ALLIANCE 284 Sheffield Street, Mountainside New Jersey 07092 NEW JERSEY LAB ID#: 20012: NEW YORK LAB ID#: 11376

GC/MS SEMI-VOLATILE ANALYSIS CONFORMANCE/NON-CONFORMANCE SUMMARY

CHEM	TECH PROJECT NUMBER: Q1215	MATRIX: TCLP		
METH	OD: 8270E/3510/1311			
1. 2.	Chromatograms Labeled/Compounds Identified. (Field samples GC/MS Tuning Specifications. DFTPP Meet Criteria. (NOTE THAT THERE ARE DIFFERENT CRITERIA FOR NY CLP AND NJ)		NO	YES ✓
3.	GC/MS Tuning Frequency - Performed every 24 hours for 600 s 8000 Series.	eries and 12 hours for		✓
4.	GC/MS Calibration - Initial Calibration performed within 30 day analysis and continuing calibration performed within 24 hours of 600 series and 12 hours for 8000 series.			✓
5.	GC/MS Calibration Requirements.			✓
	The Initial Calibration met the requirements . The Continuous Calibration File ID BF141336.D met the require Pentachlorophenol , Associated samples has no hit for this comp corrective action was required.			
6.	Blank Contamination - If yes, list compounds and concentrations	s in each blank:	✓	
7.	Surrogate Recoveries Meet Criteria If not met, list those compounds and their recoveries which fall or ranges.	outside the acceptable		~
8.	Matrix Spike/Matrix Spike Duplicate Recoveries Meet Criteria If not met, list those compounds and their recoveries which fall or range.	putside the acceptable		✓

The Blank Spike met requirements for all samples .

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<u>GC/MS SEMI-VOLATILE ANALYSIS CONFORMANCE/NON-CONFORMANCE SUMMARY</u> (CONTINUED)

		NA	NO	YES
9.	Internal Standard Area/Retention Time Shift Meet Criteria			\checkmark
	Comments:			
10.	Extraction Holding Time Met			\checkmark
	If not met, list number of days exceeded for each sample:			
11.	Analysis Holding Time Met			\checkmark
	If not met, list number of days exceeded for each sample:			

ADDITIONAL COMMENTS:

The Form 6 is not included in the data package because the Initial Calibration was performed using 7 points. Please use %D calculated based on Avg RF and CCRF for all compounds using Average Response Factor when the %RSD value for a compound is <20% for the Initial Calibration curve and use %D calculated based on Amount added and Calculated amount for all compounds using Linear Regression when the %RSD value for a compound is > 20% for the Initial Calibration curve for SW-846 analysis.

QA REVIEW

Date