## 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

MATRIX: TCLP

CHEMTECH PROJECT NUMBER: Q1235

METHOD: 8270E/3510/1311

METH	OD: 82/0E/3510/1311			
		NIA	NO	VEC
1.	Chromatograms Labeled/Compounds Identified. (Field samples and Method Blanks)	NA	NO	YES ✓
2.	GC/MS Tuning Specifications. DFTPP Meet Criteria. (NOTE THAT THERE ARE DIFFERENT CRITERIA FOR NY ASP CLP, CLP AND NJ)			✓
3.	GC/MS Tuning Frequency - Performed every 24 hours for 600 series and 12 hours for 8000 Series.			✓
4.	GC/MS Calibration - Initial Calibration performed within 30 days before sample analysis and continuing calibration performed within 24 hours of sample analysis for 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 requirements except for Pentachlorophenol, The associate samples have no positive hit for these compounds therefore no corrective action was taken.			
6.	Blank Contamination - If yes, list compounds and concentrations in each blank:		✓	
7.	Surrogate Recoveries Meet Criteria			$\checkmark$
	If not met, list those compounds and their recoveries which fall outside the acceptable ranges.			
8.	Matrix Spike/Matrix Spike Duplicate Recoveries Meet Criteria			✓
	If not met, list those compounds and their recoveries which fall outside the acceptable range.			
	The Blank Spike met requirements for all samples .			
9.	Internal Standard Area/Retention Time Shift Meet Criteria			✓
	Comments:			

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## GC/MS SEMI-VOLATILE ANALYSIS CONFORMANCE/NON-CONFORMANCE SUMMARY (CONTINUED)

NA NO YES

10.	Extraction Holding Time Met	✓
	If not met, list number of days exceeded for each sample:	
11.	Analysis Holding Time Met	✓
	If not met, list number of days exceeded for each sample:	
ADDIT	TIONAL COMMENTS:	
The For	orm 6 is not included in the data package because the Initial Calibration was performed using 7 I	points.
Please u	use %D calculated based on Avg RF and CCRF for all compounds using Average Response Fa	ctor when the
%RSD	O value for a compound is <15% for the Initial Calibration curve and use %D calculated based or	n Amount added
and Cal	alculated amount for all compounds using Linear Regression when the %RSD value for a compo	ound is > 15% for
the Initi	itial Calibration curve for SW-846 analysis.	
QA RE	EVIEW Date	