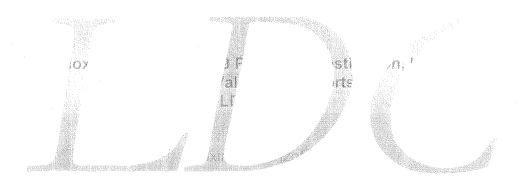
Tronox LLC Facility, 2009 Phase B Investigation, Henderson Data Validation Reports LDC #21495

Dioxins/Dibenzofurans



LDC Report# 21495C21

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name:

Tronox LLC Facility, 2009 Phase B Investigation,

Henderson, Nevada

Collection Date:

June 1 through June 4, 2009

LDC Report Date:

November 2, 2009

Matrix:

Soil

Parameters:

Dioxins/Dibenzofurans

Validation Level:

Stage 2B

Laboratory:

Columbia Analytical Services, Inc.

Sample Delivery Group (SDG): R0903051

Sample Identification

RSA12-0.5B

RSA12-0.5BDL

RSAI3-0.5B

RSAJ5-0.5BDL RSAJ5-0.5B

RSAJ5-0.5BDL

RSAK5-0.5B

RSAK5-0.5BDL

SA76-0.5B

SA76-0.5BDL

SA76009-0.5B

SA76009-0.5BDL

RSAL3-0.5B

RSAL3-0.5BDL

SA100-0.5B

RSAM3-0.5B

RSAM2-0.5B

SA189-0.5B

SA189-0.5BDL

SA88-0.5B

SA88-0.5BDL

SA152-0.5B

SA152009-0.5B

RSAJ2-0.5B

RSAJ2-0.5BDL

RSAJ3-0.5B

SA202-0.5B

Introduction

This data review covers 27 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8290 for Polychlorinated Dioxins/Dibenzofurans.

This review follows the Standard Operating Procedures (SOP) 40, Data Review/Validation (BRC 2009), the Quality Assurance Project Plan Tronox LLC Facility, Henderson, Nevada (June 2009), NDEP guidance (May 2006), and USEPA Contract Laboratory Program National Functional Guidelines for Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) as there are no current guidelines for the method stated above.

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.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XIV.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- J+ Data are qualified as estimated, with a high bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J- Data are qualified as estimated, with a low bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J Data are qualified as estimated; it is not possible to assess the direction of the potential bias. False positives or false negatives are unlikely to have been reported.
- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- R Data are qualified as rejected. There is a significant potential for the reporting of false negatives or false positives.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- B The analytical result may be a false positive totally attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JB The analytical result may be biased high and partially attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JK The analytical result is an estimated maximum possible concentration (EMPC).
- X The analytical result is not used for reporting because a more accurate and precise result is reported in its place.
- J-TDS The analytical result is estimated based on failure of the Total Dissolved Solids (TDS) correctness check performed in accordance with the Standard Method 1030E.
- J-CAB The analytical result is estimated based on failure of the cation-anion balance correctness check performed in accordance with Standard Method 1030E.
- J-TDS & CAB The analytical result is unreliable based on the failure of the cation-anion balance and TDS correctness check performed in accordance with standard Method 1030E.
- 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.

Retention time windows were established for all homologues. The chromatographic resolution between 2,3,7,8-TCDD and peaks representing any other unlabeled TCDD isomer was less than or equal to 25%.

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 30.0% for labeled compounds.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

IV. Routine Calibration (Continuing)

Routine calibration was performed at the required frequencies.

All of the routine calibration percent differences (%D) between the initial calibration RRF and the routine calibration RRF were less than or equal to 20.0% for unlabeled compounds and less than or equal to 30.0% for labeled compounds.

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
EQ0900197-01	6/10/09	1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,6,7,8-HpCDF OCDF Total HpCDD	0.449 ng/Kg 3.26 ng/Kg 0.193 ng/Kg 0.399 ng/Kg 0.449 ng/Kg	RSA12-0.5B RSA12-0.5BDL RSA13-0.5B RSA13-0.5BDL RSAJ5-0.5BDL RSAJ5-0.5BDL RSAK5-0.5BDL SA76-0.5BDL SA76-0.5BDL SA76009-0.5B SA76009-0.5BDL RSAL3-0.5B RSAL3-0.5BDL SA100-0.5B RSAM3-0.5B RSAM3-0.5B RSAM2-0.5B RSAM2-0.5B SA88-0.5B SA88-0.5BDL SA152-0.5B RSAJ2-0.5B RSAJ2-0.5B RSAJ2-0.5B
EQ0900210-01	6/16/09	1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,6,7,8-HpCDF OCDF Total HpCDD Total HpCDF	0.283 ng/Kg 1.52 ng/Kg 0.268 ng/Kg 0.468 ng/Kg 0.283 ng/Kg 0.268 ng/Kg	SA189-0.5B SA189-0.5BDL SA152009-0.5B RSAJ3-0.5B

Sample concentrations were compared to concentrations detected in the method blanks as required by the QAPP. No sample data was qualified with the following exceptions:

Sample	Compound	Reported Concentration	Modified Final Concentration
SA100-0.5B	OCDD	14.3 ng/Kg	14.3U ng/Kg
RSAM2-0.5B	1,2,3,4,6,7,8-HpCDD OCDD	1.94 ng/Kg 5.08 ng/Kg	1.94U ng/Kg 5.08U ng/Kg

Sample FB072109-SO (from SDG R0904016) was identified as a field blank. No polychlorinated dioxin/dibenzofuran contaminants were found in this blank with the following exceptions:

Field Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB072109-SO	7/21/09	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF Total HpCDD Total HxCDF Total HpCDF	8.37 pg/L 21.8 pg/L 6.33 pg/L 3.57 pg/L 4.04 pg/L 43.3 pg/L 19.6 pg/L 205 pg/L 8.37 pg/L 18.1 pg/L 72.1 pg/L	All samples in SDG R0903051

Sample concentrations were compared to concentrations detected in the field blanks as required by the QAPP. No sample data was qualified.

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 (LCS)

Laboratory control samples were reviewed for each matrix as applicable. Although the LCSD percent recoveries (%R) and relative percent differences (RPD) were not within QC limits for some compounds, the LCS or LCSD percent recoveries (%R) were within QC limits and no data were qualified.

VIII. Regional Quality Assurance and Quality Control

Not applicable.

IX. Internal Standards

All internal standard recoveries were within QC limits with the following exceptions:

Sample	Internal Standards	%R (Limits)	Compound	Flag	A or P
RSA12-0.5B	¹³ C-2,3,7,8-TCDF	19 (40-135)	2,3,7,8-TCDF Total TCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р
RSA12-0.5BDL	¹³ C-2,3,7,8-TCDF (DB-225)	156 (40-135)	2,3,7,8-TCDF (DB-225)	J (all detects) UJ (all non-detects)	P

Sample	Internal Standards	%R (Limits)	Compound	Flag	A or P
SA76-0.5B	¹³ C-2,3,7,8-TCDF	20 (40-135)	2,3,7,8-TCDF Total TCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р
SA76009-0.5B	¹³ C-2,3,7,8-TCDF (DB-225) ¹³ C-2,3,7,8-TCDF (DB-5)	142 (40-135) 17 (40-135)	2,3,7,8-TCDF (DB-225) 2,3,7,8-TCDF (DB-5) Total TCDF	J (all detects) UJ (all non-detects)	Р
SA76009-0.5BDL	¹³ C-2,3,7,8-TCDF ¹³ C-1,2,3,4,7,8-HxCDF	17 (40-135) 38 (40-135)	2,3,7,8-TCDF 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	J (all detects) UJ (all non-detects)	Р
SA202-0.5B	¹³ C-2,3,7,8-TCDF	38 (40-135)	2,3,7,8-TCDF Total TCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р

X. Target Compound Identifications

Raw data were not reviewed for this SDG.

*XI. Project Quantitation Limit

All project quantitation limits were within validation criteria with the following exceptions:

Sample	Compound	Finding	Criteria	Flag	A or P
RSA12-0.5B SA76009-0.5B SA76-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HyCDD 0CDD 1,2,3,4,6,7,8-HyCDD 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,7,8,9-HyCDF 0CDF Total TCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A

Sample RSAI3-0.5B	Compound 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 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	Finding Sample result exceeded calibration range.	Criteria Reported result should be within callibration range.	J (all detects)	A or P
RSAJ5-0.5B SA88-0.5B RSAJ2-0.5B	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 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A
RSAK5-0.5B	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 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A
*RSAL3-0.5B	1,2,3,4,6,7,8-HpCDD 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	А

Sample	Compound	Finding	Criteria	Flag	A or P
SA189-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A
RSAJ3-0.5B	1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	Р

^{*}Changed associated compounds for sample noted above.

All compounds reported below the PQL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903051	All compounds reported below the PQL.	J (all detects)	А

All compounds reported as EMPC were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903051	All compounds reported as estimated maximum possible concentration (EMPC).	JK (all detects)	А

Raw data were not reviewed for this SDG.

XII. System Performance

Raw data were not reviewed for this SDG.

*XIII. Overall Assessment of Data

The overall assessment of data was acceptable. In the case where more than one result was reported for an individual sample, the least technically acceptable results were rejected as follows:

Sample	Compound	Flag	A or P
RSA12-0.5B SA76009-0.5B SA76-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 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 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X X X	A
RSA12-0.5BDL SA76009-0.5BDL SA100-0.5B RSAM3-0.5B RSAM2-0.5B SA152-0.5B SA152009-0.5B RSAJ3-0.5B SA202-0.5B *SA76-0.5BDL	2,3,7,8-TCDF (from DB-5)	x	А
RSAI3-0.5B	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 1,2,3,4,6,7,8-HpCDF 2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X X	A
RSAI3-0.5BDL	All TCL compounds except 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-225)	x	А

Sample	Compound	Flag	A or P
RSAJ5-0.5B SA88-0.5B RSAJ2-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x x x x	A
RSAJ5-0.5BDL SA88-0.5BDL RSAJ2-0.5BDL	All TCL compounds except 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-225)	x	А
RSAK5-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X X X	Α

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Sample	Compound	Flag	A or P
RSAK5-0.5BDL	All TCL compounds except 1,2,3,7,8-PeCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-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 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 0CDF 2,3,7,8-TCDF (DB-225)	X	
*RSAL3-0.5B	1,2,3,4,6,7,8-HpCDD 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X	А
*RSAL3-0.5BDL	All TCL compounds except 1,2,3,4,6,7,8-HpCDD 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF	X	A
SA189-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X	А
SA189-0.5BDL	All TCL compounds except 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF	X	A

^{*}Changed SA76-0.5B to SA76-0.5BDL in table above and changed associated compounds for samples noted above.

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

Samples SA76-0.5B and SA76009-0.5B, samples SA76-0.5BDL and SA76009-0.5BDL, and samples SA152-0.5B and SA152009-0.5B were identified as field duplicates. No polychlorinated dioxins/dibenzofurans were detected in any of the samples with the following exceptions:

	Concentrati	on (ng/Kg)		D:#		
Compound	SA76-0.5B	SA76009-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P
2,3,7,8-TCDD	433	676	44 (≤50)	-	-	-
1,2,3,7,8-PeCDD	2460	3870	45 (≤50)	-	-	-
1,2,3,4,7,8-HxCDD	1860	3070	49 (≤50)	-	-	_
1,2,3,6,7,8-HxCDD	3330	5260	45 (≤50)	-	-	
1,2,3,7,8,9-HxCDD	4020	6580	48 (≤50)	-	-	-
1,2,3,4,6,7,8-HpCDD	14400	17200	18 (≤50)	-	-	-
OCDD	13200	20800	45 (≤50)	-	-	-
2,3,7,8-TCDF	81400	138000	52 (≤50)	-	J (all detects)	А
1,2,3,7,8-PeCDF	29900	51900	54 (≤50)	-	J (all detects)	А
2,3,4,7,8-PeCDF	14500	26700	59 (≤50)	-	J (all detects)	А
1,2,3,4,7,8-HxCDF	77200	120000	43 (≤50)	-	-	-
1,2,3,6,7,8-HxCDF	47000	77900	49 (≤50)	-	-	-
1,2,3,7,8,9-HxCDF	9010	16400	58 (≤50)	-	J (all detects)	А
2,3,4,6,7,8-HxCDF	9770	16400	51 (≤50)	-	J (all detects)	А
1,2,3,4,6,7,8-HpCDF	95300	132000	32 (≤50)	-	-	-
1,2,3,4,7,8,9-HpCDF	84900	121000	35 (≤50)	-	-	-

	Concentrati	on (ng/Kg)				
Compound	SA76-0.5B	SA76009-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P
OCDF	197000	271000	32 (≤50)	-	-	_
Total TCDD	25100	35700	35 (≤50)	-	-	-
Total PeCDD	31200	48800	44 (≤50)	-	-	-
Total HxCDD	26900	43600	47 (≤50)	-	-	-
Total HpCDD	22400	26700	18 (≤50)	•	-	-
Total TCDF	983000	1630000	50 (≤50)	-	-	-
Total PeCDF	234000	283000	19 (≤50)	-	-	-
Total HxCDF	366000	536000	38 (≤50)	-	-	-
Total HpCDF	296000	431000	37 (≤50)	-	-	-
2,3,7,8-TCDF (DB-225)	12900	15100	16 (≤50)	-	-	-

	Concentrati	ion (ng/Kg)				
Compound	SA76-0.5BDL	SA76009-0.5BDL	RPD (Limits)	Difference (Limits)	Flag	A or P
2,3,7,8-TCDD	259	376	-	117 (≤199)	-	-
1,2,3,7,8-PeCDD	1040	1190	-	150 (≤497)	-	-
1,2,3,4,7,8-HxCDD	824	1040	-	216 (≤497)	-	-
1,2,3,6,7,8-HxCDD	1920	2460	-	540 (≤497)	J (all detects)	А
1,2,3,7,8,9-HxCDD	2060	2640	-	580 (≤497)	J (all detects)	А
1,2,3,4,6,7,8-HpCDD	8140	7640	6 (≤50)	-	-	-
OCDD	8370	10600	24 (≤50)	-	-	-
2,3,7,8-TCDF	12400	103000	157 (≤50)	-	J (all detects)	А

	Concentrati	on (ng/Kg)		D	 	
Compound	SA76-0.5BDL	SA76009-0.5BDL	RPD (Limits)	Difference (Limits)	Flag	A or P
1,2,3,7,8-PeCDF	12600	21400	52 (≤50)	-	J (all detects)	А
2,3,4,7,8-PeCDF	5050	10400	69 (≤50)	-	J (all detects)	А
1,2,3,4,7,8-HxCDF	62800	88200	34 (≤50)	-	-	-
1,2,3,6,7,8-HxCDF	37100	54500	38 (≤50)	•	-	-
1,2,3,7,8,9-HxCDF	4280	7060	49 (≤50)	-	-	-
2,3,4,6,7,8-HxCDF	8640	11700	30 (≤50)	-	-	-
1,2,3,4,6,7,8-HpCDF	135000	167000	21 (≤50)	-	-	-
1,2,3,4,7,8,9-HpCDF	62300	80600	26 (≤50)	-	-	-
OCDF	305000	397000	26 (≤50)	-	-	-
2,3,7,8-TCDF (DB-225)	6470	6240	4 (≤50)	-	-	<u>-</u>

	Concentration (ng/Kg) RPD Difference		D			
Compound	SA152-0.5B	SA152009-0.5B	(Limits)	(Limits)	Flag	A or P
2,3,7,8-TCDD	1.11	1.90U	-	0.79 (≤1.90)	-	-
1,2,3,7,8-PeCDD	3.79	4.75U	-	0.96 (≤4.75)	-	-
1,2,3,4,7,8-HxCDD	1.97	4.75U	-	2.78 (≤4.75)	-	-
1,2,3,6,7,8-HxCDD	4.33	4.75U	-	0.42 (≤4.75)	-	-
1,2,3,7,8,9-HxCDD	5.90	0.605	•	5.295 (≤4.75)	J (all detects)	Α
1,2,3,4,6,7,8-HpCDD	13.7	1.50	-	12.2 (≤4.75)	J (all detects)	А
OCDD	15.9	9.17	-	6.73 (≤9.50)	-	-
2,3,7,8-TCDF	62.4	0.565	•	61.835 (≤1.90)	J (all detects)	А

	Concentra	tion (ng/Kg)				
Compound	SA152-0.5B	SA152009-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P
1,2,3,7,8-PeCDF	52.0	4.75U	-	47.25 (≤4.75)	J (all detects) UJ (all non-detects)	А
2,3,4,7,8-PeCDF	27.9	4.75U	-	23.15 (≤4.75)	J (all detects) UJ (all non-detects)	А
1,2,3,4,7,8-HxCDF	107	0.952	-	106.048 (≤4.75)	J (all detects)	А
1,2,3,6,7,8-HxCDF	63.0	0.634	-	62.366 (≤4.75)	J (all detects)	А
1,2,3,7,8,9-HxCDF	10.0	4.75U	-	5.25 (≤4.75)	J (all detects) UJ (all non-detects)	А
2,3,4,6,7,8-HxCDF	15.9	4.75U	-	11.15 (≤4.75)	J (all detects) UJ (all non-detects)	А
1,2,3,4,6,7,8-HpCDF	187	2.52	-	184.48 (≤4.75)	J (all detects)	А
1,2,3,4,7,8,9-HpCDF	92.2	0.941	•	91.259 (≤4.75)	J (all detects)	А
OCDF	518	6.45	-	511.55 (≤9.50)	J (all detects)	А
Total TCDD	26.3	1.90U	-	24.4 (≤1.90)	J (all detects) UJ (all non-detects)	А
Total PeCDD	37.4	4.75U	-	32.65 (≤4.75)	J (all detects) UJ (all non-detects)	А
Total HxCDD	33.4	0.605	-	32.795 (≤4.75)	J (all detects)	А
Total HpCDD	21.4	1.50	-	19.9 (≤4.75)	J (all detects)	А
Total TCDF	471	0.565	-	470.435 (≤1.90)	J (all detects)	А
Total PeCDF	500	1.12	-	498.88 (≤4.75)	J (all detects)	А
Total HxCDF	400	0.952	-	399.048 (≤4.75)	J (all detects)	А
Total HpCDF	405	3.08	*	401.92 (≤4.75)	J (all detects)	А
2,3,7,8-TCDF (DB-225)	11.2	3.35	_	7.85 (≤1.90)	J (all detects)	А

*Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Data Qualification Summary - SDG R0903051

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SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903051	RSA12-0.5B SA76-0.5B SA202-0.5B	2,3,7,8-TCDF Total TCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р	Internal standards (area) (i)
R0903051	RSA12-0.5BDL	2,3,7,8-TCDF (DB-225)	J (all detects) UJ (all non-detects)	Р	Internal standards (area)
R0903051	SA76009-0.5B	2,3,7,8-TCDF (DB-225) 2,3,7,8-TCDF (DB-5) Total TCDF	J (all detects) UJ (all non-detects)	Р	Internal standards (area) (i)
R0903051	SA76009-0.5BDL	2,3,7,8-TCDF 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	J (all detects) UJ (all non-detects)	Р	Internal standards (area) (i)
R0903051	RSA12-0.5B SA76009-0.5B SA76-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HyCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 1,2,3,4,7,8,9-HpCDF Total TCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903051	RSAI3-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903051	RSAJ5-0.5B SA88-0.5B RSAJ2-0.5B	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 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	А	Project Quantitation Limit (e)
R0903051	RSAK5-0.5B	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 1,2,3,7,8,9-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
*R0903051	RSAL3-0.5B	1,2,3,4,6,7,8-HpCDD 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	А	Project Quantitation Limit (e)
R0903051	SA189-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903051	RSAJ3-0.5B	1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5)	J (all detects)	Р	Project Quantitation Limit (e)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903051	RSA12-0.5B RSA12-0.5BDL RSA13-0.5BDL RSA13-0.5BDL RSAJ5-0.5BDL RSAK5-0.5BDL SA76-0.5BDL SA76-0.5BDL SA76-0.5BDL SA76009-0.5B SA76009-0.5BDL RSAL3-0.5B RSAL3-0.5BDL SA100-0.5B RSAM3-0.5B RSAM3-0.5B SA189-0.5B SA189-0.5B SA189-0.5B SA189-0.5B SA152-0.5B SA152-0.5B SA152-0.5B SA152-0.5B SA152-0.5B SA152-0.5B RSAJ2-0.5B RSAJ2-0.5B RSAJ2-0.5B RSAJ2-0.5B RSAJ2-0.5B RSAJ2-0.5BDL	All compounds reported below the PQL.	J (all detects)	A	Project Quantitation Limit (sp)
R0903051	RSA12-0.5B RSA12-0.5BDL RSAI3-0.5B RSAI3-0.5BDL RSAJ5-0.5B RSAJ5-0.5BDL RSAK5-0.5BDL SA76-0.5BDL SA76-0.5BDL SA76-0.5BDL SA76009-0.5B SA76009-0.5BDL RSAL3-0.5BDL SA100-0.5B RSAM3-0.5BDL SA100-0.5B RSAM3-0.5B RSAM3-0.5B SA189-0.5B SA189-0.5B SA189-0.5B SA189-0.5BDL SA152-0.5B SA152-0.5B SA8-0.5BDL SA152-0.5B SA152-0.5B SA152-0.5B SA152-0.5B SA152-0.5B RSAJ2-0.5BDL SA152-0.5B RSAJ2-0.5B RSAJ2-0.5BDL RSAJ2-0.5BDL RSAJ2-0.5BDL RSAJ2-0.5BDL RSAJ2-0.5B RSAJ2-0.5BDL	All compounds reported as EMPC	JK (all detects)	A	Project Quantitation Limit (k)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903051	RSA12-0.5B SA76009-0.5B SA76-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8,8-TCDF (DB-5 & DB-225)	X X X X X X X X X X	A	Overall assessment of data (o)
*R0903051	RSA12-0.5BDL SA76009-0.5BDL SA100-0.5B RSAM3-0.5B RSAM2-0.5B SA152-0.5B SA152009-0.5B RSAJ3-0.5B SA202-0.5B SA76-0.5BDL	2,3,7,8-TCDF (from DB-5)	X	A	Overall assessment of data (o)
R0903051	RSAI3-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X	А	Overall assessment of data (o)
R0903051	RSAI3-0.5BDL	All TCL compounds except 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 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-225)	X	А	Overall assessment of data (o)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903051	RSAJ5-0.5B SA88-0.5B RSAJ2-0.5B	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 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x x x x x	A	Overall assessment of data (o)
R0903051	RSAJ5-0.5BDL SA88-0.5BDL RSAJ2-0.5BDL	All TCL compounds except 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 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-225)	X	A	Overall assessment of data (o)
R0903051	RSAK5-0.5B	1,2,3,7,8-PeCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-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 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X X	A	Overall assessment of data (o)

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SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903051	RSAK5-0.5BDL	All TCL compounds except 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-225)	X	A	Overall assessment of data (o)
*R0903051	RSAL3-0.5B	1,2,3,4,6,7,8-HpCDD 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x	A	Overall assessment of data (o)
*R0903051	RSAL3-0.5BDL	All TCL compounds except 1,2,3,4,6,7,8-HpCDD 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF	x	A	Overall assessment of data (o)
R0903051	SA189-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X	A	Overall assessment of data (o)
R0903051	SA189-0.5BDL	All TCL compounds except 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF	X	A	Overall assessment of data (o)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903051	SA76-0.5B SA76009-0.5B	2,3,7,8-TCDF 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,7,8,9-HxCDF 2,3,4,6,7,8-HxCDF	J (all detects)	A	Field duplicates (RPD) (fd)
R0903051	SA76-0.5BDL SA76009-0.5BDL	1,2,3,6,7,8-HxCDD 1,2,3,7,8,9-HxCDD	J (all detects) J (all detects)	Α	Field duplicates (RPD) (fd)
R0903051	SA76-0.5BDL SA76009-0.5BDL	2,3,7,8-TCDF 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF	J (all detects) J (all detects) J (all detects)	А	Field duplicates (Difference) (fd)
R0903051	SA152-0.5B SA152009-0.5B	1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDD 2,3,7,8-TCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF Total HxCDD Total HpCDD Total TCDF Total PeCDF Total HxCDF Total HxCDF Total HxCDF Total HxCDF Total HyCDF	J (all detects)	A	Field duplicates (Difference) (fd)
R0903051	SA152-0.5B SA152009-0.5B	1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,7,8,9-HxCDF 2,3,4,6,7,8-HxCDF Total TCDD Total PeCDD	J (all detects) UJ (all non-detects)	А	Field duplicates (Difference) (fd)

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Laboratory Blank Data Qualification Summary - SDG R0903051

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
R0903051	SA100-0.5B	OCDD	14.3U ng/Kg	A	bl
R0903051	RSAM2-0.5B	1,2,3,4,6,7,8-HpCDD OCDD	1.94U ng/Kg 5.08U ng/Kg	A	bl

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Field Blank Data Qualification Summary - SDG R0903051

No Sample Data Qualified in this SDG

		Trollox Hortilgate Heriderson	
LDC #:_	21495C21	VALIDATION COMPLETENESS WORKSHEET	
SDG #:	R0903051	Stage 2B	
Laborato	ory: <u>Columbia</u>	Analytical Services	R 2nd B
			'/na ₩

2nd Reviewer

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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
1.	Technical holding times	*	Sampling dates: 6/1-4/09
II.	HRGC/HRMS Instrument performance check	4	,
III.	Initial calibration	4	
IV.	Routine calibration/I	B	
V.	Blanks	M	
VI.	Matrix spike/Matrix spike duplicates	N	ainst Deried
VII	Laboratory control samples	W	2C5 D
VIII.	Regional quality assurance and quality control	N .	\
IX.	Internal standards	m	
X.	Target compound identifications	N	
XI.	Compound quantitation and CRQLs	ÁN.	
XII.	System performance	N	
XIII.	Overall assessment of data	w	
XIV.	Field duplicates	W	D=9+11.10+1=, 22+23
XV.	Field blanks	2	D=9+11.10+12, 22+23 FB072109-50(\$0904016)

A = Acceptable

N = Not provided/applicable SW = See worksheet

R = Rinsate

ND = No compounds detected

FB = Field blank

D = Duplicate TB = Trip blank EB = Equipment blank

Validated Samples:

7 RSA	12-0.5B	117	SA76009-0.5B	21/2	SA88-0.5BDL	31	ZR0900197-01
RSA	12-0.5BDL	123	SA76009-0.5BDL	22/	SA152-0.5B	32	ZQ0900210-0/
/3 RSA	13-0.5B	13/5	RSAL3-0.5B	23/	SA152009-0.5B	33	P103460 >
A RSA	13-0.5BDL	14	RSAL3-0.5BDL	24/3	RSAJ2-0.5B	34	>103553
RSA	J5-0.5B	15/5	SA100-0.5B	25/4	RSAJ2-0.5BDL	35	P112515 x
RSA RSA	J5-0.5BDL	6 ₁₆ /5	RSAM3-0.5B	26/-	RSAJ3-0.5B	36	D202729 x
/> RSA	K5-0.5B	17/5	RSAM2-0.5B	27/2	SA202-0.5B	37	7103529 >
2 RSA	K5-0.5BDL	181	SA189-0.5B	28		386	P103504 +
SA76	6-0.5B	19	SA189-0.5BDL	29		397	P103583
10 SA76	6-0.5BDL	20/5	SA88-0.5B	30		408	P20275 8, 972086

VALIDATION FINDINGS WORKSHEET

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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	a. 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	1. 1,2,3,7,8-PeCDF	N. 1,2,3,7,8,9-HxCDF	S. Total PeCDD	X. Total HxCDF
) }	J. 2,3,4,7,8-PeCDF	O. 1,2,3,4,6,7,8-HpCDF	T. Total HxCDD	Y. Total HpCDF

Notes:

SDG#:28 LDC #2

VALIDATION FINDINGS WORKSHEET

2nd Reviewer._ Page: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Were all samples associated with a method blank?

Was a method blank analyzed for each matrix?

W N/A Was the blank contaminated? If yes, please see qualification below. Blank extraction date: 100 Blank analysis date:

٢

Conc. units: バシ/<<		,	Associa	ted Samples:	7//-/	Associated Samples: /- //・・20~ 22・ 24 - />・ 2 /	١٥٥ / ١٥٨	
Compound	Blank ID	:			Sam	Sample Identification		
2,80	10-16100 0XZ	5	2	17	NA		ofus VAX	
<i>,</i>	0,449	l	1	1946.1	17251			
4	3.26	14.3/U		1/41.6 US0:5	9.18th			
9	0.193	ı		1	<u> </u>			
\forall	0399			l	F			
И	0.449		,		1.90g.			
	/				<u> </u>			
					:			
	THE RESERVE THE PERSON NAMED IN COLUMN TWO IS NOT THE PERSON NAMED IN		-	The second secon				

23.26 61-81 Associated Samples: Blank extraction date: 6/16/19 Blank analysis date: 62719 Conc. units: ハクトラ

Compound	Blank ID			ats	Sample Identification	ion		
Z\$1.9		43	53					
, ±	0283	<i>/+</i>	igg.					
4	25.1.	/ +						
9	895.0	Н						
À	8950	+						
П	1.383	7						
×	0.28	+						
)							

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".

FIDE # STORES

SDG #:See Cover

VALIDATION FINDINGS WORKSHEET

Field Blanks

Reviewer: 2

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

V N N/A Were field blanks identified in this SDG?

ng/Kg Associated sample units:_ Blank units: pg/L

Sampling date: 7/21/09

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

X 6 <) W Associated Samples:

Sample Identification 0.04185 0.03165 0.01785 0.04185 0.3605 0.0202 0.2165 0.0905 0.109 0.098 1.025 EB072109-SO Blank ID 21.8 6.33 43.3 19.6 8.37 3.57 4.04 205 8.37 18.1 72.1 Compound

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

SDG # 120 colun LDC#249822

VALIDATION FINDINGS WORKSHEET Laboratory Control Samples (LCS)

2nd Reviewer: Reviewer: 9 Page:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Was a LCS required? Y N/A

Was a LCS analyzed every 20 samples for each matrix or whenever a sample extraction was performed?

YiN N/A Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?

#	# Date		Compound	LCS Lab ID/Reference Compound %R (Limits)	LCSD %R (Limits)	RPD (Limits)	Associated Samples	Qualifications
		120.79.009.02	1	(84-60 22	()	1-17.00-22.	No Gual
		(37.50)		()	187 (80-130)	625157	Cz.5c-42	
Γ			l	()		()	0-161006082	(105 m)
				()	()	()		
		•		()	()	()		
				()	(()		
		2x0900210-12168	0)		(25) (5	18-19,23.26	No and
			1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1		_		200900 210-01	(70 K W)
		950760/	Q5	()	()	()	,	
	20.00			()	()	()		
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)	()	()		
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)	()	()		
	Section (1994) (1994) and the section of the sectio	i i i i i i i i i i i i i i i i i i i)	()	()		
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	The second secon			J	() (()		
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LDC #: 249502 SDG #: 26, COMP

VALIDATION FINDINGS WORKSHEET Internal Standards

Page: ZofZ 2nd Reviewer: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)
Plegse see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A". Y W N/A

Are all internal standard recoveries were within the 40-135% criteria? Was the S/N ratio all internal standard peaks \geq 10?

*	Ş	Concession of the 1	Consideration of the state of t	76 %	9, Daconson (1 mlt: 40.1389.)	Ouslifications
			<i>y</i>	6	1 751 17	(/ [/ 4 (/)]
T)		
1			1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	13)		(一种好女)
1			# (No 2007) #	90		
)	
		# 4	4	20	((¼ · ¼)
					()	
		- A	(See-80) A	4	()	(525- 80m H)
					()	
		4	*	<u>}</u>	()	(士)
			7	38	((k- \(\)
					(/
		27	XX	38	(/)	(/ · //) /
					(
		()	4 (DB-S)	<u> </u>	(40-135)	1/W/P (4.V)
			,			
					()	
					(
					(
					(
					()	
		Internal Standards	Check Standard Used		Internal Standards	Check Standard Used
Ą	13C-2,3,7,8-TCDF	CDF		1. 43C-OCDD		
69	13C-2,3,7,8-TCDD	coo		K. 13C-1,2,3,4-TCDD	rcdd	
O	¹³ C-1,2,3,7,8-PeCDF	PecoF		L. 1 13C-1,2,3,7,8,9-HxCDD	3,9-HxCDD	
Ö	13C-1,2,3,7,8-PeCDD	Pecoo		W		
ui	¹³ C-1,2,3,4,7,8-HxCDF	8-HxCDF		ż		
'n.	¹³ C-1,2,3,6,7,8-HxCDD	8-HxCDD		Ö		
ე		7,8-HpCDF		ď.		7
Ï		7,8-HpCDD				

LDC #: 2495c= | SDG #: 261 COWEN

VALIDATION FINDINGS WORKSHEET Compound Quantitation and Reported CRQLs

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound? Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary).

Qualifications	4 TK (K)			J65/4 (6)	II .							
Associated Samples	119	:		1,11.9		4	12 60 2 X	7	A THE REAL PROPERTY AND A STREET	J		8 - 30
Finding	ZWPC USULTS	Gods > adab cano		A-8.VIHONDBS&DB-205)		S-F I-B. H. B-5 208-20	2-47. 4m 2B5 K DR+25	R. 2-4. I-R	A control of the cont	下本 1 大人 0-18 19	185 & OB-225	H. K. L. O-B H DN DR-C & DS-C
Sample ID			The state of the s	Per [1], 9		N	5, DD, 24			20042000000000000000000000000000000000	A secondarion and the second of the second o	
# Date	5 5	 The state of the s					2 A Company of the Co		·	** Management Common and Statement (CA)		

Comments: See sample calculation verification worksheet for recalculations

1DC # 349603 SDG # 326 COUNTY

VALIDATION FINDINGS WORKSHEET Compound Quantitation and Reported CRQLs

Page: 2 of 2 Reviewer: 4

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound? Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary). Y N N/A

*	Date	Sample ID	and rated unga	Associated Samples	Qualifications
		K	K.O.D. & H W.D. S. B. 25	Ħ	(a) d/49(F)
	٠				
				-	

Comments: See sample calculation verification worksheet for recalculations

SDG #: 2495CEND

VALIDATION FINDINGS WORKSHEET Overall Assessment of Data

Page: of E-

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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.

(Y) N/A Was the overall quality and usability of the data acceptable?

L					
*	Date	Sample ID	Finding	Associated Samples	Qualifications
	The space of	. 1, 11, 9	4-8 (4008-5208-25)	5 111 3	(0) X
			,		
	187	Ez-17, 27, 51, E	35 H on DB-5	Ec. 22, 71-21-51	75-92
		3	2-4. I-A. HOW/B-5 & DEOX	ex 3	
		4	All 1xcept 2-4. 1-B	4	
			ر ا		
		5, 20, 24	2-4.1-8.4m 18-5 & DB-2	12,05, 20,24	
		55, 5,	\$1 pount 10-4. I-R. Han DB-2x	18-2x 6,2/x	
		-	B. D. F. L. A. H. M. D. S. G. D. B.	3-275 7	
			i l		
		90	21. 2xcat B-1. 3-6. 1-18. 4 dr 28-2x	1 DB-225 8	Ą
Comments:	ents:				

C * SOT

VALIDATION FINDINGS WORKSHEET Overall Assessment of Data

Page:

2nd Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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.

Y N N/A

Was the overall quality and usability of the data acceptable?

*	Date	Sample ID	Finding	Associated Samples	Qualifications
		(3)	F* 5. K. L. 0-B.	13	× (0)
			Han 3B-5 82 225		
		*	A1 UXGD+ F-X. H-I.	4	
			K-2.0-18 / "		
		8	2.4.4.0-8. 4 MOBS & RADS	81 2528	
		61	41. Broot & H-I. K-L	19	
					\rightarrow
Comments:	nents:				

LDC#:<u>21495C21</u> SDG#:<u>See Cover</u>

VALIDATION FINDINGS WORKSHEET Field Duplicates

Page:_	[of_>
Reviewer:	, Q
2nd Reviewer:	<u> </u>

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

YN NA YN NA Were field duplicate pairs identified in this SDG? Were target analytes detected in the field duplicate pairs?

	Concentration	on (ng/Kg)	(≤50)	(ng/Kg)	(ng/Kg)	Qualifications	
Compound	9	11	RPD	Difference	Limits	(Parent Only)	
A	433	676	44				
В	2460	3870	45				
С	1860	3070	49				
D	3330	5260	45				
E	4020	6580	48				
F	14400	17200	18				
G	13200	20800	45				
н	81400	138000	52			1 dets/x(fd
ı	29900	51900	54			1 4 1	
J	14500	26700	59			V	
К	77200	120000	43				
L	47000	77900	49				
N	9010	16400	58			Jots/A	F
М	9770	16400	51			V /	
0	95300	132000	32			-	
Р	84900	121000	35				
Q	197000	271000	32				
R	25100	35700	35				
S	31200	48800	44				
Т	26900	43600	47				
บ	22400	26700	18				
V	983000	1630000	50				
w	234000	283000	19				
x	366000	536000	38				
Υ	296000	431000	37				
Н (DB-225)	12900	15100	16				

LDC#:<u>21495C21</u> SDG#:<u>See Cover</u>

VALIDATION FINDINGS WORKSHEET Field Duplicates

Page: 2 of 3
Reviewer: 9
2nd Reviewer:/

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

YN NA YN NA Were field duplicate pairs identified in this SDG? Were target analytes detected in the field duplicate pairs?

	Concentration	on (ng/Kg)	(≤50)	(ng/Kg)	(ng/Kg)	Qualifications	
Compound	10	12	RPD	Difference	Limits	(Parent Only)	
Α	259	376		117	(≤199)		
В	1040	1190		150	(<u><</u> 497)		
С	824	1040		216	(<u><</u> 497)		
D	1920	2460		540	(<497)	idets/A1	fd
E	2060	2640		580	(<u>≤</u> 497)	V	
F	8140	7640	6				
G	8370	10600	24				
Н	12400	103000	157		*	Idets/A	(40
	12600	21400	52				
J	5050	10400	69			V	
К	62800	88200	34				
L	37100	54500	38				
N	4280	7060	49				
M	8640	11700	30				
0	135000	167000	21				
P	62300	80600	26				
Q	305000	397000	26		·		
H (DB-225)	6470	6240	4				

	Concentration (ng/Kg)		(≤50)	(ng/Kg)	(ng/Kg)	Qualifications	
Compound	22	23	RPD	Difference	Limits	(Parent Only)	
А	1.11	1.90U		0.79	(<u><</u> 1.90)		
В	3.79	4.75U		0.96	(≤4.75)		
С	1.97	4.75U		2.78	(≤4.75)		
D	4.33	4.75U		0.42	(≤4.75)		
E	5.90	0.605	, w	5.295	(≤4.75)	Ndets/A	
F	13.7	1.50	1	12.2	(≤4.75)	I V	

Hd)

LDC#:<u>21495C21</u> SDG#:<u>See Cover</u>

VALIDATION FINDINGS WORKSHEET Field Duplicates

Page:	3 63
Reviewer:_	Y
2nd Reviewer:_	N.

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

YN NA YN NA Were field duplicate pairs identified in this SDG? Were target analytes detected in the field duplicate pairs?

	Concentration	n (ng/Kg)	(≤50)	(ng/Kg)	(ng/Kg)	Qualifications	
Compound	22	23	RPD	Difference	Limits	(Parent Only)	
G	15.9	9.17		6.73	(<u><</u> 9.50)		
Н	62.4	0.565		61.835	(<u>≤</u> 1.90)	Utots/A	(+
l	52.0	4.75U		47.25	(<u><</u> 4.75)	NW/A	1+
J ·	27.9	4.75U		23.15	(<u>≤</u> 4.75)	V	`
К	107	0.952		106.048	(<u><</u> 4.75)	Jacks A	17
L	63.0	0.634		62.366	(<u><</u> 4.75)	↓ √	
N ·	10.0	4.75U		5.25	(<u><</u> 4.75)	VINA	
М	15.9	4.75U		11.15	(<u><</u> 4.75)	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	
0	187	2.52	·	184.48	(<u>≤</u> 4.75)	Jdets/10	
Р	92.2	0.941		91.259	(<u>≤</u> 4.75)		
Q	518	6.45		511.55	(<u><</u> 9.50)	V	
R	26.3	1.90U		24.4	(≤1.90)	varo	
s	37.4	4.75U		32.65	(≤4.75)	\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	
Т	33.4	0.605		32.795	(≤4.75)	Ndet3/0	
U	21.4	1.50		19.9	(<u>≤</u> 4.75)		
V	471	0.565		470.435	(≤1.90)		
W	500	1.12		498.88	(<u><</u> 4.75)		
х	400	0.952		399.048	(<u><</u> 4,75)		
Y	405	3.08		401.92	(<u><</u> 4.75)		
H (DB-225)	11.2	3.35		7.85	(<1.90)		

LDC Report# 21495F21

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name:

Tronox LLC Facility, 2009 Phase B Investigation,

Henderson, Nevada

Collection Date:

June 5 through June 11, 2009

LDC Report Date:

November 2, 2009

Matrix:

Soil

Parameters:

Dioxins/Dibenzofurans

Validation Level:

Stage 2B

Laboratory:

Columbia Analytical Services, Inc.

Sample Delivery Group (SDG): R0903184

Sample Identification

SA127-0.5B SA127-0.5BDL RSAJ6-0.5B RSAJ6-0.5BDL RSAK6-0.5B RSAK8-0.5B RSAK8-0.5BDL RSAL7-0.5B

RSAL8-0.5BDL SA35-0.5B

SA55-0.5B SA56-0.5B

SA176-0.5B

RSAO3-0.5B

RSAO3-0.5BDL

SA182-0.5B

SA182-0.5BDL

SA201-0.5B SA201-0.5BDL SA134-0.5B SA134-0.5BDL SA127-0.5BMS SA127-0.5BMSD

SA166-0.5B

RSAK4-0.5B

RSAK4-0.5BDL

RSAK4009-0.5B

RSAK4009-0.5BDL

Introduction

This data review covers 29 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8290 for Polychlorinated Dioxins/Dibenzofurans.

This review follows the Standard Operating Procedures (SOP) 40, Data Review/Validation (BRC 2009), the Quality Assurance Project Plan Tronox LLC Facility, Henderson, Nevada (June 2009), NDEP guidance (May 2006), and USEPA Contract Laboratory Program National Functional Guidelines for Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) as there are no current guidelines for the method stated above.

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.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XIV.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- J+ Data are qualified as estimated, with a high bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J- Data are qualified as estimated, with a low bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J Data are qualified as estimated; it is not possible to assess the direction of the potential bias. False positives or false negatives are unlikely to have been reported.
- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- R Data are qualified as rejected. There is a significant potential for the reporting of false negatives or false positives.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- B The analytical result may be a false positive totally attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JB The analytical result may be biased high and partially attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JK The analytical result is an estimated maximum possible concentration (EMPC).
- X The analytical result is not used for reporting because a more accurate and precise result is reported in its place.
- J-TDS The analytical result is estimated based on failure of the Total Dissolved Solids (TDS) correctness check performed in accordance with the Standard Method 1030E.
- J-CAB The analytical result is estimated based on failure of the cation-anion balance correctness check performed in accordance with Standard Method 1030E.
- J-TDS & CAB The analytical result is unreliable based on the failure of the cation-anion balance and TDS correctness check performed in accordance with standard Method 1030E.
- 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.

Retention time windows were established for all homologues. The chromatographic resolution between 2,3,7,8-TCDD and peaks representing any other unlabeled TCDD isomer was less than or equal to 25%.

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 30.0% for labeled compounds.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

IV. Routine Calibration (Continuing)

Routine calibration was performed at the required frequencies.

All of the routine calibration percent differences (%D) between the initial calibration RRF and the routine calibration RRF were less than or equal to 20.0% for unlabeled compounds and less than or equal to 30.0% for labeled compounds.

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
EQ0900212-01	6/17/09	OCDD OCDF	1.17 ng/Kg 1.44 ng/Kg	All samples in SDG R0903184

Sample concentrations were compared to concentrations detected in the method blanks as required by the QAPP. No sample data was qualified with the following exceptions:

Sample	Compound	Reported Concentration	Modified Final Concentration
RSAK6-0.5B	OCDD	5.07 ng/Kg	5.07U ng/Kg
RSAL7-0.5B	OCDD	3.92 ng/Kg	3.92U ng/Kg
SA55-0.5B	OCDD OCDF	1.15 ng/Kg 3.17 ng/Kg	1.15U ng/Kg 3.17U ng/Kg
SA176-0.5B	OCDD	1.07 ng/Kg	1.07U ng/Kg
SA166-0.5B	OCDD	5.03 ng/Kg	5.03U ng/Kg
RSAK4-0.5BDL (100X)	OCDD	527 ng/Kg	527U ng/Kg
RSAK4009-0.5BDL (100X)	OCDD	558 ng/Kg	558U ng/Kg

Sample FB072109-SO (from SDG R0904016) was identified as a field blank. No polychlorinated dioxin/dibenzofuran contaminants were found in this blank with the following exceptions:

Field Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB072109-SO	7/21/09	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF Total HpCDD Total HxCDF Total HpCDF	8.37 pg/L 21.8 pg/L 6.33 pg/L 3.57 pg/L 4.04 pg/L 43.3 pg/L 19.6 pg/L 205 pg/L 8.37 pg/L 18.1 pg/L 72.1 pg/L	All samples in SDG R0903184

Sample concentrations were compared to concentrations detected in the field blanks as required by the QAPP. No sample data was qualified.

VI. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Although the MS/MSD percent recoveries (%R) and relative percent differences (RPD) were not within QC limits for some compounds, the LCS percent recovery (%R) was within QC limits and no data were qualified.

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 recoveries were within QC limits with the following exceptions:

Sample	Internal Standards	%R (Limits)	Compound	Flag	A or P
RSAJ6-0.5B	¹³ C-2,3,7,8-TCDF ¹³ C-1,2,3,7,8-PeCDF ¹³ C-1,2,3,7,8-PeCDD ¹³ C-1,2,3,4,6,7,8-HpCDF ¹³ C-2,3,7,8-TCDF (DB-225)	4 (40-135) 173 (40-135) 168 (40-135) 35 (40-135) 140 (40-135)	2,3,7,8-TCDF Total TCDF 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF Total PeCDF 1,2,3,7,8-PeCDD Total PeCDD 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF Total HpCDF 2,3,7,8-TCDF (DB-225)	J (all detects) UJ (all non-detects)	Р
RSAJ6-0.5BDL	¹³ C-2,3,7,8-TCDF (DB-225)	4 (40-135)	2,3,7,8-TCDF (DB-225)	J (all detects) UJ (all non-detects)	Р
RSAK8-0.5BDL	¹³ C-1,2,3,4,7,8-HxCDF	39 (40-135)	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	J (all detects) UJ (all non-detects)	Р
*SA55-0.5B	¹³ C-OCDD	38 (40-135)	OCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р
*SA56-0.5B	¹³ C-OCDD	39 (40-135)	OCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р

*Corrected associated compounds for SA55-0.5B and SA56-0.5B

X. Target Compound Identifications

Raw data were not reviewed for this SDG.

*XI. Project Quantitation Limit

All project quantitation limits were within validation criteria with the following exceptions:

Sample	Compound	Finding	Criteria	Flag	A or P
*SA127-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A
RSAJ6-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,4,7,8-HxCDD 1,2,3,4,6,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,7,8-PeCDF 2,3,4,7,8-HxCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 1,2,3,4,7,8,9-HpCDF Total TCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A
RSAK8-0.5B	1,2,3,4,6,7,8-HpCDD 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A

Sample	Compound	Finding	Criteria	Flag	A or P
RSAL8-0.5B RSAO3-0.5B SA56-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	А
SA182-0.5B SA201-0.5B SA134-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 1,2,3,4,7,8,9-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	Α
RSAK4-0.5B *RSAK4009-0.5B	1,2,3,4,6,7,8-HpCDD 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 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	Α

^{*}Changed compounds for RSAK4009-0.5B in table above

All compounds reported below the PQL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903184	All compounds reported below the PQL.	J (all detects)	Α

All compounds reported as EMPC were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903184	All compounds reported as estimated maximum possible concentration (EMPC).	JK (all detects)	А

Raw data were not reviewed for this SDG.

XII. System Performance

Raw data were not reviewed for this SDG.

*XIII. Overall Assessment of Data

The overall assessment of data was acceptable. In the case where more than one result was reported for an individual sample, the least technically acceptable results were rejected as follows:

Sample	Compound	Flag	A or P
*SA127-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225) 2,3,4,6,7,8-HxCDF	X X X X X X	А
*SA127-0.5BDL	All TCL compounds except 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-225) 2,3,4,6,7,8-HxCDF	X	Α

		NUMBER 1	
Sample	Compound	Flag	A or P
RSAJ6-0.5B SA182-0.5B SA201-0.5B SA134-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-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 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x x x x x x	A
RSAJ6-0.5BDL	2,3,7,8-TCDF (DB-225)	х	А
RSAK6-0.5B RSAL7-0.5B SA35-0.5B SA56-0.5B SA182-0.5BDL SA201-0.5BDL SA166-0.5B SA134-0.5BDL	2,3,7,8-TCDF (DB-5)	X	A
RSAK8-0.5B	1,2,3,4,6,7,8-HpCDD 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x x	A
RSAK8-0.5BDL	All TCL compounds except 1,2,3,4,6,7,8-HpCDD 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	x	A

Sample	Compound	Flag	A or P
RSAL8-0.5B RSAO3-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x	A
RSAL8-0.5BDL RSAO3-0.5BDL	All TCL compounds except 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	×	A
*RSAK4-0.5B *RSAK4009-0.5B	1,2,3,4,6,7,8-HpCDD 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 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X	A
*RSAK4009-0.5BDL *RSAK4-0.5BDL	All TCL compounds except 1,2,3,4,6,7,8-HpCDD 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 0CDF 2,3,7,8-TCDF	X	A

^{*}Changed compounds for samples noted above.

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

Samples RSAK4-0.5B and RSAK4009-0.5B and samples RSAK4-0.5BDL and RSAK4009-0.5BDL were identified as field duplicates. No polychlorinated dioxins/dibenzofurans were detected in any of the samples with the following exceptions:

	Concentrat	DDD	D:#			
Compound	RSAK4-0.5B	RSAK4009-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P
2,3,7,8-TCDD	32.1	34.5	7 (≤50)	-	~	-
1,2,3,7,8-PeCDD	118	127	7 (≤50)	-	-	-
1,2,3,4,7,8-HxCDD	100	109	9 (≤50)	-	-	-
1,2,3,6,7,8-HxCDD	178	186	4 (≤50)	-	-	-
1,2,3,7,8,9-HxCDD	221	233	5 (≤50)	-	-	-
1,2,3,4,6,7,8-HpCDD	729	773	6 (≤50)	-	-	-
OCDD	767	811	6 (≤50)	-	-	-
2,3,7,8-TCDF	1850	2010	8 (≤50)	•	-	-
1,2,3,7,8-PeCDF	1700	1840	8 (≤50)	•	-	-
2,3,4,7,8-PeCDF	837	903	8 (≤50)	•	-	-
1,2,3,4,7,8-HxCDF	4280	4610	7 (≤50)	-	-	-
1,2,3,6,7,8-HxCDF	2730	2930	7 (≤50)	•	-	-
1,2,3,7,8,9-HxCDF	469	447	5 (≤50)	-	-	-
2,3,4,6,7,8-HxCDF	618	669	8 (≤50)	-	-	-
1,2,3,4,6,7,8-HpCDF	9680	10100	4 (≤50)	-	-	-
1,2,3,4,7,8,9-HpCDF	4760	5190	9 (≤50)	-	-	-
OCDF	20700	22000	6 (≤50)	•	-	-
Total TCDD	1280	1350	5 (≤50)	-	-	-
Total PeCDD	1390	1490	7 (≤50)	<u>-</u>	-	-
Total HxCDD	1500	1580	5 (≤50)	-	-	-

	Concentrati	on (ng/Kg)	DDD	D.##		
Compound	RSAK4-0.5B	RSAK4009-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P
Total HpCDD	1150	1230	7 (≤50)	-	-	-
Total TCDF	15200	16300	7 (≤50)	-	-	-
Total PeCDF	15200	17200	12 (≤50)	-	-	-
Total HxCDF	21100	21300	1 (≤50)	-	-	-
Total HpCDF	21400	22900	7 (≤50)	-	-	-
2,3,7,8-TCDF (DB-225)	720	795	10 (≤50)	-	-	-

	Concentrati	ion (ng/Kg)	- Dan	Diff.		
Compound	RSAK4-0.5BDL	RSAK4009-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P
1,2,3,7,8-PeCDD	93.5	242U	-	148.5 (≤242)	-	-
1,2,3,4,7,8-HxCDD	80.9	66.1	-	14.8 (≤251)	-	-
1,2,3,6,7,8-HxCDD	172	159	-	13 (≤251)	-	-
1,2,3,7,8,9-HxCDD	179	183	•	4 (≤251)	-	-
1,2,3,4,6,7,8-HpCDD	514	573	•	59 (≤501)	-	-
OCDD	527	558	-	31 (≤276)	-	-
2,3,7,8-TCDF	1430	1460	2 (≤50)	-	-	-
1,2,3,7,8-PeCDF	1300	1150	-	150 (≤251)	-	-
2,3,4,7,8-PeCDF	581	620	•	39 (≤251)	-	-
1,2,3,4,7,8-HxCDF	4410	4640	5 (≤50)	-	-	-
1,2,3,6,7,8-HxCDF	3060	3170	4 (≤50)	-	-	-
1,2,3,7,8,9-HxCDF	296	321	-	25 (≤251)	-	-

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	Concentrati	on (ng/Kg)	RPD	Difference		
Compound	RSAK4-0.5BDL	RSAK4009-0.5B	(Limits)	(Limits)	Flag	A or P
2,3,4,6,7,8-HxCDF	734	684	-	50 (≤251)	-	•
1,2,3,4,6,7,8-HpCDF	10000	11000	10 (≤50)	-	-	-
1,2,3,4,7,8,9-HpCDF	3840	4110	7 (≤50)	-	-	-
OCDF	21300	23400	9 (≤50)	-	-	-

*Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Data Qualification Summary - SDG R0903184

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903184	RSAJ6-0.5B	2,3,7,8-TCDF Total TCDF 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF Total PeCDF 1,2,3,7,8-PeCDD Total PeCDD 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF Total HpCDF 2,3,7,8-TCDF (DB-225)	J (all detects) UJ (all non-detects)	Р	Internal standards (area) (i)
R0903184	RSAJ6-0.5BDL	2,3,7,8-TCDF (DB-225)	J (all detects) UJ (all non-detects)	Р	Internal standards (area) (i)
R0903184	RSAK8-0.5BDL	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	J (all detects) UJ (all non-detects)	Р	Internal standards (area) (i)
*R0903184	SA55-0.5B SA56-0.5B	OCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р	Internal standards (area) (i)
*R0903184	SA127-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903184	RSAJ6-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD CCDD 1,2,3,4,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF Total TCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903184	RSAK8-0.5B	1,2,3,4,6,7,8-HpCDD 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	А	Project Quantitation Limit (e)
R0903184	RSAL8-0.5B RSAO3-0.5B SA56-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903184	SA182-0.5B SA201-0.5B SA134-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 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 1,2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
*R0903184	RSAK4-0.5B RSAK4009-0.5B	1,2,3,4,6,7,8-HpCDD 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 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903184	SA127-0.5B SA127-0.5BDL RSAJ6-0.5BDL RSAJ6-0.5BDL RSAK6-0.5B RSAK8-0.5BDL RSAL8-0.5BDL RSAL8-0.5BDL SA35-0.5B SA55-0.5B SA55-0.5B SA56-0.5B SA176-0.5B RSAO3-0.5BDL SA182-0.5BDL SA182-0.5BDL SA182-0.5BDL SA182-0.5BDL SA201-0.5BDL SA201-0.5BDL SA201-0.5BDL SA201-0.5BDL SA166-0.5B RSAC4-0.5BDL SA166-0.5B RSAK4-0.5BDL RSAK4-0.5BDL RSAK4009-0.5BDL SA134-0.5BDL SA134-0.5BDL	All compounds reported below the PQL.	J (all detects)	A	Project Quantitation Limit (sp)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903184	SA127-0.5B SA127-0.5BDL RSAJ6-0.5B RSAJ6-0.5BDL RSAK6-0.5B RSAK8-0.5B RSAK8-0.5BDL RSAL8-0.5B RSAL8-0.5BDL RSAL8-0.5B RSAL8-0.5B RSAL8-0.5B RSAL8-0.5B SA55-0.5B SA55-0.5B SA56-0.5B RSAC9-0.5BDL SA182-0.5BDL SA182-0.5BDL SA182-0.5BDL SA201-0.5BDL SA201-0.5BDL SA201-0.5BDL SA182-0.5BDL SA134-0.5BDL SA134-0.5BDL SA134-0.5BDL	All compounds reported as EMPC	JK (all detects)	A	Project Quantitation Limit (k)
*R0903184	SA127-0.5B	1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225) 2,3,4,6,7,8-HxCDF	X X X X X X X	A	Overall assessment of data (o)
*R0903184	SA127-0.5BDL	All TCL compounds except 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-225) 2,3,4,6,7,8-HxCDF	X	A	Overall assessment of data (o)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903184	RSAJ6-0.5B	2,3,7,8-TCDD	Х	A	Overall assessment of
	SA182-0.5B	1,2,3,7,8-PeCDD	x		data (o)
	SA201-0.5B	1,2,3,4,7,8-HxCDD	x		""" (")
	SA134-0.5B	1,2,3,6,7,8-HxCDD	x		
	071104 0.05	1,2,3,7,8,9-HxCDD	x		
		1,2,3,4,6,7,8-HpCDD	x		
		OCDD	x		
		1,2,3,7,8-PeCDF	x		
		2,3,4,7,8-PeCDF	X		
		1,2,3,4,7,8-HxCDF	X		
		1,2,3,6,7,8-HxCDF	X		
	ŀ	2,3,4,6,7,8-HxCDF	Х		
		1,2,3,7,8,9-HxCDF	X		
		1,2,3,4,6,7,8-HpCDF	X		
		1,2,3,4,7,8,9-HpCDF	Χ		
		OCDF	Χ		
		2,3,7,8-TCDF (DB-5 & DB-225)	Χ		
				ļ	
R0903184	RSAJ6-0,5BDL	2,3,7,8-TCDF (DB-225)	х	A	Overall assessment of data (o)
R0903184	RSAK6-0.5B RSAL7-0.5B SA35-0.5B SA56-0.5B SA182-0.5BDL SA201-0.5BDL SA166-0.5B SA134-0.5BDL	2,3,7,8-TCDF (DB-5)	х	A	Overall assessment of data (o)
R0903184	RSAK8-0.5B	8-0.5B 1,2,3,4,6,7,8-HpCDD 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)		A	Overall assessment of data (o)
R0903184	RSAK8-0.5BDL	All TCL compounds except 1,2,3,4,6,7,8-HpCDD 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	Х	A	Overall assessment of data (o)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903184	RSAL8-0.5B RSAO3-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X	A	Overall assessment of data (o)
R0903184	RSAL8-0.5BDL RSAO3-0.5BDL	All TCL compounds except 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X	A	Overall assessment of data (o)
*R0903184	RSAK4-0.5B RSAK4009-0.5B	1,2,3,4,6,7,8-HpCDD 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 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X	A	Overall assessment of data (o)
*R0903184	RSAK4009-0.5BDL RSAK4-0.5BDL	All TCL compounds except 1,2,3,4,6,7,8-HpCDD 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 1,2,3,4,6,7,8-HpCDF 0CDF 2,3,7,8-TCDF	X	A	Overall assessment of data (o)

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Laboratory Blank Data Qualification Summary - SDG R0903184

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
R0903184	RSAK6-0.5B	OCDD	5.07U ng/Kg	А	bl
R0903184	RSAL7-0.5B	OCDD	3.92U ng/Kg	А	bl

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
R0903184	SA55-0.5B	OCDD OCDF	1.15U ng/Kg 3.17U ng/Kg	A	bl
R0903184	SA176-0.5B	OCDD	1.07U ng/Kg	А	bl
R0903184	SA166-0.5B	OCDD	5.03U ng/Kg	А	bl
R0903184	RSAK4-0.5BDL (100X)	OCDD	527U ng/Kg	А	bl
R0903184	RSAK4009-0.5BDL (100X)	OCDD	558U ng/Kg	Α	bl

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Field Blank Data Qualification Summary - SDG R0903184

No Sample Data Qualified in this SDG

Trongy Northgate Henderson

		Tronox Horalgate Trenderson	
LDC #:_	21495F21	VALIDATION COMPLETENESS WORKSHEET	Date: 9/17/0=
SDG #:_	R0903184	Stage 2B	Page: / of // Reviewer:
Laborato	ory: Columbia Analytica	al Services	Reviewer:
			2nd Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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
1.	Technical holding times	A	Sampling dates: 6/5-11/09
11.	HRGC/HRMS Instrument performance check	4	, , ,
ł11.	Initial calibration	1	
IV.	Routine calibration/lev	A	
V.	Blanks	aw	
VI.	Matrix spike/Matrix spike duplicates	W	2859: 70 Fand FPD out - No bual (2051)
VII.	Laboratory control samples	A	10-
VIII.	Regional quality assurance and quality control	N	
IX.	Internal standards	$\langle m \rangle$	·
X.	Target compound identifications	N	
XI.	Compound quantitation and CRQLs	ŹN_	
XII.	System performance	N	
XIII.	Overall assessment of data	w	
XIV.	Field duplicates	aw	D=22+24,23+25
XV.	Field blanks	5W	D=22+24,23+25 FB072109-50(R0904016)

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 5	·		
1	SA127-0.5B	116 SA35-0.5B	21 SA166-0.5B	31 ZR0900-12-01
2 -	SA127-0.5BDL	12 SA55-0.5B	22/5 RSAK4-0.5B	32
3	RSAJ6-0.5B	13/5 SA56-0.5B	23/5 RSAK4-0.5BDL	33
4	RSAJ6-0.5BDL	14 SA176-0.5B	24 RSAK4009-0.5B	34 11132041
5	RSAK6-0.5B	15/3 RSA03-0.5B	25 RSAK4009-0.5BDL	35 U132087
6	RSAK8-0.5B	16 RSA03-0.5BDL	26 SA134-0.5B	363 U132073
7	RSAK8-0.5BDL	17 SA182-0.5B	27/4 SA134-0.5BDL	37 >202989
85/	RSAL7-0.5B	18 SA182-0.5BDL	28 SA127-0.5BMS	385 UZ2057/
96/	RSAL8-0.5B	19/5 SA201-0.5B	29 SA127-0.5BMSD	39
101	RSAL8-0.5BDL	20/4 SA201-0.5BDL	30	40

Notes:	1103559	1/2 >20	2924	3.7103612	1 421	9519	5 > 20 2 9	103/
6	P1035	73' TPI	035831	8 P2026	369	, , , , , , , , , , , , , , , , , , ,		

VALIDATION FINDINGS WORKSHEET

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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	1, 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:

1DC #1 SDG #

VALIDATION FINDINGS WORKSHEET Blanks

2nd Reviewer: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Were all samples associated with a method blank?

Was a method blank analyzed for each matrix? N N X

W N N/A Was the blank contaminated? If yes, please see qualification below. Blank extraction date: 6/23/2

251100 258 23(100x) 127 Sample Identification 5.03 7 2/201 Associated Samples: 1/2/1 V m 9 **%** 4 201 Z & 02/2-0 Blank ID Sonc. units: Notes Compound

₩

Associated Samples: Blank analysis date:___ Blank extraction date:_ Conc. units:

Compound	Blank ID	Sample Identification	

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".

SDG #:See Cover LDC#:0

VALIDATION FINDINGS WORKSHEET

Field Blanks

2nd Reviewer: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

X N/A Were field blanks identified in this SDG?

Blank units: <u>pg/L</u> Associated sample units:__Sampling date: 7/21/09

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

) H	Blank ID	Blank ID	Sample Identification	
EB072109.SO 5X	2X			
0	0.04185			
21.8 0.109	0.109			
6.33 0.03165	0.03165			
	0.01785	1		
4.04 0.0202	0.0202			
43.3 0.2165	0.2165	_		
19.6 0.098	0.098			
205 1.025	1.025			
8.37 0.04185	0.04185			
	0.0905			
	0.3605	L		
		1		
a de la companya de l		<u> </u>		

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

SDG #: 48 COS LDC # 3445

VALIDATION FINDINGS WORKSHEET Internal Standards

Page: __of _ Reviewer: _____ 2nd Reviewer: _____

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".	A Are all internal standard recoveries were within the 40-135% criteria?	A Was the S/N ratio all internal standard peaks > 10?
Please se	X (N/N/A	A/N N/A

**	Date	Lab ID/Reference	Internal Standard	6	% Recovery (Limit: 40-135%)	135%)		Cocle: n Qualifications
		4)	4	`	4)	20-135	7	Q3 4 (H.V)
-			> \$	17	173 (-		(M · r · #)
			Ð	10) 2 &		_	(B.S)
				38	٧(>	_	A
H			(5 <c-8\$) a<="" th=""><th>140</th><th>) 0</th><th>//</th><th>^</th><th>* (F. 8× 8 B-5×</th></c-8\$)>	140) 0	//	^	* (F. 8× 8 B-5×
-)		^	
-		4	(255-8K) A O	4	Ĵ	40-135	7	Ay 6 CH ON 213-2-25
)	-	7	
Н					—		+	- 1
)	8	30	J	^	K	and (K-N)
-					J		+	
		7	H	38	-)	40-135	7	/W P (F. &)
)	+	1	
		(3	t	000	`	4	 	<u> </u>
)			
		28 (MS)	∌	30	,	40-135		Do and
			H	380			$\frac{1}{1}$	
		(asn) be	F)	300	~	7	7	<i>\</i>
<u> </u>			₩	34	_		7	
				W W)	/	7	Y
		Internal Standards	Check Standard Used		Intern	Internal Standards		Check Standard Used
∢	¹³ C-2,3,7,8-TCDF	ЭF			¹³ C-OCDD			
æ	¹³ C-2,3,7,8-TCDD	Q.		×	13C-1,2,3,4-TCDD			
ن	¹³ C-1,2,3,7,8-PeCDF	SCDF		ند	¹³ C-1,2,3,7,8,9-HxCDD			
o.	13C-1,2,3,7,8-PeCDD	ecdd		Σ				
uj	¹³ C-1,2,3,4,7,8-HxCDF	HXCDF		ż				
'n.	¹³ C-1,2,3,6,7,8-HxCDD	HXCDD		ö				
Ö	¹³ C-1,2,3,4,6,7,8-HpCDF	8-нрсDF		a:				
ī	13C-1,2,3,4,6,7,8-HpCDD	8-HpCDD						

1DC #:3/495 2 | SDG #: 3ec 60 MOV

VALIDATION FINDINGS WORKSHEET Compound Quantitation and Reported CRQLs

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Y N/N/A

Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound? Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary).

*	Date	Sample ID	As > calls lange Finding	Associated Samples	Qualifications
			H-6.0-8		1 2 4 ()
			CH on back (BS SQDR >> X)		}
		W	A-Q.V(Howbath	(1)	
	•		3-5-5		
		9	#.#-4.0-8	9	
			(4 or 08-5 & DB-2>5)		
			i		
		9,15	H. = . K. L. 0-8	9, 15-	
			(# & DB-5 B DB-2×5)		7
		13 /	# 7. K. C. O - &	8	Joeth Fro
			(How DB-5 & DB-225)		ł
		17,19,26	4-8 CHM 28-5 & UB-225)	17,19,26	14th/410)

Comments: See sample calculation verification worksheet for recalculations

LDC # 3/495 F-3 SDG #: 46 (16)

Compound Quantitation and Reported CRQLs VALIDATION FINDINGS WORKSHEET

Page:

2nd Reviewer: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Y M N/A

Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound? Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary).

Qualifications	2 det (2)			(F)					
- Associated Samples	+x / 4x	0 70 (*		114					
Finding	P. K. HER (HANDB-S			ZM7c Mswlts					
Sample ID	75, 24	***		-≱ [
# Date									

Comments: See sample calculation verification worksheet for recalculations

1DC #: 2149 EF 2

VALIDATION FINDINGS WORKSHEET Overall Assessment of Data

Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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.

Was the overall quality and usability of the data acceptable?

*	Date	Sample ID	Finding	Associated Samples	Qualifications
			H-41 8 (# on		(o) ×
			buth DB S & B-225)		
		4	安女 Uxcer+ エーや	d	
			8-0-8, Hon B-25	1	
		3, 17, 19,26	X-A. SCHONDER BIR	(# 0x 0B 5 2 08 235) 2/ 3/ 17.19	
		4	35= H ON 2B-225 (15 / 1944	7 (22	
	27, 21,	5,8,11,13,18,20	H ON BB-5	1-0-181 815/K	
	-				
		9	F. H-L. 0-B	9	
			(2 cz - 8 d. 2 - 3 d m /)		
			A11 except F. H-L. O-B	>	
Comments:	ents:				

DC #: = \$95 F2

VALIDATION FINDINGS WORKSHEET Overall Assessment of Data

Page: 2 of 2 Reviewer:

Reviewer: _____

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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	Sample ID	Finding	Associated Samples	Qualifications
		9, 15	H.I.K. L. O- X	51'6	(0) X
)	
		91,01	All except about	91'01	
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Somr	Comments:				

3

LDC#:<u>21495F21</u> SDG#:<u>See Cover</u>

VALIDATION FINDINGS WORKSHEET Field Duplicates

Page: lof 2
Reviewer: 0
2nd Reviewer: 4

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

YN NA YN NA Were field duplicate pairs identified in this SDG?
Were target analytes detected in the field duplicate pairs?

	Concentration	on (ng/Kg)	(≤50)	(ng/Kg)	(ng/Kg)	Qualifications
Compound	22	24	RPD	Difference	Limits	(Parent Only)
А	32.1	34.5	7			
В	118	127	7			
С	100	109	9			
D	178	186	4			
Ε .	221	233	5			
F	729	773	6			
G	767	811	6			
Н	1850	2010	8			
ł	1700	1840	8			
J	837	903	8			
К	4280	4610	7			
L	2730	2930	7			
N	469	447	5			
M	618	669	8			
0	9680	10100	4			
ρ	4760	5190	9			
Q .	20700	22000	6			
R	1280	1350	5			
s	1390	1490	7			
Т	1500	1580	5			
υ	1150	1230	7			
V	15200	16300	7			
w	15200	17200	12	-		
х	21100	21300	1			
Υ	21400	22900	7			
H (DB-225)	720	795	10			

LDC#:<u>21495I21</u> SDG#: <u>See Cover</u>

VALIDATION FINDINGS WORKSHEET Field Duplicates

Page: 2 of 2 Reviewer: 1 2nd Reviewer: 1

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

YN NA YN NA Were field duplicate pairs identified in this SDG? Were target analytes detected in the field duplicate pairs?

	Concentration	on (ng/Kg)	(≤50)	(ng/Kg)	(ng/Kg)	Qualifications
Compound	23	25	RPD	Difference	Limits	(Parent Only)
В	93.5	242U		148.5	(<u><</u> 242)	
С	80.9	66.1		14.8	(<u><</u> 251)	
D	172	159		13	(<u><</u> 251)	
E	179	183		4	(<u><</u> 251)	
F	514	573		59	(<u><</u> 501)	
G	527	558		31	(≤276)	
Н	1430	1460	2			
ı	1300	1150		150	(<u><</u> 251)	
J	581	620		39	(≤251)	
к	4410	4640	5			
L	3060	3170	4			
N ·	296	321		25	(<u><</u> 251)	
М	734	684		50	(<u><</u> 251)	
0	10000	11000	10			
Р	3840	4110	7			
Q	21300	23400	9			

LDC Report# 21495G21

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name:

Tronox LLC Facility, 2009 Phase B Investigation,

Henderson, Nevada

Collection Date:

June 19 through June 24, 2009

LDC Report Date:

November 2, 2009

Matrix:

Soil

Parameters:

Dioxins/Dibenzofurans

Validation Level:

Stage 2B

Laboratory:

Columbia Analytical Services, Inc.

Sample Delivery Group (SDG): R0903443

Sample Identification

SA197-0.5B SA198-0.5B

SA198-0.5BDL

SA64-0.5B

SA104-0.5B

SA104-0.5BDL

SA129-0.5B

SA129-0.5BDL

SA70-0.5B

SA70-0.5BDL

SA60-0.5B

SA60-0.5BDL

SA150-0.5B

SA150-0.5BDL

RSAN5-0.5B

SA53-0.5B

SA43009-0.5B

SA40-0.5B

SA200-0.5B

SA200-0.5BDL

RSAO6-0.5B

SA51-0.5B

SA51-0.5BDL

SA43-0.5B

SA150-0.5BMS

SA150-0.5BMSD

Introduction

This data review covers 26 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8290 for Polychlorinated Dioxins/Dibenzofurans.

This review follows the Standard Operating Procedures (SOP) 40, Data Review/Validation (BRC 2009), the Quality Assurance Project Plan Tronox LLC Facility, Henderson, Nevada (June 2009), NDEP guidance (May 2006), and USEPA Contract Laboratory Program National Functional Guidelines for Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) as there are no current guidelines for the method stated above.

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.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XIV.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- J+ Data are qualified as estimated, with a high bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J- Data are qualified as estimated, with a low bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J Data are qualified as estimated; it is not possible to assess the direction of the potential bias. False positives or false negatives are unlikely to have been reported.
- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- R Data are qualified as rejected. There is a significant potential for the reporting of false negatives or false positives.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- B The analytical result may be a false positive totally attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JB The analytical result may be biased high and partially attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JK The analytical result is an estimated maximum possible concentration (EMPC).
- X The analytical result is not used for reporting because a more accurate and precise result is reported in its place.
- J-TDS The analytical result is estimated based on failure of the Total Dissolved Solids (TDS) correctness check performed in accordance with the Standard Method 1030E.
- J-CAB The analytical result is estimated based on failure of the cation-anion balance correctness check performed in accordance with Standard Method 1030E.
- J-TDS & CAB The analytical result is unreliable based on the failure of the cation-anion balance and TDS correctness check performed in accordance with standard Method 1030E.
- 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.

Retention time windows were established for all homologues. The chromatographic resolution between 2,3,7,8-TCDD and peaks representing any other unlabeled TCDD isomer was less than or equal to 25%.

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 30.0% for labeled compounds.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

IV. Routine Calibration (Continuing)

Routine calibration was performed at the required frequencies.

All of the routine calibration percent differences (%D) between the initial calibration RRF and the routine calibration RRF were less than or equal to 20.0% for unlabeled compounds and less than or equal to 30.0% for labeled compounds.

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
EQ0900235-01	6/29/09	1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,6,7,8-HpCDF OCDF Total HpCDF	0.199 ng/Kg 3.11 ng/Kg 0.218 ng/Kg 0.955 ng/Kg 0.218 ng/Kg	All samples in SDG R0903443

Sample concentrations were compared to concentrations detected in the method blanks as required by the QAPP. No sample data was qualified with the following exceptions:

Sample	Compound	Reported Concentration	Modified Final Concentration
SA53-0.5B	1,2,3,4,6,7,8-HpCDD OCDD	0.502 ng/Kg 1.70 ng/Kg	0.502U ng/Kg 1.70U ng/Kg
RSAN5-0.5B	OCDD	7.58 ng/Kg	7.58U ng/Kg
RSAO6-0.5B	OCDD	12.6 ng/Kg	12.6U ng/Kg

Sample FB072109-SO (from SDG R0904016) was identified as a field blank. No polychlorinated dioxin/dibenzofuran contaminants were found in this blank with the following exceptions:

Field Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB072109-SO	7/21/09	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF Total HpCDD Total HxCDF Total HpCDF	8.37 pg/L 21.8 pg/L 6.33 pg/L 3.57 pg/L 4.04 pg/L 43.3 pg/L 19.6 pg/L 205 pg/L 8.37 pg/L 18.1 pg/L 72.1 pg/L	All samples in SDG R0903443

Sample concentrations were compared to concentrations detected in the field blanks as required by the QAPP. No sample data was qualified.

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 the QC limits with the following exceptions:

Spike ID (Associated Samples)	Compound	MS (%R) (Limits)	MSD (%R) (Limits)	RPD (Limits)	Flag	A or P
SA150-0.5BMS/MSD (SA150-0.5B SA150-0.5BDL)	OCDF	-	-	28 (≤20)	J (all detects)	А

VII. Laboratory Control Samples (LCS)

Laboratory control samples were reviewed for each matrix as applicable. The percent recoveries (%R) were within the QC limits with the following exceptions:

LCS ID	Compound	%R (Limits)	Associated Samples	Flag	A or P
EQ0900235-02LCS	OCDF	148 (70-130)	All samples in SDG R0903443	J+ (all detects)	Р

VIII. Regional Quality Assurance and Quality Control

Not applicable.

IX. Internal Standards

All internal standard recoveries were within QC limits with the following exceptions:

Sample	Internal Standards	%R (Limits)	Compound	Flag	A or P
SA129-0.5B	¹³ C-1,2,3,6,7,8-HxCDD ¹³ C-OCDD	32 (40-135) 17 (40-135)	1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,7,8,9-HxCDD Total HxCDD OCDD OCDF	J (all detects) UJ (all non-detects)	Р
SA129-0.5BDL	¹³ C-1,2,3,6,7,8-HxCDD ¹³ C-OCDD	37 (40-135) 17 (40-135)	1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,7,8,9-HxCDD OCDD OCDF	J (all detects) UJ (all non-detects)	Р
SA60-0.5BDL	¹³ C-OCDD	38 (40-135)	OCDD	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р
SA150-0,5BDL	¹³ C-OCDD	39 (40-135)	OCDD	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р

X. Target Compound Identifications

Raw data were not reviewed for this SDG.

XI. Project Quantitation Limit

All project quantitation limits were within validation criteria with the following exceptions:

Sample	Compound	Finding	Criteria	Flag	A or P
SA197-0.5B	1,2,3,4,6,7,8-HpCDF OCDF	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects) J (all detects)	Р
SA198-0.5B SA70-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	А
SA64-0.5B	1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	Р
SA104-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A
SA129-0.5B	OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A

Sample	Compound	Finding	Criteria	Flag	A or P
SA60-0.5B SA150-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A
SA43009-0.5B SA43-0.5B	1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	Р
SA40-0.5B	1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	Р
SA200-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	А
SA51-0,5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A

All compounds reported below the PQL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903443	All compounds reported below the PQL.	J (all detects)	А

All compounds reported as EMPC were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903443	All compounds reported as estimated maximum possible concentration (EMPC).	JK (all detects)	А

Raw data were not reviewed for this SDG.

XII. System Performance

Raw data were not reviewed for this SDG.

XIII. Overall Assessment of Data

The overall assessment of data was acceptable. In the case where more than one result was reported for an individual sample, the least technically acceptable results were rejected as follows:

Sample	Compound	Flag	A or P
SA197-0.5B SA64-0.5B RSAN5-0.5B SA53-0.5B SA43009-0.5B SA40-0.5B RSAO6-0.5B SA43-0.5B	2,3,7,8-TCDF (from DB-5)	X	A
SA198-0.5B SA70-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X	A
SA198-0.5BDL SA70-0.5BDL	All TCL compounds except 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF	X	A

Sample	Compound	Flag	A or P
SA104-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X	A
*SA104-0.5BDL	All TCL compounds except 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 2,3,7,8-TCDF (DB-225)	X	A
SA129-0.5B	OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X	A
SA129-0.5BDL	All TCL compounds except OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF	х	A
SA60-0.5B SA150-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X X	A

Sample	Compound	Flag	A or P
*SA60-0.5BDL *SA150-0.5BDL	All TCL compounds except 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 0CDF 2,3,7,8-TCDF (DB-225)	X	A
SA200-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X	A
SA200-0.5BDL	All TCL compounds except 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF	х	A
SA51-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X	Α
SA51-0.5BDL	All TCL compounds except 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 2,3,7,8-TCDF	x	А

^{*}Changed associated compounds for samples noted above

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

Samples SA43009-0.5B and SA43-0.5B were identified as field duplicates. No polychlorinated dioxins/dibenzofurans were detected in any of the samples with the following exceptions:

	Concentrati	on (ng/Kg)		Diff		
Compound	SA43009-0.5B	SA43-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P
2,3,7,8-TCDD	8.27	8.38	1 (≤50)	-	-	-
1,2,3,7,8-PeCDD	27.3	26.9	1 (≤50)	-	-	-
1,2,3,4,7,8-HxCDD	18.1	18.6	3 (≤50)	-	-	-
1,2,3,6,7,8-HxCDD	32.7	32.2	2 (≤50)	-	-	_
1,2,3,7,8,9-HxCDD	36,9	37.9	3 (≤50)	-	-	-
1,2,3,4,6,7,8-HpCDD	159	176	10 (≤50)	-	-	-
OCDD	419	572	31 (≤50)	-	-	-
2,3,7,8-TCDF	488	475	3 (≤50)	-	-	-
1,2,3,7,8-PeCDF	374	372	1 (≤50)	-	-	-
2,3,4,7,8-PeCDF	185	181	2 (≤50)	-	-	-
1,2,3,4,7,8-HxCDF	786	789	0 (≤50)	-	-	_
1,2,3,6,7,8-HxCDF	470	478	2 (≤50)	-	-	-
1,2,3,7,8,9-HxCDF	83.9	75.8	10 (≤50)	-	-	-
2,3,4,6,7,8-HxCDF	278	278	0 (≤50)	-	-	•
1,2,3,4,6,7,8-HpCDF	1660	1660	0 (≤50)	-	-	•
1,2,3,4,7,8,9-HpCDF	811	767	6 (≤50)	-	-	-

	Concentrati	on (ng/Kg)	RPD	Difference		
Compound	SA43009-0.5B	SA43-0.5B	(Limits)	(Limits)	Flag	A or P
OCDF	5830	5730	2 (≤50)	-	-	-
Total TCDD	350	326	7 (≤50)	-	-	-
Total PeCDD	310	302	3 (≤50)	-	-	-
Total HxCDD	268	271	1 (≤50)	-	-	-
Total HpCDD	263	296	12 (≤50)	-	-	-
Total TCDF	3220	3090	4 (≤50)	-	-	-
Total PeCDF	2980	2910	2 (≤50)	-	-	-
Total HxCDF	3450	3650	6 (≤50)	-	-	-
Total HpCDF	3590	3510	2 (≤50)	-	-	-
2,3,7,8-TCDF (DB-225)	233	231	1 (≤50)	-	-	-

*Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Data Qualification Summary - SDG R0903443

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903443	SA150-0.5B SA150-0.5BDL	OCDF	J (all detects)	A	Matrix spike/Matrix spike duplicates (RPD) (ld)
R0903443	SA197-0.5B SA198-0.5B SA198-0.5BDL SA64-0.5B SA104-0.5B SA104-0.5BDL SA129-0.5B SA129-0.5BDL SA70-0.5B SA70-0.5BDL SA60-0.5BDL SA60-0.5BDL SA150-0.5B SA150-0.5B SA150-0.5B SA150-0.5B SA200-0.5B SA43009-0.5B SA43009-0.5B SA200-0.5B SA200-0.5B SA200-0.5B SA200-0.5B SA200-0.5B SA200-0.5B SA200-0.5B SA200-0.5B SA51-0.5B SA51-0.5B	OCDF	J+ (all detects)	P	Laboratory control samples (%R) (I)
R0903443	SA129-0.5B	1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,7,8,9-HxCDD Total HxCDD OCDD OCDF	J (all detects) UJ (all non-detects)	Р	Internal standards (area) (i)
R0903443	SA129-0.5BDL	1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,7,8,9-HxCDD OCDD OCDF	J (all detects) UJ (all non-detects)	Р	Internal standards (area) (i)
R0903443	SA60-0.5BDL SA150-0.5BDL	OCDD	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р	internal standards (area) (i)
R0903443	SA197-0.5B	1,2,3,4,6,7,8-HpCDF OCDF	J (all detects) J (all detects)	Р	Project Quantitation Limit (e)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903443	SA198-0.5B SA70-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903443	SA64-0.5B	1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	Р	Project Quantitation Limit (e)
R0903443	SA104-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903443	SA129-0.5B	OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903443	SA60-0.5B SA150-0.5B	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 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903443	SA43009-0.5B SA43-0.5B	1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	Р	Project Quantitation Limit (e)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903443	SA40-0.5B	1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	Р	Project Quantitation Limit (e)
R0903443	SA200-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903443	SA51-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	А	Project Quantitation Limit (e)
R0903443	SA197-0.5B SA198-0.5B SA198-0.5BDL SA64-0.5B SA104-0.5B SA104-0.5BDL SA129-0.5B SA129-0.5BDL SA70-0.5BDL SA70-0.5BDL SA60-0.5BDL SA60-0.5BDL SA150-0.5BDL SA150-0.5B SA150-0.5B SA150-0.5B SA150-0.5B SA3-0.5B SA43009-0.5B SA200-0.5B SA200-0.5B SA200-0.5B SA200-0.5B SA200-0.5B SA200-0.5B SA51-0.5B SA51-0.5B SA51-0.5B SA51-0.5B	All compounds reported below the PQL.	J (all detects)	A	Project Quantitation Limit (sp)

<u> </u>		1			
SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903443	SA197-0.5B SA198-0.5B SA198-0.5BDL SA64-0.5B SA104-0.5BDL SA104-0.5BDL SA129-0.5BDL SA70-0.5BDL SA70-0.5BDL SA60-0.5BDL SA60-0.5BDL SA150-0.5BDL SA150-0.5B SA150-0.5BS SA150-0.5BS SA43009-0.5B SA43009-0.5B SA43009-0.5B SA200-0.5B SA200-0.5BDL RSA06-0.5B SA200-0.5BDL RSA06-0.5B SA51-0.5B SA51-0.5B SA51-0.5B	All compounds reported as EMPC	JK (all detects)	A	Project Quantitation Limit (k)
R0903443	SA197-0.5B SA64-0.5B RSAN5-0.5B SA53-0.5B SA43009-0.5B SA40-0.5B RSAO6-0.5B SA43-0.5B	2,3,7,8-TCDF (from DB-5)	X	A	Overall assessment of data (o)
R0903443	SA198-0.5B SA70-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x	А	Overall assessment of data (o)
R0903443	SA198-0.5BDL SA70-0.5BDL	All TCL compounds except 1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF	x	A	Overall assessment of data (o)

67.5		0	Eler	A or P	Possen (Cada)
SDG	Sample	Compound	Flag	AOPP	Reason (Code)
R0903443	SA104-0.5B	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 1,2,3,4,6,7,8-HpCDF 0CDF	x x x x x x x x	A	Overall assessment of data (o)
*R0903443	SA104-0.5BDL	2,3,7,8-TCDF (DB-5 & DB-225) All TCL compounds except 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	×	A	Overall assessment of data (o)
D0000440	01400 0 FD	2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-225)	X	A	Overall assessment of
R0903443	SA129-0.5B	OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X		data (o)
R0903443	SA129-0.5BDL	All TCL compounds except OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF	х	A	Overall assessment of data (o)
R0903443	SA60-0.5B SA150-0.5B	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 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X	A	Overall assessment of data (o)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
*R0903443	SA60-0.5BDL SA150-0.5BDL	All TCL compounds except 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 0CDF 2,3,7,8-TCDF (DB-225)	X	A	Overall assessment of data (o)
R0903443	SA200-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X	A	Overall assessment of data (o)
R0903443	SA200-0.5BDL	All TCL compounds except 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF	X	A	Overall assessment of data (o)
R0903443	SA51-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X	A	Overall assessment of data (o)
R0903443	SA51-0.5BDL	All TCL compounds except 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 2,3,7,8-TCDF	X	A	Overall assessment of data (o)

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Laboratory Blank Data Qualification Summary - SDG R0903443

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
R0903443	SA53-0.5B	1,2,3,4,6,7,8-HpCDD OCDD	0.502U ng/Kg 1.70U ng/Kg	A	bl
R0903443	RSAN5-0.5B	OCDD	7,58U ng/Kg	А	bl
R0903443	RSAO6-0.5B	OCDD	12.6U ng/Kg	А	bl

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Field Blank Data Qualification Summary - SDG R0903443

No Sample Data Qualified in this SDG

Tronox Northgate Henderson T

LDC #:	21495G21	VALIDATION COMPLETENESS WORKSHEE
SDG #:_	R0903443	Stage 2B
Laborato	ry: Columbia /	Analytical Services

Date: 9/18/09
Page:_/of_/
Reviewer:
2nd Reviewer: A

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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
l.	Technical holding times	A	Sampling dates: 6/19 - 24/09
II.	HRGC/HRMS Instrument performance check	4	, ,
111.	Initial calibration	₫.	
IV.	Routine calibration/IOV	₩	
V.	Blanks	W	
VI.	Matrix spike/Matrix spike duplicates	w	
VII.	Laboratory control samples	W	105
VIII.	Regional quality assurance and quality control	N,	
IX.	Internal standards	W	
X.	Target compound identifications	N	
XI.	Compound quantitation and CRQLs	√N_	
XII.	System performance	N	
XIII.	Overall assessment of data	-\$₩_	
XIV.	Field duplicates	W	D=16+17-17+24
XV.	Field blanks	SN	B07=10980 (R0904016)

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:

M 50:19			
1 SA197-0.5B	11/2 SA60-0.5B	21/5 RSAO6-0.5B	31 220900235-01
2 SA198-0.5B	12/4 SA60-0.5BDL	22 SA51-0.5B	32 /
3 SA198-0.5BDL	13 /2 SA150-0.5B	23 SA51-0.5BDL	33 DIO3737
4 /2 SA64-0.5B	14/4 SA150-0.5BDL	24 SA43-0.5B	34 HO381T
5 /2 SA104-0.5B	15 RSAN5-0.5B	25 SA150-0.5BMS	35 0103753
6/2 SA104-0.5BDL	16 X SA53-0.5B	26 SA150-0.5BMSD	36 P1038+0
7/2 SA129-0.5B	17 SA43009-0.5B	27	37 P1038-6
8 SA129-0.5BDL	18/5 SA40-0.5B	28	38 421965
9 SA70-0.5B	19'5 SA200-0.5B	29	39 P103791 1
7 10 SA70-0.5BDL	20 SA200-0.5BDL	30	408 P103864
<u> </u>			94103874

Notes: \$20306	*P203094	· 3 > 20315T	47203139	57203(20	7103864
			1		P703803

VALIDATION FINDINGS WORKSHEET

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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	a. 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	1. 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

2nd Reviewer: _______ Reviewer:

Blanks

Please see qualifications below for all questions answered "N", Not applicable questions are identified as "N/A". METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 8290)

Were all samples associated with a method blank?

Was a method blank analyzed for each matrix?

Was the blank contaminated? If yes, please see qualification below. In date: Blank extraction date: Y/N N/A

Sample Identification Associated Samples: 12.0 d 200 199110502 2 955 10-582 01/2 Blank ID Į. Compound Conc. units: 1/0/

Blank analysis date:_ Blank extraction date:_

Associated Samples: Conc. units:

_		 	 	 	
	llon				
	Sample Identification				
	Ø,				
	Blank ID				
	Compound				
	Comp				

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".

LDC #: 2|40824 SDG #:See Cover

VALIDATION FINDINGS WORKSHEET

Field Blanks

Page: of 2nd Reviewer: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

ng/Ka

Sampling date: 7/21/09

 $(\times S \times) \setminus M$ Sample Identification Associated Samples: Field blank type: (circle one) Field Blank / Rinsate / Other: 0.04185 0.03165 0.01785 0.04185 0.0202 0.2165 0.0905 0.3605 0.109 0.098 1.025 K EB072109-SO Blank ID 8.37 21.8 6.33 43.3 4.04 19.6 3.57 8.37 18.1 205 72.1 Compound

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

V:\Pei\Tronox\FB_SO_DIOXINS.wpd

LDC #:2495K

VALIDATION FINDINGS WORKSHEET Matrix Spike/Matrix Spike Duplicates

2nd Reviewer: Page: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

N/A

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.

N NA N NA

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?

Qualifications			X No gual YN TO R	1.2											1200 m						
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Date																					

SDG #: LECCOLL LDC #.249

VALIDATION FINDINGS WORKSHEET Internal Standards

Page: Cof L 2nd Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)
Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Are all internal standard recoveries were within the 40-135% criteria?

Y N/A Was the S/N ratio all internal standard peaks > 10?

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*	Date	l ah ID/Reference	Internal Standard	%	% Recovery (Limit: 40-135%)	0-135%)	Qualifications	lions
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					4			
					J			
					J			
))		
		Internal Standards	Check Standard Used		Inter	Internal Standards	Check S	Check Standard Used
Ą	¹³ C-2,3,7,8-TCDF	CDF			13C-OCDD			
ω̈	13C-2,3,7,8-TCDD	CDD		χ.	13C-1,2,3,4-TCDD			
ن	¹³ C-1,2,3,7,8-PeCDF	-PeCDF		ij	¹³ C-1,2,3,7,8,9-HxCDD	Q		
Ö	13C-1,2,3,7,8-PeCDD	-PeCDD		Σ				
ш	13C-1,2,3,4,7,8-HxCDF	,8-HxCDF		ż				
u.	¹³ C-1,2,3,6,7,8-HxCDD	,8-HxCDD		Ö				
Ö	¹³ C-1,2,3,4,6,7,8-HpCDF	,7,8-HpCDF		ai				
Ï	¹³ C-1,2,3,4,6,7,8-HpCDD	,7,8-HpCDD						

LDC #: 2149SE

VALIDATION FINDINGS WORKSHEET Compound Quantitation and Reported CRQLs

Page: of 4

Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Y MINIA

Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound? Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary).

Qualifications	(무)		1 deta (e)		12th 14 (0)	•		(4-4/47)	(de tratal 1 e)		
Associated Samples	ad							※シン 4	B5818-225 S	8-2%	
Finding	ente usulta.	de > calto lange	0.78		H.K.C. D. D. & 4. 9.	01.1 TR-22C		KAN X IN TO TO DE SYC	75-318-54 W # 8-2 N. N-1-1- 1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-1-	4.I-K.1-R, Han 185 & 18-23	
Sample ID	≱ ![2,9			4	2		
Date				·				•			
*											

Comments: See sample calculation verification worksheet for recalculations

LDC #:24405 SDG # 220000

Compound Quantitation and Reported CRQLs VALIDATION FINDINGS WORKSHEET

Page:

2nd Reviewer: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A",

Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound? Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary).

*	Date	Sample ID	spols > caliborange	Associated Samples	Qualifications
		51 11	F-8 (400 03520835)	=	-184/A1E)
					2
		17,24	K.O. 4. 8. 4 m 185 & DR-2X	18-2x 17 34	1 Start 10)
					1
		(8	K. L. OD Q, H W DBS RUB->X	8-22	
		B 19	T. K.L. M. D. D. B. H W NR-5 ROR-2X	5 ROB-2% , 4	1 1 th 1 6 0)
		22	7-14.0-28. 4 00 BS GOB-225	500 VV	

Comments: See sample calculation verification worksheet for recalculations

SDG #: 24 CO WAY

VALIDATION FINDINGS WORKSHEET Overall Assessment of Data

Page: of Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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.

(Y)N N/A Was the overall quality and usability of the data acceptable?

*	Date	Sample ID	Finding	Associated Samples	Qualifications
		1,4,815-18,7	H ex 123 -5	1,4,8,15-18,3	(0) X
		70		** **********************************	
		y , 4	I K. L. O. P. B. and Hm	2,4	
			75 21 BB-225		
		3,10	41 except 4.1.10. 6.0.7.8	3,10	
		19	F-4. I-M. 0-8. HMDBS&RDB-3>S	085x5 5	
			1		
		9	41 Wat F.F. I.M.O. A. B	9	
			255 4 or 78-255		
			-		
		_	4. 5-K. 0-R. How DB-5 & B-25	18-25 T	
		×	A1 except 4. H-K. 0-8	B	
Comments:	ınts:				

SDG #: 26 CONE

VALIDATION FINDINGS WORKSHEET Overall Assessment of Data

Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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.

(M) N/A Was the overall quality and usability of the data acceptable?

*	Date	Sample ID	Finding	Associated Samples	Qualifications
		શ /	F- & 1 HON (BS & DR-2X)	<u></u>	(o) ×
		L, (4	411 except FG 3. 1-8	4. 4	
			Han BB-4-25		
]					
		# 19	KT. K. L.H. 0-R.	0	
			Hr 08-5 808-25		
		R	AIL OXCEST H, I. K L. M	20	
			0.0.		
		x 22	I-M.O-R. HONDS-S & DB-2>5	な タ	
		4 X	All except H-H. O-B	45	
Comments:	ınts:				

LDC #: 24953 SDG #74 ZZ

VALIDATION FINDINGS WORKSHEET Laboratory Control Samples (LCS)

2nd Reviewer:

Page: Reviewer:__

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290) Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Was a LCS required?
Was a LCS analyzed every 20 samples for each matrix or whenever a sample extraction was performed?
Were the LCS percent recoveries (%R) and relative percent difference (RPD) within the QC limits?

		-	_			- 17	· · · · · ·	- T			11	,	- T	1	- 1	- 1 1			<u> </u>	- 7	7	 1			T	T	\neg
Qualifications	126ts 4 1)																										
Associated Samples	M+W																										
RPD (Limits)		,		^ }	()	()	()	()	()	()	()	()	(`)	()	()	()	((()	())	()	()
LCSD %R (Limits)	,			(()	()	()	(()	()	()		(()	())	()	()	().	()	()	())	()	()
LCS %R (1 imits)	CL 11. 871	140 (177)		()	()	(1 ((((_		((()	()	()	(()	()	()	()	()	()
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3	Lab iDineielelice	-Boronsoux	,																								
	# Date																										

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name:

Tronox LLC Facility, 2009 Phase B Investigation,

Henderson, Nevada

Collection Date:

June 29 through June 30, 2009

LDC Report Date:

September 29, 2009

Matrix:

Soil

Parameters:

Dioxins/Dibenzofurans

Validation Level:

Stage 2B

Laboratory:

Columbia Analytical Services, Inc.

Sample Delivery Group (SDG): R0903615

Sample Identification

SA45-0.5B

SA452009-0.5B

SA187-0.5B

SA187-0.5BDL

SA153-0.5B

SA186-0.5B

SA185-0.5B

RSA05-0.5B

SA50-0.5B

SA54-0.5B

SA106-0.5B

SA106-0.5BDL

SA102-0.5B

SA109-0.5B

SA106-0.5BMS

SA106-0.5BMSD

Introduction

This data review covers 16 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8290 for Polychlorinated Dioxins/Dibenzofurans.

This review follows the Standard Operating Procedures (SOP) 40, Data Review/Validation (BRC 2009), the Quality Assurance Project Plan Tronox LLC Facility, Henderson, Nevada (June 2009), NDEP guidance (May 2006), and USEPA Contract Laboratory Program National Functional Guidelines for Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) as there are no current guidelines for the method stated above.

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.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XIV.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- J+ Data are qualified as estimated, with a high bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J- Data are qualified as estimated, with a low bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J Data are qualified as estimated; it is not possible to assess the direction of the potential bias. False positives or false negatives are unlikely to have been reported.
- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- R Data are qualified as rejected. There is a significant potential for the reporting of false negatives or false positives.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- B The analytical result may be a false positive totally attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JB The analytical result may be biased high and partially attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JK The analytical result is an estimated maximum possible concentration (EMPC).
- X The analytical result is not used for reporting because a more accurate and precise result is reported in its place.
- J-TDS The analytical result is estimated based on failure of the Total Dissolved Solids (TDS) correctness check performed in accordance with the Standard Method 1030E.
- J-CAB The analytical result is estimated based on failure of the cation-anion balance correctness check performed in accordance with Standard Method 1030E.
- J-TDS & CAB The analytical result is unreliable based on the failure of the cation-anion balance and TDS correctness check performed in accordance with standard Method 1030E.
- 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.

Retention time windows were established for all homologues. The chromatographic resolution between 2,3,7,8-TCDD and peaks representing any other unlabeled TCDD isomer was less than or equal to 25%.

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 30.0% for labeled compounds.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

IV. Routine Calibration (Continuing)

Routine calibration was performed at the required frequencies.

All of the routine calibration percent differences (%D) between the initial calibration RRF and the routine calibration RRF were less than or equal to 20.0% for unlabeled compounds and less than or equal to 30.0% for labeled compounds.

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
EQ0900252-01	7/8/09	1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF Total HpCDD Total HpCDF	0.189 ng/Kg 0.496 ng/Kg 0.358 ng/Kg 0.609 ng/Kg 1.55 ng/Kg 0.213 ng/Kg 0.609 ng/Kg	All samples in SDG R0903615

Sample concentrations were compared to concentrations detected in the method blanks as required by the QAPP. No sample data was qualified with the following exceptions:

Sample	Compound	Reported Concentration	Modified Final Concentration
SA45-0.5B	OCDD	1.97 ng/Kg	1.97U ng/Kg
SA452009-0.5B	OCDD	2.16 ng/Kg	2.16U ng/Kg
SA153-0.5B	OCDD	3.02 ng/Kg	3.02U ng/Kg
SA54-0.5B	1,2,3,4,6,7,8-HpCDD OCDD Total HpCDD	0.419 ng/Kg 0.900 ng/Kg 0.419 ng/Kg	0.419U ng/Kg 0.900U ng/Kg 0.419U ng/Kg

No field blanks were identified in this SDG.

VI. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Although the MS/MSD percent recoveries (%R) and relative percent differences (RPD) were not within QC limits for some compounds, the LCS percent recoveries (%R) were within QC limits and no data were qualified.

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 recoveries were within QC limits with the following exceptions:

Sample	Internal Standards	%R (Limits)	Compound	Flag	A or P
RSAO5-0.5B	¹³ C-OCDD	34 (40-135)	OCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р
SA50-0.5B	¹³ C-OCDD	26 (40-135)	OCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р
SA54-0.5B	¹³ C-OCDD	29 (40-135)	OCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р

X. Target Compound Identifications

Raw data were not reviewed for this SDG.

XI. Project Quantitation Limit

All project quantitation limits were within validation criteria with the following exceptions:

Sample	Compound	Finding	Criteria	Flag	A or P
SA187-0.5B	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 1,2,3,4,6,7,8-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 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	А

Sample	Compound	Finding	Criteria	Flag	A or P
SA106-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HxCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A
SA106-0.5BDL	OCDF	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	А

All compounds reported below the PQL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903615	All compounds reported below the PQL.	J (all detects)	А

All compounds reported as EMPC were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903615	All compounds reported as estimated maximum possible concentration (EMPC).	JK (all detects)	А

Raw data were not reviewed for this SDG.

XII. System Performance

Raw data were not reviewed for this SDG.

XIII. Overall Assessment of Data

The overall assessment of data was acceptable. In the case where more than one result was reported for an individual sample, the least technically acceptable results were rejected as follows:

Sample	Compound	Flag	A or P
SA187-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X	А
SA187-0.5BDL	All TCL compounds except 1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-HxCDF 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 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-225)	X	A
SA45-0.5B SA452009-0.5B SA153-0.5B SA186-0.5B SA185-0.5B RSAO5-0.5B SA50-0.5B SA54-0.5B SA102-0.5B SA109-0.5B SA109-0.5B	2,3,7,8-TCDF (DB-5)	X	A
SA106-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-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 1,2,3,4,7,8-HxCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X X	A

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

Samples SA45-0.5B and SA452009-0.5B were identified as field duplicates. No polychlorinated dioxins/dibenzofurans were detected in any of the samples with the following exceptions:

	Concentra	tion (ng/Kg)		_			
Compound	SA45-0.5B	SA452009-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P	
2,3,7,8-TCDD	0.189	0.204	-	0.015 (≤0.952)	-	-	
1,2,3,7,8-PeCDD	0.277	0.255	-	0.022 (≤2.38)	-	•	
1,2,3,4,7,8-HxCDD	0.135	0.114	-	0.021 (≤2.38)	-	-	
1,2,3,6,7,8-HxCDD	0.413	0.410	-	0.003 (≤2.38)	-	-	
1,2,3,7,8,9-HxCDD	0.462	0.480	-	0.018 (≤2.38)	-	-	
1,2,3,4,6,7,8-HpCDD	1.10	1.14	-	0.04 (≤2.38)	-	-	
OCDD	1.97	2.16	-	0.19 (≤4.76)	-	-	
2,3,7,8-TCDF	2.79	3,42	•	0.63 (≤0.952)	-	-	
1,2,3,7,8-PeCDF	1.96	2.27	-	0.31 (≤2.38)	-	-	
2,3,4,7,8-PeCDF	1.12	1.23	•	0.11 (≤2.38)	-	_	
1,2,3,4,7,8-HxCDF	3.52	3.67	-	0.15 (≤2.38)	-	-	
1,2,3,6,7,8-HxCDF	2.28	2.46	•	0.18 (≤2.38)	-	-	
1,2,3,7,8,9-HxCDF	0.461	0.483	•	0.022 (≤2.38)	-	-	
2,3,4,6,7,8-HxCDF	1.55	1.43	-	0.12 (≤2.38)	-	-	
1,2,3,4,6,7,8-HpCDF	7.03	6.76		0.27 (≤2.38)	-	-	
1,2,3,4,7,8,9-HpCDF	3.16	3.12	-	0.04 (≤2.38)	-	-	
OCDF	19.3	19.2	_	0.1 (≤4.76)	-	÷	

	Concentration (ng/Kg)		, ppp	Difference		
Compound	SA45-0.5B	SA452009-0.5B	RPD (Limits)	(Limits)	Flag	A or P
Total TCDD	0.695	1.45	-	0.755 (≤0.952)	-	-
Total PeCDD	0.542	1.03	-	0.488 (≤2.38)	+	-
Total HxCDD	1.01	2.12	-	1.11 (≤2.38)	-	-
Total HpCDD	2.21	2.18	-	0.03 (≤2.38)	*	-
Total TCDF	18.4	21.5	16 (≤50)	-	-	-
Total PeCDF	17.2	18.3	6 (≤50)	•	-	-
Total HxCDF	16.4	17.4	6 (≤50)	-	-	<u>-</u>
Total HpCDF	14.6	14.3	2 (≤50)	-	-	-
2,3,7,8-TCDF (DB-225)	1.01	2.10	-	1.09 (≤0.952)	•	-

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Data Qualification Summary - SDG R0903615

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903615	RSAO5-0.5B SA50-0.5B SA54-0.5B	OCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р	Internal standards (%R) (i)
R0903615	SA187-0.5B	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 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903615	SA106-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HxCDD 1,2,3,7,8-PeCDF 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 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	A	Project Quantitation Limit (e)
R0903615	SA106-0.5BDL	OCDF	J (all detects)	Α	Project Quantitation Limit (e)
R0903615	SA45-0.5B SA452009-0.5B SA187-0.5B SA187-0.5BDL SA153-0.5B SA186-0.5B SA185-0.5B RSAO5-0.5B SA50-0.5B SA54-0.5B SA106-0.5B SA106-0.5B SA109-0.5B	All compounds reported below the PQL.	J (all detects)	A	Project Quantitation Limit (sp)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903615	SA45-0.5B SA452009-0.5B SA187-0.5B SA187-0.5BDL SA153-0.5B SA186-0.5B SA185-0.5B RSA05-0.5B SA50-0.5B SA50-0.5B SA106-0.5B SA106-0.5B SA106-0.5B SA106-0.5B	All compounds reported as EMPC	JK (all detects)	A	Project Quantitation Limit (k)
R0903615	SA187-0.5B	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 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X	A	Overall assessment of data (o)
R0903615	SA187-0.5BDL	All TCL compounds except 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 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-225)	X	A	Overall assessment of data (o)
R0903615	SA45-0.5B SA452009-0.5B SA153-0.5B SA186-0.5B SA185-0.5B RSAO5-0.5B SA50-0.5B SA54-0.5B SA102-0.5B SA109-0.5B SA109-0.5B	2,3,7,8-TCDF (DB-5)	X	A	Overall assessment of data (o)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903615	SA106-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HxCDD 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 1,2,3,6,7,8-HxCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x x x x x x	A	Overall assessment of data (o)

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Laboratory Blank Data Qualification Summary - SDG R0903615

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
R0903615	SA45-0.5B	OCDD	1.97U ng/Kg	А	bl
R0903615	SA452009-0.5B	OCDD	2.16U ng/Kg	А	bl
R0903615	SA153-0.5B	OCDD	3.02U ng/Kg	А	bl
R0903615	SA54-0.5B	1,2,3,4,6,7,8-HpCDD OCDD Total HpCDD	0.419U ng/Kg 0.900U ng/Kg 0.419U ng/Kg	А	bl

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Field Blank Data Qualification Summary - SDG R0903615

No Sample Data Qualified in this SDG

Tronox Northgate Henderson SHEET

LDC #: 21495H21	VALIDATION COMPLETENESS WORKS
SDG #: R0903615	Stage 2B
Laboratory: Columbia Analytical	Services

Date: 9/17/29	=
Page:/of/	
Reviewer:	
2nd Reviewer:	

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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	
1.	Technical holding times	A	Sampling dates: 6/29-30/0 9	
11.	HRGC/HRMS Instrument performance check	4	′ ′ ′	
III.	Initial calibration	4		
IV.	Routine calibration/I	1		
V.	Blanks	áW	+	
VI.	Matrix spike/Matrix spike duplicates	W	7. E and RDO and IN 15/16 - No Canal 1605,	m
VII.	Laboratory control samples	A	109	
VIII.	Regional quality assurance and quality control	N N		
IX.	Internal standards	W		
X.	Target compound identifications	N	·	
XI.	Compound quantitation and CRQLs	1/W		
XII.	System performance	N		
XIII.	Overall assessment of data	av		
XIV.	Field duplicates	W	7=1+2 ,	
XV.	Field blanks	XN	FB07=109 50(10904016) C	

Note:

A = Acceptable

N = Not provided/applicable

SW = See worksheet

ND = No compounds detected

R = Rinsate

D = Duplicate

TB = Trip blank

FB = Field blank

EB = Equipment blank

Validated Samples:

<u> </u>	15015							
1	SA45-0.5B	11	1/2	SA106-0.5B	21 ³	20900252-01	31	U219752
1	SA452009-0.5B	12	ST.	SA106-0.5BDL	22	` /	32 -	P103991
3/2	SA187-0.5 B		1/2	-SA102-0.5B	23		33	
4/3	SA187-0.50LBDL	14		SA109-0.5B	24		34	P103952
5 />	SA153-0.5B	15	, ;	SA106-0.5BMS	25		35	
67/4	SA186-0.5B	16	3	SA106-0.5BMSD	26		36	
73/4	SA185-0.5B	17	,		27	P203448	37	
1 3 6	∕RSAO5-0.5B	18	3		28	D2133(1	38	
9/2	SA50-0.5B	19)		29 ³	703360	39	
10/2	≻SA54-0.5B	20)		$_{30}\psi$	P203327	40	

Notes:		

VALIDATION FINDINGS WORKSHEET

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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	1. 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:

LDC #21495 SDG #: 22

VALIDATION FINDINGS WORKSHEET Blanks

Page:

2nd Reviewer:_ Reviewer:_

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A". METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 8290)

Were all samples associated with a method blank? N N N

Was a method blank analyzed for each matrix?

 \overline{V} N/A Was the blank contaminated? If yes, please see qualification below. Blank extraction date: \overline{Z} Blank analysis date: \overline{Z}

Sample Identification 4/9 1900 1 Associated Samples: 6 9 W Ì d Ŋ 4 9 189 .35B 600 496 人のなかのかり Blank ID Compound Sonc. units: 1/25 4

Associated Samples: Blank analysis date: Blank extraction date: Conc. units:

609 M.

2

5	_	-	_	_		
	Ę					
	Sample Identification					
	San					
including maintain						
	Blank ID					
	Compound					

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

Page: of A

2nd Reviewer:__

Internal Standards

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A". METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Are all internal standard recoveries were within the 40-135% criteria? Was the S/N ratio all internal standard peaks > 10?

Y M N/A

							/	
#	Date	Lab ID/Reference	Internal Standard		% Recovery (Limit: 40-135%)	:: 40-135%)	Qualifications	/ tions
		X	A		አ	140-130) 4/ NY/ \	(★,★)
T		3				(_	
		В	4	d	26			
		0/	+	29	7	(//	Λ	
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		Internal Standards	Check Standard Used			Internal Standards	Check	Check Standard Used
ά	¹³ C-2,3,7,8-TCDF	CDF			13C-OCDD			
ю	¹³ C-2,3,7,8-TCDD	CDD		Υ.	¹³ C-1,2,3,4-TCDD			
ن	 	PeCDF			¹³ C-1,2,3,7,8,9-HxCDD	хсрр		
۵	¹³ C-1,2,3,7,8-PeCDD	PeCDD		Σ				
шi		8-HxCDF		z				
ıı.		8-HxCDD		이				
Ö	¹³ C-1,2,3,4,6,7,8-HpCDF	7,8-HpCDF		a:			7	
Ξ	13C-1,2,3,4,6,7,8-HpCDD	,7,8-HpCDD						

LDC #: 26/54/2/ SDG #: 566 COND

Compound Quantitation and Reported CRQLs VALIDATION FINDINGS WORKSHEET

Page: Reviewer:

2nd Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Y N N/A

Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound? Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary).

	Qualifications	Lt. /4 (0)						\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\				
	Associated Samples	7 m				<u> </u>		17				
	ods > cally range	F-> & (+) an booth	1B5 R 1B-225)	4-8 (How both DB-5	and DB-22S)	«		ZMPC MSW AS				
	Sample ID	M		1		4						
	# Date				٠			-				
L										·	<u> </u>	<u></u>

Comments: See sample calculation verification worksheet for recalculations

LDC #:>495#>

VALIDATION FINDINGS WORKSHEET Overall Assessment of Data

Page: /of /
Reviewer: &

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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.

(Y) N/A Was the overall quality and usability of the data acceptable?

*	Date	Sample ID	Finding	Associated Samples	Qualifications
		M	F> & CHON bath	Ŋ	(o) X
			(25-3C X 2-3C)		
		4	All 2x cept F. G.	1	
			2 -> & Hon BB-225		
		1-2. \$ 5-10, 13-14	\$ H ON DB-S	11-2.5-1	
		,			
			A- & (# or both 28-5 &		
			\$\$-25E)		
		4	H 5N OB-5	7	
Comments:	ents:				

LDC#:21495H21 SDG#: See Cover

VALIDATION FINDINGS WORKSHEET Field Duplicates

Page:_/of_/
Reviewer:______
2nd Reviewer:______

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

YN NA YN NA Were field duplicate pairs identified in this SDG?
Were target analytes detected in the field duplicate pairs?

	Concentrati	on (ng/Kg)	(≤50)	(ng/Kg)	(ng/Kg)	Qualifications
Compound	1	2	RPD	Difference	Limits	(Parent Only)
Α	0.189	0.204		0.015	(<u><</u> 0.952)	
В	0.277	0.255		0.022	(≤2.38)	
С	0.135	0.114		0.021	(<u><</u> 2.38)	
D	0.413	0.410		0.003	(<u><</u> 2.38)	
E	0.462	0.480		0.018	(≤2.38)	
F	1.10	1.14		0.04	(≤2.38)	
G	1.97	2.16		0.19	(<u><</u> 4.76)	
Н	2.79	3.42		0.63	(≤0.952)	
I.	1.96	2.27		0.31	(<u>≤</u> 2.38)	
J	1.12	1.23		0.11	(≤2.38)	
к	3.52	3.67		0.15	(<u>≤</u> 2.38)	
L	2.28	2.46		0.18	(<u>≤</u> 2.38)	
N	0.461	0.483		0.022	(≤2.38)	
M	1.55	1.43		0.12	(<u>≤</u> 2.38)	
0	7.03	6.76		0.27	(<u><</u> 2.38)	
P	3.16	3.12		0.04	(<u>≤</u> 2.38)	
Q	19.3	19.2		0.1	(<u>≤</u> 4.76)	
R	0.695	1.45		0.755	(<u><</u> 0.952)	
s	0.542	1.03		0.488	(<u><</u> 2.38)	
Т	1.01	2.12		1.11	(<u>≤</u> 2.38)	
U	2.21	2.18		0.03	(≤2.38)	
V	18.4	21.5	16			
w	17.2	18.3	6			
х	16.4	17.4	6			
Υ	14.6	14.3	2			
H (DB-225)	1.01	2.10		1.09	(≤0.952)	

LDC Report# 21495l21

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name:

Tronox LLC Facility, 2009 Phase B Investigation,

Henderson, Nevada

Collection Date:

July 1 through July 2, 2009

LDC Report Date:

October 21, 2009

Matrix:

Soil

Parameters:

Dioxins/Dibenzofurans

Validation Level:

Stage 2B

Laboratory:

Columbia Analytical Services, Inc.

Sample Delivery Group (SDG): R0903678

Sample Identification

SA114-0.5B

SA114-0.5BDL

SA114009-0.5B

SA114009-0.5BDL

RSAN6-0.5B

SA82-0.5B

SA82-0.5BDL

RSAK3-0.5B

RSAK3-0.5BDL

SA82-0.5BMS

SA82-0.5BMSD

Introduction

This data review covers 11 soil samples listed on the cover sheet including dilutions and reanalysis as applicable. The analyses were per EPA SW 846 Method 8290 for Polychlorinated Dioxins/Dibenzofurans.

This review follows the Standard Operating Procedures (SOP) 40, Data Review/Validation (BRC 2009), the Quality Assurance Project Plan Tronox LLC Facility, Henderson, Nevada (June 2009), NDEP guidance (May 2006), and USEPA Contract Laboratory Program National Functional Guidelines for Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) as there are no current guidelines for the method stated above.

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.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XIV.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- J+ Data are qualified as estimated, with a high bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J- Data are qualified as estimated, with a low bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J Data are qualified as estimated; it is not possible to assess the direction of the potential bias. False positives or false negatives are unlikely to have been reported.
- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- R Data are qualified as rejected. There is a significant potential for the reporting of false negatives or false positives.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- B The analytical result may be a false positive totally attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JB The analytical result may be biased high and partially attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JK The analytical result is an estimated maximum possible concentration (EMPC).
- X The analytical result is not used for reporting because a more accurate and precise result is reported in its place.
- J-TDS The analytical result is estimated based on failure of the Total Dissolved Solids (TDS) correctness check performed in accordance with the Standard Method 1030E.
- J-CAB The analytical result is estimated based on failure of the cation-anion balance correctness check performed in accordance with Standard Method 1030E.
- J-TDS & CAB The analytical result is unreliable based on the failure of the cation-anion balance and TDS correctness check performed in accordance with standard Method 1030E.
- 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.

Retention time windows were established for all homologues. The chromatographic resolution between 2,3,7,8-TCDD and peaks representing any other unlabeled TCDD isomer was less than or equal to 25%.

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 30.0% for labeled compounds.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

IV. Routine Calibration (Continuing)

Routine calibration was performed at the required frequencies.

All of the routine calibration percent differences (%D) between the initial calibration RRF and the routine calibration RRF were less than or equal to 20.0% for unlabeled compounds and less than or equal to 30.0% for labeled compounds.

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
EQ0900252-01	7/8/09	1,2,3,4,6,7,8-HpCDD OCDD 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF Total HpCDD Total HpCDF	0.189 ng/Kg 0.496 ng/Kg 0.358 ng/Kg 0.609 ng/Kg 1.55 ng/Kg 0.213 ng/Kg 0.609 ng/Kg	All samples in SDG R0903678

Sample concentrations were compared to concentrations detected in the method blanks as required by the QAPP. No sample data was qualified.

Sample FB072109-SO (from SDG R0904016) was identified as a field blank. No polychlorinated dioxin/dibenzofuran contaminants were found in this blank with the following exceptions:

Field Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB072109-SO	7/21/09	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-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF Total HpCDD Total HxCDF	8.37 pg/L 21.8 pg/L 6.33 pg/L 3.57 pg/L 4.04 pg/L 43.3 pg/L 19.6 pg/L 205 pg/L 8.37 pg/L 18.1 pg/L 72.1 pg/L	SA82-0.5B SA82-0.5BDL RSAK3-0.5B RSAK3-0.5BDL

Sample concentrations were compared to concentrations detected in the field blanks as required by the QAPP. No sample data was qualified.

VI. Matrix Spike/Matrix Spike Duplicates

Matrix spike (MS) and matrix spike duplicate (MSD) samples were reviewed for each matrix as applicable. Although the MS/MSD percent recoveries (%R) and relative percent differences (RPD) were not within QC limits for some compounds, the LCS percent recoveries (%R) were within QC limits and no data were qualified.

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 recoveries were within QC limits with the following exceptions:

Sample	Internal Standards	%R (Limits)	Compound	Flag	A or P
SA114-0.5BDL	13C-OCDD	39 (40-135)	OCDD OCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р
SA114009-0.5B	¹³ C-OCDD	37 (40-135)	OCDD	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р
SA114009-0.5BDL	¹³ C-OCDD	29 (40-135)	OCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р
SA82-0.5BDL	¹³ C-OCDD	37 (40-135)	OCDD	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р
RSAK3-0.5B	¹³ C-2,3,7,8-TCDF ¹³ C-2,3,7,8-TCDD ¹³ C-1,2,3,4,6,7,8-HpCDF ¹³ C-OCDD	11 (40-135) 29 (40-135) 30 (40-135) 32 (40-135)	2,3,7,8-TCDF Total TCDF 2,3,7,8-TCDD Total TCDD 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF Total HpCDF OCDD OCDF	J (all detects) UJ (all non-detects)	P
RSAK3-0.5BDL	¹³ C-2,3,7,8-TCDF ¹³ C-2,3,7,8-TCDD ¹³ C-1,2,3,4,7,8-HxCDF ¹³ C-1,2,3,6,7,8-HxCDD ¹³ C-1,2,3,4,6,7,8-HpCDF ¹³ C-OCDD	11 (40-135) 27 (40-135) 34 (40-135) 36 (40-135) 30 (40-135) 23 (40-135)	2,3,7,8-TCDF 2,3,7,8-TCDD 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDD OCDF	J (all detects) UJ (all non-detects)	Р

^{*}Corrected compounds for RSAK3-0.5BDL

X. Target Compound Identifications

Raw data were not reviewed for this SDG.

XI. Project Quantitation Limit

All project quantitation limits were within validation criteria with the following exceptions:

Sample	Compound	Finding	Criteria	Flag	A or P
SA114-0.5B SA114009-0.5B	1,2,3,4,6,7,8-HpCDD OCDD 2,3,7,8-TCDF 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	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	А
RSAN6-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A
SA82-0.5B	2,3,7,8-TCDF 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	А
RSAK3-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HyCDD 1,2,3,4,6,7,8-HpCDD OCDD 2,3,7,8-TCDF 1,2,3,7,8-PeCDF 1,2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HyCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	A

All compounds reported below the PQL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903678	All compounds reported below the PQL.	J (all detects)	А

All compounds reported as EMPC were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903678	All compounds reported as estimated maximum possible concentration (EMPC).	JK (all detects)	А

Raw data were not reviewed for this SDG.

XII. System Performance

Raw data were not reviewed for this SDG.

*XIII. Overall Assessment of Data

The overall assessment of data was acceptable. In the case where more than one result was reported for an individual sample, the least technically acceptable results were rejected as follows:

Sample	Compound	Flag	A or P
SA114-0.5B SA114009-0.5B	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 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 1,2,3,4,7,8,9-HpCDF 2,3,7,8,1-TCDF (DB-5 & DB-225)	x x x x x x x x	А

Sample	Compound	Flag	A or P
SA114-0.5BDL SA114009-0.5BDL	All TCL compounds except 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 0CDF 2,3,7,8-TCDF (DB-225)	x	A
SA82-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x	А
SA82-0.5BDL	All TCL compounds except 2,3,7,8-TCDF 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	X	А
RSAK3-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 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 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X X X	A
RSAN6-0.5B RSAK3-0.5BDL	2,3,7,8-TCDF (DB-5)	x	А

^{*}Corrected compounds in table above.

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

Samples SA114-0.5B and SA114009-0.5B and samples SA114-0.5BDL and SA114009-0.5BDL were identified as field duplicates. No polychlorinated dioxins/dibenzofurans were detected in any of the samples with the following exceptions:

	Concentrati	on (ng/Kg)	RPD	Difference		
Compound	SA114-0.5B	SA114009-0.5B	(Limits)	(Limits)	Flag	A or P
2,3,7,8-TCDD	73.5	57.7	24 (≤50)	•	-	-
1,2,3,7,8-PeCDD	263	195	30 (≤50)	•	-	-
1,2,3,4,7,8-HxCDD	213	152	33 (≤50)	-	-	•
1,2,3,6,7,8-HxCDD	356	256	33 (≤50)	-	-	-
1,2,3,7,8,9-HxCDD	417	286	37 (≤50)	-	-	-
1,2,3,4,6,7,8-HpCDD	1240	940	28 (≤50)	-	-	-
OCDD	1360	1050	26 (≤50)	-	-	-
2,3,7,8-TCDF	3510	4190	18 (≤50)	•	-	-
1,2,3,7,8-PeCDF	2590	2860	10 (≤50)	-	-	-
2,3,4,7,8-PeCDF	1440	1530	6 (≤50)	-	-	-
1,2,3,4,7,8-HxCDF	6050	6620	9 (≤50)	-	-	-
1,2,3,6,7,8-HxCDF	3330	3950	17 (≤50)	-	-	-
1,2,3,7,8,9-HxCDF	559	725	26 (≤50)	-	-	-
2,3,4,6,7,8-HxCDF	2070	2290	10 (≤50)	-	-	•
1,2,3,4,6,7,8-HpCDF	11500	13600	17 (≤50)	-	-	-
1,2,3,4,7,8,9-HpCDF	6040	6870	13 (≤50)	-	-	-

	Concentration (ng/Kg)		D :#	Difference		
Compound	SA114-0.5B	SA114009-0.5B	RPD (Limits)	(Limits)	Flag	A or P
OCDF	32900	42800	26 (≤50)	-	- -	-
Total TCDD	3340	2310	36 (≤50)	-	-	_
Total PeCDD	2960	2200	29 (≤50)	-	-	-
Total HxCDD	2990	2230	29 (≤50)	-	+	-
Total HpCDD	1900	1500	24 (≤50)	-	-	-
Total TCDF	33400	42600	24 (≤50)	•	-	-
Total PeCDF	21100	23500	11 (≤50)	-	-	-
Total HxCDF	25500	19400	27 (≤50)	-	-	-
Total HpCDF	26300	24700	6 (≤50)	-	-	-
2,3,7,8-TCDF (DB-225)	1120	1380	21 (≤50)	-	-	-

	Concentration (ng/Kg)			D:#		
Compound	SA114-0.5BDL	SA114009-0.5BDL	RPD (Limits)	Difference (Limits)	Flag	A or P
2,3,7,8-TCDD	81.0	77.5	-	3.5 (≤65.3)	-	-
1,2,3,7,8-PeCDD	251	206	-	45 (≤138)	_	-
1,2,3,4,7,8-HxCDD	244	163	-	81 (≤138)	-	-
1,2,3,6,7,8-HxCDD	404	315	-	89 (≤138)	-	-
1,2,3,7,8,9-HxCDD	448	357	-	91 (≤138)	-	-
1,2,3,4,6,7,8-HpCDD	1250	972	25 (≤50)	-	-	<u>-</u> .
OCDD	1780	1240	-	540 (≤276)	J (all detects)	Α
2,3,7,8-TCDF	3810	5960	44 (≤50)	-	-	-

	Concentrati	on (ng/Kg)	RPD	Difference		
Compound	SA114-0.5BDL	SA114009-0.5BDL	(Limits)	(Limits)	Flag	A or P
1,2,3,7,8-PeCDF	2150	2960	32 (≤50)	-	-	-
2,3,4,7,8-PeCDF	1150	1490	26 (≤50)	-	-	-
1,2,3,4,7,8-HxCDF	6930	8090	15 (≤50)	-	-	-
1,2,3,6,7,8-HxCDF	3710	4990	29 (≤50)	-	-	-
1,2,3,7,8,9-HxCDF	651	680	-	29 (≤138)	-	-
2,3,4,6,7,8-HxCDF	2210	2670	19 (≤50)	-	-	-
1,2,3,4,6,7,8-HpCDF	14600	17700	19 (≤50)	-	-	-
1,2,3,4,7,8,9-HpCDF	6710	7480	11 (≤50)	-	-	-
OCDF	43200	55000	24 (≤50)	-	-	-
2,3,7,8-TCDF (DB-225)	881	1040	17 (≤50)	-	-	-

*Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Data Qualification Summary - SDG R0903678

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903678	SA114-0.5BDL SA114009-0.5B SA114009-0.5BDL SA82-0.5BDL	OCDD	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р	Internal standards (%R) (i)
R0903678	RSAK3-0.5B	2,3,7,8-TCDF Total TCDF 2,3,7,8-TCDD Total TCDD 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF Total HpCDF OCDD OCDF	J (all detects) UJ (all non-detects)	Р	Internal standards (%R) (i)
*R0903678	RSAK3-0.5BDL	2,3,7,8-TCDF 2,3,7,8-TCDD 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,7,8,9-HxCDD 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDD OCDF	J'(all detects) UJ (all non-detects)	Р	Internal standards (%R) (i)
R0903678	SA114-0.5B SA114009-0.5B	1,2,3,4,6,7,8-HpCDD OCDD 2,3,7,8-TCDF 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	J (all detects)	А	Project Quantitation Limit (e)
R0903678	RSAN6-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-225)	J (all detects)	A	Project Quantitation Limit (e)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903678	SA82-0.5B	2,3,7,8-TCDF 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	J (all detects)	A	Project Quantitation Limit (e)
R0903678	RSAK3-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD OCDD 2,3,7,8-TCDF 1,2,3,7,8-PeCDF 1,2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF	J (all detects)	A	Project Quantitation Limit (e)
R0903678	SA114-0.5B SA114-0.5BDL SA114009-0.5B SA114009-0.5BDL RSAN6-0.5B SA82-0.5B SA82-0.5BDL RSAK3-0.5B RSAK3-0.5BDL	All compounds reported below the PQL.	J (all detects)	A	Project Quantitation Limit (sp)
R0903678	SA114-0.5B SA114-0.5BDL SA114009-0.5B SA114009-0.5BDL RSAN6-0.5B SA82-0.5B SA82-0.5BDL RSAK3-0.5B RSAK3-0.5BDL	All compounds reported as EMPC	JK (all detects)	A	Project Quantitation Limit (k)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
*R0903678	SA114-0.5B SA114009-0.5B	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 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X X X	A	Overall assessment of data (o)
*R0903678	SA114-0.5BDL SA114009-0.5BDL	All TCL compounds except 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 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-225)	х	A	Overall assessment of data (o)
*R0903678	SA82-0.5B	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	X X X X X X	A	Overall assessment of data (o)
*R0903678	SA82-0.5BDL	All TCL compounds except 2,3,7,8-TCDF 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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF	X	A	Overall assessment of data (o)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
*R0903678	RSAK3-0.5B	2,3,7,8-TCDD 1,2,3,7,8-PeCDD 1,2,3,4,7,8-HxCDD 1,2,3,6,7,8-HxCDD 1,2,3,4,6,7,8-HpCDD 0CDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x x x x	Α	Overall assessment of data (o)
*R0903678	RSAN6-0.5B RSAK3-0.5BDL	2,3,7,8-TCDF (DB-5)	х	А	Overall assessment of data (o)
R0903678	SA114-0.5BDL SA114009-0.5BDL	OCDD	J (all detects)	А	Field duplicates (Difference) (fd)

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Laboratory Blank Data Qualification Summary - SDG R0903678

No Sample Data Qualified in this SDG

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Field Blank Data Qualification Summary - SDG R0903678

No Sample Data Qualified in this SDG

Tronox Northgate Henderson WORKSHEET

LDC #: 21495I21	VALIDATION COMPLETENESS
SDG #: R0903678	Stage 2B
Laboratory: Columbia Analytical	Services

	Date:	9/3/03
	Page:_	$Z_{of}Z$
	Reviewer:	W.
2nd	Reviewer:	1/

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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: 7/1 - 2 / 0-9
II.	HRGC/HRMS Instrument performance check	A	
III.	Initial calibration	A	
IV.	Routine calibration/IgV	\bigvee	
V.	Blanks	w	
VI.	Matrix spike/Matrix spike duplicates	KW	
VII.	Laboratory control samples	\forall	109
VIII.	Regional quality assurance and quality control	N,	
iX.	Internal standards	W	
Χ.	Target compound identifications	N	
XI.	Compound quantitation and CRQLs	SW	
XII.	System performance	W	
XIII.	Overall assessment of data	S W	
XIV.	Field duplicates	in	D=1+3 2+4.
XV.	Field blanks	W	D=1+3,2+4. FB07=109-50(R0904016)

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:

<u> </u>	150.19					The state of the s
12/	SA114-0.5B	7/2 11 / SA82-0.5BMSD	21	280900252-01	31	·U=1976= 1
25/2	SA114-0.5BDL ,	12	22	,	32	P103952
3 B	SA114009-0.5B	13	23		33≥	112194
4/2	SA114009-0.5BDL)	14	24		34	D103938
	RSAN6-0.5B	15	25		35	+103978
67/1	SA82-0.5B	16	26		36	
73	SA82-0.5BDL &	17	27	D203>92	37	
87/3	RSAK3-0.5B	18	حـ 28	D2022 15	38	
97	RSAK3-0.5BDL	19	293	P=03311	39	
10/1	SA82-0.5BMS	20	304	P203327	40	

Notes:		
		_
	\	

VALIDATION FINDINGS WORKSHEET

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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	11 Total Hocod
R 12378-DeCDD	2000			
	e. ocen	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 Pacing
1038 H 000	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			
U. 1,4,5,6,6,6,7,000	1. 1,4,3,7,8-PBCDF	N. 1,2,3,7,8,9-HxODF	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

lotes;

SDG #: John Box LDC #3/495/3

VALIDATION FINDINGS WORKSHEET Blanks

2nd Reviewer:_ Reviewer Page:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Were all samples associated with a method blank?

Was a method blank analyzed for each matrix?

Was the blank contaminated? If yes, please see qualification below. In date: 78/09 Blank analysis date: 7/4/09

(Y) N N/A Was the blank contaminated? If yes, please see Blank extraction date: 78/0 Blank analysis date: 7/4/4 Conc. units: NS,

Sample Identification Associated Samples: 0.189 W/4 0.358 609.0 0.496 0.609 Blank ID -x 400x2-0 55 Compound Ø

Blank analysis date: Blank extraction date:

Conc. units:

Associated Samples:

Compound	Blank ID	
	·	

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".

100 #: 2495 P SDG #:See Cover

VALIDATION FINDINGS WORKSHEET

Field Blanks

Page: of [Reviewer: A

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Associated sample units: YN N/A Were field blanks identified in this SDG?

Blank units: pg/L Associated sample units

Sampling date: 7/21/09

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

Associated Samples:

75x 16-7.8-9 (>5x Sample Identification 0.04185 0.04185 0.03165 0.01785 0.0202 0.2165 0.0905 0.3605 0.109 0.098 1.025 놝 FB072109-SD Blank ID 21.8 8.37 6.33 19.6 3.57 4.04 43.3 205 8.37 18.1 72.1 Compound CROL 0

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

LDC #: 2 SDG #:

VALIDATION FINDINGS WORKSHEET Matrix Spike/Matrix Spike Duplicates

Page: Reviewer: 2nd Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Phease see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".
| N N/A | 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.

N N/A

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?

F		T	T	T	T	T	T	T	T	7	Ţ	Ī	T	T	_		T	T	7	-	T		-		T	-
	Qualifications	10 a.s.																								
	Associated Samples	8																								
	APD (Limits)	())	()		()	())							(()	()	()	())	,		_		
MSD	%R (Limits)	25-41	()	())	()	,	()	()		, ,		, , ,		-	· ·	()	()	()					(
MS	%R (Limits)	QA.		()	()	()	()	(()))		_		,		(()	()	(_
	Compound	JOK and																								
	OI OSW/SW	11/0												-												
<u> </u>	Date																		1							
1	*										ļ		ı													

LDC #= 4495 |= | SDG #: 26.4 CONUM

VALIDATION FINDINGS WORKSHEET Internal Standards

Page: Reviewer: 2nd Revlewer:

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A". METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290) KN/N/A

Are all internal standard recoveries were within the 40-135% criteria?

Was the S/N ratio all internal standard peaks ≥ 10?

V N N/A

**	Date	Lab ID/Reference	Internal Standard		% Recovery (Limit: 40-135%)	40-135%)	:	Qualifications
		(TS() !!	tþ.		33	40-136+	11/10	19. J. J. 18.
	11.0							
		ረ	H	39	6	40-12V	1777	(X X)
П								
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		-						
		7	Н	20	6			
		-				(
		_	Н	[A]	7			<u> </u>
		8	*	1-4			1 27 7	(/1 7/ <
			R)	1	4		V	_
			4	30				(< 40)
			14	32		(-	(A. A)
						(
\neg		A	-4-	A		(
1			*	·		(
						(
						(
		Internal Standards	Check Standard Used		III	Internal Standards		Check Standard Used
¥	13C-2,3,7,8-ŤCDF	:OF			13C-OCDD			
ю	¹³ C-2,3,7,8-TCDD	000		ᅶ	13C-1,2,3,4-TCDD			
ن	¹³ C-1,2,3,7,8-PeCDF	PecDF		نـ	13C-1,2,3,7,8,9-HxCDD	QQ		
اه	¹³ C-1,2,3,7,8-PeCDD	PecdD		Σ				
ш	¹³ C-1,2,3,4,7,8-HxCDF	3-HxCDF		z			1	
u.	¹³ C-1,2,3,6,7,8-HxCDD	знхсоо		ö				
Ö	13C-1,2,3,4,6,7,8-HpCDF	,8-HpCDF		a.			7	
Ξ	{	7,8-HpCDD					-	

VALIDATION FINDINGS WORKSHEET Internal Standards

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2nd Reviewer: Reviewer;_

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

# Date Lab DiReference Internal Standard W. Recovery (Limit, 40-1354) Calonical Constitutions Calonical Constitu		T	T	T	T	T	T	T	T	Т	T	T	Ť	T	T	T	T	T	Γ		Γ	T	T	Ť	T	T	T	T	=
Date Lab Directerance Internal Standard W. Recovery (Limit: 40-135%) W. Recovery (Limit: 40-135%) W. Levis	OCC C Qualifications	これにはおし	V	(外・// //	14 119	`l.`	1 6														Charle Standard Head								
Date Lab ID/Reference Internal Standard 1/4 Recovery (Limit 1/4 1/		7 - 5%	-					A	,	, ,									(al Standards							7	
Date Lab ID/Reference Inte	% Recovery (Limit: 40-	7	7	34	36	30	23))	<u> </u>))	Interna	13C-OCDD	╄	_	<u> </u>	ż	O.	Д.	
Date 1.0.2.3.7,8-TCDF 1.0.2.3.7,8-TCDF 1.0.1.2,3.7,8-PeCDF 1.0.1.2,3.7,8-PeCDF 1.0.1.2,3.7,8-PeCDF 1.0.1.2,3.7,8-PeCDF 1.0.1.2,3.7,8-PeCDF 1.0.1.2,3.4,7,8-HxCDI 1.0.1.2,3.4,7,8-HxCDI 1.0.1.2,3.4,7,8-HxCDI 1.0.1.2,3.4,7,8-HxCDI 1.0.1.2,3.4,7,8-HxCDI 1.0.1.2,3.4,7,8-HxCDI 1.0.1.2,3.4,7,8-HxCDI 1.0.1.2,3.4,6,7,8-HpC	Internal Standard	A	Q	٧)		(—														Check Standard Used								
13C.2.3.7,8-TCD 13C.1.2.3.7,8-Pe 13C.1.2.3.7,8-Pe 13C.1.2.3.7,8-Pe 13C.1.2.3.7,8-Pe 13C.1.2.3.7,8-Pe 13C.1.2.3.7,8-Pe	Lab ID/Reference	0																			ırnal Standards					JF.	00	CDF	
																					Inte	¹³ C-2,3,7,8-TCDF	¹³ C-2,3,7,8-TCDD	¹³ C-1,2,3,7,8-PeCDF	¹³ C-1,2,3,7,8-PeCDL	¹³ C-1,2,3,4,7,8-HxCl	13C-1,2,3,6,7,8-HxCl		

V:\Validation Worksheets\Dioxin90\(\text{INTST90a.wpd}\)

1DC #: 2495|2| SDG #: 26x COUN

VALIDATION FINDINGS WORKSHEET Compound Quantitation and Reported CRQLs

Page:

Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

N N N N

Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound? Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary).

		Qualifications	1.45七/4-01	VI .								>) (c)				
	,	Associated samples	m)			3		,	9		X			7 . 4					
	Finding Lause		∀ 11		H. K. N. A &		1 av 40-23-8	O A O C V Y H T			4- X			EMPC 1/21/42					
	Sample ID	ر ر			2			9		ð	X			∯ 1					
	Date																		
L	*	١							-		<u></u>					-			

Comments: See sample calculation verification worksheet for recalculations

LDC #:3485|2| SDG #: 588-000

VALIDATION FINDINGS WORKSHEET Overall Assessment of Data

Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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.

(Y, N/A) Was the overall quality and usability of the data acceptable?

Finding Associated Samples Ousiffications	×		1xest F. 4. 2-8 2-14	HOW DB-255	. M.O.P. & 6	+ ON DB-225		1、1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			H ON DB-225 8		18-5	
# Date Sample ID	↑ 	*	41 1×8	and Ho	1 H.T. K	a = (+ 2	Z X 0x20	70 11 4	4	W.O.F	A - A	4 JAN 198	1 1 0 5 H ON D	Commente

LDC#:<u>21495I21</u> SDG#: <u>See Cover</u>

VALIDATION FINDINGS WORKSHEET Field Duplicates

Page: of Reviewer: 9

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Á	Y	N	NA
	V	N	NA

Were field duplicate pairs identified in this SDG? Were target analytes detected in the field duplicate pairs?

	Concentra	tion (ng/Kg)	(≤50)	(ng/Kg)	(ng/Kg)	Qualifications
Compound	1	3	RPD	Difference	Limits	(Parent Only)
A	73.5	57.7	24			Notal
В	263	195	30			
С	213	152	33			
D ·	356	256	33	·		
E	417	286	37			
F	1240	940	28		·	
G	1360	1050	26			
Н	3510	4190	18			
1	2590	2860	10	·		
J	1440	1530	6			
К	6050	6620	9			
L	3330	3950	17			
N	559	725	26			
М	2070	2290	10		1	
0	11500	13600	17			
P	6040	6870	13			
Q.	32900	42800	26			
R	3340	2310	36			
s	2960	2200	29			
Т	2990	2230	29			
บ	1900	1500	24			
V.	33400	42600	24			
W	21100	23500	11			
x	25500	19400	27			
Υ	26300	24700	6			
H (DB-225)	1120	1380	21	ľ		

LDC#:<u>21495l21</u> SDG#:<u>See Cover</u>

VALIDATION FINDINGS WORKSHEET Field Duplicates

Page:	<u>>of≥</u>
Reviewer:	9
2nd Reviewer:	K

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Were field duplicate pairs identified in this SDG? Were target analytes detected in the field duplicate pairs?

·	Concentra	tion (ng/Kg)	(≤50)	(ng/Kg)	(ng/Kg)	Qualifications
Compound	2	4	RPD	Difference	Limits	(Parent Only)
А	81.0	77.5		3.5	(≤55.3)	*
В	251	206	,	45	(≤138)	
С	244	163		81	(≤138)	
D	404	315	·	89	(<u><</u> 138)	
Ε	448	357		91	(≤138)	
F .	1250	972	25			
G	1780	1240		540	(≤276)	idets & 14
Н	3810	5960	44			
1.	2150	2960	32			
J	1150	1490	26			
К	6930	8090	15			
L	3710	4990	29			
N	651	680		29	(≤138)	
М	2210	2670	19			
0	14600	17700	19			
Р	6710	7480	11			
Q	43200	55000	24			
H (DB-225)	881	1040	17			

LDC Report# 21495K21

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name:

Tronox LLC Facility, 2009 Phase B Investigation,

Henderson, Nevada

Collection Date:

June 26, 2009

LDC Report Date:

October 20, 2009

Matrix:

Soil

Parameters:

Dioxins/Dibenzofurans

Validation Level:

Stage 2B

Laboratory:

Columbia Analytical Services, Inc.

Sample Delivery Group (SDG): R0903584

Sample Identification

SA188-0.5B

SA172-0.5B

SA41-0.5B

SA41-0.5BDL

SA44-0.5B

SA42-0.5B

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 8290 for Polychlorinated Dioxins/Dibenzofurans.

This review follows the Standard Operating Procedures (SOP) 40, Data Review/Validation (BRC 2009), the Quality Assurance Project Plan Tronox LLC Facility, Henderson, Nevada (June 2009), NDEP guidance (May 2006), and USEPA Contract Laboratory Program National Functional Guidelines for Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) as there are no current guidelines for the method stated above.

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.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XIV.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- J+ Data are qualified as estimated, with a high bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J- Data are qualified as estimated, with a low bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J Data are qualified as estimated; it is not possible to assess the direction of the potential bias. False positives or false negatives are unlikely to have been reported.
- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- R Data are qualified as rejected. There is a significant potential for the reporting of false negatives or false positives.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- B The analytical result may be a false positive totally attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JB The analytical result may be biased high and partially attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JK The analytical result is an estimated maximum possible concentration (EMPC).
- X The analytical result is not used for reporting because a more accurate and precise result is reported in its place.
- J-TDS The analytical result is estimated based on failure of the Total Dissolved Solids (TDS) correctness check performed in accordance with the Standard Method 1030E.
- J-CAB The analytical result is estimated based on failure of the cation-anion balance correctness check performed in accordance with Standard Method 1030E.
- J-TDS & CAB The analytical result is unreliable based on the failure of the cation-anion balance and TDS correctness check performed in accordance with standard Method 1030E.
- 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.

Retention time windows were established for all homologues. The chromatographic resolution between 2,3,7,8-TCDD and peaks representing any other unlabeled TCDD isomer was less than or equal to 25%.

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 30.0% for labeled compounds.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

IV. Routine Calibration (Continuing)

Routine calibration was performed at the required frequencies.

All of the routine calibration percent differences (%D) between the initial calibration RRF and the routine calibration RRF were less than or equal to 20.0% for unlabeled compounds and less than or equal to 30.0% for labeled compounds.

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
EQ0900248-01	7/6/09	1,2,3,4,6,7,8-HpCDD OCDD OCDF Total HpCDD	0.126 ng/Kg 0.629 ng/Kg 0.177 ng/Kg 0.136 ng/Kg	All samples in SDG R0903584

Sample concentrations were compared to concentrations detected in the method blanks as required by the QAPP. No sample data was qualified.

No field blanks were identified in this SDG.

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 (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 recoveries were within QC limits with the following exceptions:

Sample	Internal Standards	%R (Limits)	Compound	Flag	A or P
SA44-0.5B	¹³ C-OCDD	33 (40-135)	OCDD	J (all detects) UJ (all non-detects)	Р
			OCDF	J (all detects) UJ (all non-detects)	

X. Target Compound Identifications

Raw data were not reviewed for this SDG.

XI. Project Quantitation Limit

All project quantitation limits were within validation criteria with the following exceptions:

Sample	Compound	Finding	Criteria	Flag	A or P
SA188-0.5B SA44-0.5B	1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	P
SA172-0.5B	OCDF	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	Р
SA41-0.5B	1,2,3,4,6,7,8-HpCDD 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 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	А

All compounds reported below the PQL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903584	All compounds reported below the PQL.	J (all detects)	А

All compounds reported as EMPC were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903584	All compounds reported as estimated maximum possible concentration (EMPC).	JK (all detects)	Α

Raw data were not reviewed for this SDG.

XII. System Performance

Raw data were not reviewed for this SDG.

*XIII. Overall Assessment of Data

The overall assessment of data was acceptable. In the case where more than one result was reported for an individual sample, the least technically acceptable results were rejected as follows:

Sample	Compound	Flag	A or P
SA188-0.5B SA172-0.5B SA44-0.5B SA42-0.5B	2,3,7,8-TCDF (DB-5)	x	А
SA41-0.5B	1,2,3,4,6,7,8-HpCDD 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 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x x x	А
SA41-0.5BDL	All TCL compounds except 1,2,3,4,6,7,8-HpCDD 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 0CDF 2,3,7,8-TCDF (DB-225)	X	A

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.

*Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Data Qualification Summary - SDG R0903584

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903584	SA44-0.5B	OCDD OCDF	J (all detects) UJ (all non-detects) J (all detects) UJ (all non-detects)	Р	Internal standards (%R) (i)
R0903584	SA188-0.5B SA44-0.5B	1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	Р	Project Quantitation Limit (e)
R0903584	SA172-0.5B	OCDF	J (all detects)	Р	Project Quantitation Limit (e)
R0903584	SA41-0.5B	1,2,3,4,6,7,8-HpCDD 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 2,3,7,8-TCDF (DB-5 & DB-225)	J (all detects)	А	Project Quantitation Limit (e)
R0903584	SA188-0.5B SA172-0.5B SA41-0.5B SA41-0.5BDL SA44-0.5B SA42-0.5B	All compounds reported below the PQL.	J (all detects)	A	Project Quantitation Limit (sp)
R0903584	SA188-0.5B SA172-0.5B SA41-0.5B SA41-0.5BDL SA44-0.5B SA42-0.5B	All compounds reported as EMPC	JK (all detects)	А	Project Quantitation Limit (k)
R0903584	SA188-0.5B SA172-0.5B SA44-0.5B SA42-0.5B	2,3,7,8-TCDF (DB-5)	х	А	Overall assessment of data (o)

SDG	Sample	Compound	Flag	A or P	Reason (Code)
*R0903584	SA41-0.5B	1,2,3,4,6,7,8-HpCDD 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 1,2,3,7,8,9-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 0CDF 2,3,7,8-TCDF (DB-5 & DB-225)	x x x x x x x x	А	Overall assessment of data (o)
R0903584	SA41-0.5BDL	All TCL compounds except 1,2,3,4,6,7,8-HpCDD 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 0CDF 2,3,7,8-TCDF (DB-225)	X	А	Overall assessment of data (o)

^{*}Corrected 2,3,7,8-TCDF for SA41-0.5B

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Laboratory Blank Data Qualification Summary - SDG R0903584

No Sample Data Qualified in this SDG

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Field Blank Data Qualification Summary - SDG R0903584

No Sample Data Qualified in this SDG

Tronox Northgate Henderson

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LDC #: 21495K21	VALIDATION COMPLETENESS WORK	(SHEET	Date: 9/17/05
SDG #: R0903584	Stage 2B	i i i i i i i i i i i i i i i i i i i	Page:/of_//
Laboratory: Columbia Analytica	Services		Reviewer:
			2nd Reviewer:
HETHAR LIDOCALIDAD Disch	no/Dibonzofurono /EDA SM/ 846 Method 8290\		

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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
1.	Technical holding times	4	Sampling dates: 6/26/09
11.	HRGC/HRMS instrument performance check	<u> </u>	/ / /
I II.	Initial calibration	4	
IV.	Routine calibration/ICV	A	
V.	Blanks	W	
VI.	Matrix spike/Matrix spike duplicates	N	gient spirited.
VII.	Laboratory control samples	A	205/3
VIII.	Regional quality assurance and quality control	N	
IX.	Internal standards	W	
X.	Target compound identifications	N	
XI.	Compound quantitation and CRQLs	ŹN_	
χίΪ.	System performance	N	
XIII.	Overall assessment of data	w	
XIV.	Field duplicates	N,	
XV.	Field blanks	ANN.	TBA=109-50 (\$0904016) C

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:

///	120,15						
1	SA188-0.5B	11	ZD0900248-01	21	P10385/	31	P203248 /
2 /	SA172-0.5B	12	7	د 22	P103864	32	P203279
3/2	SA41-0.5B	13		233	P103962 1	334	P203327
	SA41-0.5BDL	14		24	P1038[4 C	34	<u> </u>
5/3	SA44-0.5B	15		25	1	35	
8/	SA42-0.5B	16		26		36	
7		17		27_		37	
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes:	

VALIDATION FINDINGS WORKSHEET

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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	a.ocpf	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	1. 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:

SDG #: Ser COME LDC #: 249542

VALIDATION FINDINGS WORKSHEET

2nd Reviewer. Reviewer: Page:

> Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A". METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 8290)

Were all samples associated with a method blank?

Was a method blank analyzed for each matrix? Y)N N/A

Was the blank contaminated? If yes, please see qualification below.

Blank extraction date: 7/4/0 9 Blank analysis date: 7/9/0 Conc. units: Ind.

XSXV Sample Identification Associated Samples: 629 10-870 逐 Blank 10 0 78 0.177 Compound

Blank analysis date:_ Blank extraction date:_

Conc. units:

Associated Samples:

	Sample Identification							
Blank ID								
Blank ID								
Compound								

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".

LDC#OL

VALIDATION FINDINGS WORKSHEET Internal Standards

Reviewer: 2nd Reviewer: Page:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)
Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Are all internal standard recoveries were within the 40-135% criteria? Was the S/N ratio all internal standard peaks > 10?

Y N N/A

#	Date	Lab ID/Reference	Internal Standard	% R	% Recovery (Limit: 40-135%)	Cool : P. Qualifications
\vdash		8	H	33	(40-135)	~ (M + (F. R)
					(
			•		(
					()	
П			•		(
				-	(
					(
					()	
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					()	
					()	
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					()	
					()	
					()	
					(
		Internal Standards	Check Standard Used		Internal Standards	Check Standard Used
₹	13C-2,3,7,8-TCDF	CDF		1. 13	13C-OCDD	
В.	13C-2,3,7,8-TCDD	.cop		κ.	¹³ C-1,2,3,4-TCDD	
ပ		-PeCDF		L. 13	¹³ C-1,2,3,7,8,9-HxCDD	
Ö		-PeCDD		Σ		
ui	¹³ C-1,2,3,4,7,8-HxCDF	,8-HxCDF		ż		
u:	¹³ C-1,2,3,6,7,8-HxCDD	,8-HxCDD		Ö		
Ø	13C-1,2,3,4,6,7,8-HpCDF	,7,8-HpCDF		œ.		<u></u>
ŗ		,7,8-HpCDD				

V:\Validation Worksheets\Dioxin90\inTST90a.wpd

LDC #:>L

Compound Quantitation and Reported CRQLs VALIDATION FINDINGS WORKSHEET

Page:

2nd Reviewer: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound? Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary).

#	Date Sample ID	Finding range	Associated Samples	Qualifications
	7	H. K. C. O & &	4	(0) A 40
		(How Doth OB = ad OB-2x)] -
	d	X	1	
	<u>M</u>	F. H. I.J. K. Z.M.N	M	Lots 14 (P)
		10 P. & (+ m, Lett.		
		3-5		
	₩ ₩	zwe result	118	(A)

Comments: See sample calculation verification worksheet for recalculations

LDC #:2/495K2/ SDG #:26/ COUST

VALIDATION FINDINGS WORKSHEET Overall Assessment of Data

(of / Page:

2nd Reviewer: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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.

Was the overall quality and usability of the data acceptable? (Y)N N/A

L					
*	Date	Sample ID	Finding	Associated Samples	Qualifications
		1-2. \$ 5.6	2.3.7.8-720F on DB-S	12. X-15.6	~
					3
		8	T. W. H. J. H. W. M. O.	2	
			8 + (2B->X>+DR-E)		
			A1 2xept #. 4. 1. 4. F	4	
			L.M.N.O. F. R. B. and		
			# ON DB-22S		
		-			
Comments:	nemts:				

Laboratory Data Consultants, Inc. Data Validation Report

Project/Site Name:

Tronox LLC Facility, 2009 Phase B Investigation,

Henderson, Nevada

Collection Date:

July 6 through July 7, 2009

LDC Report Date:

September 27, 2009

Matrix:

Soil

Parameters:

Dioxins/Dibenzofurans

Validation Level:

Stage 2B

Laboratory:

Columbia Analytical Services, Inc.

Sample Delivery Group (SDG): R0903729

Sample Identification

SA206-0.5B RSAL4-0.5B RSAL4009-0.5B SA69-0.5B

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 8290 for Polychlorinated Dioxins/Dibenzofurans.

This review follows the Standard Operating Procedures (SOP) 40, Data Review/Validation (BRC 2009), the Quality Assurance Project Plan Tronox LLC Facility, Henderson, Nevada (June 2009), NDEP guidance (May 2006), and USEPA Contract Laboratory Program National Functional Guidelines for Polychlorinated Dioxins/Dibenzofurans Data Review (September 2005) as there are no current guidelines for the method stated above.

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.

Blank results are summarized in Section V.

Field duplicates are summarized in Section XIV.

Raw data were not reviewed for this SDG. The review was based on QC data.

The following are definitions of the data qualifiers:

- J+ Data are qualified as estimated, with a high bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J- Data are qualified as estimated, with a low bias likely to occur. False positives or false negatives are unlikely to have been reported.
- J Data are qualified as estimated; it is not possible to assess the direction of the potential bias. False positives or false negatives are unlikely to have been reported.
- U Indicates the compound or analyte was analyzed for but not detected at or above the stated limit.
- R Data are qualified as rejected. There is a significant potential for the reporting of false negatives or false positives.
- UJ Indicates the compound or analyte was analyzed for but not detected. The sample detection limit is an estimated value.
- B The analytical result may be a false positive totally attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JB The analytical result may be biased high and partially attributable to blank contamination. This qualifier is applicable to radiochemistry analysis only.
- JK The analytical result is an estimated maximum possible concentration (EMPC).
- X The analytical result is not used for reporting because a more accurate and precise result is reported in its place.
- J-TDS The analytical result is estimated based on failure of the Total Dissolved Solids (TDS) correctness check performed in accordance with the Standard Method 1030E.
- J-CAB The analytical result is estimated based on failure of the cation-anion balance correctness check performed in accordance with Standard Method 1030E.
- J-TDS & CAB The analytical result is unreliable based on the failure of the cation-anion balance and TDS correctness check performed in accordance with standard Method 1030E.
- 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.

Retention time windows were established for all homologues. The chromatographic resolution between 2,3,7,8-TCDD and peaks representing any other unlabeled TCDD isomer was less than or equal to 25%.

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 30.0% for labeled compounds.

The ion abundance ratios for all PCDDs and PCDFs were within validation criteria.

IV. Routine Calibration (Continuing)

Routine calibration was performed at the required frequencies.

All of the routine calibration percent differences (%D) between the initial calibration RRF and the routine calibration RRF were less than or equal to 20.0% for unlabeled compounds and less than or equal to 30.0% for labeled compounds.

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
EQ0900258-01	7/10/09	OCDD 1,2,3,4,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF OCDF Total HxCDF Total HpCDF	3.72 ng/Kg 0.730 ng/Kg 2.77 ng/Kg 11.5 ng/Kg 0.730 ng/Kg 2.77 ng/Kg	All samples in SDG R0903729

Sample concentrations were compared to concentrations detected in the method blanks as required by the QAPP. No sample data was qualified with the following exceptions:

Sample	Compound	Reported Concentration	Modified Final Concentration
SA69-0.5B	OCDD	4.70 ng/Kg	4.70U ng/Kg

Sample FB072109-SO (from SDG R0904016) was identified as a field blank. No polychlorinated dioxin/dibenzofuran contaminants were found in this blank with the following exceptions:

Field Blank ID	Sampling Date	Compound	Concentration	Associated Samples
FB072109-SO	7/21/09	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 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF Total HpCDD Total HxCDF Total HpCDF	8.37 pg/L 21.8 pg/L 6.33 pg/L 3.57 pg/L 4.04 pg/L 43.3 pg/L 19.6 pg/L 205 pg/L 8.37 pg/L 18.1 pg/L 72.1 pg/L	All samples in SDG R0903729

Sample concentrations were compared to concentrations detected in the field blanks as required by the QAPP. No sample data was qualified.

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 (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 recoveries were within QC limits with the following exceptions:

Sample	Internal Standards	%R (Limits)	Compound	Flag	A or P
EQ0900258-01	¹³ C-2,3,7,8-TCDD ¹³ C-1,2,3,7,8-PeCDF ¹³ C-1,2,3,4,7,8-HxCDF ¹³ C-1,2,3,6,7,8-HpCDF ¹³ C-1,2,3,4,6,7,8-HpCDD ¹³ C-1,2,3,4,6,7,8-HpCDD	36 (40-135) 37 (40-135) 37 (40-135) 39 (40-135) 24 (40-135) 21 (40-135) 6 (40-135)	2,3,7,8-TCDD Total TCDD 1,2,3,7,8-PeCDF 2,3,4,7,8-PeCDF Total PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,7,8,9-HxCDF Total HxCDF 1,2,3,4,7,8-HxCDD Total HxCDD 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF 1,2,3,4,7,8,9-HpCDF Total HpCDF 1,2,3,4,6,7,8-HpCDF Total HpCDF 1,2,3,4,6,7,8-HpCDD Total HpCDD CCDD CCDD CCDF 1,2,3,6,7,8-HxCDD	J (all detects) UJ (all non-detects)	Р

X. Target Compound Identifications

Raw data were not reviewed for this SDG.

XI. Project Quantitation Limit

All project quantitation limits were within validation criteria with the following exceptions:

Sample	Compound	Finding	Criteria	Flag	A or P
SA206-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF OCDF 2,3,7,8-TCDF (DB-5 & DB-225)	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects)	Р
RSAL4-0.5B	2,3,7,8-TCDF 1,2,3,4,6,7,8-HpCDF OCDF	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects) J (all detects) J (all detects)	Р
RSAL4009-0.5B	1,2,3,4,6,7,8-HpCDF OCDF	Sample result exceeded calibration range.	Reported result should be within calibration range.	J (all detects) J (all detects)	Р

All compounds reported below the PQL were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903729	All compounds reported below the PQL.	J (all detects)	А

All compounds reported as EMPC were qualified as follows:

Sample	Finding	Flag	A or P
All samples in SDG R0903729	All compounds reported as estimated maximum possible concentration (EMPC).	JK (all detects)	А

Raw data were not reviewed for this SDG.

XII. System Performance

Raw data were not reviewed for this SDG.

XIII. Overall Assessment of Data

The overall assessment of data was acceptable. In the case where more than one result was reported for an individual sample, the least technically acceptable results were rejected as follows:

Sample	Compound	Flag	A or P
All samples in SDG R0903729	2,3,7,8-TCDF (DB-5)	x	Α

Data flags are summarized at the end of this report if data has been qualified.

XIV. Field Duplicates

Samples RSAL4-0.5B and RSAL4009-0.5B were identified as field duplicates. No polychlorinated dioxins/dibenzofurans were detected in any of the samples with the following exceptions:

	Concentrat	Concentration (ng/Kg)		D.//		
Compound	RSAL4-0.5B	RSAL4009-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P
2,3,7,8-TCDD	5.51	4.50	-	1.01 (≤1.04)	-	-
1,2,3,7,8-PeCDD	18.4	15.7	16 (≤50)	-	-	-

	Concentrat	tion (ng/Kg)				
Compound	RSAL4-0.5B	RSAL4009-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P
1,2,3,4,7,8-HxCDD	13.9	12.1	-	1.8 (≤2.59)	-	-
1,2,3,6,7,8-HxCDD	26.4	23.2	13 (≤50)	-	-	-
1,2,3,7,8,9-HxCDD	28.5	24.4	16 (≤50)	-	-	-
1,2,3,4,6,7,8-HpCDD	90.5	76.9	16 (≤50)	•	-	-
OCDD	104	87.2	18 (≤50)	-	•	-
2,3,7,8-TCDF	234	202	15 (≤50)	-	-	-
1,2,3,7,8-PeCDF	208	171	20 (≤50)	-	-	-
2,3,4,7,8-PeCDF	96.8	86.0	12 (≤50)	-	-	-
1,2,3,4,7,8-HxCDF	492	415	17 (≤50)	-	-	-
1,2,3,6,7,8-HxCDF	274	243	12 (≤50)	•	-	-
1,2,3,7,8,9-HxCDF	45.9	36.8	22 (≤50)	-	-	-
2,3,4,6,7,8-HxCDF	167	135	21 (≤50)	-	-	-
1,2,3,4,6,7,8-HpCDF	990	822	19 (≤50)	-	-	-
1,2,3,4,7,8,9-HpCDF	458	386	17 (≤50)	-	-	-
OCDF	3030	2520	18 (≤50)	-	-	-
Total TCDD	165	150	10 (≤50)	-	-	-
Total PeCDD	201	179	12 (≤50)	•	-	-
Total HxCDD	211	184	14 (≤50)	-	-	-
Total HpCDD	141	121	15 (≤50)	-	-	-
Total TCDF	1450	1250	15 (≤50)	-	-	-

	Concentrati	on (ng/Kg)	555	Diff		
Compound	RSAL4-0.5B	RSAL4009-0.5B	RPD (Limits)	Difference (Limits)	Flag	A or P
Total PeCDF	1840	1510	20 (≤50)	-	-	-
Total HxCDF	2200	1850	17 (≤50)	-	-	-
Total HpCDF	2120	1770	18 (≤50)	-	-	-
2,3,7,8-TCDF (DB-225)	110	93.2	17 (≤50)	-	-	-

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Data Qualification Summary - SDG R0903729

SDG	Sample	Compound	Flag	A or P	Reason (Code)
R0903729	SA206-0.5B	1,2,3,7,8-PeCDF 1,2,3,4,7,8-HxCDF 1,2,3,6,7,8-HxCDF 1,2,3,4,6,7,8-HpCDF 1,2,3,4,7,8,9-HpCDF COCDF 2,3,7,8-TCDF (DB-5 & DB-225) J (all detects) J (all detects) J (all detects) J (all detects)		Р	Project Quantitation Limit (e)
R0903729	RSAL4-0.5B	2,3,7,8-TCDF 1,2,3,4,6,7,8-HpCDF OCDF	J (all detects) J (all detects) J (all detects)	Р	Project Quantitation Limit (e)
R0903729	RSAL4009-0.5B	1,2,3,4,6,7,8-HpCDF OCDF	J (all detects) J (all detects)	Р	Project Quantitation Limit (e)
R0903729	SA206-0.5B RSAL4-0.5B RSAL4009-0.5B SA69-0.5B	All compounds reported below the PQL.	J (all detects)	А	Project Quantitation Limit (sp)
R0903729	SA206-0.5B RSAL4-0.5B RSAL4009-0.5B SA69-0.5B	All compounds reported as EMPC	JK (all detects)	A	Project Quantitation Limit (k)
R0903729	SA206-0.5B RSAL4-0.5B RSAL4009-0.5B SA69-0.5B	2,3,7,8-TCDF (DB-5)	х	A	Overall assessment of data (o)

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Laboratory Blank Data Qualification Summary - SDG R0903729

SDG	Sample	Compound	Modified Final Concentration	A or P	Code
R0903729	SA69-0.5B	OCDD	4.70U ng/Kg	А	bl

Tronox LLC Facility, 2009 Phase B Investigation, Henderson, Nevada Dioxins/Dibenzofurans - Field Blank Data Qualification Summary - SDG R0903729

No Sample Data Qualified in this SDG

Tronox Northgate Henderson

		rionex northgate rionacioch	, ,
DC #:_	21495L21	VALIDATION COMPLETENESS WORKSHEET	Date: <u>4/17/29</u>
DG #:_	R0903729	Stage 2B	Page: /of/
aborato	ory: <u>Columbia Analytic</u>	al Services	Reviewer:
			2nd Reviewer:/
	D. LIDAA/LIDMA Dis.	ina (Dibanas frances (EDA OM) 040 Mathead 0000)	

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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
1.	Technical holding times	A	Sampling dates: 7/6-7/09
II.	HRGC/HRMS Instrument performance check	A	
	Initial calibration	♦	
IV.	Routine calibration/IO	A	
V.	Blanks	W	
VI.	Matrix spike/Matrix spike duplicates	\vee	dient sertied.
VII.	Laboratory control samples	A	100 P
VIII.	Regional quality assurance and quality control	N	\
IX.	Internal standards	W	
X.	Target compound identifications	N	
XI.	Compound quantitation and CRQLs	∕N_	
XII.	System performance	N	
XIII.	Overall assessment of data	w	
XIV.	Field duplicates	W	カ=2+3·
XV.	Field blanks	5N	7B07>109-50 (R0904016)

	بمغما	
- 11	iote:	

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	SA206-0.5B S	11/	ZQ0900258-01	21/	4219779	31	P203360
2	RSAL4-0.5B	12		د 22	D108991	32	
3	RSAL4009-0.5B	13		23		33	
4	SA69-0.5B	14		24		34	
5		15		25		35	
6		16		26		36	
7		17		27		37	Δ
8		18		28		38	
9		19		29		39	
10		20		30		40	

Notes:			

VALIDATION FINDINGS WORKSHEET

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

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. ocdb	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	1. 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:

SDG# 200 OU LDC #:3495C

VALIDATION FINDINGS WORKSHEET Blanks

2nd Reviewer:

Page: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA Method 8290)

Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Were all samples associated with a method blank? Y N N/A

|Y|N|N/A Was the blank contaminated? If yes, please see qualification below. Blank extraction date: $\frac{7/6/6}{3}$ Blank analysis date: $\frac{7/15/6}{3}$ Was a method blank analyzed for each matrix? Y N N/A

Sample Identification かると Associated Samples: 87.9 S & P 20.0 470/ N 4 730 730 3.72 0-856 Blank ID તું 680 Conc. units: 1/5/43 Compound

Blank analysis date: Blank extraction date:_ Conc. units:

Associated Samples:

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".

LDC #: 24952

SDG #:See Cover

VALIDATION FINDINGS WORKSHEET

Field Blanks

2nd Reviewer: Reviewer:

METHOD: HRGC/HRMS Dioxins/Dibenzofurans (EPA SW 846 Method 8290)

| Y N N/A Were field blanks identified in this SDG? | Blank units: __pg/L____ Associated sample units

ng/Kg Associated sample units:

Sampling date: 7/21/09

Field blank type: (circle one) Field Blank PRinsate / Other:	Field Blank	Rinsate / Other:	Associated Samples:	amples:	M	メダく)	\ \ \		
Compound	Blank ID			San	Sample Identification	ation			
	FB072109-SO	5X							
Ľ	8.37	0.04185							
9	21.8	0.109							
¥	6.33	0.03165							
7	3.57	0.01785							
M	4.04	0.0202							
0	43.3	0.2165							
۵	19.6	0.098							
Ø	205	1.025				-			
n	8.37	0.04185							
×	18.1	0.0905							
,	72.1	0.3605							
								The debugged by Colonian who has been designed by	
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	ARREST CONTRACTOR OF THE PROPERTY OF THE PROPE								
	A PART OF THE PROPERTY OF THE								
CROL									

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

VALIDATION FINDINGS WORKSHEET Internal Standards

rpc#河

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Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Are all internal standard recoveries were within the 40-135% criteria? Was the S/N ratio all internal standard peaks > 10? X N/A

1		I de ID/Deference	Internal Standard		% Recovery (Limit: 40-135%)	40-135%)	€od€: (Qualifications	
#	Date	r o 2000 XFX r o 1		26		40-135	JANG (A. R.)	
\dagger		12.05/00/00					-	
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		Internal Standards	Check Standard Used		nl	Internal Standards	Check Standard Used	
4	¹³ C-2.3.7.8-TCDF	ODF		-	13C-OCDD			
ď	13C-2 3 7 8-TCDD	CDD		K.	13C-1,2,3,4-TCDD			
i c	13C-1 2 3 7 8-PeCDF	PecDF		ن	¹³ C-1,2,3,7,8,9-HxCDD	CDD		
ا ا	13C-1,2,3,7,8-PeCDD	PeCDD		Σ				
ui	¹³ C-1.2.3,4,7,8-HxCDF	8-HXCDF		ż				
Li.	¹³ C-1,2,3,6,7,8-HxCDD	8-HxCDD		Ö				
ϋ	¹³ C-1,2,3,4,6,7,8-HpCDF	7,8-HpCDF		۵				
ī	¹³ C-1,2,3,4,6,7,8-HpCDD	7,8-HpCDD						

LDC #:21/45/2

Compound Quantitation and Reported CRQLs VALIDATION FINDINGS WORKSHEET

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Please see qualifications below for all questions answered "N". Not applicable questions are identified as "N/A".

Y N N Y

Were the correct internal standard (IS), quantitation ions and relative response factors (RRF) used to quantitate the compound? Compound quantitation and CRQLs were adjusted to reflect all sample dilutions and dry weight factors (if necessary).

Comments: See sample calculation verification worksheet for recalculations

VALIDATION FINDINGS WORKSHEET Overall Assessment of Data

Page:

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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.

Was the overall quality and usability of the data acceptable? Y N N/A

*	Date	Sample ID	Finding	Associated Samples	Qualifications
		411	2.3.T.8-120F M 18-5	M	(0) X
Comments:	ents:				

LDC#:21495L21 SDG#: See Cover

VALIDATION FINDINGS WORKSHEET Field Duplicates

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	`)		
/1	Υ	N	NA
- ($\overrightarrow{\nabla}$	N	ΝΔ
١.	.4	17	141

Were field duplicate pairs identified in this SDG? Were target analytes detected in the field duplicate pairs?

	Concentration (ng/Kg)		(≤50)	(ng/Kg)	(ng/Kg)	Qualifications
Compound	2	3	RPD	Difference	Limits	(Parent Only)
A	5.51	4.50		1.01	(≤1.04)	No Ceval
В	18.4	15.7	16			
С	13.9	12.1		1.8	(≤2.59)	
D	26.4	23.2	13			
E	28.5	24.4	16			
F	90.5	76.9	16			
G	104	87.2	18			
К	234	202	15			
	208	171	20			
J	96.8	86.0	12			
К	492	415	17			
L	274	243	12			
N	45.9	36.8	22			
М	167	135	21			
0	990	822	19			
Р	458	386	17			
Q	3030	2520	18			
R	165	150	10			
s	201	179	12			
Т	211	184	14			
U	141	121	15			
.V	1450	1250	15			
w	1840	1510	20			
Х	2200	1850	17			
Υ	2120	1770	18			
H (DB-225)	110	93.2	17			