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Validating Chlorinated Herbicides GC, SW-846, Method 8151A



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	Annual Review	
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YES NO N/A 1.0 Traffic Reports and Laboratory Narrative 1.1 Are Traffic Report Forms present for all samples? \square ACTION: If no, contact lab for replacement of missing or illegible copies. 1.2 Do the Traffic Reports or SDG Narrative indicate any problems with sample receipt, condition of the samples, analytical problems or special circumstances affecting the quality of the data? \square ACTION: If any sample analyzed as a soil, other than TCLP, contains 50%-90% water, all data should be qualified as estimated (J). If a soil sample, other than TCLP, contains more than 90% water, all data should be qualified as unusable (R). ACTION: If samples were not iced (4°C) upon receipt at the laboratory, flag all positive results "J" and all non-detects "UJ". 2.0 **Holding Times** 2.1 Has the technical holding times, determined from date of sample receipt to date of extraction. been exceeded? П Note: Samples may be analyzed for herbicide ester and acid. Check Laboratory SDG Narrative. Note: Aqueous samples must be extracted within 7 days. Extracts must be analyzed within 40 days following extraction. Soil/Concentrated Waste samples must be extracted within 14 days and extracts analyzed within 40 days following extraction. ACTION: If technical holding times are exceeded, flag all positive results and non-detects(U)as estimated ("J") and document in the narrative that holding times were exceeded. Samples extracted more than 28 days from sample receipt, either on the first analysis or

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YES NO N/A

upon re-analysis, flag all positive results as Estimate ("J") and non-detects as unusable (R)

		Estimate ("J") and non-detects as unusable (R).			
3.0	Surrogate				
3.1		erbicide Surrogate Recovery Summaries quivalent) present for each of the natrices?			
	a. A	aqueous	Ц	_	_
	b. S	Soil	Ц	_	
3.2		samples listed on the appropriate Recovery Summary for each of the natrices?			
	a. A	aqueous	П	_	_
	b. S	Soil/Concentrated Waste	Ш	_	_
	ACTION:	Contact lab for explanation/resubmittals. If missing deliverables are unavailable, document effect in data assessments.			
3.3	Were outli	ers marked correctly with an asterisk?	Ц	_	_
	ACTION:	Circle all outliers with red pencil.			
	Note: reco	mmend surrogate is 2,4-Dichlorophenylacetic acid (DCAA)			
3.4	Did the laboration limits/recover	oratory provide their developed in-house QC veries?	Ц	_	_
	ACTION:	If no, use 70 -130% recovery to qualify data			
	ACTION:	No qualification is done if the surrogate is diluted out. If recovery for the surrogate is below the QC limit, but above 10%, flag all results for that sample "J". If recovery is < 10%, qualify postive results "J" and flag non-detects "R". If recovery is above the QC limits limit, qualify positive values "J".			

4.0

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YES NO N/A

	Note:	In-house QC limits must be examined for reasonableness, e.g. 10-170% may be appropriate for analytes not present in the sample.			
	Note:	Matrix effect is indicated if the LCS data are within limits but surrogate data exceeds QC limits.			
3.5		ogate retention times (RT) within the stablished during the initial 5-point analysis?	Ц	_	
	ACTION:	If the RT limits are not met, the analysis may be qualified unusable (R) for that sample on the basis of professional judgement.			
3.6		any transcription/calculation errors aw data and Form II/Equivalent?	_	Ц	
	ACTION:	If large errors exist, call lab for explanation/resubmittal. Make any necessary corrections and document effect in data assessments.			
	Matrix Spil	kes (Form III/Equivalent)			
4.1		ix Spike/Matrix Spike Duplicate Form (Form III/Equivalent) present?	Ц	_	
4.2		ix spikes analyzed at the required for each of the following matrices?			
	spike and	minimum, analysis of at least one matrix one duplicate unspiked sample or one matrix ix spike duplicate pair with each batch of amples.			
	a. A	queous	Ц	_	_
	b. S	oil/Concentrated Waste	Ц	_	_
	ACTION:	If any matrix spike data are missing, take the action specified in 3.2 above.			

			YES NO	N/A	
4.3	Did the labo	pratory provide their developed in-house ecoveries?	Ц	_	
	ACTION:	If no, use 70 -130% recovery to qualify data			
	ACTION:	No action is taken on MS/MSD data alone. However, using informed professional judgement, the data reviewer may use the matrix spike results in conjunction with other QC criteria (e.g. LCS) to determine the need for qualification of the data.			
5.0	Blanks (Fo	orm IV/Equivalent)			
5.1	Is the Meth	nod Blank Summary (Form IV) present?	Ц		_
5.2	blank beer samples o	of Analysis: has a reagent/method analyzed for each SDG or every 20 f similar matrix or concentration traction batch, whichever is more	Ш	_	
	ACTION:	If any blank data are missing, take the action specified above in 3.2. If blank data is not available, reject (R) all associated positive data. However, using professional judgement, the data reviewer may substitute field blank data for missing method blank data.			
5.3		bicide instrument blank been analyzed nning of every analytical sequence of s?	Ц	_	
AC	exp deli	ny blank data are missing, call lab for lanation/resubmittals. If missing verables are unavailable, document the ct in data assessments.			
5.4		graphy: review the blank raw data - rams, quant reports or data system			
		matographic performance (baseline or each instrument acceptable for			

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YES NO N/A

Herbicides?

 \square

ACTION: Use professional judgement to determine

the effect on the data.

Contamination 6.0

"Water blanks", "distilled water blanks" and NOTE:

"drilling water blanks" are validated like any other sample and are not used to qualify the data. Do not confuse them with the other QC

blanks discussed below.

YES NO N/A

6.1	6.1 Do any method/instrument/reagent/cleanup blanks have positive results for Herbicides? When applied as described in table below, the contaminant concentration in the method blank is multiplied by the sample dilution factor and corrected for % moisture when necessary.				
6.2	Do any fie Herbicides	ld/rinse blanks have positive s results?	_	Ц	_
	ACTION:	Prepare a list of the samples associated with each of the contaminated blanks. (Attach a separate sheet)			
	NOTE:	All field blank results associated to a particular group of samples (may exceed one per case or one per day) may be used to qualify data. Blanks may not be qualified because of contamination in another blank. Field blanks must be qualified for surrogate, calibration, or any QC problems.			
0	ACTION:	Follow the directions in the table below to qualify TCL results due to contamination. Use the largest value from all the associated blanks.			
	conc > CRC conc < CRC				
·		·			
but <	5x blank	is < 5x blank value & > 5x blank value			
	nple result	No modification			
Report C	RQL &	No qualification			
with a	a "U";	qualify "U" is needed			
NOTI	in t	ross blank contamination exists, all data he associated samples should be qualified unusable (R).			
6.3	Are there with every	field/rinse/equipment blanks associated sample?	П	_	_
ACTI		low level samples, note in data assessment there is no associated field/rinse/equipment blank.			

YES NO N/A

Exception: samples taken from a drinking water tan

		not have associated field blanks.			
7.0	Calibration				
7.1	printouts f	as Chromatograms and Data Systems or both columns present for all samples, C Check references, and matrix spikes?	Ц	_	
	ACTION:	If no, take action specified in 3.2 above.			
7.2		VI/Equivalent present and complete olumn and each analytical sequence?	Ц	_	_
	ACTION:	If no, take action specified in 3.2 above.			
7.3		any transcription/calculation errors aw data and Forms VI?	_	Ц	_
	ACTION:	If large errors exist, call lab for explanation/resubmittal, make necessary corrections and document effect in data assessments.			
7.4	average a measurem of the abs	retention time windows calculated using the bsolute retention time (at least three nents) <u>+</u> three times the standard deviation olute retention time, for each standard? Method 8000A, section 7.5).	[]		
7.5.	,	S check standard analyzed prior to		_	
		ental samples?	Ц	_	_
7.5.1	If yes, was	s the surrogate recovery >50%?	Ц	_	_
7.5.2	if surroga	CS check standard re-extracted/re-analyzed, te recovery was <50%, or any one analyte %, or two analytes < 70% ?	П	_	
	Action:	If No/' to any of the above, then qualify positive hits as estimated "J" and non-detects as rejected "R" in the original analysis of all samples in the associated analytical sequence.			

YES NO N/A 7.6 Do all standard retention times, including each Herbicides in each level of Initial Calibration fall within the windows established during the initial calibration analytical sequence? (For Initial Calibration Standards, Form VI/Equivalent - Herbicides - 1). [] ACTION: If no, all samples in the entire analytical sequence are potentially affected. Check to see if the chromatograms contain peaks within an expanded window surrounding the expected retention times. If no peaks are found and the surrogate is visible, nondetects are valid. If peaks are present and cannot be identified through pattern recognition or using a revised RT window, qualify all positive results and non-detects as unusable (R). 7.7 Are the linearity criteria for the Initial Calibration analyses within limits for both columns? (% RSD must be < 20.0% for all analytes). [] If no, qualify all associated positive ACTION: results generated during the entire analytical sequence "J" and all nondetects "UJ". When RSD >90%, flag all non-detect results for that analyte R (unusable). 7.8 Are there any transcription/calculation errors between raw data and Form VII - Herbicides-2? П ACTION: If large errors exists, call lab for explanation/resubmittal, make any necessary corrections and document effect in data assessments. 7.9 Is the resolution between any two adjacent peaks in the QC Reference Check Mixture > 60.0% for both columns? (Form VI-Herbicides- 4) \square

YES NO N/A

	ACTION:	If no, positive results for compounds that were not adequately resolved should be qualified "J". Use professional judgement to determine if non-detects which elute in areas affected by co-eluting peaks should be qualified "N" as presumptive evidence of presence or unusable (R).			
7.10		I -Continuing Calibration present and or each analytical sequence for both	Ц	_	_
	ACTION:	If no, take action as specified in 3.2 above.			
7.11		amples been injected within a 24 hr. inning with the injection of the first	П	_	
	ACTION:	If no, use professional judgement to determine the severity of the effect on the data and qualify accordingly.			
7.12	the Mid-co fall within t	yte retention times for ncentration Check standard (Form VII Herb-2) he windows established by the initial sequence?	П	_	
	ACTION:	If no, beginning with the samples which followed the last in-control standard, check to see if the chromatograms contain peaks within an expanded window surrounding the expected retention times. If no peaks are found and the surrogates are visible, non-detects are valid. If peaks are present			
		and cannot be identified through pattern recognition or using a revised RT window, qualify all positive results and non-detects as unusable (R).			

YES NO N/A 7.13 Are RPD values for all verification calibration standard compounds < 25.0% [] ACTION: The "associated samples" are those which followed the last in-control standard up to the next passing standard containing the analyte which failed the criteria. If %D is 25 -50% qualify as "J" If %D is 51-100% qualify as "NJ" If %D is >100% qualify as "R" If %D is >100% with visible interferences/qualify as "JN" 8.0 Analytical Sequence Check (Form VIII) 8.1 Is Form VIII present and complete for each column and each period of analyses? П ACTION: If no, take action specified in 3.2 above. 8.2 Was the proper analytical sequence followed for each initial calibration and subsequent analyses? (see SAS Client Request/section 8/paragraph 6) If no, use professional judgement to ACTION: determine the severity of the effect on the data and qualify it accordingly. Generally, the effect is negligible unless the sequence was grossly altered or the calibration was also out of limits. 9.0 Herbicides Identification 9.1 Is Form X complete for every sample in which a Herbicide was detected? []If no, take action specified in 3.2 above. ACTION: 9.2 Are there any transcription/calculation errors between raw data and Form X. \square

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YES NO N/A ACTION: If large errors exist, call lab for explanation/resubmittal, make necessary corrections and note errors in data assessment. 9.3 Are retention times (RT) of sample compounds within the established RT windows for both columns? [] Was GC/MS confirmation provided instead of confirmation by a second dissimilar column? Qualify as unusable (R) all Action: positive results which were not confirmed by second GC column analysis or by GC/MS. Also qualify as unusable (R) all positive results not meeting RT window unless associated standard compounds show a similar RT shift. The reviewer should use professional judgement to assign an appropriate quantitation limit. 9.4 Is the percent difference (% D) calculated for the positive sample results on the two GC columns < 25.0%? [] ACTION: If the reviewer finds neither column shows interference for the positive hits, the data should be flagged as follows: % Difference Qualifier 25-50 % J JN 50-90 % > 90 % R NOTE: The lower of the two values is reported on Form I. If using professional judgement, the reviewer determines that the higher result was more acceptable, the reviewer should replace the value and indicate the

reason for the change in the data assessment.

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YES NO N/A 9.5 Check chromatograms for false negatives. Were there any false negatives? П ACTION: Use professional judgement to decide if the compound should be reported. 10.0 Compound Quantitation and Reported Detection Limits 10.1 Are there any transcription/calculation errors in Form I results? Check at least two positive values. Were any errors found? \square NOTE: The reviewer should use professional judgement to decide whether a much larger concentration obtained on one column versus the other indicates the presence of an interfering compound. If an interfering compound is indicated, the lower of the two values should be reported and qualified as presumptively present at an approximated quantity (NJ). This necessitates a determination of an estimated concentration on the confirmation column. The narrative should indicate the presence of interferences during the evaluation of the second column confirmation. Are the CRQLs adjusted to reflect sample dilutions 10.2 and, for soils, % moisture? П ACTION: If errors are large, call lab for explanation/resubmittal, make any necessary corrections and document effect in data assessments. ACTION: When a sample is analyzed at more than one dilution, the lowest CRQLs are used (unless a QC exceedance dictates the use of the higher CRQL data from the diluted sample analysis). Replace concentrations that exceed the calibration range in the original analysis by crossing out the "E" value on the original Form I and substituting it with data from the analysis of diluted sample. Specify which Form I is to be used, then draw a red "X" across the entire page of all Form I's that should not be used, including any in the summary package.

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			YES	NO	N/A
	ACTION:	Quantitation limits affected by large, off-scale peaks should be qualified as unusable (R). If the interference is on-scale, the reviewer can provide an approximated quantitation limit (UJ) for each affected compound.			
10.3	quantitation	ata (Forms and associated chromatograms and n reports) been submitted for original, e-extraction/re-analysis samples?	Ц	_	_
11.0	Chromatog	gram Quality			
11.1	Were base	lines stable?	П	_	_
11.2	•	electropositive displacement beaks) or unusual peaks seen?	_	Ц	_
	ACTION:	Address comments under System Performance of data assessment. Explain use of professional judgement where used to qualify data.			

YES NO N/A

12.0 <u>Field Duplicates</u>

12.1 Were any field duplicates submitted for Herbicides analysis?

Ш _ _

Note: Check whether SAS Client Request required

field duplicates.

ACTION: Compare the reported results for

field duplicates and calculate the

relative percent difference.

ACTION: Any gross variation between field

duplicate results must be addressed in the reviewer narrative. However, if large differences exist, identification of field duplicates should be confirmed

by contacting the sampler.