limits)	Flag	A or P
LCS/D 19203 (All water samples in SDG PH069)	Dinoseb	-	184 (16-163)	-	J (all detects)	P

VIII. Target Compound Identification

All target compound identifications were within validation criteria.

IX. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported below the RL.	J (all detects)	A

X. System Performance

The system performance was acceptable.

XI. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XII. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Herbicides - Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH069	EB1-071713 EB2-071713	Dalapon Dinoseb	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	A	Continuing calibration (%D) (C)
PH069	SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5	Dalapon 2,4-D 2,4,5-T 2,4-DB MCPD MCPA	J (all detects) UJ (all non-detects)	A	Continuing calibration (%D) (C)
PH069	SL-514-SA8-SB-0.0-0.5	Dalapon	J (all detects)	A	Matrix spike/Matrix spike duplicate (RPD) (Q)
PH069	EB1-071713 EB2-071713	Dinoseb	J (all detects)	P	Laboratory control samples (%R) (L)
PH069	EB1-071713 EB2-071713 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Herbicides - Laboratory Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Herbicides - Field Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

LDC #: 30289D5

VALIDATION COMPLETENESS WORKSHEET

SDG #: PH069

Level IV

Laboratory: Eurofins Lancaster Laboratories

Date: 9/2/13

Page: 1 of 1

Reviewer: [Signature]

2nd Reviewer: [Signature]

METHOD: GC Herbicides (EPA SW 846 Method 8151A)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	Δ	Sampling dates: 7/17/13
II.	Initial calibration	Δ	% RSD ≤ 20
III.	Calibration verification/ICV	SW	ICV/COV = 20
IV.	Blanks	A	
V.	Surrogate recovery	A	
VI.	Matrix spike/Matrix spike duplicates	SW	
VII.	Laboratory control samples	SW	
VIII.	Target compound identification	Δ	
IX.	Compound quantitation/RL/LOQ/LODs	Δ	
X.	System Performance	Δ	
XI.	Overall assessment of data	Δ	Acceptable IS
XII.	Field duplicates	N	
XIII.	Field blanks	ND	EB = 1, 2 FB = FB-04113

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

D = Duplicate
TB = Trip blank
EB = Equipment blank

SDG PH029

Validated Samples:

SOIL + water

1	EB1-071713	w	11	PBLK 25205	21		31	
2	EB2-071713	w	12	PBLK 19203	22		32	
3	SL-514-SA8-SB-0.0-0.5		13		23		33	
4	SL-514-SA8-SB-4.0-5.0		14		24		34	
5	SL-514-SA8-SB-7.5-8.5		15		25		35	
6	#3MS		16		26		36	
7	#3MSD		17		27		37	
8			18		28		38	
9			19		29		39	
10			20		30		40	

Notes:

Method: GC HPLC

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
II. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) < 20%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a curve fit used for evaluation?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Did the initial calibration meet the curve fit acceptance criteria of > 0.990?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the RT windows properly established?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IV. Continuing calibration				
Was a continuing calibration analyzed daily?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) ≤ 20%.0 or percent recoveries 80-120%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all the retention times within the acceptance windows?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
V. Blanks				
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank analyzed for each matrix and concentration?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
If the percent recovery (%R) for one or more surrogates was out of QC limits, was a reanalysis performed to confirm samples with %R outside of criteria?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	

Validation Area	Yes	No	NA	Findings/Comments
X. Target compound identification				
Were the retention times of reported detects within the RT windows?	/			
XI. Compound quantitation/CRQLs				
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XII. System performance				
System performance was found to be acceptable.	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.		/		

VALIDATION FINDINGS WORKSHEET

METHOD: GC HPLC

8310	8330	8151	8141	8141(Con't)	8021B
A. Acenaphthene	A. HMX	A. 2,4-D	A. Dichlorvos	V. Fensulfothion	V. Benzene
B. Acenaphthylene	B. RDX	B. 2,4-DB	B. Mevinphos	W. Bolstar	CC. Toluene
C. Anthracene	C. 1,3,5-Trinitrobenzene	C. 2,4,5-T	C. Demeton-O	X. EPN	EE. Ethyl Benzene
D. Benzo(a)anthracene	D. 1,3-Dinitrobenzene	D. 2,4,5-TP	D. Demeton-S	Y. Azinphos-methyl	SSS. O-Xylene
E. Benzo(a)pyrene	E. Tetryl	E. Dinoseb	E. Ethoprop	Z. Coumaphos	RRR. MP-Xylene
F. Benzo(b)fluoranthene	F. Nitrobenzene	F. Dichlorprop	F. Naled	AA. Parathion	GG. Total Xylene
G. Benzo(g,h,i)perylene	G. 2,4,6-Trinitrotoluene	G. Dicamba	G. Sulfotep	BB. Trichloronate	
H. Benzo(k)fluoranthene	H. 4-Amino-2,6-dinitrotoluene	H. Dalapon	H. Phorate	CC. Trichlorinate	
I. Chrysene	I. 2-Amino-4,6-dinitrotoluene	I. MCPP	I. Dimethoate	DD. Trifluralin	
J. Dibenz(a,h)anthracene	J. 2,4-Dinitrotoluene	J. MCPA	J. Diazinon	EE. Def	
K. Fluoranthene	K. 2,6-Dinitrotoluene	K. Pentachlorophenol	K. Disulfoton	FF. Prowl	
L. Fluorene	L. 2-Nitrotoluene	L. 2,4,5-TP (silvex)	L. Parathion-methyl	GG. Ethion	
M. Indeno(1,2,3-cd)pyrene	M. 3-Nitrotoluene	M. Silvex	M. Ronnel		
N. Naphthalene	N. 4-Nitrotoluene		N. Malathion		
O. Phenanthrene	O.		O. Chlorpyrifos		
P. Pyrene	P.		P. Fenthion		
Q.	Q		Q. Parathion-ethyl		
R.			R. Trichloronate		
S.			S. Merphos		
			T. Stirofos		
			U. Tokuthion		

Notes: _____

LDC #: 3028905

VALIDATION FINDINGS WORKSHEET Continuing Calibration

Page: / of /

Reviewer: FT

METHOD: GC HPLC

2nd Reviewer:

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

What type of continuing calibration calculation was performed? %D or RPD

Y N N/A Were continuing calibration standards analyzed at the required frequencies?

Y N N/A Did the continuing calibration standards meet the %D / RPD validation criteria of ≤20.0%?

Level IV Only

Y N N/A Were the retention times for all calibrated compounds within their respective acceptance windows?

#	Date	Standard ID	Detector/ Column	Compound	%D / RPD (Limit ≤ 20.0)	RT (limit)	Associated Samples	Qualifications
	7/24/13	cen-056	ZB XLB	H	40	()	All water	J/U/A (C)
	10:00			E	31	()	↓	↓
			ZB 35	H	40	()	↓	↓ (C)
				E	60	()		
						()		
						()		
	7/26/13	cen-037	ZB XLB	H	28	()	All SOIL	J/U/A (C)
	1:37			A	28	()	↓	↓
				C	48	()		
				B	56	()		
			ZB 35	I	61	()		
				J	29	()	↓	↓
						()		
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LDC #: 36289DS

VALIDATION FINDINGS WORKSHEET Laboratory Control Samples (LCS)

Page: 1 of 1
Reviewer: F7
2nd Reviewer: [Signature]

METHOD: GC HPLC

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Y N N/A Were a laboratory control samples (LCS) and laboratory control sample duplicate (LCSD) analyzed for each matrix in this SDG?

Y N N/A Were the LCS percent recoveries (%R) and relative percent differences (RPD) within the QC limits?

Level IV/D Only

Y N N/A Was an LCS analyzed every 20 samples for each matrix or whenever a sample extraction was performed?

#	LCS/LCSD ID	Compound	LCS %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Associated Samples	Qualifications
	<u>lcs/d 19203</u>	<u>E</u>	()	<u>184 (16-163)</u>	()	<u>All water</u>	<u>J/Pdet (L)</u>
			()	()	()		
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LDC #: 30289 DS

VALIDATION FINDINGS WORKSHEET Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: FW
 2nd Reviewer: [Signature]

METHOD: GC ✓ HPLC _____

The calibration factors (CF) and relative standard deviation (%RSD) were recalculated using the following calculations:

CF = A/C
 Average CF = sum of the CF/number of standards
 %RSD = 100 * (S/X)

Where: A = Area of compound
 C = Concentration of compound
 S = Standard deviation of calibration factors
 X = Mean of calibration factors

#	Standard ID	Calibration Date	Compound	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				CF (<u>802</u> std)	CF (<u>802</u> std)	Ave CF (initial)	Ave CF (initial)	%RSD	%RSD
1	KAL	7/23/13	Dalapon (ZB 35)	5.01×10^{-4}	5.01×10^{-4}	4.89×10^{-4}	4.89×10^{-4}	7	7
			Dinoseb	3.59×10^{-4}	3.59×10^{-4}	3.68×10^{-4}	3.68×10^{-4}	6	6
			↓ (ZB 1B)	7.20×10^{-4}	7.20×10^{-4}	6.82×10^{-4}	6.82×10^{-4}	8	8
				6.62×10^{-4}	6.62×10^{-4}	6.27×10^{-4}	6.27×10^{-4}	6	6
2									
3									
4									

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3028905

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
 Reviewer: F7
 2nd Reviewer: [Signature]

METHOD: GC ✓ HPLC _____

The percent difference (%D) of the initial calibration average Calibration Factors (CF) and the continuing calibration CF were recalculated for the compounds identified below using the following calculation:

% Difference = 100 * (ave. CF - CF)/ave.CF
 Where: ave. CF = initial calibration average CF
 CF = continuing calibration CF
 A = Area of compound
 C = Concentration of compound

#	Standard ID	Calibration Date	Compound	Average CF(Ical)/ CCV Conc.	Reported	Recalculated	Reported	Recalculated
					CF/ Conc. CCV	CF/ Conc. CCV	%D	%D
1	ccv 10:00	7/24/13	Dalapon ZB XLB	401.00	559.77	559.77	40	40
			Dinoseb ↓	100.70	132.26	132.26	31	31
			↓ ZB 35	401.00	563.07	563.07	40	40
2			↓ ↓	100.70	160.85	160.85	60	60
3	ccv 1:37	7/26/13			287.21	287.21	28	28
					106.98	106.98	6	6
					334.68	334.68	17	17
4			↓ ↓		108.56	108.56	8	8

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3028905

VALIDATION FINDINGS WORKSHEET Surrogate Results Verification

Page: 1 of 1
Reviewer: E7
2nd reviewer: ←

METHOD: GC HPLC

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
SS = Surrogate Spiked

Sample ID: # 3

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
<u>2,4-DCAA</u>	<u>ZB35</u>	<u>66.52</u>	<u>51.226299</u>	<u>77</u>	<u>77</u>	<u>0</u>

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

VALIDATION FINDINGS WORKSHEET
Matrix Spike/Matrix Spike Duplicates Results Verification

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the matrix spike and matrix spike duplicate were recalculated for the compounds identified below using the following calculation:

%Recovery = $100 * (SSC - SC) / SA$

Where

SSC = Spiked sample concentration

SC = Sample concentration

SA = Spike added

MS = Matrix spike

MSD = Matrix spike duplicate

RPD = $((SSCMS - SSCMSD) * 2) / (SSCMS + SSCMSD) * 100$

MS/MSD samples: 6 + 7

Compound	Spike Added (ug/kg)		Sample Conc. (ug/kg)	Spike Sample Concentration (ug/kg)		Matrix spike		Matrix Spike Duplicate		MS/MSD	
	MS	MSD		MS	MSD	Percent Recovery		Percent Recovery		RPD	
						Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)											
Diesel (8015)											
Benzene (8021B)											
Methane (RSK-175)											
2,4-D (8151)	82.3	82.3	ND	80.92	76.88	98	98	93	93	5	5
Dinoseb (8151)	139	139	ND	27.77	28.98	20	20	21	21	4	4
Naphthalene (8310)											
Anthracene (8310)											
HMX (8330)											
2,4,6-Trinitrotoluene (8330)											

Comments: Refer to Matrix Spike/Matrix Spike Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET

Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

$\%Recovery = 100 * (SSC - SC) / SA$

Where SSC = Spiked sample concentration
SA = Spike added
LCS = Laboratory Control Sample

SC = Sample concentration

$RPD = ((SSCLCS - SSCLCSD) * 2) / (SSCLCS + SSCLCSD) * 100$

LCSD = Laboratory Control Sample duplicate

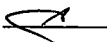
LCS/LCSD samples: LCS 19203
LCSD

Compound	Spike Added (ug/L)		Sample Conc. (ug/L)	Spike Sample Concentration (ug/L)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD		LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
						Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)											
Diesel (8015)											
Benzene (8021B)											
Methane (RSK-175)											
2,4-D (8151)	2.51	2.51	ND	2.86	2.93	114	114	117	117	2	2
Dinoseb (8151)	4.25	4.25	ND	6.45	7.8	152	152	184	184	19	19
Naphthalene (8310)											
Anthracene (8310)											
HMX (8330)											
2,4,6-Trinitrotoluene (8330)											

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30289P5

VALIDATION FINDINGS WORKSHEET Sample Calculation Verification

Page: 1 of 1
Reviewer: F7
2nd Reviewer: 

METHOD: GC HPLC

Y N N/A
Y N N/A

Were all reported results recalculated and verified for all level IV samples?
Were all recalculated results for detected target compounds within 10% of the reported results?

$$\text{Concentration} = \frac{(A)(Fv)(Df)}{(RF)(Vs \text{ or } Ws)(\%S/100)}$$

Example:

Sample ID: 10519203 Compound Name Dinoseb

- A= Area or height of the compound to be measured
- Fv= Final Volume of extract
- Df= Dilution Factor
- RF= Average response factor of the compound in the initial calibration
- Vs= Initial volume of the sample
- Ws= Initial weight of the sample
- %S= Percent Solid

$$\text{Concentration} = \frac{(3646164)(1)(10)}{(15338228)(3.68 \times 10^{-4})(1000)}$$

$$= 6.45597 \text{ ug/L}$$

#	Sample ID	Compound	Reported Concentrations ()	Recalculated Results Concentrations ()	Qualifications

Comments: _____

**Laboratory Data Consultants, Inc.
Data Validation Report**

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 23, 2013
Matrix: Soil/Water
Parameters: Hexavalent Chromium
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713
EB2-071713
SL-513-SA8-SB-0.0-0.5
SL-513-SA8-SB-4.0-5.0
SL-513-SA8-SB-8.0-9.0

Introduction

This data review covers 3 soil samples and 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 7199 for Hexavalent Chromium.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Data Review (January 2010).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Initial Calibration

All criteria for the initial calibration were met.

III. Calibration Verification

Continuing calibration frequency and analysis criteria were met.

IV. Blanks

Method blanks were reviewed for each matrix as applicable. No hexavalent chromium was found in the initial, continuing and preparation blanks.

Samples EB1-071713 and EB2-071713 were identified as equipment blanks. No hexavalent chromium was found.

Sample FB-041113 (from SDG PH029) was identified as a field blank. No hexavalent chromium was found.

V. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VI. Duplicates

The laboratory has indicated that there were no duplicate (DUP) analyses specified for the samples in this SDG, and therefore duplicate analyses were not performed for this SDG.

VII. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VIII. Sample Result Verification

All sample result verifications were acceptable.

IX. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

X. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Hexavalent Chromium - Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Hexavalent Chromium – Laboratory Blank Data Qualification Summary - SDG
PH069**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Hexavalent Chromium - Field Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

LDC #: 30289D6

VALIDATION COMPLETENESS WORKSHEET

SDG #: PH069

Level IV

Laboratory: Eurofins Lancaster Laboratories

Date: 9/4/13

Page: 1 of 1

Reviewer: [Signature]

2nd Reviewer: [Signature]

METHOD: (Analyte) Hexavalent Chromium (EPA SW846 Method 7199)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 7/17/13
Ia.	Initial calibration	A	
Iib.	Calibration verification	A	
III.	Blanks	A	
IV	Matrix Spike/Matrix Spike Duplicates	N	CS
V	Duplicates	N	↓
VI.	Laboratory control samples	A	LCS/D
VII.	Sample result verification	A	
VIII.	Overall assessment of data	A	
IX.	Field duplicates	N	
X	Field blanks	ND	EB=1,2 FB=FB-01113

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

(PH029)

Validated Samples:

soil/water

1	EB1-071713	W	11		21		31
2	EB2-071713	W	12		22		32
3	SL-513-SA8-SB-0.0-0.5	S	13		23		33
4	SL-513-SA8-SB-4.0-5.0	↓	14		24		34
5	SL-513-SA8-SB-8.0-9.0	↓	15		25		35
6			16		26		36
7			17		27		37
8			18		28		38
9			19		29		39
10			20		30		40

Notes: _____

Method: Inorganics (EPA Method See cover)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. Calibration				
Were all instruments calibrated daily, each set-up time?	/			
Were the proper number of standards used?	/			
Were all initial calibration correlation coefficients > 0.995?	/			
Were all initial and continuing calibration verification %Rs within the 90-110% QC limits?	/			
Were titrant checks performed as required? (Level IV only)			/	
Were balance checks performed as required? (Level IV only)			/	
III. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.		/		
IV. Matrix spike/Matrix spike duplicates and Duplicates				
Were a matrix spike (MS) and duplicate (DUP) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD or MS/DUP. Soil / Water.		/		
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the 75-125 QC limits? If the sample concentration exceeded the spike concentration by a factor of 4 or more, no action was taken.			/	
Were the MS/MSD or duplicate relative percent differences (RPD) ≤ 20% for waters and ≤ 35% for soil samples? A control limit of ≤ CRDL (≤ 2X CRDL for soil) was used for samples that were ≤ 5X the CRDL, including when only one of the duplicate sample values were ≤ 5X the CRDL.			/	
V. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per extraction batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the 80-120% (85-115% for Method 300.0) QC limits?	/			
VI. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?			/	
Were the performance evaluation (PE) samples within the acceptance limits?			/	

Validation Area	Yes	No	NA	Findings/Comments
VII. Sample Result Verification				
Were RLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were detection limits < RL?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
VIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IX. Field duplicates				
Field duplicate pairs were identified in this SDG.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Target analytes were detected in the field duplicates.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
X. Field blanks				
Field blanks were identified in this SDG.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Target analytes were detected in the field blanks.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	

LDC #: 3028006

Validation Findings Worksheet
Initial and Continuing Calibration Calculation Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

Method: Inorganics, Method See Cover

The correlation coefficient (r) for the calibration of Cr⁶⁺ was recalculated. Calibration date: 7/23/13

An initial or continuing calibration verification percent recovery (%R) was recalculated for each type of analysis using the following formula:

$$\%R = \frac{\text{Found} \times 100}{\text{True}}$$

Where,

Found = concentration of each analyte measured in the analysis of the ICV or CCV solution
 True = concentration of each analyte in the ICV or CCV source

Type of analysis	Analyte	Standard	Conc. (mg/L)	Area	Recalculated	Reported	Acceptable (Y/N)
					r or r ²	r or r ²	
Initial calibration	Cr ⁶⁺	s1	10	0.115518	0.9997	0.9997	Y
		s2	50	0.58082			
		s3	100	1.108824			
		s4	200	2.158982			
		s5	300	3.299498			
Calibration verification		CCV	200	201.6928	101	101	Y
Calibration verification				2105783	105	105	Y
Calibration verification							

Comments: Refer to Calibration Verification findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3028902

VALIDATION FINDINGS WORKSHEET
Level IV Recalculation Worksheet

Page: 1 of 1
Reviewer: QR
2nd Reviewer: [Signature]

METHOD: Inorganics, Method see cover

Percent recoveries (%R) for a laboratory control sample and a matrix spike sample were recalculated using the following formula:

$\%R = \frac{\text{Found}}{\text{True}} \times 100$ Where, Found = concentration of each analyte measured in the analysis of the sample. For the matrix spike calculation, Found = SSR (spiked sample result) - SR (sample result).
True = concentration of each analyte in the source.

A sample and duplicate relative percent difference (RPD) was recalculated using the following formula:

$RPD = \frac{|S-D|}{(S+D)/2} \times 100$ Where, S = Original sample concentration
D = Duplicate sample concentration

Sample ID	Type of Analysis	Element	Found / S (units)	True / D (units)	Recalculated	Reported	Acceptable (Y/N)
					%R / RPD	%R / RPD	
<u>LCS</u>	Laboratory control sample	<u>Cd⁺</u>	<u>5.07</u>	<u>5</u>	<u>101</u>	<u>101</u>	<u>Y</u>
<u>N</u>	Matrix spike sample		(SSR-SR)				
<u>N</u>	Duplicate sample						

Comments: Refer to appropriate worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3028906

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

Page: 1 of 1
 Reviewer: CR
 2nd reviewer: W

METHOD: Inorganics, Method see over

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- Y N N/A Have results been reported and calculated correctly?
- Y N N/A Are results within the calibrated range of the instruments?
- Y N N/A Are all detection limits below the CRQL?

Compound (analyte) results for Cr⁶⁺ reported with a positive detect were recalculated and verified using the following equation:

Concentration =
 $\text{Area}(91.84) - 1.415$

Recalculation:

$$\frac{0.1422(91.84) - 1.415}{0.916(26)} = 0.488 \text{ mg/kg}$$

#	Sample ID	Analyte	Reported Concentration (mg/kg)	Calculated Concentration (mg/kg)	Acceptable (Y/N)
	<u>3</u>	<u>Cr⁶⁺</u>	<u>0.49</u>	<u>0.49</u>	<u>Y</u>

Note: _____

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 23, 2013
Matrix: Soil/Water
Parameters: Total Petroleum Hydrocarbons as Gasoline
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713
EB2-071713
SL-514-SA8-SB-4.0-5.0
SL-514-SA8-SB-7.5-8.5
SL-515-SA8-SB-4.0-5.0
SL-515-SA8-SB-9.0-10.0
SL-515-SA8-SB-14.0-15.0
TB-071713
SL-513-SA8-SB-4.0-5.0
SL-513-SA8-SB-8.0-9.0

Introduction

This data review covers 7 soil samples and 3 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8015B for Total Petroleum Hydrocarbons (TPH) as Gasoline.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Initial Calibration

Initial calibration of compounds was performed as required by the method.

The percent relative standard deviations (%RSD) of calibration factors for all compounds were less than or equal to 20.0%.

III. Continuing Calibration

Continuing calibration was performed at required frequencies.

The percent differences (%D) were less than or equal to 20.0% for all compounds.

The percent differences (%D) of the second source calibration standard were less than or equal to 20.0% for all compounds.

IV. Blanks

Method blanks were reviewed for each matrix as applicable. No total petroleum hydrocarbons as gasoline contaminants were found in the method blanks.

Sample TB-071713 was identified as a trip blank. No total petroleum hydrocarbons as gasoline contaminants were found.

Samples EB1-071713 and EB2-071713 were identified as equipment blanks. No total petroleum hydrocarbons as gasoline contaminants were found.

Sample FB-041113 (from SDG PH029) was identified as a field blank. No total petroleum hydrocarbons as gasoline contaminants were found.

V. Surrogate Recovery

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VII. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VIII. Target Compound Identification

All target compound identifications were within validation criteria.

IX. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported below the RL.	J (all detects)	A

X. System Performance

The system performance was acceptable.

XI. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XII. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Gasoline - Data Qualification Summary - SDG
 PH069**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH069	EB1-071713 EB2-071713 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 TB-071713 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Gasoline - Laboratory Blank Data Qualification
 Summary - SDG PH069**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Gasoline - Field Blank Data Qualification
 Summary - SDG PH069**

No Sample Data Qualified in this SDG

METHOD: GC TPH as Gasoline (EPA SW 846 Method 8015B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 7/17/13
II.	Initial calibration	A	% RSD ≤ 20
III.	Calibration verification/ICV	A	ICV/CCV ≤ 20
IV.	Blanks	A	
V.	Surrogate recovery	A	
VI.	Matrix spike/Matrix spike duplicates	N	client specified
VII.	Laboratory control samples	A	LCSD
VIII.	Target compound identification	A	
IX.	Compound quantitation/RL/LOQ/LODs	A	
X.	System Performance	A	
XI.	Overall assessment of data	A	
XII.	Field duplicates	N	
XIII.	Field blanks	ND	EB = 1,2 TB = 8 FB = FB - 04113 (SDG PH029)

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

ND = No compounds detected
 R = Rinsate
 FB = Field blank

D = Duplicate
 TB = Trip blank
 EB = Equipment blank

Validated Samples:

soil, water

1 ²	EB1-071713	N	11	BLKLC	21		31	
2 ²	EB2-071713	N	12	BLKLJ	22		32	
3	SL-514-SA8-SB-4.0-5.0	✓	13		23		33	
4	SL-514-SA8-SB-7.5-8.5	✓	14		24		34	
5	SL-515-SA8-SB-4.0-5.0		15		25		35	
6	SL-515-SA8-SB-9.0-10.0		16		26		36	
7	SL-515-SA8-SB-14.0-15.0		17		27		37	
8 ²	TB-071713	N	18		28		38	
9	SL-513-SA8-SB-4.0-5.0		19		29		39	
10	SL-513-SA8-SB-8.0-9.0		20		30		40	

Notes: _____

Method: GC HPLC

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
II. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) < 20%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a curve fit used for evaluation?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Did the initial calibration meet the curve fit acceptance criteria of > 0.990?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the RT windows properly established?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IV. Continuing calibration				
Was a continuing calibration analyzed daily?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) < 20%.0 or percent recoveries 80-120%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all the retention times within the acceptance windows?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
V. Blanks				
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank analyzed for each matrix and concentration?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
If the percent recovery (%R) for one or more surrogates was out of QC limits, was a reanalysis performed to confirm samples with %R outside of criteria?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	

Validation Area	Yes	No	NA	Findings/Comments
X. Target compound identification				
Were the retention times of reported detects within the RT windows?	/			
XI. Compound quantitation/CRQLs				
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XII. System performance				
System performance was found to be acceptable.	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.		/		

LDC #: 3020907

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: F7
 2nd Reviewer: [Signature]

METHOD: GC HPLC

The calibration factors (CF) and relative standard deviation (%RSD) were recalculated using the following calculations:

CF = A/C
 Average CF = sum of the CF/number of standards
 %RSD = 100 * (S/X)

Where: A = Area of compound
 C = Concentration of compound
 S = Standard deviation of calibration factors
 X = Mean of calibration factors

#	Standard ID	Calibration Date	Compound	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				CF (<u>530</u> std)	CF (<u>530</u> std)	Ave CF (initial)	Ave CF (initial)	%RSD	%RSD
1	ICAL	1/12/12	GRU	6717	6717	6269	6269	9.2	9.2
				(<u>220</u>)	(<u>220</u>)				
2	ICAL	5/23/13	GRU	60597	60597	61516	61516	3	3
3									
4									

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3028907

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
 Reviewer: F7
 2nd Reviewer: [Signature]

METHOD: GC HPLC

The percent difference (%D) of the initial calibration average Calibration Factors (CF) and the continuing calibration CF were recalculated for the compounds identified below using the following calculation:

% Difference = 100 * (ave. CF - CF)/ave.CF

Where: ave. CF = initial calibration average CF
 CF = continuing calibration CF
 A = Area of compound
 C = Concentration of compound

#	Standard ID	Calibration Date	Compound	Average CF(1cal)/ CCV Conc.	Reported	Recalculated	Reported	Recalculated
					CF/ Conc. CCV	CF/ Conc. CCV	%D	%D
1	CCV 19:24	7/23/13	GRU	550.00	535.71	535.71	3	3
	CCV 17:38	7/22/13	↓	220.0	215.47	215.47	2	2
2	CCV 3:08	7/23/13	↓	220.0	206.39	206.39	6	6
3								
4								

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3028707

VALIDATION FINDINGS WORKSHEET Surrogate Results Verification

Page: 1 of 1
Reviewer: F7
2nd reviewer: [Signature]

METHOD: GC HPLC

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
SS = Surrogate Spiked *spt Added 750 (0.9728)*

Sample ID: #3

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
TFT	NS	729.60	451.5625	62	62	0

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

LDC #: 30289D7

VALIDATION FINDINGS WORKSHEET

Page: 1 of 1

Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Reviewer: FJ

2nd Reviewer: [Signature]

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

$\% \text{Recovery} = 100 * (\text{SSC} - \text{SC}) / \text{SA}$

Where SSC = Spiked sample concentration

SC = Sample concentration

$\text{RPD} = ((\text{SSCLCS} - \text{SSCLCSD}) * 2) / (\text{SSCLCS} + \text{SSCLCSD}) * 100$

SA = Spike added

LCS = Laboratory Control Sample

LCSD = Laboratory Control Sample duplicate

LCS/LCSD samples: LCSH9 / LCSDW7

Compound	Spike Added (ug/L)		Sample Conc. (ug/L)	Spike Sample Concentration (ug/L)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD		LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
						Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)	1100	1100	ND	1141.09	1085.47	104	104	99	99	5	5
Diesel (8015)											
Benzene (8021B)											
Methane (RSK-175)											
2,4-D (8151)											
Dinoseb (8151)											
Naphthalene (8310)											
Anthracene (8310)											
HMX (8330)											
2,4,6-Trinitrotoluene (8330)											

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30289P7

VALIDATION FINDINGS WORKSHEET Sample Calculation Verification

Page: 1 of 1
Reviewer: F7
2nd Reviewer: ←

METHOD: GC HPLC

Y N N/A Were all reported results recalculated and verified for all level IV samples?
Y N N/A Were all recalculated results for detected target compounds within 10% of the reported results?

$$\text{Concentration} = \frac{(A)(Fv)(Df)}{(RF)(Vs \text{ or } Ws)(\%S/100)}$$

Example:

Sample ID: # L0544 Compound Name GRO

A= Area or height of the compound to be measured
Fv= Final Volume of extract
Df= Dilution Factor
RF= Average response factor of the compound
In the initial calibration
Vs= Initial volume of the sample
Ws= Initial weight of the sample
%S= Percent Solid

$$\text{Concentration} = \frac{(22351810 - 3001740)(25)}{(61516)(1000)} = 7.9 \text{ mg/kg}$$

#	Sample ID	Compound	Reported Concentrations ()	Recalculated Results Concentrations ()	Qualifications

Comments: _____

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 23, 2013
Matrix: Soil/Water
Parameters: Total Petroleum Hydrocarbons as Extractables
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713
EB2-071713
SL-512-SA8-SB-0.0-0.5
SL-514-SA8-SB-0.0-0.5
SL-514-SA8-SB-4.0-5.0
SL-514-SA8-SB-7.5-8.5
SL-515-SA8-SB-0.0-0.5
SL-515-SA8-SB-4.0-5.0
SL-515-SA8-SB-9.0-10.0
SL-515-SA8-SB-14.0-15.0
SL-513-SA8-SB-0.0-0.5
SL-513-SA8-SB-4.0-5.0
SL-513-SA8-SB-8.0-9.0
SL-514-SA8-SB-7.5-8.5MS
SL-514-SA8-SB-7.5-8.5MSD

Introduction

This data review covers 13 soil samples and 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8015B for Total Petroleum Hydrocarbons (TPH) as Extractables.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. Initial Calibration

Initial calibration of compounds was performed as required by the method.

The percent relative standard deviations (%RSD) of calibration factors for compounds were less than or equal to 20.0%.

III. Continuing Calibration

Continuing calibration was performed at required frequencies.

The percent differences (%D) were less than or equal to 20.0% for all compounds.

The percent differences (%D) of the second source calibration standard were less than or equal to 20.0% for all compounds.

IV. Blanks

Method blanks were reviewed for each matrix as applicable. No total petroleum hydrocarbons as extractables contaminants were found in the method blanks.

Samples EB1-071713 and EB2-071713 were identified as equipment blanks. No total petroleum hydrocarbons as extractables contaminants were found.

Sample FB-041113 (from SDG PH029) was identified as a field blank. No total petroleum hydrocarbons as extractables contaminants were found.

V. Surrogate Recovery

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the following exceptions:

Spike ID (Associated Samples)	Compound	MS (%R) (Limits)	MSD (%R) (Limits)	RPD (Limits)	Flag	A or P
SL-514-SA8-SB-7.5-8.5MS/MSD (SL-514-SA8-SB-7.5-8.5)	Extractable fuel hydrocarbons (C15-C20)	162 (49-123)	138 (49-123)	-	J (all detects)	A
	Extractable fuel hydrocarbons (C21-C30)	172 (49-123)	-	28 (≤20)	J (all detects)	
	Extractable fuel hydrocarbons (C30-C40)	131 (49-123)	-	-	J (all detects)	

VII. Laboratory Control Samples

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits.

VIII. Target Compound Identification

All target compound identifications were within validation criteria.

IX. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported below the RL.	J (all detects)	A

X. System Performance

The system performance was acceptable.

XI. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XII. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Extractables - Data Qualification Summary -
 SDG PH069**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH069	SL-514-SA8-SB-7.5-8.5	Extractable fuel hydrocarbons (C15-C20) Extractable fuel hydrocarbons (C30-C40)	J (all detects) J (all detects)	A	Matrix spike/Matrix spike duplicate (%R) (Q)
PH069	SL-514-SA8-SB-7.5-8.5	Extractable fuel hydrocarbons (C21-C30)	J (all detects)	A	Matrix spike/Matrix spike duplicate (%R)(RPD) (Q)
PH069	EB1-071713 EB2-071713 SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-514-SA8-SB-7.5-8.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Extractables - Laboratory Blank Data
 Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
 Total Petroleum Hydrocarbons as Extractables - Field Blank Data Qualification
 Summary - SDG PH069**

No Sample Data Qualified in this SDG

LDC #: 30289D8

VALIDATION COMPLETENESS WORKSHEET

Date: 9/2/13

SDG #: PH069

Level IV

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: [Signature]

2nd Reviewer: [Signature]

METHOD: GC TPH as Extractables (EPA SW 846 Method 8015B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	Δ	Sampling dates: 7/17/13
II.	Initial calibration	Δ	% PSD ≤ 20
III.	Calibration verification/ICV	Δ	ICV/CCV ≤ 20
IV.	Blanks	Δ	
V.	Surrogate recovery	Δ	
VI.	Matrix spike/Matrix spike duplicates	SW	
VII.	Laboratory control samples	Δ	LCS 10
VIII.	Target compound identification	Δ	
IX.	Compound quantitation/RL/LOQ/LODs	Δ	
X.	System Performance	Δ	
XI.	Overall assessment of data	Δ	
XII.	Field duplicates	N	
XIII.	Field blanks	ND	EB = 1 EB = 2

Note: A = Acceptable
 N = Not provided/applicable
 SW = See worksheet

* ND = No compounds detected
 R = Rinsate
 FB = Field blank

* FB = FB-04113
 D = Duplicate SDG PH029
 TB = Trip blank
 EB = Equipment blank

Validated Samples:

2011 - water

12	EB1-071713	w	11	SL-513-SA8-SB-0.0-0.5	21	PBLK 15204	31	
22	EB2-071713	w	12	SL-513-SA8-SB-4.0-5.0	22	PBLK 10201	32	
3	SL-512-SA8-SB-0.0-0.5		13	SL-513-SA8-SB-8.0-9.0	23		33	
4	SL-514-SA8-SB-0.0-0.5		14	#6 MS	24		34	
5	SL-514-SA8-SB-4.0-5.0		15	#6 MS/D	25		35	
6	SL-514-SA8-SB-7.5-8.5		16		26		36	
7	SL-515-SA8-SB-0.0-0.5		17		27		37	
8	SL-515-SA8-SB-4.0-5.0		18		28		38	
9	SL-515-SA8-SB-9.0-10.0		19		29		39	
10	SL-515-SA8-SB-14.0-15.0		20		30		40	

Notes: _____

Method: GC HPLC

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
II. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent relative standard deviations (%RSD) \leq 20%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a curve fit used for evaluation?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
Did the initial calibration meet the curve fit acceptance criteria of > 0.990 ?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the RT windows properly established?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IV. Continuing calibration				
Was a continuing calibration analyzed daily?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all percent differences (%D) \leq 20%.0 or percent recoveries 80-120%?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were all the retention times within the acceptance windows?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
V. Blanks				
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a method blank analyzed for each matrix and concentration?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
VI. Surrogate spikes				
Were all surrogate %R within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
If the percent recovery (%R) for one or more surrogates was out of QC limits, was a reanalysis performed to confirm samples with %R outside of criteria?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
VII. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was a MS/MSD analyzed every 20 samples of each matrix?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input type="checkbox"/>	<input checked="" type="checkbox"/>	<input type="checkbox"/>	
VIII. Laboratory control samples				
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input checked="" type="checkbox"/>	<input type="checkbox"/>	<input type="checkbox"/>	
IX. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?	<input type="checkbox"/>	<input type="checkbox"/>	<input checked="" type="checkbox"/>	

Validation Area	Yes	No	NA	Findings/Comments
X. Target compound identification				
Were the retention times of reported detects within the RT windows?	✓			
XI. Compound quantitation/CRQLs				
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	✓			
XII. System performance				
System performance was found to be acceptable.	✓			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	✓			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		✓		
Target compounds were detected in the field duplicates.			✓	
XV. Field blanks				
Field blanks were identified in this SDG.	✓			
Target compounds were detected in the field blanks.		✓		

LDC #: 3028908

VALIDATION FINDINGS WORKSHEET Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: F7
 2nd Reviewer: [Signature]

METHOD: GC _____ HPLC _____

The calibration factors (CF) and relative standard deviation (%RSD) were recalculated using the following calculations:

CF = A/C
 Average CF = sum of the CF/number of standards
 %RSD = 100 * (S/X)

Where: A = Area of compound
 C = Concentration of compound
 S = Standard deviation of calibration factors
 X = Mean of calibration factors

#	Standard ID	Calibration Date	Compound	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				CF (288 std)	CF (288 std)	Ave CF (initial)	Ave CF (initial)	%RSD	%RSD
1	ICAL	6/28/13 (A)	TPH Ane	23751.09	23751.09	25786.45	25786.45	8.235	8.235
	FLJ179A								
	TPH J179A								
2	ICAL	6/28/13 (B)	↓	23620	23620	24187	24187	2	2
	TPH K179A								
3									
4									

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3028908

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
 Reviewer: F7
 2nd Reviewer: A

METHOD: GC HPLC

The percent difference (%D) of the initial calibration average Calibration Factors (CF) and the continuing calibration CF were recalculated for the compounds identified below using the following calculation:

% Difference = 100 * (ave. CF - CF)/ave.CF
 Where: ave. CF = initial calibration average CF
 CF = continuing calibration CF
 A = Area of compound
 C = Concentration of compound

#	Standard ID	Calibration Date	Compound	Average CF(1cal)/ CCV Conc.	Reported	Recalculated	Reported	Recalculated
					CF/ Conc. CCV	CF/ Conc. CCV	%D	%D
1	cen 20:34	7/22/13	C ₈ -C ₄₀	239.97	259.14	259.14	8	8
2	cen 16:18	7/24/13	↓	288.01	283.92	283.92	1	1
3								
4								

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 3028708

VALIDATION FINDINGS WORKSHEET
Surrogate Results Verification

Page: 1 of 1
Reviewer: F7
2nd reviewer: [Signature]

METHOD: GC HPLC

The percent recoveries (%R) of surrogates were recalculated for the compounds identified below using the following calculation:

% Recovery: SF/SS * 100

Where: SF = Surrogate Found
SS = Surrogate Spiked

Sample ID: #13

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	
chlorobenzene	ZB5	2.0	2.0408	102	102	0
ortho terphenyl	↓	2.0	2.129554	106	106	0

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

Sample ID: _____

Surrogate	Column/Detector	Surrogate Spiked	Surrogate Found	Percent Recovery	Percent Recovery	Percent Difference
				Reported	Recalculated	

LDC #: 3028908

VALIDATION FINDINGS WORKSHEET
Matrix Spike/Matrix Spike Duplicates Results Verification

Page: 1 of 1
 Reviewer: EF
 2nd Reviewer: A

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the matrix spike and matrix spike duplicate were recalculated for the compounds identified below using the following calculation:

%Recovery = $100 * (SSC - SC) / SA$

Where

SSC = Spiked sample concentration

SC = Sample concentration

SA = Spike added

RPD = $((SSCMS - SSCMSD) * 2) / (SSCMS + SSCMSD) * 100$

MS = Matrix spike

MSD = Matrix spike duplicate

MS/MSD samples: 14 & 15

Compound	Spike Added (mg/kg)		Sample Conc. (mg/kg)	Spike Sample Concentration (mg/kg)		Matrix spike		Matrix Spike Duplicate		MS/MSD	
	MS	MSD		MS	MSD	Percent Recovery		Percent Recovery		RPD	
						Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)											
Diesel (8015)											
Benzene (8021B)											
Methane (RSK-175)											
2,4-D (8151)											
Dinoseb (8151)											
Naphthalene (8310)											
Anthracene (8310)											
HMX (8330)											
2,4,6-Trinitrotoluene (8330)											
EFH (CB-C11)	1.68	1.68	ND	1.71	1.63	102	102	97	97	5	5

Comments: Refer to Matrix Spike/Matrix Spike Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30289D8

VALIDATION FINDINGS WORKSHEET

Page: 1 of 1

Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Reviewer: FJ

2nd Reviewer: [Signature]

METHOD: GC HPLC

The percent recoveries (%R) and relative percent differences (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

$\% \text{Recovery} = 100 * (\text{SSC} - \text{SC}) / \text{SA}$

Where SSC = Spiked sample concentration

SC = Sample concentration

SA = Spike added

$\text{RPD} = ((\text{SSCLCS} - \text{SSCLCSD}) * 2) / (\text{SSCLCS} + \text{SSCLCSD}) * 100$

LCS = Laboratory Control Sample

LCSD = Laboratory Control Sample duplicate

LCS/LCSD samples: LCS15204

Compound	Spike Added (mg/kg)		Sample Conc. (mg/kg)	Spike Sample Concentration (mg/kg)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD		LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
						Reported	Recalc.	Reported	Recalc.	Reported	Recalc.
Gasoline (8015)											
Diesel (8015)											
Benzene (8021B)											
Methane (RSK-175)											
2,4-D (8151)											
Dinoseb (8151)											
Naphthalene (8310)											
Anthracene (8310)											
HMX (8330)											
2,4,6-Trinitrotoluene (8330)											
EFH (CB-11)	1.68	1.68 NA	ND	1.41	NA	84	84	NA			

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicate findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30289PS

VALIDATION FINDINGS WORKSHEET Sample Calculation Verification

Page: 1 of 1
Reviewer: F7
2nd Reviewer: [Signature]

METHOD: GC HPLC

Y N N/A
Y N N/A

Were all reported results recalculated and verified for all level IV samples?

Were all recalculated results for detected target compounds within 10% of the reported results?

$$\text{Concentration} = \frac{(A)(Fv)(Df)}{(RF)(Vs \text{ or } Ws)(\%S/100)}$$

Example:

Sample ID. #3 Compound Name C30 - C40

- A= Area or height of the compound to be measured
- Fv= Final Volume of extract
- Df= Dilution Factor
- RF= Average response factor of the compound
In the initial calibration
- Vs= Initial volume of the sample
- Ws= Initial weight of the sample
- %S= Percent Solid

$$\text{Concentration} = \frac{(36268247)(1)}{(24187)(0.989)(30)}$$

51 mg/kg

#	Sample ID	Compound	Reported Concentrations ()	Recalculated Results Concentrations ()	Qualifications

Comments: _____

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 23, 2013
Matrix: Soil/Water
Parameters: Dioxins/Dibenzofurans
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories

Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713
EB2-071713
SL-512-SA8-SB-0.0-0.5
SL-514-SA8-SB-0.0-0.5
SL-514-SA8-SB-4.0-5.0
SL-515-SA8-SB-0.0-0.5
SL-515-SA8-SB-4.0-5.0
SL-515-SA8-SB-9.0-10.0
SL-515-SA8-SB-14.0-15.0
SL-513-SA8-SB-0.0-0.5
SL-513-SA8-SB-4.0-5.0
SL-513-SA8-SB-8.0-9.0

Introduction

This data review covers 10 soil samples and 2 water samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA Method 1613B for Polychlorinated Dioxins/Dibenzofurans.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and the USEPA Contract Laboratory Program National Functional Guidelines for Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. HRGC/HRMS Instrument Performance Check

Instrument performance was checked at the required daily frequency.

The chromatographic resolution between 2,3,7,8-TCDD and the peaks representing any other unlabeled TCDD isomers was resolved with a valley of less than or equal to 25%.

PFK and static resolving power were within validation criteria.

III. Initial Calibration

A five point initial calibration was performed as required by the method.

Percent relative standard deviations (%RSD) were less than or equal to 20.0% for unlabeled compounds and less than or equal to 35.0% for labeled compounds.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

The minimum S/N ratio was greater than or equal to 10 for each unlabeled compound and labeled compound.

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

All of the continuing calibration percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were within QC limits.

The percent differences (%D) of the second source calibration standard were within QC limits.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No polychlorinated dioxin/dibenzofuran contaminants were found in the method blanks with the following exceptions:

Method Blank ID	Extraction Date	Compound	Concentration	Associated Samples
BLK200001	7/19/13	1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.368 pg/L 0.364 pg/L 0.327 pg/L 0.373 pg/L 1.18 pg/L 0.465 pg/L 0.511 pg/L 0.527 pg/L 0.476 pg/L 0.238 pg/L 0.586 pg/L 0.413 pg/L 0.325 pg/L 0.838 pg/L	EB1-071713 EB2-071713
BLK206001	7/26/13	1,2,3,4,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF	0.0432 ng/Kg 0.0358 ng/Kg 0.0714 ng/Kg 0.230 ng/Kg 0.0307 ng/Kg 0.0229 ng/Kg 0.0231 ng/Kg 0.0401 ng/Kg 0.0413 ng/Kg 0.0256 ng/Kg 0.159 ng/Kg	SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0

Sample concentrations were compared to concentrations detected in the method blanks. The sample concentrations were either not detected or were significantly greater (>5X blank contaminants) than the concentrations found in the associated method blanks with the following exceptions:

Sample	Compound	Reported Concentration	Modified Final Concentration
EB1-071713	1,2,3,4,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.161 pg/L 0.413 pg/L 0.814 pg/L 0.406 pg/L 0.132 pg/L 0.222 pg/L 0.191 pg/L 0.224 pg/L 0.186 pg/L 0.238 pg/L 0.191 pg/L 0.512 pg/L	0.161U pg/L 0.413U pg/L 0.814U pg/L 0.406U pg/L 0.132U pg/L 0.222U pg/L 0.191U pg/L 0.224U pg/L 0.186U pg/L 0.238U pg/L 0.191U pg/L 0.512U pg/L
EB2-071713	1,2,3,4,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.373 pg/L 0.352 pg/L 0.797 pg/L 0.532 pg/L 0.245 pg/L 0.396 pg/L 0.359 pg/L 0.283 pg/L 0.174 pg/L 0.519 pg/L 0.733 pg/L	0.373U pg/L 0.352U pg/L 0.797U pg/L 0.532U pg/L 0.245U pg/L 0.396U pg/L 0.359U pg/L 0.283U pg/L 0.174U pg/L 0.519U pg/L 0.733U pg/L

Sample	Compound	Reported Concentration	Modified Final Concentration
SL-512-SA8-SB-0.0-0.5	1,2,3,4,7,8-HxCDD 2,3,4,6,7,8-HxCDF	0.156 ng/Kg 0.184 ng/Kg	0.156U ng/Kg 0.184U ng/Kg
SL-514-SA8-SB-0.0-0.5	1,2,3,4,7,8-HxCDD 2,3,4,6,7,8-HxCDF	0.164 ng/Kg 0.141 ng/Kg	0.164U ng/Kg 0.141U ng/Kg
SL-514-SA8-SB-4.0-5.0	1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF	0.0729 ng/Kg 0.133 ng/Kg 0.864 ng/Kg 0.0270 ng/Kg 0.0234 ng/Kg 0.0427 ng/Kg 0.0499 ng/Kg 0.0864 ng/Kg	0.0729U ng/Kg 0.133U ng/Kg 0.864U ng/Kg 0.0270U ng/Kg 0.0234U ng/Kg 0.0427U ng/Kg 0.0499U ng/Kg 0.0864U ng/Kg
SL-515-SA8-SB-0.0-0.5	1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 2,3,4,6,7,8-HxCDF	0.142 ng/Kg 0.0707 ng/Kg 0.169 ng/Kg	0.142U ng/Kg 0.0707U ng/Kg 0.169 ng/Kg
SL-515-SA8-SB-4.0-5.0	1,2,3,4,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.0544 ng/Kg 0.115 ng/Kg 0.0770 ng/Kg 0.658 ng/Kg 0.125 ng/Kg 0.0599 ng/Kg 0.0461 ng/Kg 0.0345 ng/Kg 0.0755 ng/Kg 0.0819 ng/Kg 0.135 ng/Kg	0.0544U ng/Kg 0.115U ng/Kg 0.0770U ng/Kg 0.658U ng/Kg 0.125U ng/Kg 0.0599U ng/Kg 0.0461U ng/Kg 0.0345U ng/Kg 0.0755U ng/Kg 0.0819U ng/Kg 0.135U ng/Kg
SL-515-SA8-SB-9.0-10.0	1,2,3,4,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF	0.0603 ng/Kg 0.120 ng/Kg 0.121 ng/Kg 0.562 ng/Kg 0.152 ng/Kg 0.0653 ng/Kg 0.145 ng/Kg 0.0679 ng/Kg 0.129 ng/Kg	0.0603U ng/Kg 0.120U ng/Kg 0.121U ng/Kg 0.562U ng/Kg 0.152U ng/Kg 0.0653U ng/Kg 0.145U ng/Kg 0.0679U ng/Kg 0.129U ng/Kg
SL-515-SA8-SB-14.0-15.0	1,2,3,4,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF	0.194 ng/Kg 0.121 ng/Kg 0.106 ng/Kg 0.675 ng/Kg 0.134 ng/Kg 0.0865 ng/Kg 0.116 ng/Kg	0.194U ng/Kg 0.121U ng/Kg 0.106U ng/Kg 0.675U ng/Kg 0.134U ng/Kg 0.0865U ng/Kg 0.116U ng/Kg
SL-513-SA8-SB-4.0-5.0	1,2,3,4,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF	0.0567 ng/Kg 0.0790 ng/Kg 0.151 ng/Kg 0.0433 ng/Kg 0.0453 ng/Kg 0.0507 ng/Kg 0.0828 ng/Kg	0.0567U ng/Kg 0.0790U ng/Kg 0.151U ng/Kg 0.0433U ng/Kg 0.0453U ng/Kg 0.0507U ng/Kg 0.0828U ng/Kg

Sample	Compound	Reported Concentration	Modified Final Concentration
SL-513-SA8-SB-8.0-9.0	1,2,3,7,8,9-HxCDD	0.0460 ng/Kg	0.0460U ng/Kg
	1,2,3,4,6,7,8-HpCDD	0.0671 ng/Kg	0.0671U ng/Kg
	OCDD	0.192 ng/Kg	0.192U ng/Kg
	1,2,3,4,7,8-HxCDF	0.0311 ng/Kg	0.0311U ng/Kg
	1,2,3,6,7,8-HxCDF	0.0177 ng/Kg	0.0177U ng/Kg
	2,3,4,6,7,8-HxCDF	0.0189 ng/Kg	0.0189U ng/Kg
	1,2,3,7,8,9-HxCDF	0.0393 ng/Kg	0.0393U ng/Kg
	1,2,3,4,6,7,8-HpCDF	0.0293 ng/Kg	0.0293U ng/Kg
	OCDF	0.126 ng/Kg	0.126U ng/Kg

Samples EB1-071713 and EB2-071713 were identified as equipment blanks. No polychlorinated dioxin/dibenzofuran contaminants were found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
EB1-071713	7/17/13	1,2,3,4,7,8-HxCDD	0.161 pg/L	SL-515-SA8-SB-0.0-0.5
		1,2,3,4,6,7,8-HpCDD	0.413 pg/L	SL-515-SA8-SB-4.0-5.0
		OCDD	0.814 pg/L	SL-515-SA8-SB-9.0-10.0
		1,2,3,7,8-PeCDF	0.406 pg/L	SL-515-SA8-SB-14.0-15.0
		2,3,4,7,8-PeCDF	0.132 pg/L	SL-513-SA8-SB-0.0-0.5
		1,2,3,4,7,8-HxCDF	0.222 pg/L	SL-513-SA8-SB-4.0-5.0
		1,2,3,6,7,8-HxCDF	0.191 pg/L	SL-513-SA8-SB-8.0-9.0
		2,3,4,6,7,8-HxCDF	0.224 pg/L	
		1,2,3,7,8,9-HxCDF	0.186 pg/L	
		1,2,3,4,6,7,8-HpCDF	0.238 pg/L	
		1,2,3,4,7,8,9-HpCDF	0.191 pg/L	
		OCDF	0.512 pg/L	
EB2-071713	7/17/13	1,2,3,7,8-PeCDD	0.542 pg/L	SL-512-SA8-SB-0.0-0.5
		1,2,3,4,7,8-HxCDD	0.373 pg/L	SL-514-SA8-SB-0.0-0.5
		1,2,3,4,6,7,8-HpCDD	0.352 pg/L	SL-514-SA8-SB-4.0-5.0
		OCDD	0.797 pg/L	
		1,2,3,7,8-PeCDF	0.532 pg/L	
		2,3,4,7,8-PeCDF	0.245 pg/L	
		1,2,3,6,7,8-HxCDF	0.396 pg/L	
		2,3,4,6,7,8-HxCDF	0.359 pg/L	
		1,2,3,7,8,9-HxCDF	0.283 pg/L	
		1,2,3,4,6,7,8-HpCDF	0.174 pg/L	
		1,2,3,4,7,8,9-HpCDF	0.519 pg/L	
		OCDF	0.733 pg/L	

Sample FB-041113 (from SDG PH029) was identified as a field blank. No polychlorinated dioxin/dibenzofuran contaminants were found with the following exceptions:

Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB-041113	4/11/13	1,2,3,4,7,8-HxCDD	0.125 pg/L	SL-512-SA8-SB-0.0-0.5
		1,2,3,7,8,9-HxCDD	0.134 pg/L	SL-514-SA8-SB-0.0-0.5
		1,2,3,4,6,7,8-HpCDD	0.402 pg/L	SL-514-SA8-SB-4.0-5.0
		1,2,3,7,8-PeCDF	0.398 pg/L	SL-515-SA8-SB-0.0-0.5
		2,3,4,7,8-PeCDF	0.316 pg/L	SL-515-SA8-SB-4.0-5.0
		1,2,3,4,7,8-HxCDF	0.324 pg/L	SL-515-SA8-SB-9.0-10.0
		1,2,3,6,7,8-HxCDF	0.221 pg/L	SL-515-SA8-SB-14.0-15.0
		1,2,3,7,8,9-HxCDF	0.211 pg/L	SL-513-SA8-SB-0.0-0.5
		2,3,4,6,7,8-HxCDF	0.149 pg/L	SL-513-SA8-SB-4.0-5.0
		1,2,3,4,6,7,8-HpCDF	0.254 pg/L	SL-513-SA8-SB-8.0-9.0
		OCDF	0.840 pg/L	

Sample concentrations were compared to concentrations detected in the field blanks. The sample concentrations were either not detected or were significantly greater (>5X for other contaminants) than the concentrations found in the associated field blanks.

VI. Matrix Spike/Matrix Spike Duplicates

The laboratory has indicated that there were no matrix spike (MS) and matrix spike duplicate (MSD) analyses specified for the samples in this SDG, and therefore matrix spike and matrix spike duplicate analyses were not performed for this SDG.

VII. Ongoing Precision Recovery (OPR)

Ongoing precision recovery samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Regional Quality Assurance and Quality Control

Not applicable.

IX. Internal Standards

All internal standard recoveries were within QC limits.

X. Target Compound Identifications

All target compound identifications were within validation criteria.

XI. Compound Quantitation

All compound quantitations were within validation criteria.

The 2,3,7,8-TCDF confirmation was performed with the following exceptions:

Sample	Compound	Finding	Criteria	Flag	A or P
SL-514-SA8-SB-0.0-0.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5	2,3,7,8-TCDF	2nd column confirmation was not performed for this compound.	2,3,7,8-TCDF must be confirmed on the 2nd column per the method.	None	P

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported below the RL.	J (all detects)	A

XII. System Performance

The system performance was acceptable.

XIII. Overall Assessment of Data

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Dioxins/Dibenzofurans - Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH069	SL-514-SA8-SB-0.0-0.5 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5	2,3,7,8-TCDF	None	P	Compound quantitation (no confirmation analysis)
PH069	EB1-071713 EB2-071713 SL-512-SA8-SB-0.0-0.5 SL-514-SA8-SB-0.0-0.5 SL-514-SA8-SB-4.0-5.0 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Dioxins/Dibenzofurans - Laboratory Blank Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
PH069	EB1-071713	1,2,3,4,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.161U pg/L 0.413U pg/L 0.814U pg/L 0.406U pg/L 0.132U pg/L 0.222U pg/L 0.191U pg/L 0.224U pg/L 0.186U pg/L 0.238U pg/L 0.191U pg/L 0.512U pg/L	A	B
PH069	EB2-071713	1,2,3,4,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.373U pg/L 0.352U pg/L 0.797U pg/L 0.532U pg/L 0.245U pg/L 0.396U pg/L 0.359U pg/L 0.283U pg/L 0.174U pg/L 0.519U pg/L 0.733U pg/L	A	B
PH069	SL-512-SA8-SB-0.0-0.5	1,2,3,4,7,8-HxCDD 2,3,4,6,7,8-HxCDF	0.156U ng/Kg 0.184U ng/Kg	A	B

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
PH069	SL-514-SA8-SB-0.0-0.5	1,2,3,4,7,8-HxCDD 2,3,4,6,7,8-HxCDF	0.164U ng/Kg 0.141U ng/Kg	A	B
PH069	SL-514-SA8-SB-4.0-5.0	1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF	0.0729U ng/Kg 0.133U ng/Kg 0.864U ng/Kg 0.0270U ng/Kg 0.0234U ng/Kg 0.0427U ng/Kg 0.0499U ng/Kg 0.0864U ng/Kg	A	B
PH069	SL-515-SA8-SB-0.0-0.5	1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDF 2,3,4,6,7,8-HxCDF	0.142U ng/Kg 0.0707U ng/Kg 0.169U ng/Kg	A	B
PH069	SL-515-SA8-SB-4.0-5.0	1,2,3,4,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	0.0544U ng/Kg 0.115U ng/Kg 0.0770U ng/Kg 0.658U ng/Kg 0.125U ng/Kg 0.0599U ng/Kg 0.0461U ng/Kg 0.0345U ng/Kg 0.0755U ng/Kg 0.0819U ng/Kg 0.135U ng/Kg	A	B
PH069	SL-515-SA8-SB-9.0-10.0	1,2,3,4,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF	0.0603U ng/Kg 0.120U ng/Kg 0.121U ng/Kg 0.562U ng/Kg 0.152U ng/Kg 0.0653U ng/Kg 0.145U ng/Kg 0.0679U ng/Kg 0.129U ng/Kg	A	B
PH069	SL-515-SA8-SB-14.0-15.0	1,2,3,4,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF	0.194U ng/Kg 0.121U ng/Kg 0.106U ng/Kg 0.675U ng/Kg 0.134U ng/Kg 0.0865U ng/Kg 0.116U ng/Kg	A	B
PH069	SL-513-SA8-SB-4.0-5.0	1,2,3,4,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF	0.0567U ng/Kg 0.0790U ng/Kg 0.151U ng/Kg 0.0433U ng/Kg 0.0453U ng/Kg 0.0507U ng/Kg 0.0828U ng/Kg	A	B

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
PH069	SL-513-SA8-SB-8.0-9.0	1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF	0.0460U ng/Kg 0.0671U ng/Kg 0.192U ng/Kg 0.0311U ng/Kg 0.0177U ng/Kg 0.0189U ng/Kg 0.0393U ng/Kg 0.0293U ng/Kg 0.126U ng/Kg	A	B

**Santa Susana Field Laboratory
Dioxins/Dibenzofurans - Field Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

LDC #: 30289D21
 SDG #: PH069
 Laboratory: Eurofins

VALIDATION COMPLETENESS WORKSHEET

Level IV

Date: 9-3-13
 Page: 1 of 1
 Reviewer: *OM*
 2nd Reviewer: *E*

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	A	Sampling dates: 7/17/13
II.	HRGC/HRMS Instrument performance check	A	
III.	Initial calibration	A	≤ 20/35
IV.	Continuing Calibration	A	QC limits
V.	Blanks	SW	
VI.	Matrix spike/Matrix spike duplicates	N	client
VII.	Laboratory control samples	A	OPR
VIII.	Regional quality assurance and quality control	N	
IX.	Internal standards	A	
X.	Target compound identifications	A	
XI.	Compound quantitation/RL/LOQ/LODs	SW	
XII.	System performance	A	
XIII.	Overall assessment of data	A	
XIV.	Field duplicates	N	
XV.	Field blanks	SW	EB=1,2; FB=FB-041113(PH029)

Note: A = Acceptable ND = No compounds detected D = Duplicate
 N = Not provided/applicable R = Rinsate TB = Trip blank
 SW = See worksheet FB = Field blank EB = Equipment blank

Validated Samples: 2 water / 10 Soil

1	EB1-071713	11	SL-513-SA8-SB-4.0-5.0	21		31	
2	EB2-071713	12	SL-513-SA8-SB-8.0-9.0	22		32	
3	SL-512-SA8-SB-0.0-0.5	13		23		33	
4	SL-514-SA8-SB-0.0-0.5	14		24		34	
5	SL-514-SA8-SB-4.0-5.0	15		25		35	
6	SL-515-SA8-SB-0.0-0.5	16		26		36	
7	SL-515-SA8-SB-4.0-5.0	17		27		37	
8	SL-515-SA8-SB-9.0-10.0	18		28		38	
9	SL-515-SA8-SB-14.0-15.0	19		29	BLK 2 00001	39	
10	SL-513-SA8-SB-0.0-0.5	20		30	BIK 206001	40	

Notes: _____

VALIDATION FINDINGS CHECKLIST

Method: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	/			
Cooler temperature criteria was met.	/			
II. GC/MS Instrument performance check				
Was PFK exact mass 380.9760 verified?	/			
Were the retention time windows established for all homologues?	/			
Was the chromatographic resolution between 2,3,7,8-TCDD and peaks representing any other unlabeled TCDD isomers $\leq 25\%$?	/			
Is the static resolving power at least 10,000 (10% valley definition)?	/			
Was the mass resolution adequately check with PFK?	/			
Was the presence of 1,2,8,9-TCDD and 1,3,4,6,8-PeCDF verified?	/			
III. Initial calibration				
Was the initial calibration performed at 5 concentration levels?	/			
Were all percent relative standard deviations (%RSD) $\leq 20\%$ for unlabeled compounds and $\leq 35\%$ for labeled compounds ?	/			
Did all calibration standards meet the Ion Abundance Ratio criteria?	/			
Was the signal to noise ratio for each target compound ≥ 2.5 and for each recovery and internal standard > 10 ?	/			
IV. Continuing calibration				
Was a routine calibration performed at the beginning and end of each 12 hour period?	/			
Were all the concentrations for the unlabeled compounds and labeled compounds within the QC limits (Method 1613B, Table 6)?	/			
Did all routine calibration standards meet the Ion Abundance Ratio criteria?	/			
V. Blanks				
Was a method blank associated with every sample in this SDG?	/			
Was a method blank performed for each matrix and concentration?	/			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet?	/			
VI. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.		/		
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?			/	
VII. Laboratory control samples				
Was an LCS analyzed for this SDG?	/			
Was an LCS analyzed per extraction batch?	/			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	/			

VALIDATION FINDINGS CHECKLIST

Validation Area	Yes	No	NA	Findings/Comments
VIII. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?		/		
Were the performance evaluation (PE) samples within the acceptance limits?			/	
IX. Internal standards				
Were internal standard recoveries within the 25-150% criteria?	/			
Was the minimum S/N ratio of all internal standard peaks ≥ 10 ?	/			
X. Target compound identification				
For 2,3,7,8 substituted congeners with associated labeled standards, were the retention times of the two quantitation peaks within -1 to 3 sec. of the RT of the labeled standard?	/			
For 2,3,7,8 substituted congeners without associated labeled standards, were the relative retention times of the two quantitation peaks within 0.005 time units of the RRT measured in the routine calibration?			/	
For non-2,3,7,8 substituted congeners, were the retention times of the two quantitation peaks within RT established in the performance check solution?			/	
Did compound spectra contain all characteristic ions listed in the table attached?	/			
Was the Ion Abundance Ratio for the two quantitation ions within criteria?		/		
Was the signal to noise ratio for each target compound and labeled standard ≥ 2.5 ?	/			
Does the maximum intensity of each specified characteristic ion coincide within ± 2 seconds (includes labeled standards)?	/			
For PCDF identification, was any signal ($S/N \geq 2.5$, at \pm seconds RT) detected in the corresponding PCDPE channel?		/		
Was an acceptable lock mass recorded and monitored?	/			
XI. Compound quantitation/CRQLs				
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	/			
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	/			
XII. System performance				
System performance was found to be acceptable.	/			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	/			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		/		
Target compounds were detected in the field duplicates.			/	
XV. Field blanks				
Field blanks were identified in this SDG.	/			
Target compounds were detected in the field blanks.	/			

VALIDATION FINDINGS WORKSHEET

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

A. 2,3,7,8-TCDD	F. 1,2,3,4,6,7,8-HpCDD	K. 1,2,3,4,7,8-HxCDF	P. 1,2,3,4,7,8,9-HpCDF	U. Total HpCDD
B. 1,2,3,7,8-PeCDD	G. OCDD	L. 1,2,3,6,7,8-HxCDF	Q. OCDF	V. Total TCDF
C. 1,2,3,4,7,8-HxCDD	H. 2,3,7,8-TCDF	M. 2,3,4,6,7,8-HxCDF	R. Total TCDD	W. Total PeCDF
D. 1,2,3,6,7,8-HxCDD	I. 1,2,3,7,8-PeCDF	N. 1,2,3,7,8,9-HxCDF	S. Total PeCDD	X. Total HxCDF
E. 1,2,3,7,8,9-HxCDD	J. 2,3,4,7,8-PeCDF	O. 1,2,3,4,6,7,8-HpCDF	T. Total HxCDD	Y. Total HpCDF

Notes: _____

VALIDATION FINDINGS WORKSHEET

Blanks

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- Y N N/A Were all samples associated with a method blank?
- Y N N/A Was a method blank performed for each matrix and whenever a sample extraction was performed?
- Y N N/A Was the method blank contaminated?

Blank extraction date: 07/19/13 Blank analysis date: 07/19/13

Associated samples: 1-2 Qual U (B)

Conc. units: pg/L

Compound	Blank ID	Sample Identification							
		5X	1	2					
	BLK200001								
C	0.368*	1.84	0.161*	0.373*					
D	0.364*	1.82							
E	0.327*	1.64							
F	0.373*	1.87	0.413*	0.352*					
G	1.18	5.90	0.814*	0.797*					
I	0.465	2.33	0.406	0.532					
J	0.511*	2.56	0.132*	0.245					
K	0.527*	2.64	0.222*						
L	0.476*	2.38	0.191*	0.396*					
M	0.238*	1.19	0.224	0.359*					
N	0.586*	2.93	0.186	0.283*					
O	0.413	2.07	0.238	0.174*					
P	0.325*	1.63	0.191*	0.519*					
Q	0.838*	4.19	0.512*	0.733*					

*EMPC

All contaminants within five times the method blank concentration were qualified as not detected, "U".

VALIDATION FINDINGS WORKSHEET

Blanks

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

- N N/A Were all samples associated with a method blank?
- N N/A Was a method blank performed for each matrix and whenever a sample extraction was performed?
- N N/A Was the method blank contaminated?

Blank extraction date: 07/26/13 Blank analysis date: 07/26/13

Associated samples: 3-12 Qual U (B)

Conc. units: ~~pg/L~~ ng/kg

Compound	Blank ID	Sample Identification										
		5X	3	4	5	6	7	8	9	11	12	
	BLK206001											
C	0.0432*	0.216	0.156*	0.164		0.142	0.0544*	0.0603*	0.194*	0.0567*		
E	0.0358*	0.179			0.0729		0.115*	0.120	0.121	0.0790	0.0460	
F	0.0714*	0.357			0.133*		0.0770*	0.121*	0.106*		0.0671*	
G	0.230	1.150			0.864		0.658	0.562*	0.675		0.192*	
I	0.0307*	0.154					0.125*	0.152		0.151		
K	0.0229	0.115			0.0270*	0.0707	0.0599*			0.0433*	0.0311*	
L	0.0231*	0.116					0.0461			0.0453*	0.0177*	
M	0.0401*	0.201	0.184*	0.141	0.0234	0.169*	0.0345*	0.0653*		0.0507*	0.0189*	
N	0.0413*	0.207			0.0427*		0.0755*	0.145*	0.134*	0.0828	0.0393*	
O	0.0256*	0.128			0.0499*		0.0819*	0.0679	0.0865		0.0293*	
Q	0.159	0.795			0.0864*		0.135*	0.129*	0.116*		0.126	

*EMPC

All contaminants within five times the method blank concentration were qualified as not detected, "U".

VALIDATION FINDINGS WORKSHEET

Field Blanks

METHOD: HRGC/HRMS Dioxins (EPA Method 1613B)

Blank units: pg/L **Associated sample units:** ng/kg

Sampling date: 04/11/13

Field blank type: (circle one) Field Blank / Rinsate / Other: FB Associated Samples: 3-12 >5x

Compound	Blank ID	Sample Identification							
		5X							
	FB-041113	5X							
C	0.125	0.625							
E	0.134*	0.67							
F	0.402*	2.01							
I	0.398*	1.99							
J	0.316*	1.58							
K	0.324	1.62							
L	0.221	1.105							
N	0.211*	1.055							
M	0.149	0.745							
O	0.254*	1.27							
Q	0.840*	4.2							

* EMPC
 FB-041113 (PH029)

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
 All contaminants within five times the blank concentration were qualified as not detected, "U".

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: HRGC/HRMS Dioxins (EPA Method 1613B)

Blank units: pg/L **Associated sample units:** ng/kg

Sampling date: 07/17/13

Field blank type: (circle one) Field Blank / Rinsate / Other: EB Associated Samples: 6-12 >5x

Compound	Blank ID	Sample Identification							
		5X							
	EB1-071713	5X							
C	0.161*	0.805							
F	0.413*	2.07							
G	0.814*	4.07							
I	0.406	2.03							
J	0.132*	0.660							
K	0.222*	1.11							
L	0.191*	0.955							
M	0.224	1.12							
N	0.186	0.930							
O	0.238	1.19							
P	0.191*	0.955							
Q	0.512*	2.56							

* EMPC

EB1-071713 (PH069)

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
All contaminants within five times the blank concentration were qualified as not detected, "U".

VALIDATION FINDINGS WORKSHEET
Field Blanks

METHOD: HRGC/HRMS Dioxins (EPA Method 1613B)

Blank units: pg/L **Associated sample units:** ng/kg

Sampling date: 07/17/13

Field blank type: (circle one) Field Blank / Rinsate / Other: EB Associated Samples: 3-5 >5x

Compound	Blank ID	Sample Identification							
	EB2-071713	5X							
B	0.542*	2.71							
C	0.373*	1.87							
F	0.352*	1.76							
G	0.797*	3.99							
I	0.532	2.66							
J	0.245	1.23							
L	0.396*	1.98							
M	0.359*	1.80							
N	0.283*	1.42							
O	0.174*	0.870							
P	0.519*	2.60							
Q	0.733*	3.67							

* EMPC

EB2-071713 (PH069)

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
All contaminants within five times the blank concentration were qualified as not detected, "U".

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

The Relative Response Factor (RRF), average RRF, and percent relative standard deviation (%RSD) were recalculated for the compounds identified below using the following calculations:

$$RRF = (A_x)(C_{is}) / (A_{is})(C_x)$$

average RRF = sum of the RRFs/number of standards

$$\%RSD = 100 * (S/X)$$

A_x = Area of compound,

C_x = Concentration of compound,

S = Standard deviation of the RRFs,

A_{is} = Area of associated internal standard

C_{is} = Concentration of internal standard

X = Mean of the RRFs

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Reported	Recalculated	Reported	Recalculated	Reported	Recalculated
				Average RRF (initial)	Average RRF (initial)	RRF (CS3 std)	RRF (CS3 std)	%RSD	%RSD
1	ICAL (DBSMS)	03/22/2013	2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	1.043	1.043	1.025	1.024	5.23	5.24
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	1.207	1.207	1.182	1.182	9.04	9.04
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	0.957	0.9575	0.947	0.947	2.62	2.62
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	1.048	1.048	1.028	1.028	3.46	3.47
			OCDF (¹³ C-OCDF)	0.933	0.933	0.920	0.920	5.76	5.76
2	ICAL (SP233)	3-23-13	2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	0.968	0.968	0.993	0.993	3.96	3.94
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)						
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)						
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)						
			OCDF (¹³ C-OCDF)						
3	ICAL (DBSMS)	6-19-13	2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	1.056	1.056	1.028	1.028	7.30	7.31
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	1.196	1.196	1.107	1.107	10.82	10.80
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	0.955	0.955	0.954	0.954	1.24	1.22
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	1.060	1.0605	1.034	1.034	8.10	8.10
			OCDF (¹³ C-OCDF)	0.962	0.962	0.959	0.959	4.62	4.60

Comments: Refer to Initial Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Routine Calibration Results Verification

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

$$\% \text{ Difference} = 100 * (\text{ave. RRF} - \text{RRF}) / \text{ave. RRF}$$

$$\text{RRF} = (A_x)(C_s) / (A_s)(C_x)$$

Where: ave. RRF = initial calibration average RRF
 RRF = continuing calibration RRF
 A_x = Area of compound, A_s = Area of associated internal standard
 C_x = Concentration of compound, C_s = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Spiked Conc (ng/mL)	Reported	Recalculated	Reported	Recalculated
					Conc (ng/mL)	Conc (ng/mL)	%R	%R
1	CS3CC02 (DBSMS)	7-19-13	2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	10	9.360	9.360	94	94
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	10	9.410	9.411	94	94
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	50	52.040	52.062	104	104
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	50	51.060	51.050	102	102
			OCDF (¹³ C-OCDF)	100	102.120	102.111	102	102
2	CS3CC02 (SP2331)	7-30-13	2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	10	10.970	10.962	110	110
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	10				
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	50				
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	50				
			OCDF (¹³ C-OCDF)	100				
3	CS3CC02 (DBSMS)	7-26-13	2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	10	10.000	10.000	100	100
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	10	10.250	10.243	102	102.5
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	50	51.990	51.982	104	104
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	50	49.590	49.598	99	99
			OCDF (¹³ C-OCDF)	100	99.390	99.387	99	99

Comments: Refer to Routine Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Routine Calibration Results Verification

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 1613B)

The percent difference (%D) of the initial calibration average Relative Response Factors (RRFs) and the continuing calibration RRFs were recalculated for the compounds identified below using the following calculation:

% Difference = 100 * (ave. RRF - RRF)/ave. RRF
 RRF = (A_x)(C_s)/(A_s)(C_x)

Where: ave. RRF = initial calibration average RRF
 RRF = continuing calibration RRF
 A_x = Area of compound, A_s = Area of associated internal standard
 C_x = Concentration of compound, C_s = Concentration of internal standard

#	Standard ID	Calibration Date	Compound (Reference Internal Standard)	Spiked Conc (ng/mL)	Reported	Recalculated	Reported	Recalculated
					Conc (ng/mL)	Conc (ng/mL)	%R	%R
1	CS3CC04 (DBSMS)	7-27-13	2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	10	9.370	9.368	94	94
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	10	9.960	9.956	100	100
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	50	48.620	48.610	97	97
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	50	47.850	47.857	96	96
			OCDF (¹³ C-OCDF)	100	92.730	92.731	93	93
2			2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	10				
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	10				
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	50				
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	50				
			OCDF (¹³ C-OCDF)	100				
3			2,3,7,8-TCDF (¹³ C-2,3,7,8-TCDF)	10				
			2,3,7,8-TCDD (¹³ C-2,3,7,8-TCDD)	10				
			1,2,3,6,7,8-HxCDD (¹³ C-1,2,3,6,7,8-HxCDD)	50				
			1,2,3,4,6,7,8-HpCDD (¹³ C-1,2,4,6,7,8,-HpCDD)	50				
			OCDF (¹³ C-OCDF)	100				

Comments: Refer to Routine Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name: Santa Susana Field Laboratory
Collection Date: July 17, 2013
LDC Report Date: September 23, 2013
Matrix: Soil/Water
Parameters: Perchlorate
Validation Level: Level IV
Laboratory: Eurofins Lancaster Laboratories
Sample Delivery Group (SDG): PH069

Sample Identification

EB1-071713
SL-515-SA8-SB-0.0-0.5
SL-515-SA8-SB-4.0-5.0
SL-515-SA8-SB-9.0-10.0
SL-515-SA8-SB-14.0-15.0
SL-513-SA8-SB-0.0-0.5
SL-513-SA8-SB-4.0-5.0
SL-513-SA8-SB-8.0-9.0
EB1-071713MS
EB1-071713MSD

Introduction

This data review covers 9 soil samples and one water sample listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 6850 for Perchlorate.

This review follows the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008).

A qualification summary table is provided at the end of this report if data has been qualified. Flags are classified as P (protocol) or A (advisory) to indicate whether the flag is due to a laboratory deviation from a specified protocol or is of technical advisory nature.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- A Indicates the finding is based upon technical validation criteria.
- P Indicates the finding is related to a protocol/contractual deviation.
- None Indicates the data was not significantly impacted by the finding, therefore qualification was not required.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met validation criteria.

II. LC/MS Instrument Performance Check

Instrument performance was checked as applicable. All ion abundance requirements were met.

III. Initial Calibration

Initial calibration was performed using required standard concentrations.

A curve fit, based on the initial calibration, was established for quantitation. The coefficient of determination (r^2) was greater than or equal to 0.990 .

IV. Continuing Calibration

Continuing calibration was performed at the required frequencies.

All of the continuing calibration percent differences (%D) between the initial calibration RRF and the continuing calibration RRF were less than or equal to 15.0% for unlabeled compounds and less than or equal to 50.0% for labeled compounds.

The percent differences (%D) of the second source calibration standard were less than or equal to 15.0% for unlabeled compounds and less than or equal to 50.0% for labeled compounds.

The percent differences (%D) of the limit of detection verification (LODV) calibration standard were less than or equal to 50.0% for perchlorate.

V. Blanks

Method blanks were reviewed for each matrix as applicable. No perchlorate was found in the method blanks.

Sample EB1-071713 was identified as an equipment blank. No perchlorate was found in the equipment blank.

Sample FB-041113 (from SDG PH029) was identified as a field blank. No perchlorate was found in the field blank.

VI. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD)

VII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) were within QC limits.

VIII. Regional Quality Assurance and Quality Control

Not applicable.

IX. Internal Standards

All internal standard areas and retention times were within QC limits.

X. Target Compound Identifications

All target compound identifications were within validation criteria.

XI. Compound Quantitation

All compound quantitations were within validation criteria.

All compounds reported below the RL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH069	All compounds reported below the RL.	J (all detects)	A

XII. System Performance

The system performance was acceptable.

XIII. Overall Assessment

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

No field duplicates were identified in this SDG.

**Santa Susana Field Laboratory
Perchlorate - Data Qualification Summary - SDG PH069**

SDG	Sample	Compound	Flag	A or P	Reason (Code)
PH069	EB1-071713 SL-515-SA8-SB-0.0-0.5 SL-515-SA8-SB-4.0-5.0 SL-515-SA8-SB-9.0-10.0 SL-515-SA8-SB-14.0-15.0 SL-513-SA8-SB-0.0-0.5 SL-513-SA8-SB-4.0-5.0 SL-513-SA8-SB-8.0-9.0	All compounds reported below the RL.	J (all detects)	A	Compound quantitation (Z)

**Santa Susana Field Laboratory
Perchlorate - Laboratory Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

**Santa Susana Field Laboratory
Perchlorate - Field Blank Data Qualification Summary - SDG PH069**

No Sample Data Qualified in this SDG

METHOD: LC/MS Perchlorate (EPA SW846 Method 6850)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	Δ	Sampling dates: 7/22/13 7/17/13
II.	GC/MS Instrument performance check	Δ	
III.	Initial calibration	Δ	
IV.	Continuing calibration/ICV	Δ	100/100 ^{r²} = 15/50 LODV ≤ 50
V.	Blanks	Δ	
VI.	Surrogate spikes	N	W
VII.	Matrix spike/Matrix spike duplicates	Δ	
VIII.	Laboratory control samples	A	LCS
IX.	Regional Quality Assurance and Quality Control	N	
X.	Internal standards	Δ	
XI.	Target compound identification	Δ	
XII.	Compound quantitation/RL/LOQ/LODs	Δ	
XIII.	Tentatively identified compounds (TICs)	N	W
XIV.	System performance	Δ	
XV.	Overall assessment of data	Δ	
XVI.	Field duplicates	N	
XVII.	Field blanks	ND	EB = 1 FB = FB-04113 (SDG # PH029)

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

D = Duplicate
TB = Trip blank
EB = Equipment blank

Validated Samples:

SOIL & WATER

12	EB1-071713	W	111	PBLK14201	21		31
2	SL-515-SA8-SB-0.0-0.5		122	PBLK03203	22		32
3	SL-515-SA8-SB-4.0-5.0		13		23		33
4	SL-515-SA8-SB-9.0-10.0		14		24		34
5	SL-515-SA8-SB-14.0-15.0		15		25		35
6	SL-513-SA8-SB-0.0-0.5		16		26		36
7	SL-513-SA8-SB-4.0-5.0		17		27		37
8	SL-513-SA8-SB-8.0-9.0		18		28		38
9	#1 MS		19		29		39
10	#1 MSD		20		30		40

LDC #: 30289087

VALIDATION FINDINGS CHECKLIST

Page: 1 of 2
 Reviewer: F7
 2nd Reviewer: A

Method: GC LC/MS

Validation Area	Yes	No	NA	Findings/Comments
I. Technical holding times				
All technical holding times were met.	<input checked="" type="checkbox"/>			
Cooler temperature criteria was met.	<input checked="" type="checkbox"/>			
II. LC/MS Instrument performance check				
Were the instrument performance reviewed and found to be within the specified criteria?	<input checked="" type="checkbox"/>			
Were all samples analyzed within the 12 hour clock criteria?	<input checked="" type="checkbox"/>			
III. Initial calibration				
Did the laboratory perform a 5 point calibration prior to sample analysis?	<input checked="" type="checkbox"/>			
Were all percent relative standard deviations (%RSD) < 20%?			<input checked="" type="checkbox"/>	
Was a curve fit used for evaluation?	<input checked="" type="checkbox"/>			
Did the initial calibration meet the curve fit acceptance criteria of > 0.990?	<input checked="" type="checkbox"/>			
Were the RT windows properly established?	<input checked="" type="checkbox"/>			
IV. Continuing calibration				
Was a continuing calibration analyzed daily?	<input checked="" type="checkbox"/>			
Were all percent differences (%D) < 15%?	<input checked="" type="checkbox"/>			
Were all the retention times within the acceptance windows?	<input checked="" type="checkbox"/>			
V. Blanks				
Was a method blank associated with every sample in this SDG?	<input checked="" type="checkbox"/>			
Was a method blank analyzed for each matrix and concentration?	<input checked="" type="checkbox"/>			
Was there contamination in the method blanks? If yes, please see the Blanks validation completeness worksheet.			<input checked="" type="checkbox"/>	
VI. Matrix spike/Matrix spike duplicates				
Were a matrix spike (MS) and matrix spike duplicate (MSD) analyzed for each matrix in this SDG? If no, indicate which matrix does not have an associated MS/MSD. Soil / Water.	<input checked="" type="checkbox"/>			
Was a MS/MSD analyzed every 20 samples of each matrix?	<input checked="" type="checkbox"/>			
Were the MS/MSD percent recoveries (%R) and the relative percent differences (RPD) within the QC limits?	<input checked="" type="checkbox"/>			
VII. Laboratory control samples				
Was an LCS analyzed for this SDG?	<input checked="" type="checkbox"/>			
Was an LCS analyzed per extraction batch?	<input checked="" type="checkbox"/>			
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?	<input checked="" type="checkbox"/>			
XIII. Regional Quality Assurance and Quality Control				
Were performance evaluation (PE) samples performed?			<input checked="" type="checkbox"/>	
Were the performance evaluation (PE) samples within the acceptance limits?			<input checked="" type="checkbox"/>	

Validation Area	Yes	No	NA	Findings/Comments
IX. Internal standards				
Were internal standard area counts within -50% or +100% of the associated calibration standard?	✓			
Were retention times within ± 30 seconds from the associated calibration standard?	✓			
X. Target compound identification				
Were relative retention times (RRT's) within + 0.06 RRT units of the standard?	✓			
Did compound spectra meet specified EPA "Functional Guidelines" criteria?	✓			
Were chromatogram peaks verified and accounted for?	✓			
XI. Compound quantitation/CRQLs				
Were the correct internal standard (IS), quantitation ion and relative response factor (RRF) used to quantitate the compound?	✓			
Were compound quantitation and CRQLs adjusted to reflect all sample dilutions and dry weight factors applicable to level IV validation?	✓			
XII. System performance				
System performance was found to be acceptable.	✓			
XIII. Overall assessment of data				
Overall assessment of data was found to be acceptable.	✓			
XIV. Field duplicates				
Field duplicate pairs were identified in this SDG.		✓		
Target compounds were detected in the field duplicates.			✓	
XV. Field blanks				
Field blanks were identified in this SDG.	✓			
Target compounds were detected in the field blanks.		✓		
Target compounds were detected in the field blanks. F7				

LDC#: 30289087
 SDG#: pe cover

VALIDATION FINDINGS WORKSHEET
Initial Calibration Calculation Verification

Page: 1 of 1
 Reviewer: FJ
 2nd Reviewer: A

Method: LC/MS Perchlorate (EPA SW846 Method 6850)

Calibration Date	System	Compound	Standard	(Y) Response	(X) Concentration
7/22/2013	LCMS	Perchlorate	0	0	0
			1	0.043230776	0.400
			2	0.099295788	1.000
			3	0.208311718	2.000
			4	0.441965729	4.000
			5	1.052918041	10.000
			6	2.722266856	25.000

Regression Output	Calculated	Reported
Constant	-0.006176	
Std Err of Y Est		
R Squared	0.999799	0.999500
Degrees of Freedom		
X Coefficient(s)	0.10875521	
Std Err of Coef.		
Correlation Coefficient	0.999899	
Coefficient of Determination (r ²)	0.999799	0.999500

LDC #: 30289087

VALIDATION FINDINGS WORKSHEET
Continuing Calibration Results Verification

Page: 1 of 1
 Reviewer: [Signature]
 2nd Reviewer: [Signature]

METHOD: LCMS Perchlorate (EPA SW 846 Method 6850)

The percent difference (%D) of the initial calibration average Calibration Factors (CF) and the continuing calibration CF were recalculated for the compounds identified below using the following calculation:

% Difference = 100 * (ave. CF - CF)/ave.CF

Where: ave. CF = initial calibration average CF
 CF = continuing calibration CF
 A = Area of compound
 C = Concentration of compound

#	Standard ID	Calibration Date	Compound	Average CF(Ical)/ CCV Conc.	Reported	Recalculated	Reported	Recalculated
					CF/ Conc. CCV	CF/ Conc. CCV	%D	%D
1	cew-3021	7/22/13	Perchlorate (LODV)	0.4	0.48	0.48	20	20
	- 3032	7/22/13	↓ (cew)	4.0	3.7	3.7	8	8
2	- 03043	7/23/13	↓ (LODV)	0.4	0.45	0.45	13	13
3	-3047		↓ (cew)	4	3.7	3.7	8	8 ↓ closing cew
4								

Comments: Refer to Continuing Calibration findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30289D87

VALIDATION FINDINGS WORKSHEET
Matrix Spike/Matrix Spike Duplicates Results Verification

Page: 6 / 1
 Reviewer: FJ
 2nd Reviewer: [Signature]

METHOD: LC/MS Perchlorate (EPA SW 846 Method 6850/6860)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the matrix spike and matrix spike duplicate were recalculated for the compounds identified below using the following calculation:

% Recovery = $100 * (SSC - SC) / SA$

Where: SSC = Spiked sample concentration
 SA = Spike added

SC = Sample concentration

RPD = $|MS - MSD| * 2 / (MS + MSD)$

MS = Matrix spike percent recovery

MSD = Matrix spike duplicate percent recovery

MS/MSD samples: 9 + 10

Compound	Spike Added (ug/L)		Sample Concentration (ug/L)	Spiked Sample Concentration (ug/L)		Matrix Spike		Matrix Spike Duplicate		MS/MSD	
	MS	MSD		MS	MSD	Percent Recovery		Percent Recovery		RPD	
						Reported	Recalc	Reported	Recalc	Reported	Recalculated
Perchlorate	5	5	ND	4.94	4.95	99	99	99	99	0	0

Comments: Refer to Matrix Spike/Matrix Spike Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

LDC #: 30289087

VALIDATION FINDINGS WORKSHEET

Laboratory Control Sample/Laboratory Control Sample Duplicates Results Verification

Page: 1 of 1
 Reviewer: FT
 2nd Reviewer: RA

METHOD: LC/MS Perchlorate (EPA SW 846 Method 6850/6860)

The percent recoveries (%R) and Relative Percent Difference (RPD) of the laboratory control sample and laboratory control sample duplicate were recalculated for the compounds identified below using the following calculation:

% Recovery = 100 * (SC/SA)

Where: SSC = Spike concentration
 SA = Spike added

RPD = |LCS - LCSD| * 2 / (LCS + LCSD)

LCS = Laboratory control sample percent recovery

LCSD = Laboratory control sample duplicate percent recovery

LCS/LCSD samples: LCS14201

Compound	Spike Added (<u>1 ug/kg</u>)		Spike Concentration (<u>1 ug/kg</u>)		LCS		LCSD		LCS/LCSD	
	LCS	LCSD	LCS	LCSD	Percent Recovery		Percent Recovery		RPD	
					Reported	Recalc	Reported	Recalc	Reported	Recalc
Perchlorate	100	NA	104.3	NA	104	104	NA			

Comments: Refer to Laboratory Control Sample/Laboratory Control Sample Duplicates findings worksheet for list of qualifications and associated samples when reported results do not agree within 10.0% of the recalculated results.

VALIDATION FINDINGS WORKSHEET
Sample Calculation Verification

METHOD: LCMS Perchlorate (EPA SW 846 Method 6850/6860)

Y N N/A
Y N N/A

Were all reported results recalculated and verified for all level IV samples?

Were all recalculated results for detected target compounds agree within 10.0% of the reported results?

$$\text{Concentration} = \frac{(A_x)(I_s)(V_i)(DF)(2.0)}{(A_{is})(RRF)(V_o)(V_t)(\%S)}$$

- A_x = Area of the characteristic ion (EICP) for the compound to be measured
- A_{is} = Area of the characteristic ion (EICP) for the specific internal standard
- I_s = Amount of internal standard added in nanograms (ng)
- V_o = Volume or weight of sample extract in milliliters (ml) or grams (g).
- V_i = Volume of extract injected in microliters (ul)
- V_t = Volume of the concentrated extract in microliters (ul)
- Df = Dilution Factor.
- %S = Percent solids, applicable to soil and solid matrices only.
- 2.0 = Factor of 2 to account for GPC cleanup

Example:

Sample I.D. #7, Perchlorate

$$\text{Conc.} = \frac{(\quad)(\quad)(\quad)(\quad)(\quad)}{(\quad)(\quad)(\quad)(\quad)(\quad)}$$

=

#	Sample ID	Compound	Reported Concentration ()	Calculated Concentration ()	Qualification
	<u>1508 =</u>	<u>(0.1081)(X) - 0.0029</u>			
	<u>46870</u>				
	<u>X = 0.324</u>				
	<u>final conc</u>	<u>= 0.324 x 10ml</u>	<u>= 3.605 ug/kg</u>		
		<u>(1g)(0.90)</u>			

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 8

SDG: PH070

Prepared for

CDM
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Prepared by

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September 24, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on July 18th, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Volatiles (VOAs) by EPA SW 846 Method 8260B
Semivolatiles (SVOAs) by EPA SW 846 Method 8270D
(SVOAs) by EPA SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A, and 7471B
Total Petroleum Hydrocarbon as Gasoline (TPH-G) by EPA SW 846 Method 8015M
Total Petroleum Hydrocarbons as Extractable (TPH-E) by EPA SW 846 Method 8015M
Dioxins/Dibenzofurans by EPA Method 1613B
Perchlorate by EPA method 6850
Hexavalent Chromium by EPA method 7199

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks, and field duplicate samples. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.

I. Technical Holding Times

All technical holding time requirements were met.

The chain-of-custodies were reviewed for documentation of cooler temperatures. All cooler temperatures met QC criteria.

II. Initial Calibration

Initial Calibration data were not reviewed for level III.

III. Continuing Calibration

Continuing calibration data were not reviewed for level III.

IV. Blanks

Method blanks were performed at the required frequencies. No contaminant concentrations were detected in the method or preparation blanks with the exception of three blanks for dioxins and metals. The associated sample results were qualified as non-detected (U) due to method blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the associated blanks were not qualified. The details regarding the qualification of data are provided in Enclosure I.

No contaminant concentrations were detected in the initial or continuing calibration blanks.

V. Surrogate Spikes

Surrogates were added to all samples and blanks as required by the method. All surrogate recoveries (%R) were within QC limits.

VI. ICP Interference Check Sample (ICS) Analysis

ICP interference check data were not reviewed for level III.

VII. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were performed at the required frequency. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of one MS/MSD pair for SVOAs, and TPH-E. The benzidine result in sample SL-506-SA8-SB-0.0-0.5 was qualified as rejected (R) due to MS/MSD %R grossly outside QC limits (i.e., $\leq 0\%$). The remainder of the associated sample results were qualified as detected estimated (J) as applicable. The details regarding the qualification of data are provided in Enclosure I.

VIII. Laboratory Duplicates Sample

Laboratory duplicates (DUP) sample analyses were reviewed for each matrix as applicable. Results were within QC limits.

IX. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Percent recoveries (%R) and relative percent differences (RPD) were within QC limits with the exception of several LCS/LCSD pairs for VOAs and SVOAs. The associated sample results were qualified as detected estimated (J) or non-detected estimated (UJ) as applicable. The details regarding LCS/LCSD outliers are provided in Enclosure I.

X. Internal Standards

Internal standards were reviewed for dioxins. Percent recoveries (%R) were within QC limits.

XI. ICP Serial Dilution

ICP serial dilution was not performed for this SDG.

XII. Compound Quantitation

The laboratory reporting limits were evaluated. All laboratory reporting limits met the specified requirements.

All compounds reported below the RL as detected were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG PH070	All compounds reported as detected below the RL.	J (all detects)	A

XIII. Field Duplicate Samples

No field duplicates were identified in this SDG.

XIV. Field Blank Samples

One trip blank was collected and analyzed for VOAs and TPH-G. No contaminant concentrations were found in the trip blank.

One equipment blank (from SDG PH069) was collected and analyzed for VOAs, SVOAs, PCBs, metals, perchlorate, TPH-G, TPH-E, hexavalent chromium, and dioxins. The equipment blank had detections for several VOAs, SVOAs, metals, and dioxins. All associated sample results were not detected or were significantly greater than the concentrations found in the equipment blanks and were not qualified.

One field blank (from SDG PH029) was collected and analyzed for VOAs, SVOAs, PCBs, metals, perchlorate, TPH-G, TPH-E, hexavalent chromium, and dioxins. The field blank had detections for several VOAs, SVOAs, metals, and dioxins. The associated sample results were qualified as non-detected (U) due to field blank contamination as applicable. The sample results that were not detected or were significantly greater than the concentrations found in the field blank were not qualified. The field blank outlier reports are presented in Enclosure I.

XV. Overall Assessment of Data

One SVOA result was rejected due to MS/MSD %Rs grossly outside of QC limits. These results are not useable for all purposes.

The overall assessment of QA/QC data review by automated and manual validation of this sampling event met project requirements and analytical completeness levels with the exceptions noted in the above sections. The remainder of the data are deemed useable for the intended use.

Data flags are summarized and are presented as Attachment 2.

Attachment 1
Sample Cross Reference

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
18-Jul-2013	SL-506-SA8-SB-0.0-0.5	7132364	N	3050B	6010C	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5	7132364	N	3050B	6020A	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5	7132364	N	3060A	7199	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5	7132364	N	3546	8015M	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5	7132364	N	3546	8082A	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5	7132364	N	3546	8270D	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5	7132364	N	3546	8270D SIM	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5	7132364	N	METHOD	1613B	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5	7132364	N	METHOD	6850	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5	7132364	N	METHOD	7471B	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5MSD	P132364M260558	MSD	3546	8270D SIM	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5MSD	P132364M262337	MSD	3546	8270D	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5MSD	P132364M322048A	MSD	3546	8015M	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5MS	P132364R260526	MS	3546	8270D SIM	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5MS	P132364R262312	MS	3546	8270D	III
18-Jul-2013	SL-506-SA8-SB-0.0-0.5MS	P132364R322027A	MS	3546	8015M	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	3050B	6010C	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	3050B	6020A	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	3060A	7199	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	3546	8015M	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	3546	8082A	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	3546	8270D	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	3546	8270D SIM	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	5035	8260B	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	5035A	8015M	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	METHOD	1613B	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	METHOD	6850	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0	7132365	N	METHOD	7471B	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0MSD	P132365M240232A	MSD	3546	8082A	III
18-Jul-2013	SL-506-SA8-SB-4.0-5.0MS	P132365R240213A	MS	3546	8082A	III
18-Jul-2013	TB-071813	7132370	TB	5030B	8015M	III
18-Jul-2013	TB-071813	7132370	TB	5030B	8260B	III
18-Jul-2013	SL-509-SA8-SB-0.0-0.5	7132366	N	3050B	6010C	III
18-Jul-2013	SL-509-SA8-SB-0.0-0.5	7132366	N	3050B	6020A	III
18-Jul-2013	SL-509-SA8-SB-0.0-0.5	7132366	N	3060A	7199	III
18-Jul-2013	SL-509-SA8-SB-0.0-0.5	7132366	N	3546	8015M	III
18-Jul-2013	SL-509-SA8-SB-0.0-0.5	7132366	N	3546	8082A	III
18-Jul-2013	SL-509-SA8-SB-0.0-0.5	7132366	N	3546	8270D	III
18-Jul-2013	SL-509-SA8-SB-0.0-0.5	7132366	N	3546	8270D SIM	III
18-Jul-2013	SL-509-SA8-SB-0.0-0.5	7132366	N	METHOD	1613B	III
18-Jul-2013	SL-509-SA8-SB-0.0-0.5	7132366	N	METHOD	6850	III
18-Jul-2013	SL-509-SA8-SB-0.0-0.5	7132366	N	METHOD	7471B	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	3050B	6010C	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	3050B	6020A	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	3060A	7199	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	3546	8015M	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	3546	8082A	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	3546	8270D	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	3546	8270D SIM	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	5035	8260B	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	5035A	8015M	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	METHOD	1613B	III

III = EPA Level 3 Data Review
IV = EPA Level 4 Data Validation

N = Normal Sample
FD = Field Duplicate

TB = Trip Blank
FB = Field Blank

MS = Matrix Spike
MSD = Matrix Spike Duplicate

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	METHOD	6850	III
18-Jul-2013	SL-509-SA8-SB-4.0-5.0	7132367	N	METHOD	7471B	III
18-Jul-2013	SL-601-SA8-SB-0.0-0.5	7132368	N	3050B	6010C	III
18-Jul-2013	SL-601-SA8-SB-0.0-0.5	7132368	N	3050B	6020A	III
18-Jul-2013	SL-601-SA8-SB-0.0-0.5	7132368	N	3546	8015M	III
18-Jul-2013	SL-601-SA8-SB-0.0-0.5	7132368	N	3546	8082A	III
18-Jul-2013	SL-601-SA8-SB-0.0-0.5	7132368	N	3546	8270D SIM	III
18-Jul-2013	SL-601-SA8-SB-0.0-0.5	7132368	N	METHOD	1613B	III
18-Jul-2013	SL-601-SA8-SB-0.0-0.5	7132368	N	METHOD	7471B	III
18-Jul-2013	SL-601-SA8-SB-4.0-5.0	7132369	N	3050B	6010C	III
18-Jul-2013	SL-601-SA8-SB-4.0-5.0	7132369	N	3050B	6020A	III
18-Jul-2013	SL-601-SA8-SB-4.0-5.0	7132369	N	3546	8015M	III
18-Jul-2013	SL-601-SA8-SB-4.0-5.0	7132369	N	3546	8082A	III
18-Jul-2013	SL-601-SA8-SB-4.0-5.0	7132369	N	3546	8270D SIM	III
18-Jul-2013	SL-601-SA8-SB-4.0-5.0	7132369	N	5035A	8015M	III
18-Jul-2013	SL-601-SA8-SB-4.0-5.0	7132369	N	METHOD	1613B	III
18-Jul-2013	SL-601-SA8-SB-4.0-5.0	7132369	N	METHOD	7471B	III
18-Jul-2013	SL-520-SA8-SB-0.0-0.5	7132373	N	3050B	6010C	III
18-Jul-2013	SL-520-SA8-SB-0.0-0.5	7132373	N	3050B	6020A	III
18-Jul-2013	SL-520-SA8-SB-0.0-0.5	7132373	N	3546	8015M	III
18-Jul-2013	SL-520-SA8-SB-0.0-0.5	7132373	N	3546	8082A	III
18-Jul-2013	SL-520-SA8-SB-0.0-0.5	7132373	N	3546	8270D SIM	III
18-Jul-2013	SL-520-SA8-SB-0.0-0.5	7132373	N	METHOD	1613B	III
18-Jul-2013	SL-520-SA8-SB-0.0-0.5	7132373	N	METHOD	7471B	III
18-Jul-2013	SL-519-SA8-SB-0.0-0.5	7132371	N	3050B	6010C	III
18-Jul-2013	SL-519-SA8-SB-0.0-0.5	7132371	N	3050B	6020A	III

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
18-Jul-2013	SL-519-SA8-SB-0.0-0.5	7132371	N	3546	8015M	III
18-Jul-2013	SL-519-SA8-SB-0.0-0.5	7132371	N	3546	8082A	III
18-Jul-2013	SL-519-SA8-SB-0.0-0.5	7132371	N	3546	8270D SIM	III
18-Jul-2013	SL-519-SA8-SB-0.0-0.5	7132371	N	METHOD	1613B	III
18-Jul-2013	SL-519-SA8-SB-0.0-0.5	7132371	N	METHOD	7471B	III
18-Jul-2013	SL-519-SA8-SB-3.5-4.5	7132372	N	3050B	6010C	III
18-Jul-2013	SL-519-SA8-SB-3.5-4.5	7132372	N	3050B	6020A	III
18-Jul-2013	SL-519-SA8-SB-3.5-4.5	7132372	N	3546	8015M	III
18-Jul-2013	SL-519-SA8-SB-3.5-4.5	7132372	N	3546	8082A	III
18-Jul-2013	SL-519-SA8-SB-3.5-4.5	7132372	N	3546	8270D SIM	III
18-Jul-2013	SL-519-SA8-SB-3.5-4.5	7132372	N	5035A	8015M	III
18-Jul-2013	SL-519-SA8-SB-3.5-4.5	7132372	N	METHOD	1613B	III
18-Jul-2013	SL-519-SA8-SB-3.5-4.5	7132372	N	METHOD	7471B	III
18-Jul-2013	SL-531-SA8-SB-0.0-0.5	7132374	N	3050B	6010C	III
18-Jul-2013	SL-531-SA8-SB-0.0-0.5	7132374	N	3050B	6020A	III
18-Jul-2013	SL-531-SA8-SB-0.0-0.5	7132374	N	3546	8015M	III
18-Jul-2013	SL-531-SA8-SB-0.0-0.5	7132374	N	3546	8082A	III
18-Jul-2013	SL-531-SA8-SB-0.0-0.5	7132374	N	3546	8270D SIM	III
18-Jul-2013	SL-531-SA8-SB-0.0-0.5	7132374	N	METHOD	1613B	III
18-Jul-2013	SL-531-SA8-SB-0.0-0.5	7132374	N	METHOD	7471B	III
18-Jul-2013	SL-531-SA8-SB-4.0-5.0	7132375	N	3050B	6010C	III
18-Jul-2013	SL-531-SA8-SB-4.0-5.0	7132375	N	3050B	6020A	III
18-Jul-2013	SL-531-SA8-SB-4.0-5.0	7132375	N	3546	8015M	III
18-Jul-2013	SL-531-SA8-SB-4.0-5.0	7132375	N	3546	8082A	III
18-Jul-2013	SL-531-SA8-SB-4.0-5.0	7132375	N	3546	8270D SIM	III
18-Jul-2013	SL-531-SA8-SB-4.0-5.0	7132375	N	5035A	8015M	III

III = EPA Level 3 Data Review
IV = EPA Level 4 Data Validation

N = Normal Sample
FD = Field Duplicate

TB = Trip Blank
FB = Field Blank

MS = Matrix Spike
MSD = Matrix Spike Duplicate

Sample Cross Reference

Date Collected	Field Sample ID	Lab Sample ID	Sample Type	Prep Method	Analytical Method	Review Level
18-Jul-2013	SL-531-SA8-SB-4.0-5.0	7132375	N	METHOD	1613B	III
18-Jul-2013	SL-531-SA8-SB-4.0-5.0	7132375	N	METHOD	7471B	III
18-Jul-2013	SL-531-SA8-SB-4.0-5.0DUP	P132375D220919	DUP	METHOD	7471B	III
18-Jul-2013	SL-531-SA8-SB-4.0-5.0MSD	P132375M220924	MSD	METHOD	7471B	III
18-Jul-2013	SL-531-SA8-SB-4.0-5.0MS	P132375R220922	MS	METHOD	7471B	III

Attachment 2
Overall Data Qualification Summary

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: METALS
Method: 6010C **Matrix:** SO

Sample ID: SL-506-SA8-SB-0.0-0.5 Collected: 7/18/2013 7:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.704	J	0.0704	MDL	1.05	PQL	mg/Kg	J	Z
BORON	4.93	J	0.883	MDL	10.5	PQL	mg/Kg	J	Z
CADMIUM	0.520	J	0.0799	MDL	1.05	PQL	mg/Kg	J	Z
MOLYBDENUM	0.287	J	0.179	MDL	2.10	PQL	mg/Kg	U	F
TIN	2.83	J	0.231	MDL	10.5	PQL	mg/Kg	U	B
Zirconium	4.67	J	0.883	MDL	5.25	PQL	mg/Kg	J	Z

Sample ID: SL-506-SA8-SB-4.0-5.0 Collected: 7/18/2013 7:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.677	J	0.0679	MDL	1.01	PQL	mg/Kg	J	Z
CADMIUM	0.289	J	0.0770	MDL	1.01	PQL	mg/Kg	J	Z
MOLYBDENUM	0.413	J	0.172	MDL	2.03	PQL	mg/Kg	U	F
TIN	2.86	J	0.223	MDL	10.1	PQL	mg/Kg	U	B
Zirconium	3.92	J	0.851	MDL	5.07	PQL	mg/Kg	J	Z

Sample ID: SL-509-SA8-SB-0.0-0.5 Collected: 7/18/2013 8:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.781	J	0.0735	MDL	1.10	PQL	mg/Kg	J	Z
BORON	9.98	J	0.921	MDL	11.0	PQL	mg/Kg	J	Z
CADMIUM	0.670	J	0.0833	MDL	1.10	PQL	mg/Kg	J	Z
TIN	2.80	J	0.241	MDL	11.0	PQL	mg/Kg	U	B

Sample ID: SL-509-SA8-SB-4.0-5.0 Collected: 7/18/2013 9:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.530	J	0.0691	MDL	1.03	PQL	mg/Kg	J	Z
BORON	1.26	J	0.867	MDL	10.3	PQL	mg/Kg	U	B
CADMIUM	0.425	J	0.0784	MDL	1.03	PQL	mg/Kg	J	Z
TIN	2.93	J	0.227	MDL	10.3	PQL	mg/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-519-SA8-SB-0.0-0.5 Collected: 7/18/2013 1:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.827	J	0.0712	MDL	1.06	PQL	mg/Kg	J	Z
CADMIUM	0.751	J	0.0807	MDL	1.06	PQL	mg/Kg	J	Z
MOLYBDENUM	0.377	J	0.181	MDL	2.12	PQL	mg/Kg	U	F
SODIUM	103	J	17.7	MDL	106	PQL	mg/Kg	J	Z
TIN	2.69	J	0.234	MDL	10.6	PQL	mg/Kg	U	B

Sample ID: SL-519-SA8-SB-3.5-4.5 Collected: 7/18/2013 1:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
ARSENIC	3.75	J	0.724	MDL	4.14	PQL	mg/Kg	J	Z
BERYLLIUM	0.458	J	0.0693	MDL	1.03	PQL	mg/Kg	J	Z
CADMIUM	0.339	J	0.0786	MDL	1.03	PQL	mg/Kg	J	Z
SODIUM	88.1	J	17.3	MDL	103	PQL	mg/Kg	J	Z
TIN	2.85	J	0.228	MDL	10.3	PQL	mg/Kg	U	B
Zirconium	4.37	J	0.869	MDL	5.17	PQL	mg/Kg	J	Z

Sample ID: SL-520-SA8-SB-0.0-0.5 Collected: 7/18/2013 1:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.785	J	0.0699	MDL	1.04	PQL	mg/Kg	J	Z
BORON	4.36	J	0.876	MDL	10.4	PQL	mg/Kg	U	B
CADMIUM	0.591	J	0.0792	MDL	1.04	PQL	mg/Kg	J	Z
MOLYBDENUM	0.315	J	0.177	MDL	2.09	PQL	mg/Kg	U	F
SODIUM	84.3	J	17.4	MDL	104	PQL	mg/Kg	J	Z
TIN	2.66	J	0.229	MDL	10.4	PQL	mg/Kg	U	B
Zirconium	4.76	J	0.876	MDL	5.21	PQL	mg/Kg	J	Z

Sample ID: SL-531-SA8-SB-0.0-0.5 Collected: 7/18/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.882	J	0.0712	MDL	1.06	PQL	mg/Kg	J	Z
BORON	8.08	J	0.893	MDL	10.6	PQL	mg/Kg	J	Z
CADMIUM	0.673	J	0.0808	MDL	1.06	PQL	mg/Kg	J	Z
MOLYBDENUM	0.238	J	0.181	MDL	2.13	PQL	mg/Kg	U	F
SODIUM	94.6	J	17.7	MDL	106	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6010C	Matrix: SO

Sample ID: SL-531-SA8-SB-0.0-0.5 Collected: 7/18/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
TIN	2.65	J	0.234	MDL	10.6	PQL	mg/Kg	U	B
Zirconium	4.37	J	0.893	MDL	5.31	PQL	mg/Kg	J	Z

Sample ID: SL-531-SA8-SB-4.0-5.0 Collected: 7/18/2013 2:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.944	J	0.0722	MDL	1.08	PQL	mg/Kg	J	Z
BORON	3.58	J	0.905	MDL	10.8	PQL	mg/Kg	U	B
CADMIUM	0.505	J	0.0819	MDL	1.08	PQL	mg/Kg	J	Z
MOLYBDENUM	0.343	J	0.183	MDL	2.15	PQL	mg/Kg	U	F
SODIUM	98.4	J	18.0	MDL	108	PQL	mg/Kg	J	Z
TIN	2.91	J	0.237	MDL	10.8	PQL	mg/Kg	U	B

Sample ID: SL-601-SA8-SB-0.0-0.5 Collected: 7/18/2013 10:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.790	J	0.0683	MDL	1.02	PQL	mg/Kg	J	Z
BORON	9.22	J	0.857	MDL	10.2	PQL	mg/Kg	J	Z
CADMIUM	0.599	J	0.0775	MDL	1.02	PQL	mg/Kg	J	Z
MOLYBDENUM	0.452	J	0.173	MDL	2.04	PQL	mg/Kg	U	F
SODIUM	93.6	J	17.0	MDL	102	PQL	mg/Kg	J	Z
TIN	2.63	J	0.224	MDL	10.2	PQL	mg/Kg	U	B
Zirconium	2.94	J	0.857	MDL	5.10	PQL	mg/Kg	J	Z

Sample ID: SL-601-SA8-SB-4.0-5.0 Collected: 7/18/2013 10:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BERYLLIUM	0.730	J	0.0682	MDL	1.02	PQL	mg/Kg	J	Z
BORON	4.04	J	0.855	MDL	10.2	PQL	mg/Kg	U	B
CADMIUM	0.412	J	0.0773	MDL	1.02	PQL	mg/Kg	J	Z
MOLYBDENUM	0.351	J	0.173	MDL	2.04	PQL	mg/Kg	U	F
SODIUM	92.7	J	17.0	MDL	102	PQL	mg/Kg	J	Z
TIN	2.88	J	0.224	MDL	10.2	PQL	mg/Kg	U	B
Zirconium	3.65	J	0.855	MDL	5.09	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	METALS	
Method:	6020A	Matrix: SO

Sample ID: SL-506-SA8-SB-0.0-0.5	Collected: 7/18/2013 7:35:00	Analysis Type: REA2	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.315	J	0.105	MDL	0.420	PQL	mg/Kg	J	Z

Sample ID: SL-506-SA8-SB-0.0-0.5	Collected: 7/18/2013 7:35:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0732	J	0.0273	MDL	0.210	PQL	mg/Kg	J	Z

Sample ID: SL-509-SA8-SB-0.0-0.5	Collected: 7/18/2013 8:50:00	Analysis Type: REA2	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.281	J	0.110	MDL	0.439	PQL	mg/Kg	J	Z

Sample ID: SL-509-SA8-SB-0.0-0.5	Collected: 7/18/2013 8:50:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0321	J	0.0285	MDL	0.219	PQL	mg/Kg	J	Z

Sample ID: SL-519-SA8-SB-0.0-0.5	Collected: 7/18/2013 1:30:00	Analysis Type: REA2	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.209	J	0.106	MDL	0.425	PQL	mg/Kg	J	Z

Sample ID: SL-519-SA8-SB-0.0-0.5	Collected: 7/18/2013 1:30:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0699	J	0.0276	MDL	0.212	PQL	mg/Kg	J	Z

Sample ID: SL-520-SA8-SB-0.0-0.5	Collected: 7/18/2013 1:00:00	Analysis Type: REA2	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.156	J	0.104	MDL	0.417	PQL	mg/Kg	J	Z

Sample ID: SL-520-SA8-SB-0.0-0.5	Collected: 7/18/2013 1:00:00	Analysis Type: RES	Dilution: 2						
Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0373	J	0.0271	MDL	0.209	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: METALS
Method: 6020A **Matrix:** SO

Sample ID: SL-531-SA8-SB-0.0-0.5 Collected: 7/18/2013 2:15:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.225	J	0.106	MDL	0.425	PQL	mg/Kg	J	Z

Sample ID: SL-531-SA8-SB-0.0-0.5 Collected: 7/18/2013 2:15:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0320	J	0.0276	MDL	0.213	PQL	mg/Kg	J	Z

Sample ID: SL-531-SA8-SB-4.0-5.0 Collected: 7/18/2013 2:30:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0474	J	0.0280	MDL	0.215	PQL	mg/Kg	J	Z

Sample ID: SL-601-SA8-SB-0.0-0.5 Collected: 7/18/2013 10:00:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.248	J	0.102	MDL	0.408	PQL	mg/Kg	J	Z

Sample ID: SL-601-SA8-SB-0.0-0.5 Collected: 7/18/2013 10:00:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0287	J	0.0265	MDL	0.204	PQL	mg/Kg	J	Z

Sample ID: SL-601-SA8-SB-4.0-5.0 Collected: 7/18/2013 10:30:00 Analysis Type: REA2 Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SELENIUM	0.123	J	0.102	MDL	0.407	PQL	mg/Kg	J	Z

Sample ID: SL-601-SA8-SB-4.0-5.0 Collected: 7/18/2013 10:30:00 Analysis Type: RES Dilution: 2

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
SILVER	0.0299	J	0.0265	MDL	0.204	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: METALS
Method: 7199 **Matrix:** SO

Sample ID: SL-506-SA8-SB-0.0-0.5 Collected: 7/18/2013 7:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.30	J	0.15	MDL	0.43	PQL	mg/Kg	J	Z

Sample ID: SL-509-SA8-SB-0.0-0.5 Collected: 7/18/2013 8:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.18	J	0.15	MDL	0.44	PQL	mg/Kg	J	Z

Sample ID: SL-509-SA8-SB-4.0-5.0 Collected: 7/18/2013 9:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
HEXAVALENT CHROMIUM	0.18	J	0.15	MDL	0.42	PQL	mg/Kg	J	Z

Method Category: METALS
Method: 7471B **Matrix:** SO

Sample ID: SL-531-SA8-SB-4.0-5.0 Collected: 7/18/2013 2:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
MERCURY	0.0139	J	0.0112	MDL	0.0186	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-506-SA8-SB-0.0-0.5 Collected: 7/18/2013 7:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	2.50	JB	0.0260	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0679	JB	0.0369	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0550	JQ	0.0428	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDF	0.110	JB	0.0277	MDL	5.29	PQL	ng/Kg	U	B
1,2,3,6,7,8-HxCDD	0.345	JB	0.0450	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HxCDF	0.111	JBQ	0.0275	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDD	0.394	JBQ	0.0441	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HxCDF	0.229	JB	0.0292	MDL	5.29	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.678	JB	0.0319	MDL	5.29	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-506-SA8-SB-0.0-0.5 Collected: 7/18/2013 7:35:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,3,4,6,7,8-HXCDF	0.131	JB	0.0256	MDL	5.29	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.128	JBQ	0.0298	MDL	5.29	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.102	JQ	0.0484	MDL	1.06	PQL	ng/Kg	J	Z
OCDF	4.13	J	0.0389	MDL	10.6	PQL	ng/Kg	J	Z

Sample ID: SL-506-SA8-SB-4.0-5.0 Collected: 7/18/2013 7:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.101	JBQ	0.0289	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0448	JBQ	0.00959	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0313	JB	0.0102	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0325	JBQ	0.0229	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0111	JBQ	0.0104	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0464	JB	0.0216	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0257	JBQ	0.0103	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0519	JBQ	0.0208	MDL	5.26	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0157	JBQ	0.0101	MDL	5.26	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0273	JBQ	0.0193	MDL	5.26	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.0324	JQ	0.0309	MDL	1.05	PQL	ng/Kg	J	Z
OCDD	1.07	JB	0.0302	MDL	10.5	PQL	ng/Kg	U	B
OCDF	0.150	JQ	0.0339	MDL	10.5	PQL	ng/Kg	J	Z

Sample ID: SL-509-SA8-SB-0.0-0.5 Collected: 7/18/2013 8:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.98	JB	0.0690	MDL	5.28	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.154	JBQ	0.0189	MDL	5.28	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0472	JBQ	0.0265	MDL	5.28	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0416	JQ	0.0386	MDL	5.28	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0620	JB	0.0196	MDL	5.28	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.120	JBQ	0.0422	MDL	5.28	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.145	JBQ	0.0429	MDL	5.28	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0842	JBQ	0.0214	MDL	5.28	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0696	JQ	0.0532	MDL	5.28	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.656	JBQ	0.0316	MDL	5.28	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category:	SVOA	
Method:	1613B	Matrix: SO

Sample ID: SL-509-SA8-SB-0.0-0.5 Collected: 7/18/2013 8:50:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
2,3,4,6,7,8-HXCDF	0.0198	JB	0.0192	MDL	5.28	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0517	JBQ	0.0304	MDL	5.28	PQL	ng/Kg	U	B
2,3,7,8-TCDF	0.0695	JQ	0.0518	MDL	1.06	PQL	ng/Kg	J	Z
OCDF	0.597	JQ	0.0470	MDL	10.6	PQL	ng/Kg	J	Z

Sample ID: SL-509-SA8-SB-4.0-5.0 Collected: 7/18/2013 9:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.133	JB	0.0373	MDL	5.14	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.131	JB	0.0122	MDL	5.14	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0404	JB	0.0333	MDL	5.14	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0652	JBQ	0.0320	MDL	5.14	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0682	JBQ	0.0157	MDL	5.14	PQL	ng/Kg	U	B
OCDD	1.83	JB	0.0449	MDL	10.3	PQL	ng/Kg	J	Z
OCDF	0.750	JQ	0.0431	MDL	10.3	PQL	ng/Kg	J	Z

Sample ID: SL-519-SA8-SB-0.0-0.5 Collected: 7/18/2013 1:30:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	1.92	JB	0.0231	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.161	JBQ	0.0275	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HxCDD	0.115	JQ	0.0501	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.397	JB	0.0364	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.481	JB	0.0536	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDF	0.426	JB	0.0373	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.394	JBQ	0.0514	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.122	JB	0.0324	MDL	5.35	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.134	JQ	0.0568	MDL	5.35	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	2.90	JB	0.0638	MDL	5.35	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.526	JB	0.0378	MDL	5.35	PQL	ng/Kg	J	Z
2,3,4,7,8-PECDF	1.17	JBQ	0.0642	MDL	5.35	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.519	J	0.124	MDL	1.07	PQL	ng/Kg	J	Z
OCDF	3.25	J	0.0330	MDL	10.7	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-519-SA8-SB-3.5-4.5 Collected: 7/18/2013 1:40:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0612	JBQ	0.0289	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0393	JBQ	0.00867	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0331	JBQ	0.0131	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0264	JBQ	0.0249	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0467	JBQ	0.0121	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0278	JB	0.0269	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0424	JB	0.0134	MDL	4.98	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0944	JQ	0.0406	MDL	4.98	PQL	ng/Kg	J	Z
1,2,3,7,8-PECDF	0.0864	JBQ	0.0201	MDL	4.98	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0226	JBQ	0.0126	MDL	4.98	PQL	ng/Kg	U	B
OCDD	0.151	JBQ	0.0446	MDL	9.96	PQL	ng/Kg	U	B
OCDF	0.120	J	0.0424	MDL	9.96	PQL	ng/Kg	J	Z

Sample ID: SL-520-SA8-SB-0.0-0.5 Collected: 7/18/2013 1:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.553	JB	0.0343	MDL	5.01	PQL	ng/Kg	J	Z
1,2,3,4,6,7,8-HPCDF	0.672	JB	0.0129	MDL	5.01	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0642	JBQ	0.0281	MDL	5.01	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0392	JQ	0.0246	MDL	5.01	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.213	JBQ	0.0231	MDL	5.01	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.140	JBQ	0.0272	MDL	5.01	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.480	JBQ	0.0217	MDL	5.01	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDD	0.412	JB	0.0255	MDL	5.01	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.223	JBQ	0.0250	MDL	5.01	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0872	JB	0.0224	MDL	5.01	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0360	JQ	0.0281	MDL	1.00	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.274	J	0.0702	MDL	1.00	PQL	ng/Kg	J	Z
OCDD	5.20	JB	0.0771	MDL	10.0	PQL	ng/Kg	J	Z
OCDF	0.920	J	0.0562	MDL	10.0	PQL	ng/Kg	J	Z

Sample ID: SL-531-SA8-SB-0.0-0.5 Collected: 7/18/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	1.15	JB	0.0473	MDL	5.20	PQL	ng/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-531-SA8-SB-0.0-0.5 **Collected:** 7/18/2013 2:15:00 **Analysis Type:** RES **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDF	0.486	JB	0.0129	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,4,7,8,9-HPCDF	0.0349	JBQ	0.0305	MDL	5.20	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0742	JBQ	0.0201	MDL	5.20	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.105	JBQ	0.0289	MDL	5.20	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0725	JBQ	0.0190	MDL	5.20	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.200	JB	0.0273	MDL	5.20	PQL	ng/Kg	J	Z
1,2,3,7,8,9-HXCDF	0.178	JB	0.0193	MDL	5.20	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.483	JBQ	0.0254	MDL	5.20	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0588	JBQ	0.0196	MDL	5.20	PQL	ng/Kg	U	B
2,3,4,7,8-PECDF	0.0804	JBQ	0.0286	MDL	5.20	PQL	ng/Kg	U	B
2,3,7,8-TCDD	0.0392	J	0.0384	MDL	1.04	PQL	ng/Kg	J	Z
2,3,7,8-TCDF	0.0875	JQ	0.0453	MDL	1.04	PQL	ng/Kg	J	Z
OCDF	1.44	J	0.0664	MDL	10.4	PQL	ng/Kg	J	Z

Sample ID: SL-531-SA8-SB-4.0-5.0 **Collected:** 7/18/2013 2:30:00 **Analysis Type:** RES **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0908	JB	0.0296	MDL	5.49	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0321	JBQ	0.00916	MDL	5.49	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0315	JBQ	0.0178	MDL	5.49	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0210	JQ	0.0206	MDL	5.49	PQL	ng/Kg	J	Z
1,2,3,6,7,8-HXCDD	0.0348	JBQ	0.0234	MDL	5.49	PQL	ng/Kg	U	B
1,2,3,7,8-PECDD	0.0482	JQ	0.0360	MDL	5.49	PQL	ng/Kg	J	Z
2,3,4,6,7,8-HXCDF	0.0291	JBQ	0.0106	MDL	5.49	PQL	ng/Kg	U	B
OCDD	0.484	JBQ	0.0458	MDL	11.0	PQL	ng/Kg	U	B
OCDF	0.0502	JQ	0.0465	MDL	11.0	PQL	ng/Kg	J	Z

Sample ID: SL-601-SA8-SB-0.0-0.5 **Collected:** 7/18/2013 10:00:00 **Analysis Type:** RES **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.153	JBQ	0.0401	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0548	JBQ	0.0142	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,4,7,8-HxCDD	0.0367	JQ	0.0330	MDL	5.30	PQL	ng/Kg	J	Z
1,2,3,4,7,8-HXCDF	0.0247	JBQ	0.0162	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDD	0.0569	JBQ	0.0373	MDL	5.30	PQL	ng/Kg	U	B

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 1613B **Matrix:** SO

Sample ID: SL-601-SA8-SB-0.0-0.5 **Collected:** 7/18/2013 10:00:00 **Analysis Type:** RES **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,6,7,8-HXCDF	0.0352	JB	0.0160	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.0378	JB	0.0348	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0244	JBQ	0.0173	MDL	5.30	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0513	JBQ	0.0309	MDL	5.30	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0369	JBQ	0.0156	MDL	5.30	PQL	ng/Kg	U	B
OCDD	1.01	JBQ	0.0400	MDL	10.6	PQL	ng/Kg	U	B
OCDF	0.178	JQ	0.0516	MDL	10.6	PQL	ng/Kg	J	Z

Sample ID: SL-601-SA8-SB-4.0-5.0 **Collected:** 7/18/2013 10:30:00 **Analysis Type:** RES **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1,2,3,4,6,7,8-HPCDD	0.0961	JBQ	0.0292	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,4,6,7,8-HPCDF	0.0294	JBQ	0.00794	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,4,7,8,9-HPCDF	0.0204	JBQ	0.0148	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,4,7,8-HXCDF	0.0281	JBQ	0.0132	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,6,7,8-HXCDF	0.0226	JB	0.0127	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDD	0.135	JBQ	0.0275	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,7,8,9-HXCDF	0.0890	JB	0.0143	MDL	5.26	PQL	ng/Kg	U	B
1,2,3,7,8-PECDF	0.0707	JB	0.0204	MDL	5.26	PQL	ng/Kg	U	B
2,3,4,6,7,8-HXCDF	0.0192	JBQ	0.0127	MDL	5.26	PQL	ng/Kg	U	B
OCDD	0.337	JBQ	0.0333	MDL	10.5	PQL	ng/Kg	U	B
OCDF	0.0885	J	0.0388	MDL	10.5	PQL	ng/Kg	J	Z

Method Category: SVOA
Method: 8015M **Matrix:** SO

Sample ID: SL-506-SA8-SB-0.0-0.5 **Collected:** 7/18/2013 7:35:00 **Analysis Type:** RES **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	8.5		2.1	MDL	5.4	PQL	mg/Kg	J	Q, Q

Sample ID: SL-519-SA8-SB-0.0-0.5 **Collected:** 7/18/2013 1:30:00 **Analysis Type:** RES **Dilution:** 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	2.8	J	2.2	MDL	5.4	PQL	mg/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 8015M Matrix: SO

Sample ID: SL-520-SA8-SB-0.0-0.5 Collected: 7/18/2013 1:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	3.9	J	2.1	MDL	5.2	PQL	mg/Kg	J	Z
EFH (C30-C40)	7.9	J	4.2	MDL	10	PQL	mg/Kg	J	Z

Sample ID: SL-531-SA8-SB-0.0-0.5 Collected: 7/18/2013 2:15:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C15-C20)	2.5	J	2.1	MDL	5.3	PQL	mg/Kg	J	Z

Sample ID: SL-601-SA8-SB-4.0-5.0 Collected: 7/18/2013 10:30:00 Analysis Type: REA Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
EFH (C21-C30)	4.4	J	2.1	MDL	5.3	PQL	mg/Kg	J	Z
EFH (C30-C40)	7.3	J	4.2	MDL	11	PQL	mg/Kg	J	Z

Method Category: SVOA
Method: 8082A Matrix: SO

Sample ID: SL-506-SA8-SB-0.0-0.5 Collected: 7/18/2013 7:35:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	5.0	J	4.7	MDL	18	PQL	ug/Kg	J	Z

Sample ID: SL-531-SA8-SB-0.0-0.5 Collected: 7/18/2013 2:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
AROCLOR 1254	5.5	J	4.6	MDL	18	PQL	ug/Kg	J	Z

Method Category: SVOA
Method: 8270D Matrix: SO

Sample ID: SL-506-SA8-SB-0.0-0.5 Collected: 7/18/2013 7:35:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZIDINE	1800	U	750	MDL	1800	PQL	ug/Kg	R	Q

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-506-SA8-SB-0.0-0.5 Collected: 7/18/2013 7:35:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(K)FLUORANTHENE	1.7	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	6.6	J	6.4	MDL	19	PQL	ug/Kg	J	Z
CHRYSENE	1.1	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	0.74	J	0.71	MDL	1.8	PQL	ug/Kg	U	F

Sample ID: SL-509-SA8-SB-0.0-0.5 Collected: 7/18/2013 8:50:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(B)FLUORANTHENE	0.78	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(K)FLUORANTHENE	1.1	J	0.72	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	1.1	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	0.85	J	0.72	MDL	1.8	PQL	ug/Kg	U	F

Sample ID: SL-519-SA8-SB-0.0-0.5 Collected: 7/18/2013 1:30:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
1-METHYLNAPHTHALENE	1.1	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
ANTHRACENE	0.38	J	0.36	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(A)ANTHRACENE	0.96	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(A)PYRENE	1.4	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
BENZO(E)PYRENE	4.0	J	3.5	MDL	18	PQL	ug/Kg	J	Z
BIS(2-ETHYLHEXYL)PHTHALATE	8.9	J	6.4	MDL	19	PQL	ug/Kg	J	Z
INDENO(1,2,3-CD)PYRENE	1.7	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z

Sample ID: SL-520-SA8-SB-0.0-0.5 Collected: 7/18/2013 1:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(K)FLUORANTHENE	0.92	J	0.69	MDL	1.7	PQL	ug/Kg	J	Z
CHRYSENE	0.43	J	0.35	MDL	1.7	PQL	ug/Kg	J	Z

Sample ID: SL-531-SA8-SB-0.0-0.5 Collected: 7/18/2013 2:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
BENZO(K)FLUORANTHENE	1.5	J	0.71	MDL	1.8	PQL	ug/Kg	J	Z
CHRYSENE	1.0	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method Category: SVOA
Method: 8270D SIM **Matrix:** SO

Sample ID: SL-531-SA8-SB-0.0-0.5 Collected: 7/18/2013 2:15:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
NAPHTHALENE	0.82	J	0.71	MDL	1.8	PQL	ug/Kg	U	F

Sample ID: SL-601-SA8-SB-0.0-0.5 Collected: 7/18/2013 10:00:00 Analysis Type: RES-BASE/NEUTRAL Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
CHRYSENE	0.73	J	0.35	MDL	1.8	PQL	ug/Kg	J	Z
NAPHTHALENE	0.93	J	0.70	MDL	1.8	PQL	ug/Kg	J	Z

Method Category: VOA
Method: 8260B **Matrix:** AQ

Sample ID: TB-071813 Collected: 7/18/2013 8:00:00 Analysis Type: RES Dilution: 1

Analyte	Lab Result	Lab Qual	DL	DL Type	RL	RL Type	Units	Data Review Qual	Reason Code
Chlorotrifluoroethylene	5	U	2	MDL	5	PQL	ug/L	UJ	L

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

Data Qualifier Summary

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Reason Code Legend

<i>Reason Code</i>	<i>Description</i>
B	Method Blank Contamination
E	Laboratory Duplicate Precision
E	Matrix Spike Precision
F	Field Blank Contamination
L	Laboratory Control Spike Lower Estimation
L	Laboratory Control Spike Upper Estimation
Q	Laboratory Duplicate Precision
Q	Matrix Spike Lower Rejection
Q	Matrix Spike Precision
Q	Matrix Spike Upper Estimation
Z	Reporting Limit Trace Value

* denotes a non-reportable result

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Enclosure I
EPA Level III ADR Outliers
(Including Manual Review Outliers)

Quality Control Outlier Reports

PH070

Method Blank Outlier Report

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
BLK2070B371809	7/29/2013 6:09:00 PM	1,2,3,4,6,7,8-HPCDD 1,2,3,4,6,7,8-HPCDF 1,2,3,4,7,8,9-HPCDF 1,2,3,4,7,8-HXCDF 1,2,3,6,7,8-HXCDD 1,2,3,6,7,8-HXCDF 1,2,3,7,8,9-HXCDD 1,2,3,7,8,9-HXCDF 1,2,3,7,8-PECDF 2,3,4,6,7,8-HXCDF 2,3,4,7,8-PECDF OCDD	0.0394 ng/Kg 0.0286 ng/Kg 0.0272 ng/Kg 0.0219 ng/Kg 0.0352 ng/Kg 0.0208 ng/Kg 0.0341 ng/Kg 0.0358 ng/Kg 0.0485 ng/Kg 0.0291 ng/Kg 0.0474 ng/Kg 0.233 ng/Kg	SL-506-SA8-SB-0.0-0.5 SL-506-SA8-SB-4.0-5.0 SL-509-SA8-SB-0.0-0.5 SL-509-SA8-SB-4.0-5.0 SL-519-SA8-SB-0.0-0.5 SL-519-SA8-SB-3.5-4.5 SL-520-SA8-SB-0.0-0.5 SL-531-SA8-SB-0.0-0.5 SL-531-SA8-SB-4.0-5.0 SL-601-SA8-SB-0.0-0.5 SL-601-SA8-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-506-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0679 ng/Kg	0.0679U ng/Kg
SL-506-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.110 ng/Kg	0.110U ng/Kg
SL-506-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.131 ng/Kg	0.131U ng/Kg
SL-506-SA8-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.128 ng/Kg	0.128U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDD	0.101 ng/Kg	0.101U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0448 ng/Kg	0.0448U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HXCDF	0.0313 ng/Kg	0.0313U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDD	0.0325 ng/Kg	0.0325U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDF	0.0111 ng/Kg	0.0111U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDD	0.0464 ng/Kg	0.0464U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDF	0.0257 ng/Kg	0.0257U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.0519 ng/Kg	0.0519U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0157 ng/Kg	0.0157U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	2,3,4,7,8-PECDF	0.0273 ng/Kg	0.0273U ng/Kg
SL-506-SA8-SB-4.0-5.0(RES)	OCDD	1.07 ng/Kg	1.07U ng/Kg
SL-509-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0472 ng/Kg	0.0472U ng/Kg
SL-509-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0620 ng/Kg	0.0620U ng/Kg
SL-509-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.120 ng/Kg	0.120U ng/Kg
SL-509-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.145 ng/Kg	0.145U ng/Kg
SL-509-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.0842 ng/Kg	0.0842U ng/Kg
SL-509-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0198 ng/Kg	0.0198U ng/Kg
SL-509-SA8-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.0517 ng/Kg	0.0517U ng/Kg
SL-509-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDD	0.133 ng/Kg	0.133U ng/Kg
SL-509-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.131 ng/Kg	0.131U ng/Kg
SL-509-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDD	0.0404 ng/Kg	0.0404U ng/Kg
SL-509-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDD	0.0652 ng/Kg	0.0652U ng/Kg
SL-509-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDF	0.0682 ng/Kg	0.0682U ng/Kg
SL-519-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.122 ng/Kg	0.122U ng/Kg
SL-519-SA8-SB-3.5-4.5(RES)	1,2,3,4,6,7,8-HPCDD	0.0612 ng/Kg	0.0612U ng/Kg
SL-519-SA8-SB-3.5-4.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0393 ng/Kg	0.0393U ng/Kg
SL-519-SA8-SB-3.5-4.5(RES)	1,2,3,4,7,8-HXCDF	0.0331 ng/Kg	0.0331U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-519-SA8-SB-3.5-4.5(RES)	1,2,3,6,7,8-HXCDD	0.0264 ng/Kg	0.0264U ng/Kg
SL-519-SA8-SB-3.5-4.5(RES)	1,2,3,6,7,8-HXCDF	0.0467 ng/Kg	0.0467U ng/Kg
SL-519-SA8-SB-3.5-4.5(RES)	1,2,3,7,8,9-HXCDD	0.0278 ng/Kg	0.0278U ng/Kg
SL-519-SA8-SB-3.5-4.5(RES)	1,2,3,7,8,9-HXCDF	0.0424 ng/Kg	0.0424U ng/Kg
SL-519-SA8-SB-3.5-4.5(RES)	1,2,3,7,8-PECDF	0.0864 ng/Kg	0.0864U ng/Kg
SL-519-SA8-SB-3.5-4.5(RES)	2,3,4,6,7,8-HXCDF	0.0226 ng/Kg	0.0226U ng/Kg
SL-519-SA8-SB-3.5-4.5(RES)	OCDD	0.151 ng/Kg	0.151U ng/Kg
SL-520-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0642 ng/Kg	0.0642U ng/Kg
SL-520-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.140 ng/Kg	0.140U ng/Kg
SL-520-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0872 ng/Kg	0.0872U ng/Kg
SL-531-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8,9-HPCDF	0.0349 ng/Kg	0.0349U ng/Kg
SL-531-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0742 ng/Kg	0.0742U ng/Kg
SL-531-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.105 ng/Kg	0.105U ng/Kg
SL-531-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0725 ng/Kg	0.0725U ng/Kg
SL-531-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.178 ng/Kg	0.178U ng/Kg
SL-531-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0588 ng/Kg	0.0588U ng/Kg
SL-531-SA8-SB-0.0-0.5(RES)	2,3,4,7,8-PECDF	0.0804 ng/Kg	0.0804U ng/Kg
SL-531-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDD	0.0908 ng/Kg	0.0908U ng/Kg
SL-531-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0321 ng/Kg	0.0321U ng/Kg
SL-531-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0315 ng/Kg	0.0315U ng/Kg
SL-531-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDD	0.0348 ng/Kg	0.0348U ng/Kg
SL-531-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0291 ng/Kg	0.0291U ng/Kg
SL-531-SA8-SB-4.0-5.0(RES)	OCDD	0.484 ng/Kg	0.484U ng/Kg
SL-601-SA8-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDD	0.153 ng/Kg	0.153U ng/Kg
SL-601-SA8-SB-0.0-0.5(RES)	1,2,3,4,6,7,8-HPCDF	0.0548 ng/Kg	0.0548U ng/Kg
SL-601-SA8-SB-0.0-0.5(RES)	1,2,3,4,7,8-HXCDF	0.0247 ng/Kg	0.0247U ng/Kg
SL-601-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDD	0.0569 ng/Kg	0.0569U ng/Kg
SL-601-SA8-SB-0.0-0.5(RES)	1,2,3,6,7,8-HXCDF	0.0352 ng/Kg	0.0352U ng/Kg
SL-601-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDD	0.0378 ng/Kg	0.0378U ng/Kg
SL-601-SA8-SB-0.0-0.5(RES)	1,2,3,7,8,9-HXCDF	0.0244 ng/Kg	0.0244U ng/Kg
SL-601-SA8-SB-0.0-0.5(RES)	1,2,3,7,8-PECDF	0.0513 ng/Kg	0.0513U ng/Kg
SL-601-SA8-SB-0.0-0.5(RES)	2,3,4,6,7,8-HXCDF	0.0369 ng/Kg	0.0369U ng/Kg
SL-601-SA8-SB-0.0-0.5(RES)	OCDD	1.01 ng/Kg	1.01U ng/Kg
SL-601-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDD	0.0961 ng/Kg	0.0961U ng/Kg
SL-601-SA8-SB-4.0-5.0(RES)	1,2,3,4,6,7,8-HPCDF	0.0294 ng/Kg	0.0294U ng/Kg
SL-601-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8,9-HPCDF	0.0204 ng/Kg	0.0204U ng/Kg
SL-601-SA8-SB-4.0-5.0(RES)	1,2,3,4,7,8-HXCDF	0.0281 ng/Kg	0.0281U ng/Kg
SL-601-SA8-SB-4.0-5.0(RES)	1,2,3,6,7,8-HXCDF	0.0226 ng/Kg	0.0226U ng/Kg
SL-601-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDD	0.135 ng/Kg	0.135U ng/Kg
SL-601-SA8-SB-4.0-5.0(RES)	1,2,3,7,8,9-HXCDF	0.0890 ng/Kg	0.0890U ng/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Method Blank Outlier Report

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-601-SA8-SB-4.0-5.0(RES)	1,2,3,7,8-PECDF	0.0707 ng/Kg	0.0707U ng/Kg
SL-601-SA8-SB-4.0-5.0(RES)	2,3,4,6,7,8-HXCDF	0.0192 ng/Kg	0.0192U ng/Kg
SL-601-SA8-SB-4.0-5.0(RES)	OCDD	0.337 ng/Kg	0.337U ng/Kg

Method: 6010C
Matrix: SO

Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
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P20337CB222014	7/23/2013 8:14:00 PM	BORON CALCIUM TIN ZINC	0.920 mg/Kg 12.9 mg/Kg 1.46 mg/Kg 0.353 mg/Kg	SL-506-SA8-SB-0.0-0.5 SL-506-SA8-SB-4.0-5.0 SL-509-SA8-SB-0.0-0.5 SL-509-SA8-SB-4.0-5.0 SL-519-SA8-SB-0.0-0.5 SL-519-SA8-SB-3.5-4.5 SL-520-SA8-SB-0.0-0.5 SL-531-SA8-SB-0.0-0.5 SL-531-SA8-SB-4.0-5.0 SL-601-SA8-SB-0.0-0.5 SL-601-SA8-SB-4.0-5.0
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The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-506-SA8-SB-0.0-0.5(RES)	TIN	2.83 mg/Kg	2.83U mg/Kg
SL-506-SA8-SB-4.0-5.0(RES)	TIN	2.86 mg/Kg	2.86U mg/Kg
SL-509-SA8-SB-0.0-0.5(RES)	TIN	2.80 mg/Kg	2.80U mg/Kg
SL-509-SA8-SB-4.0-5.0(RES)	BORON	1.26 mg/Kg	1.26U mg/Kg
SL-509-SA8-SB-4.0-5.0(RES)	TIN	2.93 mg/Kg	2.93U mg/Kg
SL-519-SA8-SB-0.0-0.5(RES)	TIN	2.69 mg/Kg	2.69U mg/Kg
SL-519-SA8-SB-3.5-4.5(RES)	TIN	2.85 mg/Kg	2.85U mg/Kg
SL-520-SA8-SB-0.0-0.5(RES)	BORON	4.36 mg/Kg	4.36U mg/Kg
SL-520-SA8-SB-0.0-0.5(RES)	TIN	2.66 mg/Kg	2.66U mg/Kg
SL-531-SA8-SB-0.0-0.5(RES)	TIN	2.65 mg/Kg	2.65U mg/Kg
SL-531-SA8-SB-4.0-5.0(RES)	BORON	3.58 mg/Kg	3.58U mg/Kg
SL-531-SA8-SB-4.0-5.0(RES)	TIN	2.91 mg/Kg	2.91U mg/Kg
SL-601-SA8-SB-0.0-0.5(RES)	TIN	2.63 mg/Kg	2.63U mg/Kg
SL-601-SA8-SB-4.0-5.0(RES)	BORON	4.04 mg/Kg	4.04U mg/Kg
SL-601-SA8-SB-4.0-5.0(RES)	TIN	2.88 mg/Kg	2.88U mg/Kg

Method Blank Outlier Report

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6020A				
Matrix: SO				
Method Blank Sample ID	Analysis Date	Analyte	Result	Associated Samples
P20337CB220738A	7/24/2013 7:38:00 AM	STRONTIUM	0.0744 mg/Kg	SL-506-SA8-SB-0.0-0.5 SL-506-SA8-SB-4.0-5.0 SL-509-SA8-SB-0.0-0.5 SL-509-SA8-SB-4.0-5.0 SL-519-SA8-SB-0.0-0.5 SL-519-SA8-SB-3.5-4.5 SL-520-SA8-SB-0.0-0.5 SL-531-SA8-SB-0.0-0.5 SL-531-SA8-SB-4.0-5.0 SL-601-SA8-SB-0.0-0.5 SL-601-SA8-SB-4.0-5.0

Field Blank Outlier Report

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PH070_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(REA2)	4/11/2013 3:00:00 PM	COPPER MOLYBDENUM	0.0036 mg/L 0.0036 mg/L	SL-506-SA8-SB-0.0-0.5 SL-506-SA8-SB-4.0-5.0 SL-509-SA8-SB-0.0-0.5 SL-509-SA8-SB-4.0-5.0 SL-519-SA8-SB-0.0-0.5 SL-519-SA8-SB-3.5-4.5 SL-520-SA8-SB-0.0-0.5 SL-531-SA8-SB-0.0-0.5 SL-531-SA8-SB-4.0-5.0 SL-601-SA8-SB-0.0-0.5 SL-601-SA8-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-506-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.287 mg/Kg	0.287U mg/Kg
SL-506-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.413 mg/Kg	0.413U mg/Kg
SL-519-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.377 mg/Kg	0.377U mg/Kg
SL-520-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.315 mg/Kg	0.315U mg/Kg
SL-531-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.238 mg/Kg	0.238U mg/Kg
SL-531-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.343 mg/Kg	0.343U mg/Kg
SL-601-SA8-SB-0.0-0.5(RES)	MOLYBDENUM	0.452 mg/Kg	0.452U mg/Kg
SL-601-SA8-SB-4.0-5.0(RES)	MOLYBDENUM	0.351 mg/Kg	0.351U mg/Kg

Method: 8270D SIM
Matrix: SO

Field Blank Sample ID	Collected Date	Analyte	Result	Associated Samples
FB-041113(RES)	4/11/2013 3:00:00 PM	1-METHYLNAPHTHALENE 2-METHYLNAPHTHALENE BIS(2-ETHYLHEXYL)PHTHALATE Diethylphthalate Di-n-butylphthalate NAPHTHALENE	0.019 ug/L 0.024 ug/L 0.082 ug/L 0.18 ug/L 0.17 ug/L 0.17 ug/L	SL-506-SA8-SB-0.0-0.5 SL-506-SA8-SB-4.0-5.0 SL-509-SA8-SB-0.0-0.5 SL-509-SA8-SB-4.0-5.0 SL-519-SA8-SB-0.0-0.5 SL-519-SA8-SB-3.5-4.5 SL-520-SA8-SB-0.0-0.5 SL-531-SA8-SB-0.0-0.5 SL-531-SA8-SB-4.0-5.0 SL-601-SA8-SB-0.0-0.5 SL-601-SA8-SB-4.0-5.0

The following samples and their listed target analytes were qualified due to contamination reported in this blank

Sample ID	Analyte	Reported Result	Modified Final Result
SL-506-SA8-SB-0.0-0.5(RES)	NAPHTHALENE	0.74 ug/Kg	1.8U ug/Kg
SL-509-SA8-SB-0.0-0.5(RES)	NAPHTHALENE	0.85 ug/Kg	1.8U ug/Kg
SL-531-SA8-SB-0.0-0.5(RES)	NAPHTHALENE	0.82 ug/Kg	1.8U ug/Kg

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Matrix Spike/Matrix Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PH070_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8015M

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-506-SA8-SB-0.0-0.5MS	EFH (C15-C20)	154	203	49.00-123.00	28 (20.00)	EFH (C15-C20)	J (all detects)
SL-506-SA8-SB-0.0-0.5MSD	EFH (C21-C30)	135	258	49.00-123.00	32 (20.00)	EFH (C21-C30)	EFH (C30-C40)
(SL-506-SA8-SB-0.0-0.5)	EFH (C30-C40)	182	449	49.00-123.00	36 (20.00)	EFH (C30-C40)	No Qual, >4X

Method: 8270D SIM

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-506-SA8-SB-0.0-0.5MSD	Diethylphthalate	-	128	76.00-127.00	-	Diethylphthalate	J(all detects)
(SL-506-SA8-SB-0.0-0.5)							

Method: 8270D

Matrix: SO

QC Sample ID (Associated Samples)	Compound	MS %R	MSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
SL-506-SA8-SB-0.0-0.5MS	BENZIDINE	0	0	35.00-141.00	-	BENZIDINE	J(all detects)
SL-506-SA8-SB-0.0-0.5MSD							R(all non-detects)
(SL-506-SA8-SB-0.0-0.5)							

Lab Duplicate Outlier Report

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 7471B

Matrix: SO

QC Sample ID (Associated Sample ID)	Analyte	Sample RPD	eQAPP RPD	Flag
SL-531-SA8-SB-4.0-5.0DUP (TOT) (SL-506-SA8-SB-0.0-0.5 SL-506-SA8-SB-4.0-5.0 SL-509-SA8-SB-0.0-0.5 SL-509-SA8-SB-4.0-5.0 SL-519-SA8-SB-0.0-0.5 SL-519-SA8-SB-3.5-4.5 SL-520-SA8-SB-0.0-0.5 SL-531-SA8-SB-0.0-0.5 SL-531-SA8-SB-4.0-5.0 SL-601-SA8-SB-0.0-0.5 SL-601-SA8-SB-4.0-5.0)	MERCURY	200	20.00	No Qual, OK by Difference

Lab Control Spike/Lab Control Spike Duplicate Outlier Report

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PrepPH070_v1

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8260B
Matrix: AQ

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
LCSY77Q211110A LCSY77Y211131A (TB-071813)	Chlorotrifluoroethylene	44	43	47.00-120.00	-	Chlorotrifluoroethylene	J(all detects) UJ(all non-detects)

Method: 8270D SIM
Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P3LDLCSQ260423 (SL-506-SA8-SB-0.0-0.5 SL-506-SA8-SB-4.0-5.0 SL-509-SA8-SB-0.0-0.5 SL-509-SA8-SB-4.0-5.0 SL-519-SA8-SB-0.0-0.5 SL-519-SA8-SB-3.5-4.5 SL-520-SA8-SB-0.0-0.5 SL-531-SA8-SB-0.0-0.5 SL-531-SA8-SB-4.0-5.0 SL-601-SA8-SB-0.0-0.5 SL-601-SA8-SB-4.0-5.0)	Diethylphthalate	128	-	68.00-125.00	-	Diethylphthalate	J (all detects)

Method: 8270D
Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
P3LELCSQ262136 (SL-506-SA8-SB-0.0-0.5 SL-506-SA8-SB-4.0-5.0 SL-509-SA8-SB-0.0-0.5 SL-509-SA8-SB-4.0-5.0)	1,2-Diphenylhydrazine/Azobenz	115	-	77.00-111.00	-	1,2-Diphenylhydrazine/Azobenz	J(all detects)

Method: 8260B
Matrix: SO

QC Sample ID (Associated Samples)	Compound	LCS %R	LCSD %R	%R Limits	RPD (Limits)	Affected Compounds	Flag
LCSB68Q212141A LCSB68Y212203A (SL-506-SA8-SB-4.0-5.0 SL-509-SA8-SB-4.0-5.0)	VINYL ACETATE	-	-	10.00-160.00	-	VINYL ACETATE	J(all detects)

Reporting Limit Outliers

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PH070_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-506-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	2.50	5.29	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JB	0.0679	5.29	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0550	5.29	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.110	5.29	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.345	5.29	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.111	5.29	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.394	5.29	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.229	5.29	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	0.678	5.29	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.131	5.29	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.128	5.29	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.102	1.06	PQL	ng/Kg	
	OCDF	J	4.13	10.6	PQL	ng/Kg	
SL-506-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.101	5.26	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0448	5.26	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.0313	5.26	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.0325	5.26	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0111	5.26	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.0464	5.26	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.0257	5.26	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0519	5.26	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0157	5.26	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0273	5.26	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0324	1.05	PQL	ng/Kg	
	OCDD	JB	1.07	10.5	PQL	ng/Kg	
	OCDF	JQ	0.150	10.5	PQL	ng/Kg	
SL-509-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	1.98	5.28	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.154	5.28	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0472	5.28	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0416	5.28	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.0620	5.28	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.120	5.28	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.145	5.28	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.0842	5.28	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0696	5.28	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.656	5.28	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.0198	5.28	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0517	5.28	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0695	1.06	PQL	ng/Kg	
OCDF	JQ	0.597	10.6	PQL	ng/Kg		
SL-509-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	0.133	5.14	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.131	5.14	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.0404	5.14	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.0652	5.14	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.0682	5.14	PQL	ng/Kg	
	OCDD	JB	1.83	10.3	PQL	ng/Kg	
	OCDF	JQ	0.750	10.3	PQL	ng/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PH070_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-519-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDF	JB	1.92	5.35	PQL	ng/Kg	J (all detects)
	1,2,3,4,7,8,9-HPCDF	JBQ	0.161	5.35	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.115	5.35	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JB	0.397	5.35	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JB	0.481	5.35	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.426	5.35	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JBQ	0.394	5.35	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.122	5.35	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.134	5.35	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JB	2.90	5.35	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.526	5.35	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	1.17	5.35	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.519	1.07	PQL	ng/Kg	
	OCDF	J	3.25	10.7	PQL	ng/Kg	
SL-519-SA8-SB-3.5-4.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.0612	4.98	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0393	4.98	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0331	4.98	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.0264	4.98	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0467	4.98	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.0278	4.98	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.0424	4.98	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0944	4.98	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0864	4.98	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0226	4.98	PQL	ng/Kg	
	OCDD	JBQ	0.151	9.96	PQL	ng/Kg	
OCDF	J	0.120	9.96	PQL	ng/Kg		
SL-520-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	0.553	5.01	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.672	5.01	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0642	5.01	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0392	5.01	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.213	5.01	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.140	5.01	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.480	5.01	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.412	5.01	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.223	5.01	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JB	0.0872	5.01	PQL	ng/Kg	
	2,3,7,8-TCDD	JQ	0.0360	1.00	PQL	ng/Kg	
	2,3,7,8-TCDF	J	0.274	1.00	PQL	ng/Kg	
	OCDD	JB	5.20	10.0	PQL	ng/Kg	
	OCDF	J	0.920	10.0	PQL	ng/Kg	
SL-531-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JB	1.15	5.20	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JB	0.486	5.20	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0349	5.20	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0742	5.20	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDD	JBQ	0.105	5.20	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JBQ	0.0725	5.20	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDD	JB	0.200	5.20	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JB	0.178	5.20	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.483	5.20	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0588	5.20	PQL	ng/Kg	
	2,3,4,7,8-PECDF	JBQ	0.0804	5.20	PQL	ng/Kg	
	2,3,7,8-TCDD	J	0.0392	1.04	PQL	ng/Kg	
	2,3,7,8-TCDF	JQ	0.0875	1.04	PQL	ng/Kg	
	OCDF	J	1.44	10.4	PQL	ng/Kg	

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Reporting Limit Outliers

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PH070_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 1613B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-531-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JB	0.0908	5.49	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0321	5.49	PQL	ng/Kg	
	1,2,3,4,7,8,9-HPCDF	JBQ	0.0315	5.49	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0210	5.49	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.0348	5.49	PQL	ng/Kg	
	1,2,3,7,8-PECDD	JQ	0.0482	5.49	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0291	5.49	PQL	ng/Kg	
	OCDD	JBQ	0.484	11.0	PQL	ng/Kg	
	OCDF	JQ	0.0502	11.0	PQL	ng/Kg	
SL-601-SA8-SB-0.0-0.5	1,2,3,4,6,7,8-HPCDD	JBQ	0.153	5.30	PQL	ng/Kg	J (all detects)
	1,2,3,4,6,7,8-HPCDF	JBQ	0.0548	5.30	PQL	ng/Kg	
	1,2,3,4,7,8-HxCDD	JQ	0.0367	5.30	PQL	ng/Kg	
	1,2,3,4,7,8-HXCDF	JBQ	0.0247	5.30	PQL	ng/Kg	
	1,2,3,6,7,8-HxCDD	JBQ	0.0569	5.30	PQL	ng/Kg	
	1,2,3,6,7,8-HXCDF	JB	0.0352	5.30	PQL	ng/Kg	
	1,2,3,7,8,9-HxCDD	JB	0.0378	5.30	PQL	ng/Kg	
	1,2,3,7,8,9-HXCDF	JBQ	0.0244	5.30	PQL	ng/Kg	
	1,2,3,7,8-PECDF	JBQ	0.0513	5.30	PQL	ng/Kg	
	2,3,4,6,7,8-HXCDF	JBQ	0.0369	5.30	PQL	ng/Kg	
	OCDD	JBQ	1.01	10.6	PQL	ng/Kg	
	OCDF	JQ	0.178	10.6	PQL	ng/Kg	
	SL-601-SA8-SB-4.0-5.0	1,2,3,4,6,7,8-HPCDD	JBQ	0.0961	5.26	PQL	
1,2,3,4,6,7,8-HPCDF		JBQ	0.0294	5.26	PQL	ng/Kg	
1,2,3,4,7,8,9-HPCDF		JBQ	0.0204	5.26	PQL	ng/Kg	
1,2,3,4,7,8-HXCDF		JBQ	0.0281	5.26	PQL	ng/Kg	
1,2,3,6,7,8-HXCDF		JB	0.0226	5.26	PQL	ng/Kg	
1,2,3,7,8,9-HxCDD		JBQ	0.135	5.26	PQL	ng/Kg	
1,2,3,7,8,9-HXCDF		JB	0.0890	5.26	PQL	ng/Kg	
1,2,3,7,8-PECDF		JB	0.0707	5.26	PQL	ng/Kg	
2,3,4,6,7,8-HXCDF		JBQ	0.0192	5.26	PQL	ng/Kg	
OCDD		JBQ	0.337	10.5	PQL	ng/Kg	
OCDF		J	0.0885	10.5	PQL	ng/Kg	

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-506-SA8-SB-0.0-0.5	BERYLLIUM	J	0.704	1.05	PQL	mg/Kg	J (all detects)
	BORON	J	4.93	10.5	PQL	mg/Kg	
	CADMIUM	J	0.520	1.05	PQL	mg/Kg	
	MOLYBDENUM	J	0.287	2.10	PQL	mg/Kg	
	TIN	J	2.83	10.5	PQL	mg/Kg	
	Zirconium	J	4.67	5.25	PQL	mg/Kg	
SL-506-SA8-SB-4.0-5.0	BERYLLIUM	J	0.677	1.01	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.289	1.01	PQL	mg/Kg	
	MOLYBDENUM	J	0.413	2.03	PQL	mg/Kg	
	TIN	J	2.86	10.1	PQL	mg/Kg	
	Zirconium	J	3.92	5.07	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PH070_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-509-SA8-SB-0.0-0.5	BERYLLIUM	J	0.781	1.10	PQL	mg/Kg	J (all detects)
	BORON	J	9.98	11.0	PQL	mg/Kg	
	CADMIUM	J	0.670	1.10	PQL	mg/Kg	
	TIN	J	2.80	11.0	PQL	mg/Kg	
SL-509-SA8-SB-4.0-5.0	BERYLLIUM	J	0.530	1.03	PQL	mg/Kg	J (all detects)
	BORON	J	1.26	10.3	PQL	mg/Kg	
	CADMIUM	J	0.425	1.03	PQL	mg/Kg	
	TIN	J	2.93	10.3	PQL	mg/Kg	
SL-519-SA8-SB-0.0-0.5	BERYLLIUM	J	0.827	1.06	PQL	mg/Kg	J (all detects)
	CADMIUM	J	0.751	1.06	PQL	mg/Kg	
	MOLYBDENUM	J	0.377	2.12	PQL	mg/Kg	
	SODIUM	J	103	106	PQL	mg/Kg	
	TIN	J	2.69	10.6	PQL	mg/Kg	
SL-519-SA8-SB-3.5-4.5	ARSENIC	J	3.75	4.14	PQL	mg/Kg	J (all detects)
	BERYLLIUM	J	0.458	1.03	PQL	mg/Kg	
	CADMIUM	J	0.339	1.03	PQL	mg/Kg	
	SODIUM	J	88.1	103	PQL	mg/Kg	
	TIN	J	2.85	10.3	PQL	mg/Kg	
	Zirconium	J	4.37	5.17	PQL	mg/Kg	
SL-520-SA8-SB-0.0-0.5	BERYLLIUM	J	0.785	1.04	PQL	mg/Kg	J (all detects)
	BORON	J	4.36	10.4	PQL	mg/Kg	
	CADMIUM	J	0.591	1.04	PQL	mg/Kg	
	MOLYBDENUM	J	0.315	2.09	PQL	mg/Kg	
	SODIUM	J	84.3	104	PQL	mg/Kg	
	TIN	J	2.66	10.4	PQL	mg/Kg	
	Zirconium	J	4.76	5.21	PQL	mg/Kg	
SL-531-SA8-SB-0.0-0.5	BERYLLIUM	J	0.882	1.06	PQL	mg/Kg	J (all detects)
	BORON	J	8.08	10.6	PQL	mg/Kg	
	CADMIUM	J	0.673	1.06	PQL	mg/Kg	
	MOLYBDENUM	J	0.238	2.13	PQL	mg/Kg	
	SODIUM	J	94.6	106	PQL	mg/Kg	
	TIN	J	2.65	10.6	PQL	mg/Kg	
	Zirconium	J	4.37	5.31	PQL	mg/Kg	
SL-531-SA8-SB-4.0-5.0	BERYLLIUM	J	0.944	1.08	PQL	mg/Kg	J (all detects)
	BORON	J	3.58	10.8	PQL	mg/Kg	
	CADMIUM	J	0.505	1.08	PQL	mg/Kg	
	MOLYBDENUM	J	0.343	2.15	PQL	mg/Kg	
	SODIUM	J	98.4	108	PQL	mg/Kg	
	TIN	J	2.91	10.8	PQL	mg/Kg	
SL-601-SA8-SB-0.0-0.5	BERYLLIUM	J	0.790	1.02	PQL	mg/Kg	J (all detects)
	BORON	J	9.22	10.2	PQL	mg/Kg	
	CADMIUM	J	0.599	1.02	PQL	mg/Kg	
	MOLYBDENUM	J	0.452	2.04	PQL	mg/Kg	
	SODIUM	J	93.6	102	PQL	mg/Kg	
	TIN	J	2.63	10.2	PQL	mg/Kg	
	Zirconium	J	2.94	5.10	PQL	mg/Kg	

Reporting Limit Outliers

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PH070_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 6010C
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-601-SA8-SB-4.0-5.0	BERYLLIUM	J	0.730	1.02	PQL	mg/Kg	J (all detects)
	BORON	J	4.04	10.2	PQL	mg/Kg	
	CADMIUM	J	0.412	1.02	PQL	mg/Kg	
	MOLYBDENUM	J	0.351	2.04	PQL	mg/Kg	
	SODIUM	J	92.7	102	PQL	mg/Kg	
	TIN	J	2.88	10.2	PQL	mg/Kg	
	Zirconium	J	3.65	5.09	PQL	mg/Kg	

Method: 6020A
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-506-SA8-SB-0.0-0.5	SELENIUM	J	0.315	0.420	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0732	0.210	PQL	mg/Kg	
SL-509-SA8-SB-0.0-0.5	SELENIUM	J	0.281	0.439	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0321	0.219	PQL	mg/Kg	
SL-519-SA8-SB-0.0-0.5	SELENIUM	J	0.209	0.425	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0699	0.212	PQL	mg/Kg	
SL-520-SA8-SB-0.0-0.5	SELENIUM	J	0.156	0.417	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0373	0.209	PQL	mg/Kg	
SL-531-SA8-SB-0.0-0.5	SELENIUM	J	0.225	0.425	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0320	0.213	PQL	mg/Kg	
SL-531-SA8-SB-4.0-5.0	SILVER	J	0.0474	0.215	PQL	mg/Kg	J (all detects)
SL-601-SA8-SB-0.0-0.5	SELENIUM	J	0.248	0.408	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0287	0.204	PQL	mg/Kg	
SL-601-SA8-SB-4.0-5.0	SELENIUM	J	0.123	0.407	PQL	mg/Kg	J (all detects)
	SILVER	J	0.0299	0.204	PQL	mg/Kg	

Method: 7199
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-506-SA8-SB-0.0-0.5	HEXAVALENT CHROMIUM	J	0.30	0.43	PQL	mg/Kg	J (all detects)
SL-509-SA8-SB-0.0-0.5	HEXAVALENT CHROMIUM	J	0.18	0.44	PQL	mg/Kg	J (all detects)
SL-509-SA8-SB-4.0-5.0	HEXAVALENT CHROMIUM	J	0.18	0.42	PQL	mg/Kg	J (all detects)

Method: 7471B
Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-531-SA8-SB-4.0-5.0	MERCURY	J	0.0139	0.0186	PQL	mg/Kg	J (all detects)

Project Name and Number: 1204-002-001-AL - SSFL Area IV Phase 3

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Reporting Limit Outliers

Lab Reporting Batch ID: PH070

Laboratory: LL

EDD Filename: PH070_v1.

eQAPP Name: CDM_SSFL_130808_Lan

Method: 8015M

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-519-SA8-SB-0.0-0.5	EFH (C15-C20)	J	2.8	5.4	PQL	mg/Kg	J (all detects)
SL-520-SA8-SB-0.0-0.5	EFH (C21-C30)	J	3.9	5.2	PQL	mg/Kg	J (all detects)
	EFH (C30-C40)	J	7.9	10	PQL	mg/Kg	J (all detects)
SL-531-SA8-SB-0.0-0.5	EFH (C15-C20)	J	2.5	5.3	PQL	mg/Kg	J (all detects)
SL-601-SA8-SB-4.0-5.0	EFH (C21-C30)	J	4.4	5.3	PQL	mg/Kg	J (all detects)
	EFH (C30-C40)	J	7.3	11	PQL	mg/Kg	J (all detects)

Method: 8082A

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-506-SA8-SB-0.0-0.5	AROCLOR 1254	J	5.0	18	PQL	ug/Kg	J (all detects)
SL-531-SA8-SB-0.0-0.5	AROCLOR 1254	J	5.5	18	PQL	ug/Kg	J (all detects)

Method: 8270D SIM

Matrix: SO

SampleID	Analyte	Lab Qual	Result	Reporting Limit	RL Type	Units	Flag
SL-506-SA8-SB-0.0-0.5	BENZO(K)FLUORANTHENE	J	1.7	1.8	PQL	ug/Kg	J (all detects)
	BIS(2-ETHYLHEXYL)PHTHALATE	J	6.6	19	PQL	ug/Kg	
	CHRYSENE	J	1.1	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	0.74	1.8	PQL	ug/Kg	
SL-509-SA8-SB-0.0-0.5	BENZO(B)FLUORANTHENE	J	0.78	1.8	PQL	ug/Kg	J (all detects)
	BENZO(K)FLUORANTHENE	J	1.1	1.8	PQL	ug/Kg	
	CHRYSENE	J	1.1	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	0.85	1.8	PQL	ug/Kg	
SL-519-SA8-SB-0.0-0.5	1-METHYLNAPHTHALENE	J	1.1	1.8	PQL	ug/Kg	J (all detects)
	ANTHRACENE	J	0.38	1.8	PQL	ug/Kg	
	BENZO(A)ANTHRACENE	J	0.96	1.8	PQL	ug/Kg	
	BENZO(A)PYRENE	J	1.4	1.8	PQL	ug/Kg	
	BENZO(E)PYRENE	J	4.0	18	PQL	ug/Kg	
	BIS(2-ETHYLHEXYL)PHTHALATE	J	8.9	19	PQL	ug/Kg	
	INDENO(1,2,3-CD)PYRENE	J	1.7	1.8	PQL	ug/Kg	
SL-520-SA8-SB-0.0-0.5	BENZO(K)FLUORANTHENE	J	0.92	1.7	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	0.43	1.7	PQL	ug/Kg	
SL-531-SA8-SB-0.0-0.5	BENZO(K)FLUORANTHENE	J	1.5	1.8	PQL	ug/Kg	J (all detects)
	CHRYSENE	J	1.0	1.8	PQL	ug/Kg	
	NAPHTHALENE	J	0.82	1.8	PQL	ug/Kg	
SL-601-SA8-SB-0.0-0.5	CHRYSENE	J	0.73	1.8	PQL	ug/Kg	J (all detects)
	NAPHTHALENE	J	0.93	1.8	PQL	ug/Kg	

LDC #: 30289E4

VALIDATION COMPLETENESS WORKSHEET

Date: 9/4/13

SDG #: PH070

ADR

Page: 1 of 1

Laboratory: Eurofins Lancaster Laboratories

Reviewer: [Signature]

2nd Reviewer: [Signature]

METHOD: Metals (EPA SW 846 Method 6010C/6020A/7000)

The samples listed below were reviewed for each of the following validation areas. Validation findings are noted in attached validation findings worksheets.

	Validation Area		Comments
I.	Technical holding times	-	Sampling dates: 7/18/13
II.	ICP/MS Tune	-	
III.	Calibration	-	
IV.	Blanks	A	
V.	ICP Interference Check Sample (ICS) Analysis	-	
VI.	Matrix Spike Analysis	A	MSPD (Hg only)
VII.	Duplicate Sample Analysis	SW A	Dup ↓ Hg RPD=200 ok by difference
VIII.	Laboratory Control Samples (LCS)	A	LCS
IX.	Internal Standard (ICP-MS)	N	
X.	Furnace Atomic Absorption QC	N	
XI.	ICP Serial Dilution	N	
XII.	Sample Result Verification	N	
XIII.	Overall Assessment of Data	N	
XIV.	Field Duplicates	-	
XV.	Field Blanks	SW	EB = EB1-071713 FB = FB-041113

Note: A = Acceptable
N = Not provided/applicable
SW = See worksheet

ND = No compounds detected
R = Rinsate
FB = Field blank

(P17069)
D = Duplicate
TB = Trip blank
EB = Equipment blank

(P17029)

Validated Samples:

soil

1	SL-506-SA8-SB-0.0-0.5	11	SL-531-SA8-SB-4.0-5.0	21		31	
2	SL-506-SA8-SB-4.0-5.0	12	(X11) MS (trg)	22		32	
3	SL-509-SA8-SB-0.0-0.5	13	MSPD	23		33	
4	SL-509-SA8-SB-4.0-5.0	14	DUP	24		34	
5	SL-601-SA8-SB-0.0-0.5	15		25		35	
6	SL-601-SA8-SB-4.0-5.0	16		26		36	
7	SL-519-SA8-SB-0.0-0.5	17		27		37	
8	SL-519-SA8-SB-3.5-4.5	18		28		38	
9	SL-520-SA8-SB-0.0-0.5	19		29		39	
10	SL-531-SA8-SB-0.0-0.5	20		30		40	

Notes: _____

VALIDATION FINDINGS WORKSHEET Field Blanks

METHOD: Trace Metals (EPA SW846 6010B/7000)

Blank units: mg/L **Associated sample units:** mg/Kg Reason: F

Sampling date: 4/11/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification										
	FB-041113 (SDG: PH029)	Action Limit	1	2	5	6	7	9	10	11		
Cu	0.0036	1.8										
Mo	0.0036	1.8	0.29	0.41	0.45	0.35	0.38	0.31	0.24	0.34		

Sampling date: 7/17/13 Soil factor applied 100x

Field blank type: (circle one) Field Blank / Rinsate / Other: _____ Associated Samples: All

Analyte	Blank ID	Sample Identification										
	EB1-071713 (SDG: PH069)	Action Limit	No Qualifiers									
Sb	0.0069	3.45										

CIRCLED RESULTS WERE NOT QUALIFIED. ALL RESULTS NOT CIRCLED WERE QUALIFIED BY THE FOLLOWING STATEMENT:
Samples with analyte concentrations within five times the associated field blank concentration are listed above, these sample results were qualified as not detected, "U".

**Data Validation Report
Santa Susana Field Laboratory**

Subarea 8

SDG: PH071

Prepared for

CDM
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Prepared by

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September 24, 2013

INTRODUCTION

This Data Validation Report (DVR) presents Level III data validation results for samples collected on July 19th, 2013. Data validation was performed in accordance with the Quality Assurance Project Plan for Santa Susana Field Laboratory (SSFL), RCRA Facility Investigation, Surficial Media Operable Unit (March 2009, Revision 4) and a modified outline of the USEPA Contract Laboratory Program National Functional Guidelines for Superfund Organic Methods Data Review (June 2008), Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005), and for Inorganic Data Review (January 2010). Where specific guidance is not available, the data has been evaluated in a conservative manner consistent with industry standards using professional experience.

The analyses were performed by the following methods:

Semivolatiles (SVOAs) by EPA SW 846 Method 8270D utilizing Selective Ion Monitoring (SIM)
Polychlorinated Biphenyls (PCBs) by EPA SW 846 Method 8082A
Metals by EPA SW 846 Method 6010C, 6020A, and 7471B
Total Petroleum Hydrocarbon as Gasoline (TPH-G) by EPA SW 846 Method 8015M
Total Petroleum Hydrocarbons as Extractable (TPH-E) by EPA SW 846 Method 8015M
Dioxins/Dibenzofurans by EPA Method 1613B
Hexavalent Chromium by EPA method 7199

The sample identification and methods of analyses performed on each sample is presented in Attachment 1. Overall data qualification summary is presented in Attachment II. Level III Automated Data Review outliers are presented in Enclosure I.

All sample results were subjected to Level III data validation, which comprises an evaluation of quality control (QC) summary results for sample holding times, initial and continuing calibration blanks (ICB/CCBs), surrogates, internal standards (dioxins only), matrix spike/matrix spike duplicates (MS/MSD), laboratory duplicates (DUP), laboratory control sample/laboratory control sample duplicates (LCS/LCSD), ICP serial dilutions, method blanks, trip blanks, equipment blanks, field blanks, and field duplicate samples. No samples in this SDG were subjected to Level IV evaluation.

Automated data review was performed on all QC summary results using the Automated Data Review (ADR) software program (LDC, 2013) with exception of the ICB/CCBs and ICP serial dilutions, which were validated manually. Quality assurance (QA)/QC criteria specified in the QAPP and CLPNFGs were incorporated with the program's reference library to assess compliance with project requirements.

The following are definitions of the data qualifiers:

- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- J Indicates an estimated value.
- R Quality control indicates the data is not usable.
- NJ Presumptive evidence of presence of the compound at an estimated quantity.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.