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FEDERAL EXPRESS



Mr. John E. Kieling Chief Hazardous Waste Bureau New Mexico Environment Department 2905 Rodeo Park Dr. East, Bldg 1 Santa Fe, New Mexico 87505

Subject: Submittal of Updated Reference Documents Cited in the Mixed Waste Landfill Long-Term

Monitoring and Maintenance Plan for Sandia National Laboratories, New Mexico,

Environmental Protection Agency Identification Number NM5890110518

Dear Mr. Kieling:

The Department of Energy/National Nuclear Security Administration and National Technology and Engineering Solutions of Sandia, LLC are submitting the enclosed updated reference document to the New Mexico Environment Department. This submittal is required within 30 days of the effective date of the updated documents, which is June 19, 2017.

This submittal is comprised of one procedure used by personnel to perform validation of analytical data for the Mixed Waste Landfill. The updated reference document is:

AOP 00-03 Data Validation Procedure for Chemical and Radiochemical Data

Revisions include updates to keep the reference document current and to reflect ongoing modifications and improvements in industry practices.

If you have questions, please contact Steven Black of our staff at (505) 845-6885.

Sincerely

James W. Todd

Assistant Manager for Engineering

Enclosure cc: See Page 2

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Submittal of Updated Reference Document Cited in the Mixed Waste Landfill Long-Term Monitoring and Maintenance Plan

Sandia National Laboratories Albuquerque, New Mexico EPA ID No. NM5890110518

CERTIFICATION STATEMENT

I certify under penalty of law that this document and all attachments were prepared under my direction or supervision according to a system designed to ensure that qualified personnel properly gather and evaluate the information submitted. Based on my inquiry of the person or persons who manage the system, or those persons directly responsible for gathering the information, the information submitted is, to the best of my knowledge and belief, true, accurate, and complete. I am aware that there are significant penalties for submitting false information, including the possibility of fine or imprisonment for knowing violations.

Jaime L. Moya, Director Chief of Safety

National Technology & Engineering Solutions of Sandia, LLC

Albuquerque, New Mexico

Operator

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U.S. Department of Energy

National Nuclear Security Administration

Sandia Field Office (SFO)

Owner

Date signed 2017

Enclosure A

Updated Reference Document Cited in the Mixed Waste Landfill Long-Term Monitoring and Maintenance Plan

AOP 00-03 Data Validation Procedure for Chemical and Radiochemical Data

June 2017

Sandia National Laboratories EPA ID No. NM5890110518

		Procedure No:	AOP 00-03
Sand	lia National Laboratories	Issue No: Effective Date:	05 06/19/2017
DATA VALI	DATION PROCEDURE FOR CHEN DATA	MICAL AND RAD	IOCHEMICAL
	SAMPLE MANAGEMEN	T OFFICE	
Revised by:	May Donward M. Donivan Analytical Quality Associates, Inc.		06/08/2017 Date
Reviewed by:	K. Lambert CRM Quality Assurance Specialis	t	06-13 2017 Date
Reviewed by:	M. Moore Waste Characterization Project Le	ader	6/13/2017 Date
Reviewed by:	C. White SMO Operations Manager		06-13-17 Date
Owner and	5-187		1 /4/-

Approved by:

S. Shanks

ASP Program Leader

Table of Contents

			O ABBREVIATIONS	
ME.	ASURE	EMENT	S AND SYMBOLS	xiii
1	וווס	DDASE	<u> </u>	1
2			ND OWNERSHIP	
4	2.1			
		_	rship	
•			A	
3			SIBLE INDIVIDUALS AND ORGANIZATIONS	
	3.1		am Leader	
	3.2		Operations Manager	
	3.3	Comp	liance & Requirements Management (CRM) Staff	2
	3.4	SMO	Project Coordinator	2
	3.5	SMO	Staff	3
	3.6	Labora	atory Oversight/Data Validation Contractor	3
4			U RÉ	
			al	
		4.1.1	CVR	
		4.1.2	QC Exemptions	
		4.1.3	Holding Times	
		4.1.4	Preservation (chemical and temperature)	
		4.1.5	Calibration Points	
		4.1.6	Calibration for QC Samples	
		4.1.7	Blank Hierarchy	
		4.1.8	Blank Normalization	
		4.1.9	Field Duplicates	
		4.1.10	Sample-Specific External Standard Recovery	10
		4.1.11 4.1.12	Rounding Rules and Significant Figures	ll
		4.1.12	Special Laboratory Flags	
		4.1.13	Calculations	
		4.1.15	Reanalysis	
		4.1.16	Manual Integration	
		4.1.17	Failed Batch QC	
		4.1.18	MS/MSD, LCS/LCSD and Replicates	
		4.1.19	MS/MSD	
		4.1.20	MS/MSD with Elevated Analyte Concentration Requirements	14

IMPORTANT NOTICE:

	4.1.21	Blank Qualification with QC Failures	14
	4.1.22	Initial Dilutions	
	4.1.23	Reporting Limit Verification (RLV)	
	4.1.24	Filtered Samples	
	4.1.25	Sample Contamination	
4.2	Proced	ure for Gas Chromatography/Mass Spectrometry (GC/MS)	
			16
, ,,,	4.2.1	Instrument Tuning for GC/MS	
	4.2.2	Calibration	
	4.2.3	Calibration Verification	
	4.2.4	Blanks	
	4.2.5	Surrogate Recovery	
	4.2.6	IS Performance	
	4.2.7	MS/MSD	28
	4.2.8	Replicate	30
	4.2.9	LCS	31
	4.2.10	Sample Carry-over	33
	4.2.11	Dilutions	34
	4.2.12	Mass Spectra Acceptability	35
	4.2.13	TICs	36
	4.2.14	Method-specific Analytical Requirements-Organic GC/MS	37
4.3	Proced	ure for GC and HPLC Validation	64
	4.3.1	Calibration	64
	4.3.2	Calibration Verification	67
	4.3.3	Blanks	70
	4.3.4	Surrogate Recovery	72
	4.3.5	Internal Standard Performance	74
	4.3.6	MS/MSD	75
	4.3.7	Replicate	77
	4.3.8	LCS	77
	4.3.9	TAL Compound Identification	79
	4.3.10	Sample Carry-over	80
	4.3.11	Dilutions	
	4.3.12	Quantification and Confirmation	
	4.3.13	Method-specific Analytical Requirements-Organic GC and HPLC	85
4.4	Proced	ure for Liquid Chromatography/Mass Spectrometry/Mass	
Spe	ctrometr	y (LC/MS/MS) Validation	90
•	4.4.1	Instrument Calibration for LC/MS/MS	
	4.4.2	Calibration Verification	92
	4.4.3	RLV	95

		4.4.4	Blanks	96
		4.4.5	Surrogate Recovery – HE analysis only	98
		4.4.6	IS Performance	100
		4.4.7	MS/MSD	101
		4.4.8	Replicate	103
		4.4.9	LCS	104
		4.4.10	Sample Carry-over	105
		4.4.11	Dilutions	106
		4.4.12	Perchlorate Chlorine Ratios	
		4.4.13	Perchlorate Interference Check Standard	
		4.4.14	Method-specific Analytical Requirements – Organic LC/MS/MS	
	4.5		ation Guidelines for Confirmation by LC/MS/MS	
		4.5.1	Required LC/MS/MS Data	
		4.5.2	LC/MS/MS QC are Acceptable	
		4.5.3	Method Blank	
		4.5.4	Continuing Calibration	112
		4.5.5	PS/CRI	113
		4.5.6	IS Performance	
	4.6	Proced	dure for Inorganic Data Validation	114
		4.6.1	Initial Calibration	
		4.6.2	CCV	116
		4.6.3	Blanks	117
		4.6.4	MS	119
		4.6.5	Replicate	120
		4.6.6	LCS	121
		4.6.7	RLV	
		4.6.8	Method-specific analytical requirements (inorganic)	123
	4.7	Proced	dure for Radiochemical Analyses Validation	136
		4.7.1	Quantification	136
		4.7.2	Blanks	137
		4.7.3	Sample-Specific Chemical/Tracer Recovery	138
		4.7.4	MS	139
		4.7.5	Replicate	140
		4.7.6	LCS	141
		4.7.7	Instrument Control Charts	
		4.7.8	Method-Specific Analytical Requirements – Radiochemical	147
5	DA	TA VA	LIDATION REPORTS	149
	5.1	Sampl	e Findings Summary and Validation EDD Files	149
	5.2		Validation Narrative (format may vary by project)	
	5.3		Qualification Summary	

IMPORTANT NOTICE:

151
151
151
151
152
153
155
159
161

List of Appendices

Appendix A: Sample Preservation and Holding Times

Appendix B: Data Reporting Requirements

Appendix C: Surrogate Recovery Limits

Appendix D: GC/MS Internal Standards

Appendix E: Laboratory Control Limits

Appendix F: Mass Spectra Acceptability

Revision History

Revision	Effective Date	Summary of Changes
0	12/21/1999	New document
1	12/08/2003	Update to reflect current administrative changes
2	7/16/2007	Update to reflect current administrative changes
3	5/16/2011	EPA Method updates and references. Process changes for the automatic upload of the data validation qualifiers and the necessary QC steps associated with the new data processing procedure.
4	6/16/2014	Update to responsibilities of SMO personnel, and EPA methods and references. Clarified data validation requirements associated with mass spectra acceptability and TIC evaluation. Added language that states an MS/MSD is not required for an isotope dilution analysis. Included additional method-specific analytical requirements for Organic LC/MS/MS (PPCP) and Inorganic Anions by Ion Chromatography.
		Acronym list updated.
		A global change from RF to RRF was made throughout the document.
5		Section 4.1.3; Current NFG criteria for VOCs were updated, no marginal exceedance (>1X but ≤2X) will be allowed. "Gross exceedance" defined as >2X HT infraction for other organic parameters. The qualification for pH remains "J."
		Section 4.1.4; Cyanide, SVOCs, pesticides and PCBs added to table for specific temperature infractions. Numeric temperature infractions specified.
	06/19/2017	Section 4.1.23, Reporting Limit Verification; Added "reprocessing the RLV against the new curve."
		Section 4.2.2; RRF minimum limits identified as 0.010 and 0.050 (instead of 0.01 and 0.05). Linear Curves; added an additional intercept calculation. The coefficient of determination QC limit was changed from \geq 0.99 to \geq 0.990.
	The state of the s	Section 4.2.2, Linear Curves, added an additional intercept calculation.
		Section 4.2.4, Blanks: Scenarios removed and table added that reflects current NFG criteria.
		Section 4.2.5, Surrogate Recovery: Only 1 SVOC surrogate outlier per fraction is necessary for qualification, per NFG.

IMPORTANT NOTICE:

Section 4.2.6, Internal Standards; 20% (formerly 25%) is the lower limit for qualifying non-detects "R," per NFG.

Section 4.2.7, MS/MSD; Non-detects are now rejected for MS/MSD recoveries <20%, per NFG.

Section 4.2.9, LCS; Non-detects are now rejected for LCS recoveries <20%, per NFG. Also, clarified the criteria for LCS reanalysis.

Section 4.2.11, Dilutions; Replaced instrument response with concentration.

Section 4.2.14.2; Added program manager notification if extract cleanup was not performed.

Section 4.2.14.3; 8280B method references updated from those in 8280A. Internal Standards, 20% (formerly 25%) is the lower limit for qualifying non-detects "R," per NFG.

Section 4.2.14.4, 8290A method references updated from those in 8290. CCV section was revised for clarity based on QSM and input from Cape Fear Analytical. Added client notification if extract cleanup was not performed.

Section 4.2.14.5; removed references to TO-14A. Corrected RRF evaluation criteria for TO-15 and added discussion of surrogate recovery.

Section 4.2.14.6; added 1668C. OPR lower recovery limit <20%, per NFG. Added program manager notification if % solids, particle size reduction or extract cleanup was not performed, also if mass spectrometer performance was not evaluated at required frequency. Removed the bias from qualifiers for labeled cleanup standard infractions.

Section 4.2.14.7; LCS and QC check sample lower recovery limit <20%, per NFG. Added program manager notification if extract cleanup was not performed or mass spectrometer performance was not evaluated at required frequency.

Section 4.2.14.8; Added program manager notification if extract cleanup was not performed, mass spectrometer performance was not evaluated at required frequency, GC column performance was not evaluated at required frequency or 2,3,7,8-TCDF was detected and not confirmed.

Section 4.3.1, Calibration; Removed discussion of intercepts when results are reported at PQL. Added RRF as an ICAL option.

Section 4.3.2, Calibration Verification; Added acceptance criteria for RRFs.

Section 4.3.3, Blanks; Scenarios removed and table added that reflects current NFG criteria.

Section 4.3.5, NEW SECTION; Added Internal Standard

IMPORTANT NOTICE:

Performance. All subsequent sections of 4.3 were renumbered.

Section 4.3.6, MS/MSD; Non-detects are now rejected for MS/MSD recoveries <20%, per NFG.

Section 4.3.8, LCS; Non-detects are now rejected for LCS recoveries <20%, per NFG. Also, clarified the criteria for LCS reanalysis.

Section 4.3.11, Dilutions; Replaced instrument response with concentration.

Section 4.3.12, Quantitation and Confirmation; Added caveat that confirmation results shall not be reported from the secondary column in HPLC HE analysis. Information regarding the use of alternate columns was also added.

Section 4.3.13.4, 8310 PAH; Replaced "two columns" with "two detectors."

Section 4.4, LC/MS/MS Validation; Updated to reflect acceptance criteria for HE by LC/MS/MS using Method 8330B.

Section 4.4.1, Calibration; Removed discussion of intercepts when results are reported at PQL. RRF minimum limits identified as 0.010 and 0.050 (instead of 0.01 and 0.05).

Section 4.4.2, Calibration Verification; Added use of professional judgment when qualifying for CCV frequency infractions.

Section 4.4.3, RLV; Added discussion of the IRA, which has been added to HE 8330B analysis.

Section 4.4.4, Blanks; Scenarios removed and table added that reflects current NFG criteria.

Section 4.4.7, MS/MSD; Non-detects are now rejected for MS/MSD recoveries <20%, per NFG.

Section 4.4.9, LCS; Non-detects are now rejected for LCS recoveries <20%, per NFG. Also, clarified the criteria for LCS reanalysis.

Section 4.4.11, Dilutions; Replaced instrument response with concentration.

Section 4.4.12; Perchlorate isotope ratio acceptance criteria clarified.

Section 4.6.1, Initial Calibration; List of specific metals for axialview ICP-AES was removed. Clarified the qualification of multiple calibration curves.

Section 4.6.2, CCV; ICV/CCV acceptance criteria for Hg and cyanide is now 90-110% to reflect QSM criteria.

Section 4.6.3, Blanks; Revised table to reflect current NFG criteria.

Section 4.6.4, Inorganic MS; Updated post-digestion spike

IMPORTANT NOTICE:

recovery limits, per NFG.

Section 4.6.5, Inorganic Replicate; Added the option of qualifying results with an RPD >100%, per NFG.

Section 4.6.7, Reporting Limit Verification; LLCCV analyzed for metals at PQL concentration with 80-120% acceptance criteria per QSM; no longer required for cyanide.

Section 4.6.8.1, ICS; Added qualification for elements not present in the ICS AB solution.

Section 4.6.8.3, Total Cyanide; CCV acceptance criteria updated to 90-110%, per QSM. Qualification criteria for negative blank infractions updated.

Section 4.6.8.6 was amended to include retention time criteria for Perchlorate by Ion Chromatography.

Section 4.7.8.2, Gross alpha beta; Added preferred approach for flaming planchets.

Section 5.3, Data Qualification Summary; In RLV bullet point, both instances of CRA/CRDL were replaced with LLCCV.

Section 6.2 was renamed "Sample Detection/Quantification Limits."

Section 6.3, Formulas; The linear curve and %RSD formulas were modified.

Appendix A and associated references deleted due to revision of VOC HT qualifiers. Subsequent appendices renamed (A through F) and references within text were updated.

Update to Appendix E, Laboratory Control Limits, to reflect those in Appendix C of QSM. Added limits for TO-15.

A global change of RRF minimum limits from 0.01 and 0.05 to 0.010 and 0.050, respectively was made throughout the document.

A global change of the coefficient of determination QC limit from ≥ 0.99 to ≥ 0.990 was made throughout the document.

Effective Date: 06/19/2017

Procedure No: AOP 00-03 Issue No: 05

ACRONYMS AND ABBREVIATIONS

SNL acronyms that do not need to be defined in SNL Memos

AA Atomic Absorption

ARCOC Analysis Request and Chain of Custody

BOD Biological Oxygen Demand

CAS Chemical Abstract Service CB Chlorinated Biphenyl

Continuing Calibration Blank **CCB** Continuing Calibration Verification **CCV**

CF Calibration Factor

Contract Laboratory Program **CLP** Common Laboratory Contaminant CLC RLV for LC/MS/MS Method CRI

Compliance & Requirements Management **CRM**

Contract Verification Review **CVR**

DL **Detection Limit**

Dichlorodiphenyldichloroethane **DDD** Dichlorodiphenyldichloroethylene DDE Dichlorodiphenyltrichloroethane **DDT** U.S. Department of Energy DOE Diesel Range Organics DRO

 $\mathbf{E}\mathbf{B}$ Equipment Blank

Electronic Data Deliverable **EDD Extracted Ion Current Profile EICP**

EPA U.S. Environmental Protection Agency

FB Field Blank

Full-width Half-maximum **FWHM**

GC Gas Chromatography

Gas Chromatography/Mass Spectrometry GC/MS

GPC Gel Permeation Chromatography

Gasoline Range Organics GRO

HE High Explosive

High-Performance Liquid Chromatography **HPLC** High Resolution Gas Chromatography **HRGC HRMS** High Resolution Mass Spectrometry

IMPORTANT NOTICE:

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

ACRONYMS AND ABBREVIATIONS (continued)

Ion Chromatography IC **Initial Calibration ICAL** Initial Calibration Blank **ICB**

Inductively Coupled Plasma-Atomic Emission Spectroscopy **ICP-AES**

Inductively Coupled Plasma-Mass Spectrometry **ICP-MS**

ICS Interference Check Standard

Interference Check Sample Solution A ICS A **ICS AB** Interference Check Sample Solution AB

Initial Calibration Verification **ICV**

Re-analysis of low-level standard following calibration for HE by LC/MS/MS IRA

Internal Standard IS

Liquid Chromatography/Mass Spectrometry/Mass Spectrometry LC/MS/MS

LCS Laboratory Control Sample

Laboratory Control Sample Duplicate LCSD

Low Level Continuing Calibration Verification LLCCV

LOC Level of Chlorination

Low Resolution Mass Spectrometry LRMS

Method Blank MB

MDA Minimum Detectable Activity Minimum Detectable Concentration **MDC**

Method Detection Limit MDL Manual Integration Review MIR

Matrix Spike MS

MSD Matrix Spike Duplicate

MSDC Mass Spectrometry Data Centre

National Bureau of Standards **NBS** NFG National Functional Guidelines NIH National Institute of Health

National Institute of Standards and Technology **NIST** National Nuclear Security Administration **NNSA**

OP Operating Procedure

OPR Ongoing Precision and Recovery

Polyaromatic Hydrocarbon PAH Polychlorinated Biphenyl **PCB** Polychlorinated Dibenzodioxin **PCDD** Polychlorinated Dibenzofuran **PCDF**

IMPORTANT NOTICE:

ACRONYMS AND ABBREVIATIONS (concluded)

PPCP Pharmaceuticals and Personal Care Products

PQL Practical Quantitation Limit

PS Post-digestion spike

QC Quality Control

QSM DoD/DOE Consolidated Quality Systems Manual for Environmental

Laboratories

RER Replicate Error Ratio

RL Reporting Limit

RLV Reporting Limit Verification
RPD Relative Percent Difference
RRF Relative Response Factor
RRT Relative Retention Time
RSD Relative Standard Deviation

RT Retention Time

SD Serial Dilution

SDG Sample Delivery Group SMO Sample Management Office

SNL/NM Sandia National Laboratories/New Mexico

SOW Statement of Work

SVOC Semivolatile Organic Compound SW Solid Waste (EPA procedure number)

TAL Target Analyte List

TB Trip Blank

TCDF Tetrachlorodibenzofuran

TIC Tentatively Identified Compound

TKN Total Kjeldahl Nitrogen **TOC** Total Organic Carbon

TPH Total Petroleum Hydrocarbon
TPU Total Propagated Uncertainty

VOC Volatile Organic Compound

Effective Date: 06/19/2017

Procedure No: AOP 00-03 Issue No: 05

MEASUREMENTS AND SYMBOLS

°C	degrees Centigrade
----	--------------------

> greater than

greater than or equal to ≥

< less than

less than or equal to ≤

% percent

%D percent difference %R percent recovery

%RSD percent relative standard deviation

plus or minus

X times sigma σ

AMU atomic mass units correlation coefficient r r^2 coefficient of determination

kilo electron volt kev

kilogram kg L liter

m/e mass/charge ratio microgram μg milligram mg mLmilliliter(s) parts per billion ppb

ppbv parts per billion by volume Effective Date: 06/19/2017

1 **PURPOSE**

The purpose of data validation is to identify, through the evaluation of supporting documentation, those data that do not meet the expected precision and accuracy of an analytical method. This procedure presents the guidelines used to evaluate chemical (organic and inorganic) and/or radiochemical analytical data acquired in support of environmental and waste management activities. The purpose of the procedure is to consistently qualify data using defined criteria; however, it is not intended to eliminate the need for professional judgment in evaluating the data quality. The data validator may be more or less stringent in evaluating the results based on experience and familiarity with the analytical techniques, historical data, sample matrices, or intended use of the data. The product of this procedure is a data validation report that includes information regarding the overall quality of the data and the resulting data qualifiers. When variations in the application of data qualifiers are warranted, the justification and rationale will be explained in the data validation report.

Procedure No: AOP 00-03

Issue No: 05

SCOPE AND OWNERSHIP

2.1 Scope

This procedure specifically covers the validation of chemical or radiochemical analytical results from environmental methods required for Sandia National Laboratories/New Mexico (SNL/NM) Sample Management Office (SMO) decisions but may be used by other organizations as appropriate. The format is based on analytical techniques, standard reporting protocols used by the laboratories, and the general format followed by the U.S. Environmental Protection Agency (EPA) Contract Laboratory Program (CLP) national functional guidelines (NFG). Additions and modifications were made to address analyses requested by the SNL/NM SMO customers. Any apparent redundancies between sections, is stylistically intentional for the sake of completeness and accuracy. Qualification of data performed under this procedure does not replace any data usability review for specific project use.

2.2 Ownership

The SNL/NM SMO owns this operating procedure (OP). The SMO is responsible for maintaining and revising this OP as necessary. Any comments or suggestions for improvement should be forwarded to the SMO.

During the lifetime of this OP, it is anticipated that some of the requirements and procedures of the SMO may change. This OP will be updated every three years, and changes provided to appropriate customers as required for information, review, concurrence, or approval. The SMO owns this document. The SMO is responsible for preparing, revising, and distributing this document as necessary.

RESPONSIBLE INDIVIDUALS AND ORGANIZATIONS 3

This section describes the responsibilities of SNL/NM personnel and contractors regarding this OP.

Procedure No: AOP 00-03 Effective Date: 06/19/2017

3.1 Program Leader

Responsible for providing programmatic guidance leading to the development of this OP and the following:

- Reviewing and approving the OP.
- Acting as liaison to the U.S. Department of Energy (DOE) and the National Nuclear Security Administration (NNSA)/Sandia Field Office on data validation issues.

Issue No: 05

• Ensuring that resources are available to perform tasks in compliance with this OP.

3.2 SMO Operations Manager

Responsible for the operations and activities conducted within the SMO. The principal responsibilities of the SMO operations manager include but are not limited to the following:

- Updating this OP.
- Managing the validation contract, acting as the Sandia Delegated Representative, reviewing routine performance assessments, and conducting general contract oversight.
- Providing oversight of the data review and validation process.
- Ensuring this OP is implemented for review and validation of analytical data provided by the contract laboratories when data validation is requested.
- Developing and maintaining processes that ensure the necessary documentation, to perform data review and validation, is made available to the laboratory oversight/data validation contractor.

3.3 Compliance & Requirements Management (CRM) Staff

CRM staff is responsible for:

- Providing project data quality assurance guidance.
- Ensuring that this procedure is distributed to the appropriate personnel for project/program
- Ensuring that sufficient quality checks are in place to maintain the integrity of the SMO sample information management and analytical result database.
- Documenting non-conformances and corrective actions in accordance with the applicable SMO-QAPP.
- Interfacing with the Records Management Coordinator for maintenance of project documentation and to resolve record management concerns for storage and maintenance of sampling and analysis records.
- Reviews updates to regulatory methods, SNL corporate requirements and DOE guidance documents, such as the DOE QSM, and implements changes as necessary to remain current and to assure efficient program development.

3.4 SMO Project Coordinator

The SMO project coordinator is responsible for coordinating efforts associated with SMO analytical services. The principal responsibilities of the SMO project coordinator include but are not limited to the following:

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Acting as a point of contact between Task/Project Leaders, the analytical laboratories, and the laboratory oversight/data validation contractor.

- Performing the Contract Verification Review (CVR) and completing form SMO 2012-CVR on analytical data packages from the contract laboratories pursuant to "Procedure for Completing the Contract Verification Review," (SMO-05-03) and the SNL/NM Contract Statement of Work (SOW) for Analytical Laboratories (contract SOW).
- Transmitting and tracking electronically the complete analytical data package along with the CVR form to the laboratory oversight/data validation contractor.
- Implementing and follow-up of all nonconformances and corrective actions with the contract analytical laboratories and laboratory oversight/data validation contractor.
- Processing the EDD that includes the data validation qualifiers, into the Environmental Data Management System (EDMS), pursuant to "Procedure for Electronic Data Deliverable (EDD) Processing," (SMO-05-04).
- Performing quality control (QC) checks on all data validation results by reviewing the report and comparing the results to the data validation qualifiers captured on the EDD.
- Transmitting the complete analytical data package to SMO project/data management staff for final archiving.

3.5 SMO Staff

The SMO project/data management staff is responsible for:

- Ensuring compliance with the "SMO Data Management Plan," (AOP 95-44).
- Receiving and processing analytical data packages
- Managing data flow and data storage, including both hardcopy paper records from field activities and analytical laboratories, and electronic data relating to sample tracking or analytical results.
- Forwarding the complete and final analytical data package and electronic data to the SNL/NM Records Center for archiving.

3.6 Laboratory Oversight/Data Validation Contractor

The Laboratory Oversight/Data Validation Contractor is responsible for:

- Performing data validation in accordance with this OP.
- Requesting data corrections or additional information needed from the contract analytical laboratories and notifying SMO of the request.
- Notifying SMO of all data determined as rejected ("R" coded) according to this OP.
- Communicating non-compliance issues to the SMO technical lead and/or SMO project coordinator(s) and ensuring that nonconformances (e.g., incorrect or missing analytical information) are adequately addressed.
- Completing the data validation report including checklist(s) and if applicable generating validation EDD files (see Section 5.1).
- Communicating with the SMO customer or designated representative when data review and validation is complete and returning the complete data package to the SMO.

Data Validation Procedure for Chemical and Radiochemical Data Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

- Verifying implementation of laboratory corrective action plans.
- Performing laboratory oversight as directed by the SMO.

PROCEDURE

Data are evaluated using common quality parameters from QC measurements specified in the methods and the SNL/NM contract SOW. These parameters are compared to statistically derived or regulatory method criteria to estimate the quality of the results. The quality parameters are measures of the analytical precision and accuracy, potential contamination both from the field and from the laboratory, sample matrix effects, and sample inhomogeneity. The laboratory may define the acceptance criteria as long as they meet or exceed those specifically defined within the method or contract. The appropriateness of acceptance criteria generated by the laboratory should be evaluated periodically by the SMO.

Qualification is based on minimal reporting requirements and does not address method or contract compliance requirements, except within the context of QC data. Complete method and contract compliance cannot generally be performed using only the laboratory data package and should be done during on-site assessments at the laboratory where all supporting documentation is available.

If any QC element for a method is not provided, the data validation report must document that the QC data are missing and any qualification is at the discretion of the data validator. A QC failure for an analyte that results in "R" coded (unusable) data due to matrix problems (e.g., matrix interference that cannot be alleviated by acceptable clean-up procedures) brings the appropriateness of the analytical method into question. As a result, the data validation report must document that analysis by another acceptable method or modification of the existing method may be necessary.

4.1 General

This section provides the portions of the method for reviewing QC data that are pertinent to both chemical (organic and inorganic) and radiochemical analytical data.

4.1.1 CVR

The SMO is responsible for conducting a CVR of analytical data packages delivered from the contract laboratories using the SMO "Procedure for Completing the Contract Verification Review," (SMO-05-03).

Criteria: A CVR form shall be included with the analytical data package that specifically

addresses the Analysis Request and Chain of Custody (ARCOC), receipt of samples by the laboratory, and the technical, OC, and reporting requirements imposed upon the analytical laboratory through the contract SOW.

IMPORTANT NOTICE:

Effective Date: 06/19/2017

Evaluation	Action
	Action
The CVR form should be checked to confirm:	Report any discrepancies and/or anomalies associated with the CVR
ARCOC (SMO 2012-ARCOC) and laboratory login information have been reviewed,	form to SMO.
missing samples and sample container irregularities are discussed,	
preservation and hold time deficiencies are indicated,	
appropriate target analyte lists (TALs) and contract-required laboratory qualifiers are used,	
results are reported, in correct units, for all analytes requested,	
all radiochemistry results include the calculated total propagated uncertainty (TPU),	
the required detection limits (DL) are reported and clearly defined or an explanation of why they were not met is given,	
all outstanding reporting issues are resolved,	
any request for an amended report from the laboratory has been received, and	
signatures and dates are present indicating CVR was completed.	

Procedure No: AOP 00-03

Issue No: 05

4.1.2 QC Exemptions

Various filter materials may be submitted for analysis. Matrix spike (MS) and replicate sample analysis requirements shall not apply to filter materials because representative splits of these samples are generally not obtainable. All other QC criteria shall apply to the analysis of filters.

The requirements for reanalysis for QC failures are waived when insufficient sample remains. A detailed discussion of that condition shall be included in the laboratory case narrative when it is encountered.

Acidity, alkalinity, biological oxygen demand (BOD), color, corrosivity, dissolved oxygen, gravimetric oil and grease, hardness, ignitability, pH, titrimetric sulfide, conductivity, all of the solids methods, and turbidity analyses are generally exempt from the general inorganic QC requirements.

Criteria: The analyses referenced directly above shall be controlled according to the method QC and/or the laboratory's QC policies. In general, one or more of the following should be included:

- Blank; result less than (<) the method detection limit (MDL).
- Laboratory control sample (LCS); measured value within plus or minus (±) 20 percent (%) of known value.
- Duplicate; relative percent difference (RPD) <25%.
- Independent calibration check standard; result within \pm 10% of true value.

Note: Blanks (method blank [MB]/field blank [FB])/equipment blank [EB]) are not applicable for acidity by titration, alkalinity, conductivity, flash point, pH, and specific gravity. In the Blanks section of the data validation report, document that the blank result was reported but not assessed for data validation.

Evaluation	Action
If there are any QC failures for any of the analyses listed above,	qualify sample results associated with QC failures according to the appropriate requirements in Section 4.6, Procedure for Inorganic Data Validation.
	Note: Sample results shall not be qualified due to the lack of QC data. QC exemptions shall be discussed in the data validation report.

4.1.3 Holding Times

Samples must be extracted and analyzed within EPA-specified holding times for results to be considered reflective of total concentrations. Analytical data generated outside of the specified holding time criteria must be considered suspect. Holding times must be evaluated to ascertain the validity of results based on the holding time of the sample from time of collection to time of analysis.

Solid materials, such as soils, that are being analyzed for radioisotopes or metals are generally exempt from qualification for exceeded holding times. The reviewer should evaluate the stability of the analyte and half-life, if applicable, and qualify based on professional judgment.

Updates to the VOC holding time data qualification guidance herein are based on strict adherence to the 2017 revision of the NFG. For all other organic analyses, the use of professional judgment in

qualifying sample data when extraction/analysis is performed >1X but \leq 2X the holding time is addressed in the 2017 NFG in sub-Section E (Action) of the Preservation and Holding Time chapters. Based upon that language, and the explicit definition of "gross exceedance" found in the 1999 revision of the NFG and reiterated by the USACE in EM 200-1-10, the current NFG term "gross exceedance" is defined as >2X the holding time for the purposes of this procedure.

Criteria: All samples will be extracted and analyzed within specified holding times, per Appendix A.

Evaluation	Action
If a holding time infraction is <5% of the holding time criteria,	sample results may be accepted without qualification based on professional judgment. Note: Consideration should be given to the relevant holding time requirement; for example, "days" versus "hours."
If holding times are exceeded and preservation requirements are not met (see Section 4.1.4),	qualify all associated detects as "J-" and all associated non-detects as "R."
If samples for VOC analysis were analyzed after their holding time had expired,	qualify all associated detects as "J-" and all associated non-detects as "R."
If samples for analysis other than VOC were analyzed after their holding time had expired but within 2 times (X) the specified holding time,	qualify all associated detects as "J-" and all associated non-detects as "UJ." Infractions for pH will be qualified "J."
If samples were analyzed beyond 2X the specified holding time,	qualify all associated detects as "J-" and all associated non-detects as "R." Infractions for pH will be qualified "J."
If samples were analyzed within holding time and reanalyzed out of holding time due to a QC failure and	
the original and reanalysis calibration, sample, and QC data are provided and the sample results are similar,	accept the results of the reanalysis without holding time qualification.
the original or reanalysis calibration, sample, and QC data are not provided, or the sample results of the original analysis and the reanalysis are not similar,	qualify all associated detects as "J" and all associated non-detects as "UJ."

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Preservation (chemical and temperature)

Samples must be preserved according to EPA-specified criteria for results to be considered reflective of actual concentrations. Analytical data generated for samples that were outside of the specified preservation criteria must be considered suspect. The data validation report shall include a discussion of any preservation violations and a discussion supporting any qualifications.

Many organic compounds, and most metals and radioisotopes, are not affected by temperature variations up to ambient temperature and are generally not qualified. VOCs and mercury are subject to analyte loss at elevated temperatures.

Criteria: All samples shall be preserved and shipped under conditions specified in Appendix A.

> Samples for metals or radiochemical analysis that were received without the required chemical preservation but that were preserved by the laboratory after receipt generally do not require qualification if the samples were allowed to equilibrate at least 16 hours before a sample aliquot is taken.

Evaluation	Action
If samples were received outside the temperature criteria,	all associated detects may be qualified as "J" and all associated non-detects may be qualified as "UJ" or "R" using professional judgment (see below).
If temperature violations of >6°C but ≤10°C occur for VOCs and/or mercury (soil/sediment) or cyanide,	qualify all associated detects as "J" and all associated non-detects as "UJ" based on professional judgment.
If temperature violations of >10°C occur for VOCs, mercury (soil/sediment) or cyanide,	qualify all associated detects as "J-" and all associated non-detects as "R" based on professional judgment.
If temperature violations of >10°C occur for SVOCs, pesticides or PCBs,	qualify all associated detects as "J" and all associated non-detects as "UJ" based on professional judgment.
If samples preserved by the laboratory upon receipt were not allowed to equilibrate after laboratory preservation, or if no documentation shows the samples were allowed to equilibrate,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If samples were received without the required preservation and were not preserved by the laboratory after receipt,	qualify all associated detects as "J" and all associated non-detects as "R."

Effective Date: 06/19/2017

Procedure No: AOP 00-03 Issue No: 05

4.1.5 Calibration Points

Generally, it is not acceptable to remove points from the calibration curve unless the points are at the high or low ends of the curve. For the purpose of meeting calibration criteria, if a point is removed from the low end, the practical quantitation limit (PQL) must be adjusted accordingly. If a point is removed from the high end, the calibration range must be adjusted accordingly. Whenever a point is removed, it must be clearly documented in the instrument log. All initial calibration (ICAL) points must be analyzed without any changes to instrument conditions, and all points must be analyzed within 24 hours.

The laboratory may remove ICAL data points that are not the low or high points of the average or linear/quadratic curve, if the reason can be clearly documented. Acceptable reasons include misinjection of the standard or minor instrument failure for the particular data point. Notify the laboratory project manager if no such documentation is present.

4.1.6 Calibration for QC Samples

If any QC samples are analyzed using a different ICAL than that of the field samples, the laboratory must include a calibration report from the calibration affecting the QC samples. This calibration data shall only be used to evaluate the QC samples and only if the QC samples fail to meet recovery or RPD acceptance criteria. The laboratory is not required to report calibration data associated with OC samples from another sample delivery group (SDG).

4.1.7 Blank Hierarchy

The general hierarchy for application of qualifiers due to blank contamination is 1) instrument blank, 2) preparation blank or MB, and 3) FB, EB, or trip blank (TB). As a general guideline, if the instrument blank is contaminated, then associated detected results in field samples, MB, FBs, EBs, and TBs that are analyzed in the same analytical run may be qualified. If the preparation blank is contaminated, all associated detected results in samples prepared with that blank may be qualified even if the samples are analyzed in different runs. If an FB or EB is contaminated, all associated detected results in samples collected during the same sampling event may be qualified. If a TB is contaminated, all associated detected results in samples transported in the same container (cooler) may be qualified. Professional judgment must be employed to determine the effect of multiple blank contaminations upon the quality of field sample data.

4.1.8 **Blank Normalization**

Because sample aliquot values (masses or volumes) seldom vary significantly within a batch, the laboratory generally assigns a representative aliquot value to the MB. When a sample has a significantly different aliquot size than that of the MB, a detected MB result must be normalized to the detected sample result before a comparison can be performed for blank assessment. The blank data are normalized to the sample results using the following equation:

Normalized blank concentration = (blank concentration) X (blank aliquot value/sample aliquot value)

It should be noted that the blank analyses might not involve the same weights, volumes, and/or dilution factors as the associated samples. These factors must be taken into consideration when applying the 5X and 10X criteria, such that the total amount of contamination is actually compared.

4.1.9 **Field Duplicates**

Field duplicate samples may be collected and analyzed as an indication of overall precision. These analyses measure both field and laboratory precision; therefore, the results may have more variability than laboratory duplicates which measure only laboratory performance. It is expected that solid or waste duplicate results will have a greater variance than water matrices due to inhomogeneity.

If samples are identified as field duplicates, document the occurrence in the data validation report and state that there are no "required" review criteria for field duplicate analyses comparability.

4.1.10 Sample-Specific External Standard Recovery

In lieu of an internal standard (IS) addition, an addition of a known quantity of material to a second sample aliquot may be used to calculate sample results. To evaluate external standard recovery (standard addition), the spike amount and spike recovery must be reported.

Criteria: Recovery guidelines for external standard recovery shall be 50% to 105%. The quantity of external standard used should be adequate to provide a reasonable confidence level in the measured recovery; that is, the spike level should be greater than (>) the indigenous level.

> **Note:** For samples that require dilution the evaluation uses the concentration of the diluted result not the corrected result.

Evaluation	Action
If the measured sample result is >2X the external standard spike added,	qualify all associated results as "J."
If the measured sample result is >4X the external standard spike added,	qualify all associated results as "R."
If the recovery is >105% but less than or equal to (\le) 125%,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
If the recovery is >125%,	qualify all associated non-detects as "R." Associated detects may be qualified "J-" or "R" based on professional judgment.
If the recovery is <50% but greater than or equal to (≥) 20%,	qualify all associated detects as "J+" and non-detects as "UJ."
If the recovery is <20%,	qualify all associated detects as "J+" and non-detects as "UJ" or "R" based on professional judgment.

Data Validation Procedure for Chemical and Radiochemical Data Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

4.1.11 Rounding Rules and Significant Figures

If the figure is ≥5, round up; otherwise, round down. For example, 11.443 is rounded down to 11.44, and 11.455 is rounded up to 11.46. If a series of multiple operations is to be performed (i.e., add, subtract, divide, and/or multiply), all figures are carried through the calculations. The final answer is rounded to the proper number of significant figures. Before evaluating a number for being in control or out of control of a certain limit, the number evaluated shall be rounded using these rounding rules to the significance reported for that limit. For example, if the acceptance limit is $\pm 10\%$ of the true value, then a calculated percent recovery (%R) of 110.46 shall be reported as 110%, which is within the acceptance limits of 90% to 110%. On the other hand, a calculated %R of 110.50 shall be reported as 111%, which is not within the 90% to 110% acceptance limits.

Blank qualifications with an associated numerical value should be recorded with no more than three significant figures for values ≥100, and no more than two significant figures for values <100 in the data validation report; for example, 125U, 18U, 9.9U, 0.32U or 0.032U.

4.1.12 Special Laboratory Flags

"X" Flags

Criteria: The laboratory or analyst may have reason to believe that the result for a specific analysis has a high probability of being a false positive due to interferences. In this case, the laboratory shall qualify the result as "X" and narrate the justification for the flag. Generally, use of the "X" flag is restricted to use in conjunction with additional data such as spectral matching or results from another analytical technique. The raw data and case narrative should be reviewed to determine if they support the identification of a false positive.

Evaluation	Action
When evaluating the "X" qualifier, if it is	qualify detects determined to be
determined that the interference is the most	primarily false positives as "R" and
significant source of the instrument	detects determined to have very high bias
response (i.e., if the detect is primarily a	as "NJ+." Include a thorough discussion
false positive or if it is a detect with a very	supporting the qualification in the data
high bias),	validation report.

4.1.13 Analytical Methods

The laboratory shall follow the requirements specified in the analytical methods and those specified in the contract SOW. When these requirements are not met, reanalysis is required. In those cases where reanalysis cannot occur, the failure to reanalyze will be discussed in the case narrative. This discussion should also be included in the data validation report. See Appendix B for data reporting requirements.

Procedure No: AOP 00-03 Data Validation Procedure for Chemical and Radiochemical Data Issue No: 05 Effective Date: 06/19/2017

4.1.14 Calculations

Criteria: Laboratories will generally use commercial software whenever possible. Spreadsheets and laboratory developed software shall be verified and uniquely identified, and shall include a revision number (i.e., be under version control). Reverification of commercial software and other software is not routinely required. Hand-calculated data or data calculated from a spreadsheet or other software not under version control must be verified by the random recalculation of some of the results. Hand calculated results and spreadsheets should have all required formulas and data included in the package. In addition, any spreadsheet that is not under version control should be brought to the attention of the SMO.

Evaluation	Action
If results cannot be regenerated using the reported data,	require a formal corrective action by the laboratory.
If results are verified by recalculation using reported data,	discuss the recalculation in the data validation report.

Criteria: Laboratories are required to calculate the RPD between the MS and matrix spike duplicate (MSD) using the actual results (Solid Waste [SW]-846 Method 8000D). CLP and some other programs use calculation routines, which calculate the RPD using the %Rs.

$$RPD = (MS \%R - MSD \%R)/[(MS \%R + MSD \%R)/2]$$

This does not give an equivalent result to that obtained using the SW-846 formula (see Section 6.3 below) when the sample contains indigenous analyte. When the RPD is calculated using %Rs, the results should be recalculated before the evaluation is performed.

Evaluation	Action
If results are recalculated using the correct data,	discuss the recalculation in the data validation report.

4.1.15 Reanalysis

The laboratory may perform a reanalysis on one or more samples because of QC failures. This may occur because of MS failures, or it may occur because a small subset such as the acid fraction in semivolatile organic compound (SVOC) analysis had QC failures for the first analysis and the second analysis was performed outside the method-specific holding time. Based on professional judgment the laboratory is to report only the best data set on the certificate of analysis (COA). All supporting documentation concerning a reanalysis will be provided in the miscellaneous data section of the analytical data package.

IMPORTANT NOTICE:

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

4.1.16 Manual Integration

Manual integration review (MIR) is typically outside the scope of routine validation. When MIR is required by the program, it must be performed in accordance with standard operating procedures.

Manual integration is used to correct improper integration performed by the instrument software, not for the purpose of meeting QC criteria. While MIR is not normally required for data validation, manually integrated peaks may be reviewed based on professional judgment or whenever QC problems indicate it may be necessary

Evaluation	Action
If a manual integration was not documented correctly or was performed incorrectly, or does not meet one or more of the applicable criteria,	request reprocessed data from the laboratory. Data may be qualified as "J" or "R" based on professional judgment.

4.1.17 Failed Batch QC

Occasionally the batch QC sample (i.e., MS, LCS, blank, etc.) will fail and the individual QC sample will also fail sample-specific QC parameters (i.e., ISs, surrogates, etc.) The usefulness of the QC data from these batch QC samples is based on professional judgment for minor excursions. However, significant failures where the QC sample fails both sample parameters (i.e., surrogates, etc.) and batch parameters (e.g., analyte) require that the batch QC data be rejected and the batch be treated as if it did not include the batch QC sample. That is, the samples are qualified as if no QC sample was run with the batch.

4.1.18 MS/MSD, LCS/LCSD and Replicates

Occasionally the laboratory may analyze for replicates, matrix spike/matrix spike duplicate (MS/MSD) pairs, and/or LCS/laboratory control sample duplicate (LCSD) pairs, presenting more than one measure of precision. If the sample has little or no indigenous analyte, the MS/MSD RPD is the best indicator of precision. If the sample has significant indigenous analyte, the replicate is the best indicator of precision. As a general rule, the replicate precision is used if the indigenous analyte is >2X or 3X the MS spike concentration. The LCS/LCSD RPD should only be used as a measure of precision in the absence of both MS/MSD and replicate analyses.

More than one measurement of precision is not assessed for the same sample/analyte. The data validation report should include a discussion on which measure of precision was used for assessment and why.

4.1.19 MS/MSD

Occasionally the laboratory may dilute before spiking or may run the MS/MSD pairs at a reduced volume. For example, the sample aliquot will be 1000 milliliters (mL) while the MS aliquot is 500 mL. If the extract volume is the same for both the sample aliquot and the MS aliquot, the RPD is still

IMPORTANT NOTICE:

Procedure No: AOP 00-03 Data Validation Procedure for Chemical and Radiochemical Data Effective Date: 06/19/2017 Issue No: 05

a good measure of precision, but the %R is not a good measure of accuracy and matrix effect. At a minimum, this issue should be noted in the data validation report. If the final volume of the MS aliquot was adjusted for sample size (in this case, adjusted to half the sample extract volume), it should be noted that the laboratory may have adjusted the extract volume to account for a smaller sample aliquot. In this case, the MS %R is a good measure of accuracy and matrix effect.

MS samples that require dilution due to matrix should not be used to evaluate associated sample data unless the relative dilution factor between the MS and the field samples is ≤5, in which case there may still be significant sample matrix similarity between the MS and field samples. If the MS sample is not used to evaluate sample data, the sample results should be qualified for lack of accuracy and/or precision data, as applicable, if specified by the program.

Evaluation	Action
If the MS/MSD relative dilution factor is >5 compared to the samples,	qualify all associated detects as "J" and all associated non-detects as "UJ."

4.1.20 MS/MSD with Elevated Analyte Concentration Requirements

When the sample used for the MS/MSD has an analyte concentration >4X the analyte spike concentration and the MS and/or MSD %R is out of limits, sample results should be qualified due to a lack of matrix-specific accuracy data. Matrix-specific precision can still be assessed using the MS/MSD RPD. If a post-digestion spike (PS) is also performed, it can be used to assess matrixspecific accuracy data for the analytes not evaluated using the MS/MSD. The 4X rule also applies to the PS; however, its analyte spike concentration may be higher than that of the MS/MSD. The PS recovery limits are usually narrower than the MS recovery limits. The MS and MSD results may be used in conjunction with other QC results to determine the need for qualification of the data.

Evaluation	Action
If the sample used for MS/MSD has an analyte concentration >4X the analyte spike concentration and the MS and/or MSD %R for that analyte is out of limits,	qualify all associated detects as "J" and all associated non-detects as "UJ."

4.1.21 Blank Qualification with QC Failures

Data may be qualified as a non-detect (U) based on blank contamination and have other QC failures. While the general approach is to qualify the sample result as a non-detect with no further qualification, other quality issues should be considered to determine whether additional qualification is warranted. For example, if the LCS had very low recovery, the actual sample result may be below the blank result because of poor recovery, not just because of blank contamination. In this case, the result may be qualified "UJ" rather than "U." In general, samples with results that are qualified "U" or "UJ" due to blank contamination are not rejected. Justification for additional qualification must be explained in the data validation report.

IMPORTANT NOTICE:

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

4.1.22 Initial Dilutions

Initial dilutions may be required due to high indigenous analyte concentrations. For multi-analyte determinations where initial dilutions are required to keep from saturating the detector, the DLs and reporting limits (RL) must be adjusted for the initial dilution. In addition, the matrix effect of the over-range analyte on the other analytes being measured cannot be determined.

Evaluation	Action
If all target analytes are reported from the same diluted run,	all associated detects may be qualified as "J" and all associated non-detects may be qualified as "UJ" based on professional judgment.

4.1.23 Reporting Limit Verification (RLV)

Data from independent RLV standards may be used to additionally evaluate the intercept. Acceptable RLVs may be used to minimize qualification based on professional judgment. An acceptable curve with a low standard at the RL does not meet this requirement. The RLV must be the measurement of an independent standard or the low standard reprocessed against the new curve.

4.1.24 Filtered Samples

Water samples may be submitted as both field filtered and unfiltered fractions. When it is evident that both a filtered and an unfiltered sample are submitted, both results will be reviewed. The analyte concentrations for the filtered portion should be \leq the unfiltered portion.

Evaluation	Action
If the analyte concentrations of the filtered portion are generally > that of the unfiltered portion,	contact the laboratory to determine whether a sample mix-up has occurred.
If the analyte concentrations of the filtered portion are generally > that of the unfiltered portion and the reason cannot be identified,	document the problem and contact the technical data support for further direction.

4.1.25 Sample Contamination

There may be instances where little or no contamination is present in the associated blank, but qualification of the sample due to contamination is deemed necessary. Contamination introduced in a diluent is one example. Although it is not always possible to determine, evidence of this occurrence can be identified when contaminants are found in the diluted sample result but are absent in the undiluted sample.

Evaluation	Action
If it is determined that the sample contamination is from a source not identified in the blank,	qualify the results for that analyte as "R" and discuss such circumstances in the data validation report.

4.2 Procedure for Gas Chromatography/Mass Spectrometry (GC/MS) Validation

The requirements addressed within this section are applicable to all GC/MS analytical techniques.

4.2.1 Instrument Tuning for GC/MS

Tuning and performance criteria are established to ensure mass resolution, identification; and, to some degree, sensitivity. These criteria are not sample-specific. Conformance is determined using standard materials. Therefore, these criteria should be met in all circumstances.

Criteria: The GC/MS tune shall be evaluated daily. The relative abundance criteria listed in the appropriate method must be met.

Evaluation	Action
If tunes are not run daily or if all abundance criteria are not met,	contact the laboratory for immediate corrective action and use professional judgment to determine which data should be used. The following actions are suggested:
	qualify all associated detects as "J" and all associated non-detects as "UJ."
If multiple QC failures also occurred,	qualify all results as "R."

4.2.2 Calibration

Initial Calibration

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing acceptable qualitative and quantitative data for compounds on the TAL. Initial calibration demonstrates that the instrument is capable of acceptable performance at the beginning of the analytical run and of producing an adequate curve. In the absence of, or in addition to, method-specific calibration acceptance criteria, the following general calibration acceptance criteria should be applied.

The laboratory may establish a calibration curve using either the regression approach or the average relative response factor (RRF) approach. If both approaches are used to quantify and report target analytes within the same data package, calibration is to be assessed on an analyte-by-analyte basis.

Criteria: GC/MS instrument calibration shall be performed using a minimum of five calibration standards unless otherwise specified by the method. If calibration curves are used, five standards are required for a linear (first-order) calibration model, six standards are required for a quadratic (second-order) model, and seven standards are required for a third-order polynomial. Higher order curves (second order and higher) should not normally be used. If the laboratory uses a higher-order equation to establish a calibration curve, it should be evaluated for appropriate application.

Evaluation	Action
If an insufficient number of calibration standards was used,	qualify all associated detects as "J" and all associated non-detects as "UJ."

RFs

Criteria: RRFs are a measure of the slope of the calibration relationship and assumes that the curve passes through the origin. Under ideal conditions, the factors will not vary with the concentration of the standard that is injected into the instrument. In practice, some variation is to be expected.

When the variation, measured as the percent relative standard deviation (%RSD), is \leq 15%, the use of the linear model is appropriate and the calibration curve can be assumed to be linear and to pass through the origin. This criterion is derived from SW-846 GC/MS Methods 8260B/8260C and 8270C/8270D.

As a general rule, the amount of IS should produce an instrument response (e.g., area counts) that is no more than 100X that produced by the lowest concentration of the least responsive target compound associated with the IS. This should result in a minimum RRF of not <0.010 for the least responsive target compound.

The %RSD for the RRFs obtained from the five ICAL standards must be ≤15% and the average RRF shall be ≥ the method-specified minimum RRF for each compound. The minimum RRFs_for the system performance check compounds per method SW-846 8260B/8260C (VOC) are:

 Bromoform 	0.10
 Chlorobenzene 	0.30
 Chloromethane 	0.10
• 1,1-Dichloroethane	0.10
• 1,1,2,2-Tetrachloroethane	0.30

RRFs for compounds (VOC and SVOC) without specified minimum RRFs will be >0.050.

Evaluation	Action
If the average RRF for any target compound is < the specified minimum	qualify all associated detects as "J" and all associated non-detects as "UJ" if the
RRF, or <0.050 if no minimum is	average RRF is ≥0.010 and as "R" if the
specified,	average RRF is <0.010.
If the %RSD for any target compound is	
>15% but ≤40%,	qualify all associated detects as "J" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>40% but ≤60%,	qualify all associated detects as "J" and all associated non-detects as "UJ."
>60%,	qualify all associated detects as "J" and all associated non-detects as "R."

Linear Curves

Criteria: The coefficient of determination (r2) of the ICAL curve shall be ≥0.990 and have a slope ≥ the method-specified minimum RRF for each compound. Compounds without method-specified minimum RRFs shall have a slope ≥ 0.050 . The absolute value of the intercept shall be $\leq 3X$ the MDL.

Note: The intercept reported in the instrument calibration report may not be in appropriate units. When the intercept is not in appropriate units, the instrument conversion routine may be needed to evaluate the intercept.

Depending on the laboratory instrumentation, the intercept will be calculated differently. The instrument software used for determining calibration curves often differs from the usual y = mx+b linear equation. When this occurs, it is up to the validator to determine the actual equation used and the corresponding slope and intercept.

For calibrations using most commercial data system software, the intercept in concentration units is calculated using one of the following equations:

Concentration Intercept = $(b)(C_{IS})$

Concentration Intercept = $(-b/m)(C_{IS})$

Where:

b = reported intercept

m = slope of the curve

 C_{IS} = concentration of IS (on-column conc. on quant. report)

Evaluation	Action
If the slope for any target compound is < the minimum RRF, or <0.050 if no minimum is specified,	qualify all associated detects as "J" and all associated non-detects as "UJ" if the slope is ≥0.010 and as "R" if the slope is <0.010.
If the r ² for any target compound is	
<0.990 but ≥0.90,	qualify all associated detects as "J" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
$<0.90 \text{ but } \ge 0.80,$	qualify all associated detects as "J" and all associated non-detects as "UJ."
<0.80,	qualify all associated detects as "J" and all associated non-detects as "R."
If the intercept for any target compound is positive and > the MDL,	qualify all associated detects <3X the intercept as "J+."
When results are reported at the MDL:	
If the intercept for any target compound is negative with an absolute value	
> the MDL but $\leq 3X$ the MDL,	qualify all associated detects <3X the absolute value of the intercept as "J-" and all associated non-detects as "UJ."
>3X the MDL,	qualify all associated detects <3X the absolute value of the intercept as "J-" and all associated non-detects as "R."

IMPORTANT NOTICE:

4.2.3 Calibration Verification

Compliance requirements for satisfactory initial and continuing instrument calibration verification are established to ensure that the instrument is capable of producing acceptable qualitative and quantitative data for compounds on the TAL. Initial calibration verification (ICV) independently verifies the calibration and continuing calibration verification (CCV) establishes the 12-hour relative RRFs on which the quantitations are based and checks satisfactory performance of the instrument on a day-to-day basis.

Criteria: An ICV standard must be analyzed immediately following an ICAL.

The ICV standard analysis results are not required to be reported in the data package unless the samples in the SDG were analyzed after the ICAL standard but before a CCV standard analysis was performed. In this case, the ICV percent difference (%D) is assessed according to the calibration verification criteria described below for the associated samples. If a CCV is analyzed prior to samples and ICV data are also reported in the package, both the ICV %D and the appropriate CCV %D are to be assessed as described below. If both ICV %D and CCV %D infractions occur, the worst infraction should be evaluated for result qualification. A CCV standard must be analyzed:

- (1) if analysis continues for longer than 12 hours, and
- (2) at the beginning of each additional 12-hour period.

The laboratory is allowed to perform corrective action and reanalyze the CCV once after a failure. If more than two CCVs were analyzed to obtain a passing CCV, then the calibration was not verified and the calibration verification frequency criteria were not met.

Evaluation	Action
If the ICV and CCV standards were not analyzed at the proper frequency, or if either a required ICV or CCV was not analyzed, or if all target compounds were not present in any ICV or CCV standard,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If all required ICVs and CCVs were not analyzed,	qualify all associated detects as "J" and all associated non-detects as "R."

RRFs

Criteria: The %D between the ICV and/or CCV RRFs and the average RRFs obtained from the ICAL shall be calculated according to the formula in Section 6.3 and must be ≤20%.

IMPORTANT NOTICE:

Evaluation Action If the %D was reported with the wrong sign document the occurrence in the data (e.g., + %D for a negative bias), validation report and assess any infractions using the correct sign. If the %D between an ICAL RRF and an ICV or CCV RRF for any target compound is... qualify all associated detects as "J+." >20% and positive (high bias), qualify all associated detects as "J-" and, >20% but $\le 40\%$ and negative (low bias), if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ." qualify all associated detects as "J-" and >40% but $\leq 60\%$ and negative, all associated non-detects as "UJ." qualify all associated detects as "J-" and >60% and negative, all associated non-detects as "R."

Procedure No: AOP 00-03

Issue No: 05

Linear Curves

Criteria: The %D (see Section 6.3) between the ICV and/or CCV standard concentrations and their true values must be $\leq 20\%$.

Evaluation	Action
If the %D was reported with the wrong sign (e.g., +%D for negative bias),	document the occurrence in the data validation report and assess any infractions using the correct sign.
If the %D between the measured ICV and/or CCV concentrations and their true values for any target compound is	
>20% and positive (high bias),	qualify all associated detects as "J+."
>20% but ≤40% and negative (low bias),	qualify all associated detects as "J-" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ.

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home page.

Evaluation (concluded)	Action (concluded)
>40% but ≤60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
>60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "R."

4.2.4 Blanks

The purpose of laboratory (or field) blank analysis is to determine the nature and magnitude of contamination resulting from laboratory (or field) activities.

The criteria for evaluation of blanks apply to any blank associated with the samples and include MBs, and, if submitted, EBs, FBs, and TBs. Action in the case of unsuitable blank results depends on the circumstances and origin of the blank. In instances where more than one blank is associated with a given sample, qualification should be based upon a comparison with the associated blank having the highest concentration of a contaminant. For purposes of evaluating multiple blanks, each preparation batch may be considered an independent event in evaluating MBs, and each sampling event may be considered an independent event for evaluating EBs, FBs, and TBs.

Criteria: The concentration of each target analyte found in the blank must be < the associated MDL. The sample results must not be corrected by subtracting any blank value. If QC problems with any blank exist, all data associated with the case must be carefully evaluated to determine whether there is an inherent bias in the data, or if the problem is an isolated occurrence not affecting other data.

Evaluation	Action
If a compound detected in a blank is also detected in a field sample,	qualify the sample result for that compound in accordance with the scenarios given below.
If gross contamination (e.g., saturated peaks by GC/MS) exists,	qualify results for all affected compounds as "R" due to interference.
If inordinate numbers of target compounds are found at low levels in the blank(s),	Discuss the presence of these compounds in the data validation report as it may be indicative of a problem at the laboratory. Note: Similar consideration should be given to tentatively identified compounds (TICs) that are found in both the sample and its associated blank(s) (see Section 4.2.13).

IMPORTANT NOTICE:

Blank Type	Blank Result	Sample Result	Action
Method, Storage,	Detect	Non-detect	No qualification.
Field, Trip, Equipment	≤PQL	≤PQL	Qualify as non-detect "U" at PQL.
		>PQL but ≤5X the blank value or >PQL but ≤10X the blank value for CLCs*	Qualify "J+."
	>PQL	≤PQL	Qualify as non-detect "U" at PQL.
		>PQL but ≤2X blank value	Qualify as non-detect "U" at sample result and request corrective action from laboratory.
		>PQL and >2X blank value but ≤5X blank value or > PQL but ≤10X blank value for CLCs*	Qualify "J+" and request corrective action from laboratory.
	Gross Contamination	Detect	Report at sample result and qualify "J+" or "R", based on professional judgment and request corrective action from laboratory.

*CLCs – common laboratory contaminants – Acetone, Methylene chloride, Toluene, 2-Butanone and common phthalate esters [e.g., bis(2-ethylhexyl)phthalate and di-n-octyl phthalate]. It should be noted that toluene and the phthalate esters are no longer included in the NFG revisions as CLCs but are included in this document due to the possibility of contamination from these analytes occurring during field sampling.

Gross contamination is not specifically defined but will be evaluated using professional judgment on a case by case basis. The infraction may be a high concentration of a single analyte or low-level contamination involving several analytes in the blank.

NOTE: In some instances, the laboratory may adjust the MDLs to account for low-level common laboratory contaminants. In these cases, it may be possible to have a low-level detection in a blank that would be considered a non-detect when compared to the adjusted MDL, resulting in the blank data being reported as a non-detect (PQL U). This may result in sample results that are above the MDL but <5X or 10X the actual blank concentration not being qualified. In instances where it is believed that there is low-level contamination of common laboratory contaminants that are not identified in the blank, the sample results may be qualified as "NJ" based on professional judgment and narrated in the summary.

IMPORTANT NOTICE:

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

4.2.5 Surrogate Recovery

Laboratory performance for individual samples is evaluated by means of surrogate spikes. All samples are spiked with surrogate compounds prior to sample preparation. The evaluation of the results of these surrogate spikes is not necessarily straightforward. The sample itself may produce effects due to such factors as interference and high concentrations of analytes. Because the effects of the sample matrix are frequently outside the control of the laboratory and may present relatively unique problems, the review and validation of data based on specific surrogate results is frequently subjective and demands analytical experience and professional judgment. In addition, surrogate recoveries can be influenced by the success in recoveries of the ISs. The evaluation of surrogate recoveries and ISs should be done concurrently. Accordingly, this section consists primarily of guidelines, in some cases with several optional approaches suggested.

Criteria: Sample and blank surrogate recoveries must be within limits specified by the laboratory. Surrogate compound recoveries shall be calculated using the procedure described in SW-846 Method 8000C. Reported recoveries shall be accompanied by the applicable acceptance limits. No qualification with respect to surrogate recovery is placed on data unless one or more of the following occurs:

- 1) one surrogate is out of specification in the base/neutral fraction or acid fraction (SVOC analysis),
- 2) one surrogate is out of specification in the volatile fraction (VOC analysis), or
- 3) any surrogate has < 10 %R.

Under these three conditions, there should be a reanalysis.

Note: The common acid fraction analytes (SVOC) are all phenols; all cresols; benzoic acid; dichlorophenoxyacetic acid; dinoseb; and hexachlorophene.

Note: When there are unacceptable surrogate recoveries followed by successful reanalysis, the laboratories are required to report only the successful run.

See Appendix C for general guidelines for surrogate recovery limits.

Note: Results from spiked or replicate QC samples that have surrogate recoveries < 10% cannot be used to evaluate associated sample results. Sample results should be qualified for lack of accuracy and/or precision data, as applicable, if specified by the program.

Evaluation	Action
If surrogate recovery acceptance criteria were not reported in the data package,	request amended data from the laboratory.

Data Validation Procedure for Chemical and Radiochemical Data Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation (continued)	Action (continued)
If, based on professional judgment, the laboratory's internal acceptance criteria are excessively wide or biased,	notify the program manager.
If an initial dilution was performed on any sample and at least one surrogate recovery is < the lower acceptance limit but ≥10%, or all surrogate recoveries are <10% and the results for one or more compounds are ≥ the PQL,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
If an initial dilution was performed on any sample, all surrogate recoveries are <10%, and all results are non-detect,	qualify all associated sample results as "R."
If there are two or more analyses for a particular fraction at the same dilution,	determine which analysis contains the best data to report using the considerations below, qualify all data from the rejected analysis as "R," and document the reason for rejecting data from one analysis in the data validation report. Considerations should include: 1. surrogate recovery (marginal vs. gross deviation); 2. holding times; 3. comparison of the values of the target analytes reported in each fraction; and 4. performance of ISs.
For surrogate recoveries out of specification, the following approaches are suggested based on a review of all data from the batch, especially considering the apparent complexity of the sample matrix:	
if at least one surrogate in the base/neutral fraction, one surrogate in the acid fraction, or one surrogate in the volatile fraction, are out of specification low but have recoveries ≥ 10%,	qualify all detects for that fraction as "J-" and all non-detects for that fraction as "UJ."

Evaluation (concluded)	Action (concluded)
if any surrogate recovery in a fraction is <10%,	qualify all detects for that fraction as "J-" and all non-detects for that fraction as "R."
if at least one surrogate in the base/neutral, one surrogate in the acid fraction, or one surrogate in the volatile fraction, are out of specification high,	qualify all detects for that fraction as "J+."

Criteria: In the case of a blank analysis with surrogates out of specification, special consideration must be given to the validity of associated sample data. The basic concern is whether the blank problems represent an isolated problem with the blank alone or whether there is a fundamental problem with the analytical

process.

If one or more samples in the batch show acceptable surrogate recoveries, the blank problem may be considered an isolated occurrence. However, even if this judgment allows some use of the affected data, analytical problems remain that must be corrected by the laboratory.

Evaluation	Action
If surrogate recovery in the blank does not meet acceptance criteria,	all detects < the PQL in all samples associated with the blank may be qualified as "J" and all non-detects in all samples associated with the blank may be qualified as "UJ."

4.2.6 IS Performance

IS criteria ensure that GC/MS sensitivity and response are stable and acceptable during each analysis.

Criteria: Sample and blank IS results must be within limits given in the specific SW-846 method.

> IS area counts must not vary by more than a factor of two (50% to 200%) from the average of those obtained from the calibration standards.

The retention time (RT) of the IS must not vary more than ± 30 seconds from that of the associated calibration standard.

When qualification of sample results is warranted due to failure of an IS to meet RT or area count acceptance criteria, results of all target compounds associated with that IS are qualified.

Refer to Appendix D for IS/target compound correlation guidelines.

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation	Action
If there are two analyses for a particular fraction and internal standard failures occur,	determine which analysis contains the best data to report using the considerations below, qualify all data from the rejected analysis as "R," and document the reason for rejecting the data from one analysis in the data validation report.
	Considerations should include: 1. magnitude of the RT shift; 2. holding times; 3. comparison of the values of the target compounds reported in each fraction; and 4. surrogate recovery.
If any IS area count is <50% of the average of that obtained from the calibration standards,	qualify all associated detects as "J+" and all associated non-detects as "UJ." Non-detects may be qualified as "R" based on professional judgment if the IS area counts are <20% of that of the average obtained from the calibration standards. Note: If extremely low area counts are reported, or if performance exhibits a major abrupt drop-off, then a severe loss of sensitivity is indicated.
If the IS area count is >200% of the average of that obtained from the calibration standards,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
If the IS RT varies by more than ±30 seconds from that of the associated CCV standard,	qualify all associated detects as "N" or "R" and all associated non-detects as "R."

4.2.7 MS/MSD

Data for MS/MSD are generated to determine long-term precision and accuracy of the analytical method on samples of various matrices and to demonstrate acceptable compound recovery by the laboratory at the time of sample analysis.

Criteria: The MS/MSD data shall not be used to qualify field sample results unless the MS/MSD sample was from the same client and of similar matrix.

> An MS and MSD sample shall be analyzed at a frequency of once per data package, once per 20 samples of similar matrix, or once per sample matrix, whichever is more frequent.

The laboratory shall not use FBs, EBs, or TBs to satisfy these requirements, if the laboratory can identify these blanks.

Unless otherwise stated in the specific method, the MS and MSD accuracy and precision acceptance criteria shall be those calculated by the laboratory using the procedure given in SW-846 Method 8000C. If the acceptance criteria are not given, recovery limits of 70% to 130% and ±30% RPD should be used as the criteria. It may be appropriate to use wider default recovery acceptance criteria for SVOC analysis based on professional judgment. For solid and waste samples, it may be appropriate to accept up to a 40% RPD based on the professional judgment. The MS and MSD %R must be within the acceptance limits, unless the sample concentration is > 4X the spike concentration (see Section 4.1.20).

The MS and MSD analyses must meet all sample analysis acceptance criteria. An effort to determine to what extent the results of the MS/MSD affect the associated data should be made. This determination should be made considering the MS/MSD sample matrix, the surrogate recoveries, and the LCS results.

Professional judgment should be used to determine whether MS/MSD failure warrants qualification of only the results for the failed compounds, or if results for all the compounds associated with the failed MS compound and its associated IS are affected. Generally, unless evidence exists to warrant qualification of other compounds, only the compounds in the MS spiking mixture shall be qualified.

For programs that require application of one final qualifier to sample results, if a recovery (accuracy) infraction is identified in one or both of the MS samples along with an RPD (precision) infraction between the MS and MSD, the sample is qualified for the accuracy infraction. For example, if a compound has low MS recovery and the RPD is not within criteria, the data are qualified as "J-."

Evaluation	Action
If the MS/MSD analysis was from another client or of a dissimilar matrix; if the frequency of the MS/MSD did not meet specified criteria; if no MS/MSD was analyzed; or if FB, EB, or TB samples were used for MS/MSD purposes,	qualify all detects as "J" and all non- detects as "UJ."

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation (concluded)	Action (concluded)
If no other measure of precision (i.e., LCSD or replicate) is available,	qualify all detects as "J" and all non- detects as "UJ."
If the surrogate, IS, and LCS %Rs are within the required acceptance criteria and either the MS or MSD %R for any target compound is > the upper acceptance limit,	qualify all associated detects as "J+."
either the MS or MSD %R for any target compound is < the lower acceptance limit,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the recovery is ≥20% and as "R" if the recovery is <20%.
If the RPD for any target compound does not meet the acceptance criteria or %Rs fail both high and low,	qualify all associated detects for that compound as "J" and all associated non-detects as "UJ."

Note: The laboratory may analyze TBs in a separate batch than that of soil samples due to differences in sample matrices. In this situation, the laboratory may not analyze an MS/MSD for the batch associated with the TBs. The TB results should then be assessed for accuracy and precision using an LCS/LCSD.

4.2.8 Replicate

Replicate analyses are indicators of laboratory precision based on each sample matrix. If a replicate was performed instead of an MSD, the following criteria are applied. If insufficient sample was submitted to analyze an MS/MSD or replicate, the laboratory may run a LCSD to measure precision. LCSD precision shall be assessed as described in Section 4.2.7.

Criteria: A replicate sample shall be analyzed at a frequency of once per data package, once per 20 samples of similar matrix, or once per sample matrix, whichever is more frequent. All sample acceptance criteria must be met in the replicate analysis.

> Samples identified as FBs, EBs, or TBs should not be used for replicate sample analysis.

> Unless otherwise stated in the specific method, the replicate precision acceptance criteria shall be those calculated by the laboratory using the procedure given in SW-846 Method 8000C. When no laboratory-derived control limits are reported, a control limit of 30% for the RPD shall be used for

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

> sample values $\geq 5X$ the PQL. For solid and waste samples, it may be appropriate to accept up to a 40% RPD based on the professional judgment.

> A control limit of \pm the PQL shall be used for sample values < 5X the PQL, including the case when only one of the replicate sample values is <5X the PQL.

No precision criteria apply when both replicate sample values are < the PQL.

Evaluation	Action
If no replicate sample, no MSD, and no LCSD were analyzed for each matrix or for each data package,	qualify all associated detects of the same matrix as "J" and all associated non- detects of the same matrix as "UJ."
If an FB, EB, or TB was used for the replicate analysis and no MSD or LCSD was run,	qualify all associated detects of the same matrix as "J" and all associated non- detects of the same matrix as "UJ."
If the original result and replicate result for target compound are both ≥5X the PQL, and the RPD exceeds the appropriate control limit,	qualify all associated detects of the same matrix as "J" and all associated non-detects of the same matrix as "UJ."
If the original and/or replicate result for any target compound is <5X the PQL (including non-detects) and the difference between the original result and replicate result is > the PQL,	qualify all associated detects of the same matrix as "J" and all associated non-detects of the same matrix as "UJ."

4.2.9 LCS

Data for LCSs are generated to provide information on the accuracy of the analytical method and the overall laboratory performance, including sample preparation.

Criteria: An LCS should be analyzed for all methods at a frequency of once per data package, once per matrix, or once per 20 analytical samples, whichever is most frequent. The LCS should have recoveries for all target analytes; however, for very large analyte lists or for known poor performers, the laboratory may have received an exemption for one or more analytes.

> The LCS must meet all sample acceptance criteria. If the MS/MSD and the samples meet all QC acceptance criteria, but the surrogate and/or IS acceptance criteria are not met in the LCS analysis, the LCS must be reanalyzed. The LCS should meet all method-specific LCS requirements and acceptance criteria. If the recovery acceptance criteria are not reported, the reviewer should use the criteria in Appendix E, or 70% to 130% for evaluation.

IMPORTANT NOTICE:

If the laboratory analyzed an LCS/LCSD as a measure of precision both the LCS and LCSD must meet recovery acceptance criteria.

General laboratory precision and accuracy can be evaluated using the LCS acceptance criteria and the interlaboratory comparison data given in Appendix E. Individual LCS recoveries may be evaluated against the criteria in Appendix E if the laboratory's criteria are significantly different from those in the tables.

For volatile organics in an aqueous matrix, a successful **second source** CCV meets the LCS requirements.

Evaluation	Action
If, based on professional judgment, the laboratory's internal acceptance criteria are excessively wide or significantly biased,	notify the program manager.
If the frequency of the LCS did not meet the specified criteria,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If results are reported for target compounds that are not in the LCS,	detects for those compounds may be qualified as "J" and non-detects for those compounds may be qualified as "UJ" based on professional judgment. Compounds missing under an exemption may be qualified based on professional judgment.

If the LCS criteria were not met and reanalysis was not performed, then laboratory performance and method accuracy are in question. Professional judgment should be used to determine whether data should be qualified for all target compounds or just those compounds associated with the failed LCS compound and its associated IS. The following may be used as guidance in qualifying data.

If a full or large TAL LCS is analyzed, the following criteria may be used for LCS %Rs which fall outside reported acceptance criteria but are >10%:

70 to 74 compounds	\leq 5 LCS fall outside acceptance criteria - no qualification
60 to 69 compounds	≤ 4 LCS fall outside acceptance criteria - no qualification
50 to 59 compounds	≤ 3 LCS fall outside acceptance criteria - no qualification
40 to 49 compounds	≤2 LCS fall outside acceptance criteria - no qualification
30 to 39 compounds	≤ 1 LCS fall outside acceptance criteria - no qualification
< 30 compounds	No LCS fall outside acceptance criteria - no qualification

Evaluation	Action
If the LCS %R is > the upper acceptance limit,	qualify all associated detects as "J+."
If the LCS %R is < the lower acceptance limit,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the %R is ≥20% and as "R" if %R is <20%.
If %Rs for more than half of the compounds in the LCS analysis are below the acceptance range,	qualify all detects as "J-" and all non- detects as "UJ" if the failures are marginally low and as "R" if %Rs are significantly below acceptance limits. Note: If recoveries of more than half of the compounds in the LCS analysis are below the acceptance range, the laboratory has not shown that it can actually meet program required DLs.
If %Rs for more than half of the compounds in the LCS analysis are above the acceptance range,	qualify all detects as "J+."
If %Rs for more than half of the compounds in the LCS analysis are outside the acceptance range, both above and below, or if an LCS/LCSD pair was analyzed and recoveries of any target compound are both above and below acceptance criteria,	qualify all detects in all associated samples as "J" and all non-detects in all associated samples as "UJ."

4.2.10 Sample Carry-over

Sample carry-over may occur when a high-concentration sample is analyzed immediately prior to another field sample. Steps must be taken to avoid introduction of false positive results in the second sample analysis due to instrument contamination.

Criteria: The absence of sample carry-over must be determined and verified. If examination of the run logs indicates that any samples in the analytical run of interest required dilution, and there is no documentation of a rinse or blank analysis immediately following the original undiluted analysis then sample carry-over may be suspected in the subsequent sample.

following sample at a concentration <5X

Evaluation Action If any target compound found in the qualify the result for that compound in sample requiring dilution exceeded the the second sample as "R" or "NJ" based high calibration standard and was also on professional judgment. found in the following sample at a concentration $\leq 5X$ the PQL, qualify the results for that compound in If no data are available for the sample that required dilution and the laboratory has not the second sample as "N." documented that carry-over was evaluated, and the compound was also found in the

Procedure No: AOP 00-03

Issue No: 05

4.2.11 Dilutions

the PQL,

Criteria: The PQLs must be adjusted to reflect all sample dilutions, concentrations, splits, clean-up activities, and dry weight factors that are not accounted for by the method.

Samples must be diluted and reanalyzed when any analytes exceed the calibration range.

Data from original samples should be included when any sample requires dilution due to one or more compounds exceeding the calibration range.

The original undiluted results document the actual MDLs for non-detects.

Evaluation	Action
If the PQLs have not been properly adjusted,	request an amended report from the laboratory.
In some cases, initial dilutions are required because of expected high concentrations of non-target analytes, or because one or more target analyte is expected to greatly exceed the instrument working range. In these instances, the laboratory may not be able to analyze the undiluted sample.	note the dilution and elevated MDLs in the data validation report.
If any target compound exceeds the calibration range and	
the original undiluted sample result was reported,	qualify all detects from the undiluted analysis that exceeded the calibration range as "J."

Data Validation Procedure for Chemical and Radiochemical Data Effective Date: 06/19/2017

Evaluation (concluded)	Action (concluded)
the sample was diluted and reanalyzed, and the diluted sample data were reported,	qualify all non-detects from the diluted analysis as "UJ."
the original undiluted sample data were not provided,	request this information from the laboratory.
If data from the original sample run are unavailable,	refer to Section 4.2.5 for assessment of initially diluted samples with low surrogate recovery.

Criteria: The laboratory shall strive to make dilutions in such a way that the final concentration is measured in the mid-range of the calibration curve and that results are not reported from measurements below the lowest concentration standard.

Evaluation	Action
If the concentration (reported result /	qualify all associated detects from the
dilution factor) from a diluted sample is <	diluted analysis as "J."
that of the lowest concentration standard,	

Criteria: The extraction efficiency for extremely high concentrations of analytes has generally not been determined for most methods. If the analysis requires an extraction and dilutions of >100,000:1 the efficiency of the extraction may be suspect.

Evaluation	Action
If dilutions of >100,000:1 was required,	qualify all associated detects as "J"

4.2.12 Mass Spectra Acceptability

Mass spectra review is typically outside the scope of routine data validation and should not be performed unless it is specifically requested by the SMO. When mass spectra review is required by the program, it must be performed by a validator experienced in the interpretation of mass spectra.

The laboratory is to identify mass spectra using either the National Bureau of Standards (NBS) /EPA/Mass Spectrometry Data Centre (MSDC) library or the National Institute of Standards and Technology (NIST)/EPA/National Institutes of Health (NIH) library. The laboratory must identity and document peaks and reference spectra for all target compounds with concentrations above the MDL. While it is not the function of the validator to determine whether the analyst correctly

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home page.

identified a compound, an evaluation of how well the analyte peak matches the reference spectra may be requested. To evaluate analyte spectra, the guidelines in Appendix F shall be used.

Evaluation	Action
If the laboratory does not identify mass spectra using a nationally recognized standard library,	notify the program manager.
If the sample spectrum does not match the reference spectrum, the extracted ion current profiles (EICPs) RT or relative retention time (RRT) does not meet criteria, or several guideline failures were observed,	qualify all associated results as "R."
If the analyte is not identifiable due to gross interference or apparent instrument instability,	qualify all associated results as "R."
If the analyte was misidentified by the laboratory,	request an amended report from the laboratory.
If identification of the analyte was hampered by interferences such that it is not certain that a positive identification could be made,	qualify all associated results as "N" based on professional judgment or request additional data from the laboratory.

4.2.13 TICs

Chromatographic peaks that are not target analytes, surrogates, or ISs are potential TICs. TIC evaluation is typically outside the scope of routine data validation and should not be performed unless specifically requested by the SMO. When TIC evaluation is required by the program, it must be performed by validators with experience in mass spectra interpretation.

Criteria: For each sample, the laboratory may be requested to conduct a mass spectral search of either the NBS/EPA/MSDC library or the NIST/EPA/NIH library. The laboratory may report the possible identity for up to 20 of the largest VOC fraction peaks and the 20 largest SVOC fraction peaks which are not surrogate, IS, or target compounds, but which have an area/height >10% of the size of the area/height of the nearest IS.

It should be noted that common laboratory artifacts/contaminants and their sources (i.e., aldol products, solvent preservatives/reagent contaminants, etc.) may be present in blanks and not reported as sample TICs.

Examples:

Common laboratory contaminants: CO₂ (mass/charge ratio (m/e) 44), siloxanes (m/e73), diethyl ether, hexane, certain freons (1,1,2-trichloro-1,2,2-trifluoroethane or fluoro-trichloromethane), phthalates at levels < 100 micrograms per liter (ug/L) or 4,000 micrograms per kilogram (ug/kg).

- Solvent preservatives: cyclohexene is a methylene chloride preservative. Related by-products include cyclohexanone, cyclohexanone, cyclohexanol, cyclohexenol, chlorocyclohexene, chlorocyclohexanol.
- Aldol reaction products of acetone include 4-hydroxy-4-methyl-2-pentanone, 4-methyl-2-penten-2-one, 5,5-dimethyl-2(5H)-furanone.

Evaluation	Action
If a low-level non-target compound that is a common artifact or laboratory contaminant is detected in a sample,	verify that TIC peaks present in samples are not found in blanks. Blank chromatograms should be examined for peaks that are <10% of the IS height but are present in the sample chromatogram at similar RRTs.
If sample TIC results are not sufficiently above the level in the blank and the results are reported,	the results may be qualified as "R" (dilutions and sample size must be taken into account when comparing the amounts present in blanks and samples).
If a result is identified as a TIC, If a compound is not found in any blanks, but is a suspected artifact or common laboratory contaminant,	qualify that result as "NJ." identify the compound as such in the data validation report. Compounds that are suspected artifacts or common laboratory contaminants result may be qualified as "R" based on professional judgment.

It should be noted that common laboratory calibration practices, along with limitations of some commercial software could result in compounds being detected and not reported in either the Form I or the TIC Summary Report. Review all quantitation reports to verify that all detected compounds are reported whenever a TIC Summary is included.

Evaluation	Action
If a compound is identified on the	request a corrected report from the
quantitation report but are not reported as	laboratory.
target detect or as a TIC,	

4.2.14 Method-specific Analytical Requirements-Organic GC/MS

The additional analytical requirements addressed below are organized by SW-846 Method. These requirements should be checked if the level of deliverable (level III or level IV) allows.

4.2.14.1 Method 8260B or 8260C, VOC Analysis by GC/MS

Criteria: The analysis of 2-chloroethyl vinyl ether in water must be performed on an unacidified sample.

Evaluation	Action
If 2-chloroethyl vinyl ether was reported	qualify all associated detects as "NJ-"
for an acidified water sample,	and all associated non-detects as "R."

4.2.14.2 Method 8270C or 8270D, SVOC Analysis by GC/MS

Criteria: Gel permeation chromatography (GPC) cleanup shall be used as necessary to eliminate interferences. In addition, all water samples containing high molecular weight compounds that interfere with the analysis of the target compounds must also undergo GPC cleanup.

Evaluation	Action
If the runlog notations, spectral data, IS %Rs, or surrogate %Rs indicate potential interferences,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If appropriate extract cleanup was not performed,	note this on the data validation report and notify the program manager.

4.2.14.3 Method 8280B, Polychlorinated Dioxins and Furans by High Resolution Gas Chromatography/Low Resolution Mass Spectrometry (HRGC/LRMS)

Sample analysis shall be performed according to the requirements listed in SW-846 Method 8280B. Evaluation of tuning reports is not required for this method.

Criteria: Initial calibration shall be performed using the five calibration solutions listed in Table 1 of the method. The %RSD for the ISs and the target compounds for the five calibration standards must be <15%.

Calibration verification shall be performed using the standards solution given in Table 4 of the method. The calibration verification analysis must meet the criteria given in Section 11.13.3.6 of the method.

Evaluation	Action
If the %RSD is >15% for any IS or target	qualify all associated detects as "J" and
compound,	all associated non-detects as "UJ."

Evaluation (concluded)	Action (concluded)
If the CCV acceptance criteria were not	qualify all associated detects as "J" and
met for any target compound,	all associated non-detects as "UJ."

Sample Analysis

Criteria: For identification of any compound, the ion abundance ratios must be within the limits specified in Table 9 of the method.

> For 2,3,7,8-substituted compounds that have an isotopically labeled IS or recovery standard present in the sample extract, the RT must be -1 to +3seconds of the isotopically labeled standard. For 2,3,7,8-subtituted compounds that do not have an isotopically labeled IS or recovery standard present in the sample extract, the RT must fall within 0.005 RRT units of the RRT measured in the continuing calibration.

> For non-2,3,7,8-substituted compounds, the RT must be within the corresponding homologous RT windows established by analyzing the column performance check solution.

Evaluation	Action
If ion abundance ratio criteria were not met for any compound,	qualify all associated results as "R."
If the RT of any compound is outside of the RT window,	qualify all associated results as "R."

Criteria: IS %R for analytical samples must be \geq 20% and <150%. IS recovery guidelines are discussed in Section 11.15.5 of the method.

> The LCS shall contain all of the target compounds at concentrations near the midpoint of the calibration range.

Evaluation	Action
If the recovery of any IS solution compound is >150%,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
If the recovery of any IS solution compound is <20%,	qualify all associated detects as "J+" and all associated non-detects as "UJ" if the recovery is ≥20% and as "R" if the recovery is <20%

Evaluation (concluded)	Action (concluded)
If results are reported for target compounds	qualify all associated detects as "J" and
that are not in the LCS,	all associated non-detects as "UJ."

GC Column Performance

Criteria: The GC column performance solution is used for defining the homologous GC RT windows and to document the chromatographic resolution. Column performance must be evaluated at the beginning of each 12-hour analytical period and must meet method acceptance criteria (see Section 11.12 of the method) before sample analysis may begin.

Evaluation	Action
If GC column performance was not evaluated at the required frequency or if method criteria were	l * *
not met,	"UJ."

Confirmation of 2,3,7,8-Tetrachlorodibenzofuran (TCDF) Detects

Criteria: The DB-5 GC column generally used for polychlorinated dibenzodioxin (PCDD) and polychlorinated dibenzofuran (PCDF) analyses does not adequately separate 2,3,7,8-TCDF from its closest eluting isomer. If 2,3,7,8-TCDF is detected in a sample, the result must be confirmed on a second column capable of separating 2,3,7,8-TCDF from all other TCDF homologues (as proven by successful analysis of the GC column performance mix with <25% valley between 2,3,7,8-TCDF and its closest eluting isomer).

Evaluation	Action
If 2,3,7,8-TCDF was detected in a sample and the result was not confirmed on a second column with successful analysis of the GC column performance mix,	qualify all associated detects as "NJ."

4.2.14.4 Method 8290A, PCDD and PCDF by HRGC/High Resolution Mass Spectrometry (HRMS)

Initial Calibration

Criteria: A 5-point calibration is prepared for each labeled and unlabeled compound. The relative response factor (RRF) %RSD for the unlabeled standards must be

IMPORTANT NOTICE:

 \leq 20%. For the labeled compounds, the %RSD must be \leq 30%. Ion abundance ratios must meet the criteria listed in Table 8 of the method.

Evaluation	Action
If the %RSD is:	
>20% for any unlabeled calibration standard or >30% for any labeled calibration standard, but \leq 40%,	qualify all associated detects as "J" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>40% but <60% for either a labeled or unlabeled calibration standard,	qualify all associated detects as "J" and all associated non-detects as "UJ."
>60% for either a labeled or unlabeled calibration standard,	qualify all associated detects as "J" and all associated non-detects as "R."
If the ion abundance criteria is not met for any compound,	qualify all associated detects as "J" and all associated non-detects as "UJ."

Continuing Calibration

Criteria: Calibration must be verified for both unlabeled and labeled compounds at the beginning and end of each 12-hour shift during which analysis is performed.

> The measured RRFs must be ≤20% of the mean values established during ICAL for unlabeled compounds and $\leq 30\%$ of the mean values established during ICAL for labeled compounds. The ion abundance must be within the limits in Table 8 of the method.

> For the calibration verification analyzed at the end of a 12-hour period, a %D of 25% for unlabeled compounds and 35% for labeled compounds is acceptable. If the criteria are exceeded for any analytes, the mean RRFs obtained from the beginning and ending daily calibration verification runs are used to calculate those analyte concentrations instead of the RRFs obtained from the initial calibration. Alternatively, if the %D of the ending calibration verification is >25% for any unlabeled compound and/or >35% for any labeled compound, then successful performance of another initial calibration may be started within two hours of sample analysis and the RRFs obtained from the new initial calibration are used to calculate analyte concentrations for samples bracketed by the failed calibration verification run.

Evaluation	Action
If the ion abundance ratio for any compound is outside of the method limits,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If the %D criteria are not met for any CCV compound at the beginning of a 12-hour shift, and	
the %D is positive,	qualify all associated detects as "J+."
the %D is negative,	qualify all associated detects as "J-" and if any other calibration criteria have been exceeded for that compound, qualify all associated as "UJ."
If the %D criteria were <u>not</u> met for any target compound at the end of a 12-hour shift, and	
the %D is positive, the %D is negative,	qualify all associated detects as "J+." qualify all associated detects as "J-" if any other calibration criteria have been
	exceeded for that compound, qualify all associated non-detects as "UJ."
If the %D criteria were not met for any target compound at the end of a 12-hour shift and a new successful ICAL was started within two hours of sample analysis,	ensure that data were quantitated using the new ICAL. Qualification of the data is not required.

Sample Preparation

Criteria: Extract cleanup shall be performed to eliminate interferences. The laboratory shall first partition the sample extract, followed by silica/alumina column cleanup and carbon column cleanup.

Evaluation	Action
If the documentation on the run log, spectra data, and/or IS or labeled compound %Rs indicate interferences and extract cleanup was not performed,	qualify all associated detects as "J" and all associated non-detects as "UJ" and notify the program manager.

IMPORTANT NOTICE:

Data Validation Procedure for Chemical and Radiochemical Data Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Sample Analysis

Criteria: For identification of any compound, the ion abundance ratios must be within the limits specified in Table 8 of the method.

> For 2,3,7,8-substituted compounds which have an isotopically labeled IS or recovery standard present in the sample extract, the RT must be -1 to +3 seconds of the isotopically labeled standard. For 2,3,7,8-substituted compounds that do not have an isotopically labeled IS or recovery standard present in the sample extract, the RT must fall within 0.005 RRT units of the RRT measured in the continuing calibration.

For non-2,3,7,8-substituted compounds, the RT must be within the corresponding homologous RT windows established by analyzing the column performance check solution.

Evaluation	Action
If ion abundance ratio criteria are not met for any compound,	qualify all associated results as "R."
If the RT of any compound is outside of the RT window,	qualify all associated results as "R."

Mass Spectrometer Performance Criteria

Criteria: Performance criteria are established to ensure mass resolution, identification and, to some degree, sensitivity. These criteria are not sample specific.

> Conformance is determined using standard materials. These criteria should be met in all circumstances. Mass spectrometer performance must be checked at the beginning and end of each analytical period in accordance with the method criteria (see Section 9.3 of the method).

Evaluation	Action
If mass spectrometer performance was not checked at the required frequency or if	qualify all associated detects as "R" and all associated non-detects as "UJ."
method criteria were not met,	

Replicate Samples

Criteria: A replicate sample should be extracted and analyzed with each batch of samples. The RPDs between results (i.e., between the recoveries for the labeled 2,3,7,8-substituted compounds and between the concentrations for the non-labeled 2,3,7,8-substituted compounds) should be ≤25%.

> **Note:** An MS/MSD is not required for this method since it is an isotope dilution analysis. A replicate sample or LCS/LCSD will suffice to demonstrate batch precision.

Evaluation	Action
If a replicate sample or LCSD were <u>not</u> analyzed for each matrix or for each data package,	qualify all detects of the same matrix as "J" and all non-detects of the same matrix as "UJ."
If the RPD between the sample (or LCS) and its replicate (or LCSD) for any compound falls outside the appropriate control window,	qualify all associated detects of the same matrix as "J" and all associated non-detects of the same matrix as "UJ."

ISs

Criteria: The laboratory must spike all samples with the sample fortification solution and all sample extracts with recovery standard solution. The %R of each compound must be within 40% to 135%.

Evaluation	Action
If the %R for any sample fortification solution compound is <40%,	qualify all detects for that sample fraction as "J+" and all non-detects for that sample fraction as "UJ" if the %R is ≥10% and as "R" if the %R is <10%.
If the %R for any sample fortification solution compound is >135%,	qualify all detects for that sample fraction as "J-" and all non-detects for that sample fraction as "UJ."

Gas Chromatography (GC) Column Performance

Criteria: The GC column performance solution is used for defining the homologous GC RT windows and to document the chromatographic resolution. Column performance must be checked at the beginning of each analytical analysis period and must meet method acceptance criteria (see Section 9.3.1 of the method) before sample analysis may begin.

IMPORTANT NOTICE:

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation	Action
If GC column performance is not checked at the required frequency or if method	qualify all associated detects as "J" and all associated non-detects as "UJ."
criteria is not met,	

Confirmation of 2,3,7,8-TCDF Detects

Criteria: The DB-5 GC column generally used for PCDD and PCDF analyses does not adequately separate 2,3,7,8-TCDF from its closest eluting isomer. If 2,3,7,8-TCDF is detected in a sample, the result must be confirmed on a second column capable of separating 2,3,7,8-TCDF from all other TCDF homologues (as proven by successful analysis of the GC column performance column mix with <25% valley between 2,3,7,8-TCDF and its closest eluting isomer).

Evaluation	Action
If 2,3,7,8-TCDF was detected in a sample and the result was not confirmed on a second column with successful analysis of the GC column performance mix,	qualify all associated detects as "NJ."

Method TO-15, VOCs in Ambient Air using GC/MS

Analysis shall be performed according to the requirements specified in EPA TO-15, "Determination of VOCs in Air Collected in Specially- Prepared Canisters and Analyzed by GC/MS." In general, validate these analyses according to Section 4.2.

Surrogates, an MS/MSD, and TICs are not required. Although they are not required by the method, surrogates are often added during the analysis and reported by the labs. The laboratory criteria will be used for validation and qualifications will be based on professional judgment.

Instrument Tuning for GC/MS

See Section 4.2.1 for tuning and performance criteria.

Initial Calibration

Criteria: Instrument calibration shall be performed using at least five standard concentration levels. In addition, a zero air certification for the sampling apparatus is to be provided.

Evaluation	Action
If an insufficient number of standards were used,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If a zero air certification is not provided,	document the occurrence in the data validation report

RRFs

Criteria: The %RSD for the RRFs must be ≤30% and the average RRF shall be ≥ the method-specified minimum RRF for each compound. Compounds without specified minimum RRFs will be >0.050.

Evaluation	Action
If the average RRF for any target compound is < the specified minimum RRF, or <0.050 if no minimum is specified,	qualify all associated detects as "J" and all associated non-detects as "UJ" if the average RRF is ≥0.010 and as "R" if the average RRF is <0.010.
If the %RSD for any target compound is	
>30% but ≤45%,	qualify all associated detects as "J" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>45% but ≤60%,	qualify all associated detects as "J" and all associated non-detects as "UJ."
>60%,	qualify all associated detects as "J" and all associated non-detects as "R."

Calibration Verification

Criteria: Prior to analysis of samples, a calibration verification standard must be analyzed immediately following an ICAL to ensure that the instrument continues to remain under control. A calibration verification standard must be analyzed:

- (1) daily and
- (2) contain all target compounds.

The laboratory is allowed to perform corrective action and reanalyze once after a failure. If more than two calibration verification standards were analyzed to obtain a passing calibration verification standard, then the calibration was not verified and the calibration verification frequency criteria was not met.

Evaluation	Action
If the calibration verification standard was not analyzed at the proper frequency, or if all target compounds were not present in any calibration verification standard,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If the required calibration verification standard was not analyzed,	qualify all associated detects as "J" and all associated non-detects as "R."

RRFs

Criteria: The %D between RRFs and the average RRFs obtained from the ICAL shall be calculated according to the formula in Section 6.3 and must be $\leq 30\%$.

Evaluation	Action
If the %D was reported with the wrong sign (e.g., + %D for a negative bias),	document the occurrence in the data validation report and assess any infractions using the correct sign.
If the %D between an ICAL RRF and continuing calibration RRF for any target compound is	
>30% and positive (high bias),	qualify all associated detects as "J+."
>30% but ≤45% and negative (low bias),	qualify all associated detects as "J-" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>45% but ≤60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
>60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "R."

IS Performance

Criteria: IS area counts must not vary by more than $\pm 40\%$ from the average of those obtained from the calibration standards.

The RT of the IS must not vary more than ± 0.33 minutes (20 sec.) from that of the associated CCV standard.

When qualification of sample results is warranted due to failure of an IS to meet RT or area count acceptance criteria, results of all target compounds associated with that IS are qualified.

Refer to Appendix D for IS/target compound correlation guidelines.

Evaluating previous CCV IS areas are not required for this method.

Evaluation	Action
If there are two analyses for a particular compound,	determine which analysis contains the best data to report using the considerations below, qualify all data from the rejected analysis as "R," and document the reason for rejecting the data from one analysis in the data validation report. Considerations should include: 1. magnitude of the RT shift; 2. holding times; 3. comparison of the values of the target compounds reported in each fraction.
If any IS area count is <40% of the average of that obtained from the calibration standards,	qualify all associated detects as "J+" and all associated non-detects as "UJ." Non-detects may be qualified as "R" based on professional judgment if the IS area counts are <20% of that of the average obtained from the calibration standards.
	Note: If extremely low area counts are reported, or if performance exhibits a major abrupt drop-off, then a severe loss of sensitivity is indicated.
If the IS area count is >140% of the average of that obtained from the calibration standards,	qualify all associated detects as "J-" and all associated non-detects as "UJ."

Data Validation Procedure for Chemical and Radiochemical Data Effective Date: 06/19/2017

Evaluation (concluded)	Action (concluded)
If the IS RT varies by more than ±0.33 minutes from that of the associated calibration verification standard,	qualify all associated detects as "N" or "R" and all associated non-detects as "R."

LCS/LCSD

See Section 4.2.9 for LCS/LCSD criteria.

Methods 1668A and 1668C, Chlorinated Biphenyl Congeners

Initial Calibration

Criteria: Isotope dilution shall be used for calibration of the toxics and beginning and ending level of chlorination (LOC) chlorinated biphenyls (CBs). A 5- or 6point calibration is prepared for each native congener. The RRF %RSD for any native toxics/LOC CBs must be <20%. If a linear curve is used for ICAL, the r^2 of the curve must be >0.990.

Evaluation	Action
If the %RSD for any target compound is	
>20% but ≤40%,	qualify all associated detects as "J" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>40% but ≤60%,	qualify all associated detects as "J" and all associated non-detects as "UJ."
>60%,	qualify all associated detects as "J" and all associated non-detects as "R."

Evaluation (concluded)	Action (concluded)
If the r ² for any target compound is	
<0.990 but ≥0.90,	qualify all associated detects as "J" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
<0.90 but ≥0.80,	qualify all associated detects as "J" and all associated non-detects as "UJ."
<0.80,	qualify all associated detects as "J" and all associated non-detects as "R."

Criteria: Calibration using ISs is used for determination of native CBs for which a labeled compound is not available. For these CBs, calibration is performed at a single point. Compounds should be quantitated using the appropriate reference IS listed in Table 2 of the method. Ion abundance ratios must meet the criteria in Table 8 of the method or must be within 15% of the theoretical ratio of the ion monitored.

Evaluation	Action
If the ion abundance criteria were not met for any calibration compound,	qualify all associated detects as "J" and all associated non-detects as "UJ."

Continuing Calibration

Criteria:

At the beginning of each 12-hour shift during which analyses are performed, calibration is verified for all native CBs and labeled compounds. The ion abundance ratios for all CBs must be within the limits in Table 8 and all compounds must meet the calibration verification recovery limits listed in Table 6 of the method.

RRTs of native CBs and labeled compounds in the calibration verification must be within $\pm 0.5\%$ of the mean RRT determined in the ICAL or most recent calibration verification standard. The diluted combined 209-congener solution must be analyzed as a final step in the calibration verification and must meet minimum analysis and resolution specifications of the method.

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation	Action
If the ion abundance ratio for any compound is outside of the method limits,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If the verification limits are not met for any calibration verification compound and	
the %R is above the verification limits,	qualify all associated detects as "J+."
the %R is below the verification limits,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the recovery is ≥10% and as "R" if the recovery is <10%
If the RRT of any compound is outside of the RRT window,	qualify all associated results as "R."

RT Calibration

Criteria: The absolute RT of CB 209 must be ≥55 minutes if the SPB-octyl column is used. If a GC column or column system alternate to the SPB-octyl column is used, the absolute RT of CB 209 must be ≥ the laboratory-established minimum RT for CB 209. If the laboratory has not established a minimum RT value for CB 209, the RT for CB 209 must be ≥55 minutes.

Evaluation	Action
If an SPB-octyl column was used, and the absolute RT of CB 209 is <55 minutes,	qualify all associated results as "R."
If a GC column or column system alternate to the SPB-octyl column was used and the absolute RT is < the laboratory-established minimum RT for CB 209, or <55 minutes if the laboratory has not established a minimum RT,	qualify all associated results as "R."

Ongoing Precision and Recovery (OPR)

Criteria: OPR must be established for every batch of samples extracted and analyzed and must meet the recovery and %RSD limits listed in Table 6 of the method. If the OPR criteria are not met and reanalysis was not performed, then the laboratory performance and method accuracy are in question.

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home page.

Issue No: 05

Evaluation	Action
If the frequency of the OPR did not meet the specified criteria,	note the deficiency in the data validation report.
If the OPR %R is > the upper acceptance limit,	qualify all associated detects as "J+."
If the OPR %R is < the lower acceptance limit,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the %R is ≥ 20% and as "R" if the %R is < 20%.
If %Rs for more than half of the compounds in the OPR analysis are below the acceptance range,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the failures are marginally low and as "R" if %Rs are significantly below acceptance limits.
	Note: If recoveries for more than half of the compounds in the OPR analysis are below the acceptance range, the laboratory has not shown that it can actually meet program required DLs.
If %Rs for more than half of the compounds in the OPR analysis are above the acceptance range,	qualify all associated detects as "J+."
If %Rs for more than half of the compounds in the OPR analysis are outside the acceptance range, both above and below,	qualify all associated detects as "J" and all associated non-detects as "UJ."

Sample Preparation

Criteria: CBs may be bound to suspended particles in aqueous samples; therefore, the preparation of aqueous samples is dependent upon the solids content of the sample. A direct extraction is used for aqueous samples containing <1% solids. For aqueous samples containing >1% solids, the sample is agitated, allowed to settle and the liquid is decanted and discarded prior to extraction of the solids. The particle size for all solid samples should be determined prior to preparation. Particle size must be 1 millimeter or less prior to sample preparation.

Evaluation	Action
If % solids and particle size were not determined prior to sample preparation or if the proper preparation method was not performed,	qualify all associated detects as "J" and all associated non-detects as "UJ" and notify the program manager.

Criteria: Extract cleanup shall be used as necessary to eliminate interferences. The

laboratory may employ GPC, acid, neutral, or base silica gel; florisil; carbopak/celite; or high-performance liquid chromatography (HPLC) cleanup methods or anthropogenic isolation column for lipids (tissue extracts only).

Evaluation	Action
If the documentation on the run log, spectra	qualify all associated detects as "J" and
data, and/or IS or labeled compound	all associated non-detects as "UJ" and
recoveries indicate interferences and	notify the program manager.
applicable cleanup was not performed,	

Sample Analysis

Criteria: For identification of any CB or labeled compound, the ion abundance ratios must be within the limits specified in Table 8 of the method or $\pm 15\%$ of the calibration verification standard. The RRT of each CB must be within $\pm 0.5\%$ of the mean RRT determined in the ICAL or ±0.5% of the RRT from the most recent calibration verification standard.

Evaluation	Action
If ion abundance ratio criteria are not met for any compound,	qualify all associated results as "R."
If the RRT of any CB is outside of the RRT window,	qualify all associated results as "R."

Mass Spectrometer Performance Criteria

Criteria: Performance criteria are established to ensure mass resolution, identification, and, to some degree, sensitivity. These criteria are not sample specific. Conformance is determined using standard materials. These criteria should

be met in all circumstances.

Evaluation	Action
If mass spectrometer performance was not	qualify all associated detects as "J" and
checked at the required frequency or if	all associated non-detects as "UJ" and
method criteria were not met,	notify the program manager.

Labeled Compounds

Criteria: To assess method performance on the sample matrix, the laboratory must spike all samples with the labeled toxics/LOC/window defining standard spiking solution and all sample extracts with the labeled cleanup standard spiking solution. The recovery of each labeled compound must be within the limits listed in Table 6 of the method.

Evaluation	Action
If the %R for any labeled toxics/LOC/window defining standard compound is below acceptance limits,	qualify all detects for that sample fraction as "J+" and all non-detects for that sample fraction as UJ" if the recovery is ≥10% and as "R if the recovery is <10%.
If the %R for any labeled toxics/LOC/window defining standard compound is above acceptance limits,	qualify all detects for that sample fraction as "J-" and all non-detects for that sample fraction as "UJ."
	NOTE: For labeled cleanup standards, sample results are qualified with no bias. Because these standards are not associated with any specific targets, all 1668 sample results are affected by any cleanup standard infraction.
If the %R for any labeled cleanup standard compound is below acceptance limits,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the recovery is ≥10% and as "R" if the recovery is <10%.
If the %R for any labeled cleanup standard compound are above acceptance limits,	qualify all associated detects as "J."

California Environmental Protection Agency Air Resources Board; Method 428, PCDD, PCDF, and Polychlorinated Biphenyl (PCB) Emissions from Stationary Sources

Initial Calibration

Criteria: A 5-point calibration is prepared for each compound (see Tables 3, 5, and 10 of

the method for standard concentrations). The RRF RSD for any compound

must be $\leq 15\%$.

Evaluation	Action
If the %RSD for any compound is >15% but ≤40%,	qualify all associated detects as "J" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>40% but ≤60%,	qualify all associated detects as "J" and all associated non-detects as "UJ."
>60%,	qualify all associated detects as "J" and all associated non-detects as "R."

Continuing Calibration

Criteria: At the beginning and end of each 12-hour shift during which analyses are performed, calibration is verified for all compounds. The measured RRFs must be $\leq 30\%$ of the mean values established during ICAL. The relative abundance must meet the requirements specified in Tables 7 and 13 of the method.

Evaluation	Action
If the mass ratio for any compound is outside of the method limits,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If the %D criteria were not met for any compound and	
the %D is positive,	qualify all associated detects for that compound as "J+."
the %D is negative,	qualify all associated detects as "J-" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."

IMPORTANT NOTICE:

GC Column Performance

Criteria: The GC column performance solution is used for defining the homologous GC RT windows, to document the chromatographic resolution, and to check relative ion abundance criteria. Column performance must be checked at the beginning and end of each 12-hour analysis period and must meet method acceptance criteria (see Sections 5.3.5 and 6.3.5 of the method). If the laboratory operates during consecutive 12-hour shifts, analysis of the performance check solution at the beginning of each 12-hour period and at the end of the final 12-hour period is sufficient.

Evaluation	Action
If GC column performance was not checked	qualify all associated detects as "J" and
at the required frequency or if method	all associated non-detects as "UJ."
criteria is not met,	

Sample Preparation

Criteria: Extract cleanup shall be performed to eliminate interferences. The laboratory shall first partition the sample extract and then follow with an appropriate cleanup procedure.

Evaluation	Action
If sample spectra and/or IS and/or	qualify all associated detects as "J" and
surrogate recoveries indicate interferences	all associated non-detects as "UJ" and
and documentation of extract cleanup was	notify the program manager.
not provided,	

Sample Analysis

Criteria: For identification of any compound, the mass ratios must be within ±15% of the mass ratios listed in Tables 7 and 13 of the method. The RRT of each compound must be within ±0.006 RRT units of the standard RRT.

Evaluation	Action
If mass ratio criteria are not met for any compound,	qualify all associated results as "R."
If the RRT of any compound is outside of the RRT window,	qualify all associated results as "R."

IMPORTANT NOTICE:

ISs

Criteria: To assess method performance on the sample matrix, the laboratory must spike all samples with known concentrations of stable isotopically labeled ISs prior to extraction.

> The laboratory must spike all samples with known concentrations of recovery ISs prior to injection. The %R of each IS must be within 40% to 120% of the known value and the absolute RTs must be within ± 10 seconds of those measured during the last previous continuing calibration check.

If IS %Rs are outside of the acceptable limits, the signal to noise ratio of the IS must be >10.

Evaluation	Action
If the %R for any IS compound is below acceptance limits,	qualify all associated detects for that sample fraction as "J+" and all associated non-detects for that sample fraction as "UJ" if the recovery is ≥10% and as "R" if the recovery is <10%.
If the %R for any IS compound is above acceptance limits,	qualify all associated detects for that sample fraction as "J-" and all associated non-detects for that sample fraction as "UJ."

Matrix Blank

Criteria: Portions of the sample matrix (resin and filter) shall be analyzed at a frequency of every extraction set of 20 or fewer samples. All samples must be associated with an uncontaminated matrix blank. An uncontaminated matrix blank is defined as not having any compounds detected at a concentration \geq the MDL. The sample results must not be corrected by subtracting blank values.

Matrix blanks should be evaluated in the same manner as an MB. Blank qualification guidelines are discussed in Section 4.2.4.

Blank Sampling Train

Criteria: There shall be a least one blank train submitted to the laboratory for each series of three or fewer test runs. For sources with air pollution control devices, there shall be at least one blank train assembled at the inlet, and one at the outlet of the air pollution control devices for each set of three or fewer runs at each location. All samples must be associated with an uncontaminated blank train. An uncontaminated blank train is defined as not having any compound detected at a concentration \geq the MDL. The sample results must not be corrected by subtracting blank values.

Blank sampling trains should be evaluated in the same manner as an MB. Blank qualification guidelines are discussed in Section 4.2.4.

LCS

Criteria: An LCS must be extracted and analyzed with every batch of 20 samples or less and it must contain at least one representative of each chlorinated class of compounds to be determined in the samples. Accuracy is considered acceptable if the %R is within 60% to 140%.

> Note: If the LCS criteria are not met and reanalysis was not performed, then the lab performance and method accuracy are in question.

Evaluation	Action
If the frequency of the LCS did not meet the specified criteria,	note the deficiency in the data validation report.
If there was not at least one compound associated with each chlorinated class of compounds,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If the %R for an LCS compound is >140%,	qualify all associated detects as "J+."
If the %R for an LCS compound is <60%,	qualify all associated detects as "J-", and all associated non-detects as "UJ" if the %R is ≥20% and as "R" if the %R is <20%.
If %Rs for more than half of the compounds in the LCS analysis are below the acceptance range,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the failures are marginally low and as "R" if %Rs are significantly below acceptance limits.
	Note: If %Rs for more than half of the compounds in the LCS analysis are below the acceptance range, the laboratory has not shown that it can actually meet program required DLs.
If %Rs for more than half of the compounds in the LCS analysis are above the acceptance range,	qualify all associated detects as "J+."

Procedure No: AOP 00-03 Issue No: 05 Effective Date: 06/19/2017

Evaluation (concluded)	Action (concluded)
If %Rs for more than half of the compounds in the LCS analysis are outside the acceptance range, both above and below.	qualify all associated detects as "J" and all associated non-detects as "UJ."

Mass Spectrometer Performance Criteria

Performance criteria are established to ensure mass resolution, identification and, to some degree, sensitivity. These criteria are not sample specific. Conformance is determined using standard materials. These criteria should be met in all circumstances.

Criteria: Mass spectrometer performance must be checked every 12 hours of analysis in accordance with the method criteria. All compounds in all ICAL and continuing calibration standards must be within the QC limits listed in Tables 7 and 13 of the method for their respective isotopic ratios.

Evaluation	Action
If mass spectrometer performance was not	qualify all associated detects as "R" and
checked at the required frequency or if method criteria are not met,	all associated non-detects as "UJ" and notify the program manager.

QC Check Sample

A QC check sample must be extracted and analyzed with every batch of 20 samples or less. Accuracy is considered acceptable if the %R is within 60% to 140% and precision is acceptable if the RPD is $\leq 30\%$.

Note: If the QC check sample criteria are not met and reanalysis was not performed, then the lab performance and method accuracy are in question.

Evaluation	Action
If the frequency of the QC check sample did not meet the specified criteria,	note the deficiency in the data validation report.
If any QC check sample RPD is > 30%,	qualify all associated detects of the same matrix as "J" and all associated non-detects as "UJ."
If the QC check sample %R is >140%,	qualify all associated detects as "J+."

Evaluation (concluded) Action (concluded) qualify all associated detects as "J-" and If the QC check sample %R is <60%, all associated non-detects as "UJ" if the %R is $\geq 20\%$ and as "R" if the %R is <20%. If %Rs for more than half of the qualify all associated detects as "J-" and all associated non-detects as "UJ" if the compounds in the QC check sample analysis are below the acceptance range, failures are marginally low and as "R" if %Rs are significantly below acceptance limits. Note: If %Rs for more than half of the compounds in the OC check sample analysis are below the acceptance range, the laboratory has not shown that it can actually meet program required DLs. qualify all associated detects as "J+." If %Rs for more than half of the compounds in the QC check sample analysis are above the acceptance range, If %Rs for more than half of the qualify all associated detects as "J" and all associated non-detects as "UJ." compounds in the QC check sample analysis are outside the acceptance range, both above and below,

Procedure No: AOP 00-03

Issue No: 05

Spiked Sampling Trains

Criteria: Surrogate standards must be spiked into each sampling train as a means of estimating the precision and accuracy of the sampling train for collecting and recovering PCDDs, PCDFs, and PCBs in the stack gas sample. Surrogate recovery is considered acceptable if the %R is within 60% to 140%.

Evaluation	Action
If the surrogate %R is >140%,	qualify all associated detects as "J+."
If the surrogate %R is <60%,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the %R is ≥10% and as "R" if the %R is <10%.

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version is located on the Sandia Restricted Network (SRN), 4100 Controlled Documents home page.

Effective Date: 06/19/2017

Procedure No: AOP 00-03 Issue No: 05

Method 1613B, Tetra- through Octa-Chlorinated Dioxins and Furans by HRGC/HRMS

Note: An MS/MSD analysis is not required for this method.

Initial Calibration

Criteria: A combined 5-point calibration is prepared for the 2,3,7,8-substituted PCDDs and PCDFs for which labeled compounds are added to the samples (isotope dilution) and for 1,2,3,7,8,9-HxCDD, OCDF, and any non-2,3,7,8substituted compounds (ISs). The RRF %RSD for the compounds calibrated using isotope dilution must be ≤20%. For the compounds calibrated using ISs, the %RSD must be ≤35%. Ion abundance ratios must meet the criteria listed in Table 9 of the method.

> The laboratory may use alternative ions for quantitation to eliminate interferences. In this case, the ion abundance ratios must meet the criteria set by the laboratory.

Evaluation	Action
If the %RSD is >20% for any compound calibrated by isotope dilution, or >35% for any compound calibrated by IS, but ≤40%,	qualify all associated detects as "J" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>40% but ≤60% for any compound,	qualify all associated detects as "J" and all associated non-detects as "UJ."
>60% for any compound,	qualify all associated detects as "J" and all associated non-detects as "R."
If the ion abundance criteria were not met for any compound,	qualify all associated results as "R."

Continuing Calibration

Criteria: At the beginning of each 12-hour period during which analysis is performed, calibration is verified for all compounds. The measured concentration of each compound must be within the limits set in Table 6 of the method. The ion abundance must be within the limits in Table 9 of the method.

The absolute RTs of the ¹³C-1,2,3,4-TCDD and ¹³C-1,2,3,7,8,9-HxCDD ISs must be within ±15 seconds of the RTs obtained during the ICAL. The RRTs of the PCDDs/PCDFs and labeled compounds must be within the limits given in Table 2 of the method.

The evaluation of RTs and subsequent qualification of sample data requires professional judgment. If RRT criteria have not been met but absolute RTs between the CCV and the ICAL and between the CCV and the sample meet criteria, qualification of data may not be necessary. If RRT criteria and absolute RT criteria are not met, this may be an indication of instrument instability warranting qualification of sample data.

Evaluation	Action
If the ion abundance ratio criteria were not met for any compound,	qualify all associated detects as "J."
If the measured concentration criteria were not met for any compound at the beginning of a 12-hour period and	
the measured concentration is > the upper acceptance limit,	qualify all associated detects as "J+."
the measured concentration is < the lower acceptance limit,	qualify all associated detects as "J-" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."

Sample Preparation

Criteria: The cleanup standard ³⁷Cl₄-2,3,7,8-TCDD shall be added to all extracts prior to cleanup to measure the efficiency of the cleanup process. The recovery of the cleanup standard shall be within the limits set in Table 7 of the method.

Evaluation	Action
If the cleanup standard was not added to a sample, MB, or QC sample extract,	qualify all associated detects as "J" and all associated non-detects as "UJ" and notify the program manager.
If the recovery of the cleanup standard is > the upper acceptance limit,	qualify all associated detects as "J."
If the recovery of the cleanup standard is < the lower acceptance limit,	qualify all associated detects as "J" and all associated non-detects as "UJ."

IMPORTANT NOTICE:

Sample Analysis

Criteria: For identification of any compound, the ion abundance ratios must be within the limits specified in Table 9 of the method.

The recoveries of the labeled compounds must be within the limits specified in Table 7 of the method.

The RRTs of the PCDDs/PCDFs and labeled compounds must be within the limits specified in Table 2 of the method.

Evaluation	Action
If the ion abundance ratio criteria were not met for any compound,	qualify all associated results as "R."
If the recovery of any labeled compound is > the upper acceptance limit,	qualify all detects for the corresponding unlabeled compound as "J."
If the recovery of any labeled compound is < the lower acceptance limit,	qualify all detects for the corresponding unlabeled compound as "J" and all non-detects for the corresponding unlabeled compound as "UJ".
If the RT of any compound is outside of the RT window,	qualify all associated results as "R."

Mass Spectrometer Performance Criteria

Criteria: Performance criteria are established to ensure mass resolution; identification; and, to some degree, sensitivity. These criteria are not sample specific. Conformance is determined using standard materials. These criteria should be met in all circumstances. System performance must be evaluated at the beginning of each 12-hour period in which analysis is performed.

Evaluation	Action
If mass spectrometer performance was not evaluated at the required frequency or if method	qualify all associated detects as "R" and all associated non-detects as "UJ" and notify the
criteria were not met,	program manager.

GC Column Performance Mix

Criteria: The GC column performance solution is used for defining the homologous GC RT windows and to document the chromatographic resolution. Column performance must be evaluated at the beginning of each 12-hour analytical period and must meet method acceptance criteria (see Section 15.4 of the method) before sample analysis may begin.

Evaluation	Action
If GC column performance was not evaluated at the required frequency or if method criteria were	qualify all associated detects as "J" and all associated non-detects as "UJ" and notify the
not met,	program manager.

Confirmation of 2,3,7,8-TCDF Detects

Criteria: The DB-5 GC column generally used for PCDD and PCDF analyses does not adequately separate 2,3,7,8-TCDF from its closest eluting isomer. If 2,3,7,8-TCDF is

> detected in a sample, the result must be confirmed on a second column capable of separating 2,3,7,8-TCDF from all other TCDF homologues (as proven by successful analysis of the GC column performance column mix with <25% valley between 2,3,7,8-TCDF and its closest eluting isomer).

Evaluation	Action
If 2,3,7,8-TCDF is detected in a sample and the	qualify all associated detects as "NJ" and notify
result is not confirmed on a second column with	the program manager.
successful analysis of the GC column	
performance mix,	

4.3 Procedure for GC and HPLC Validation

The requirements covered within this section are applicable to all GC and HPLC analytical techniques, including SW-846 Methods 8081A, 8082A, and 8330B.

4.3.1 Calibration

Initial Calibration

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing acceptable qualitative and quantitative data for compounds on the TAL. Initial calibration demonstrates that the instrument is capable of acceptable performance in the beginning of the analytical run and of producing an adequate curve.

When methods require confirmation of target analytes on a second, fully-calibrated column, the calibrations of both columns must be assessed.

The laboratory may establish a calibration curve using the linear regression (linear curve) approach, the average RRF or the calibration factor (CF) approach. If more than one approach is used to quantify and report target analytes within the same data package, calibration is to be assessed on an analyte-by-analyte basis.

Criteria: GC and HPLC instrument calibration shall be performed using a minimum of five calibration standards unless otherwise specified by the method. If calibration curves are used, five standards are required for a linear (first-order) calibration model, six standards are required for a quadratic (second-order) model, and seven standards are required for a third-order polynomial. Higherorder curves should not normally be used. If the laboratory uses a higher order equation to establish a calibration curve, it should be evaluated for appropriate application.

ISs shall not be used for quantitation.

Evaluation	Action
If an insufficient number of calibration	qualify all associated detects as "J" and
standards were used,	all associated non-detects as "UJ."

RRFs

Criteria: RRFs are a measure of the slope of the calibration relationship and assume that the curve passes through the origin. Under ideal conditions, the factors will not vary with the concentration of the standard that is injected into the instrument. In practice, some variation is to be expected.

> When the variation, measured as the percent relative standard deviation (%RSD), is \leq 20%, the use of the linear model is appropriate and the calibration curve can be assumed to be linear and to pass through the origin.

> As a general rule, the amount of internal standard should produce an instrument response (e.g., area counts) that is no more than 100X that produced by the lowest concentration of the least responsive target compound associated with the internal standard. This should result in a minimum RRF of not <0.010 for the least responsive target compound.

The %RSD for the RRFs obtained from the five initial calibration standards must be \leq 20%, and the average RRF shall be \geq the method-specified minimum RRF for each compound. Compounds without specified minimum RRFs shall be ≥0.050.

Evaluation	Action
If the average RRF for any target compound is < the specified minimum RRF, or <0.050 if no minimum is specified,	qualify all associated detects as "J" and all associated non-detects as "UJ" if the average RRF is ≥0.010 and as "R" if the average RRF is <0.010.

Evaluation (concluded)	Action (concluded)
If the %RSD for any target compound is	
>20% but ≤40%,	qualify all associated detects as "J" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."

CFs

Criteria: In the absence of or in addition to, method-specific calibration acceptance criteria, the following general calibration acceptance criteria should be applied.

The %RSD for the CFs obtained from the five ICAL standards must be $\leq 20\%$.

Evaluation	Action
If any target compound has a %RSD:	
>20% but ≤40%,	qualify all associated detects as "J" and, all associated non-detects as "UJ" if any other calibration criteria have been exceeded for that compound.
>40% but ≤60%,	qualify all associated detects as "J" and all associated non-detects as "UJ."
> 60%,	qualify all associated detects as "J" and all associated non-detects as "R."

Linear Curves

Criteria: The r^2 of the ICAL curve shall be ≥ 0.990 . The absolute value of the intercept shall be $\leq 3X$ the MDL.

Evaluation	Action
If any target compound has a r ² :	
<0.990 but ≥0.90,	qualify all associated detects as "J" and, all associated non-detects as "UJ" if any other calibration criteria have been exceeded for that compound.
<0.90 but ≥0.80,	qualify all associated detects as "J" and all associated non-detects as "UJ."
<0.80,	qualify all associated detects as "J" and all associated non-detects as "R."
When results are reported at the MDL:	
If the intercept for any target compound is negative with an absolute value	
> the MDL but ≤3X the MDL,	qualify all associated detects <3X the absolute value of the intercept as "J-" and all associated non-detects as "UJ."
>3X the MDL,	qualify all associated detects <3X the absolute value of the intercept as "J-" and all associated non-detects as "R."
If the intercept for any compound is positive and > the MDL,	qualify all associated detects <3X the intercept as "J+."

4.3.2 Calibration Verification

Compliance requirements for satisfactory initial and continuing instrument calibration are established to ensure that the instrument is capable of producing acceptable qualitative and quantitative data for compounds on the TAL. The ICV independently verifies the calibration, and the CCV establishes the relative CFs on which the quantitations are based and checks satisfactory performance of the instrument on a day-to-day basis.

Criteria: An ICV must be run immediately following an ICAL. The ICV standard analysis results are not required to be reported in the data package unless the samples in the SDG were analyzed after the ICAL standards but before a CCV

standard analysis was performed. In this case, the ICV %D is assessed according to the calibration verification criteria described below for the associated samples. If a CCV is analyzed prior to samples and ICV data are also reported in the package, both the ICV %D and the appropriate CCV %D are to be assessed as described below. If both ICV %D and CCV %D infractions occur, the worst infraction should be evaluated for result qualification.

A CCV must be run:

- (I) at the beginning of each analytical run,
- (2) at least once every 20 samples (preferably every 10), and
- (3) at the end of each analytical run.

The laboratory is allowed to perform corrective action and reanalyze the CCV once after a failure. If multiple CCVs were analyzed (more than two) to obtain a passing CCV, then the calibration was not verified and the calibration verification frequency was not met. This is applicable to both CFs and linear curves. The evaluation of CCV data applies to all CCVs that bracket samples of interest.

A closing CCV is not required for toxaphene or chlordane if these compounds are non-detect in all samples.

Evaluation	Action
If the ICV/CCV standards were not analyzed at the proper frequency, or if either a required ICV or CCV was not analyzed, or if not all target compounds were present in any ICV or CCV standard,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If not all of the required ICVs and CCVs were analyzed,	qualify all associated detects as "J" and all associated non-detects as "R."

RRFs

Criteria: The %D between the ICV and/or CCV RRFs and the average RRFs obtained from the initial calibration shall be calculated according to the formula in Section 6.3 and must

be $\le 15\%$.

Evaluation	Action
If the %D was reported with the wrong sign (e.g., +%D for a negative bias),	document the occurrence in the data validation report and assess any infractions using the correct sign.

Evaluation (concluded)	Action (concluded)
If the %D between an initial calibration RRF and an ICV or CCV RRF for any target compound is	
>15% and positive (high bias),	qualify all associated detects as "J+."
>15% but ≤40% and negative (low bias),	qualify all associated detects as "J-" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>40% but ≤60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
>60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "R."

CFs

Criteria: The %D (see Section 6.3) between the ICV and/or the daily or continuing calibration standard CFs and the average CFs obtained from the ICAL must be ≤15%.

Evaluation	Action
If the %D was reported with the wrong sign (e.g., +%D for negative bias),	document the occurrence in the data validation report and assess any infractions using the correct sign.
If the %D between ICV and/or CCV CF and the average CF obtained from the ICAL is	
>15% and positive (high bias),	qualify all associated detects as "J+."
>15% but <40% and negative (low bias),	qualify all associated detects as "J" May qualify all associated non-detects as "UJ" if any other calibration criteria have been exceeded for that compound.
>40% but ≤60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
>60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "R."

IMPORTANT NOTICE:

Linear Curves

Criteria: The %D (see Section 6.3) between the daily or continuing calibration standard concentrations and their true values must be $\leq 15\%$.

The %D shall be calculated according to the formula in Section 6.3.

Evaluation	Action
If the %D was reported with the wrong sign (e.g., +%D for negative bias),	document the occurrence in the data validation report and assess any infractions using the correct sign.
If the %D between a measured ICV and/or CCV concentration and its true value is	
>15% and positive (high bias),	qualify all associated detects as "J+."
>15% but <40% and negative (low bias),	qualify all associated detects as "J" May qualify all associated non-detects for that compound as "UJ" if any other calibration criteria have been exceeded for that compound.
>40% but ≤60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
>60% and negative,	qualify non-detects for that compound as "R."

4.3.3 Blanks

The purpose of laboratory (or field) blank analysis is to determine the nature and magnitude of contamination resulting from laboratory (or field) activities.

The criteria for evaluation of blanks apply to any blank associated with the samples and include MBs, and, if submitted, EBs, and FBs. Action in the case of unsuitable blank results depends on the circumstances and origin of the blank. In instances where more than one blank is associated with a given sample, qualification should be based upon a comparison with the associated blank having the highest concentration of a contaminant. For purposes of evaluating multiple blanks, each preparation batch may be considered an independent event in evaluating MBs, and each sampling event may be considered an independent event for evaluating FBs and EBs.

Criteria: The concentration of each target analyte found in the blank must be < the associated MDL. The sample results must not be corrected by subtracting any blank value. If QC problems exist with any blank, all data associated with the case must be carefully evaluated to determine whether there is an inherent bias in the data, or if the problem is an isolated occurrence not affecting other data.

Evaluation	Action
If a compound found in a blank is also found a sample,	qualify the sample result for that compound in accordance with the scenarios given below.
If gross contamination exists,	qualify results for all compounds affected as "R" due to interference.
If inordinate numbers of other target compounds are found at low levels in the blank(s),	discuss the presence of these compounds in the data validation report as it may be indicative of a problem at the laboratory.

Blank Type	Blank Result	Sample Result	Action
Method, Storage,	Detect	Non-detect	No qualification
Field, Trip, Equipment,	≤PQL	≤PQL	Qualify as non-detect
Instrument			"U" at PQL.
		>PQL but ≤5X the	Qualify "J+."
		blank value	
	>PQL	≤PQL	Qualify as non-detect
			"U" at PQL.
		>PQL but ≤2X blank	Qualify as non-detect
		value	"U" at sample result and
			request corrective action
			from laboratory.
		>PQL and >2X blank	Qualify "J+" and
		value but ≤5X blank	request corrective action
		value	from laboratory.
	Gross Contamination	Detect	Report at sample result
			and qualify "J+" or "R",
			based on professional
			judgment and request
			corrective action from
			laboratory.

Gross contamination is not specifically defined but will be evaluated using professional judgment on a case by case basis. The infraction may be a high concentration of a single analyte or low-level contamination involving several analytes in the blank.

IMPORTANT NOTICE:

4.3.4 Surrogate Recovery

Laboratory performance for individual samples is established by means of surrogate spikes. All samples are spiked with surrogate compounds prior to sample preparation. The evaluation of the results of these surrogate spikes is not necessarily straightforward. The sample itself may produce matrix effects due to such factors as interference and high concentrations of analytes. Because the effects of the sample matrix are frequently outside the control of the laboratory and may present relatively unique problems, the review and validation of data based on specific surrogate results is frequently subjective and demands analytical experience and professional judgment.

Criteria: Sample and blank surrogate recoveries must be within limits specified by the laboratory. Surrogate compound recoveries shall be calculated using the procedure described in SW-846 Method 8000C. Reported recoveries shall be accompanied by the applicable acceptance limits.

Note: Results from spiked or replicate QC samples that have surrogate %Rs <10% cannot be used to qualify sample results. Samples should be qualified for lack of accuracy and/or precision data, as applicable, if specified by the program.

Evaluation	Action
If surrogate recovery acceptance criteria are not reported in the packages,	request amended data from the laboratory.
If, based on professional judgment, the laboratory's internal acceptance criteria are excessively wide or biased,	notify the program manager.
If an initial dilution was performed on any sample and at least one surrogate has %R < the lower acceptance limit but ≥10%, or all surrogates have <10 %R and the results for one or more compounds were ≥ the PQL,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
If an initial dilution was performed on any sample, all surrogate %Rs are <10%, and all results are < the PQL,	qualify all associated sample results as "R."
If there are two or more analyses for a particular fraction at the same dilution,	determine which analysis contains the best data to report using the considerations below, qualify all data from the rejected analysis as "R," and document the reason for rejecting data from one analysis in the data validation report.

Evaluation (concluded)	Action (concluded)
	Considerations should include: 1. surrogate recovery (marginal vs. gross deviation); 2. holding times; and 3. comparison of the values of the TALs reported in each fraction.
For surrogate recoveries out of specification, the following approaches are suggested based on a review of all data from the case, especially considering the apparent complexity of the sample matrix.	
If any surrogate %R is out of specification low,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the recovery is ≥10% and as "R" if the recovery is <10%.
If a surrogate %R is out of specification high,	qualify all associated detects as "J+."

Criteria: In the case of a blank analysis with surrogates out of specification, special consideration must be given to the validity of associated sample data. The basic concern is whether the blank problems represent an isolated problem with the blank alone or whether there is a fundamental problem with the analytical process.

> If one or more samples in the batch show acceptable surrogate recoveries, the blank problem may be considered an isolated occurrence. However, even if this judgment allows some use of the affected data, analytical problems remain that must be corrected by the laboratory.

Evaluation	Action
If surrogate recovery in the blank does not meet acceptance criteria,	all detects < the PQL in all samples associated with the blank may be qualified as "J" and all non-detects in all samples associated with the blank may be qualified as "UJ."

Internal Standard Performance

Internal standard criteria ensure that GC or HPLC sensitivity and response are stable and acceptable during each analysis.

Criteria: Sample and blank internal standard results must be within limits given in the specific SW-846 method.

> Internal standard area counts must not vary by more than a factor of two (50% to 200%) from the average of those obtained from the calibration standards.

> The RT of the internal standard must not vary by more than ±30 seconds from that of the associated CCV standard.

> When qualification of sample results is warranted due to failure of an internal standard to meet RT or area count acceptance criteria, results of all target compounds associated with that internal standard are qualified.

Evaluation	Action
If there are two or more analyses for a particular compound,	determine which analysis contains the best data to report using the considerations below, qualify all data from the rejected analysis as "R," and document the reason for rejecting the data from one analysis in the data validation report. Considerations should include: 1. magnitude of the RT shift; 2. holding times; 3. comparison of the values of the target compounds reported in each fraction; and 4. surrogate recovery.
If any internal standard area count is <50% of the average of that obtained from the calibration standards,	qualify all associated detects as "J+" and all associated non-detects as "UJ." Non-detects may be qualified as "R" based on professional judgment if the internal standard area counts are <20% of that of the average obtained from the calibration standards. NOTE: If extremely low area counts are reported or if performance exhibits a major abrupt drop-off, then a severe loss of sensitivity is indicated.

IMPORTANT NOTICE:

Evaluation (concluded)	Action (concluded)
If the internal standard area counts are >200% of the average of that obtained from the calibration standards,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
If the internal standard RT varies by more than ±30 seconds from that of the associated CCV standard,	qualify all associated detects as "N" or "R" and all associated non-detects as "R."

4.3.6 MS/MSD

Data for MS/MSD are generated to determine long-term precision and accuracy of the analytical method on various matrices and to demonstrate acceptable compound recovery by the laboratory at the time of sample analysis.

Criteria: The MS/MSD data shall not be used to evaluate field sample results unless the MS/MSD sample was from the same client and of similar matrix.

> An MS and MSD sample shall be analyzed at a frequency of once per data package, once per 20 samples of similar matrix, or once per sample matrix, whichever is more frequent. The MS must have recoveries calculated for all single-component target compounds. The presence of multi-component target compounds in the spiking solution is recommended but not required.

The laboratory shall not use FBs or EBs to satisfy this requirement if the laboratory can identify these blanks.

Unless otherwise stated in the specific method, the MS and MSD accuracy and precision acceptance criteria shall be those calculated by the laboratory using the procedure given in SW-846 Method 8000C. If the acceptance criteria are not given, recovery limits 70% to 130% and ±30% RPD should be used as the criteria. For solid and waste samples, it may be appropriate to accept up to a 40% RPD, based on the professional judgment.

The MS %Rs must be within the limits, unless the sample concentration is >4X the spike concentration (see Section 4.1.20).

The MS and MSD analyses must meet all sample analysis acceptance criteria. The MS and MSD results may be used in conjunction with other OC results to determine the need for qualification of the data. An effort to determine to what extent the results of the MS/MSD affect the associated data should first be made. This determination should be made considering the MS/MSD sample matrix, the surrogate recoveries, and the LCS results.

Professional judgment should be used to determine whether MS/MSD failure warrants qualification of only the results for the failed compounds, or if results for all the compounds associated with the failed MS compound are affected. Generally, unless evidence exists to warrant qualification of other compounds, only the compounds in the MS spiking mixture shall be qualified.

For programs that require application of one final qualifier to sample results, if a recovery (accuracy) infraction is identified in one or both of the MS samples along with an RPD (precision) infraction between the MS and MSD, the sample is qualified for the accuracy infraction. For example, if a compound has a low MS recovery and the RPD is not within criteria, the data are qualified as "J-."

Evaluation	Action
If the program requires MS/MSD analysis for all matrices and all target compounds	
and the MS/MSD sample was from another client or of a dissimilar matrix; the frequency of the MS/MSD did not meet specified criteria; no MS/MSD was analyzed or an FB- or EB was used for MS/MSD analysis,	qualify all detects as "J" and all non- detects as "UJ."
If no other measure of precision (i.e., LCSD or replicate) is available,	qualify all detects as "J" and all non- detects as "UJ."
If results are reported for single- component target compounds that are not in the MS,	all associated detects may be qualified as "J" and all associated non-detects may be qualified as "UJ" based on professional judgment
If any multi-component target compound is missing from the MS,	note the discrepancy in the data validation report.
If the surrogate and LCS recoveries are within the required acceptance criteria and either MS or MSD %R for any target compound is > the upper acceptance limit,	qualify all associated detects as "J+."
either MS or MSD %R for any target compound is < the lower acceptance limit,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the recovery is ≥20% and as "R" if the recovery is <20%.
If the RPD for any target compound does not meet the acceptance criteria or %Rs fail both high and low,	qualify all associated detects f as "J" and all associated non-detects as "UJ."

4.3.7 Replicate

Replicate analyses are indicators of laboratory precision based on each sample matrix. If a replicate was performed instead of an MSD, the following criteria are applied. If insufficient sample was submitted to analyze an MS/MSD or replicate, the laboratory may run an LCS/LCSD to measure precision. LCSD precision will be assessed as described in Section 4.3.5.

Criteria: Replicate samples shall be analyzed at a frequency of once per data package, once per 20 samples of similar matrix, or once per sample matrix, whichever is more frequent. All sample acceptance criteria must be met in the replicate analysis.

Samples identified as FBs or EBs shall not be used for replicate sample analysis.

Unless otherwise stated in the specific method, the replicate precision acceptance criteria shall be those calculated by the laboratory using the procedure given in SW-846 Method 8000C. When no laboratory-derived control limits are reported, a control limit of 30% for the RPD shall be used for sample values >5X the PQL. For solid and waste samples, it may be appropriate to accept up to a 40% RPD, based on the professional judgment.

A control limit of \pm PQL shall be used for sample values <5X the PQL, including the case when only one of the replicate sample values is <5X the PQL.

No precision criteria apply when both replicate sample values are < the PQL.

Evaluation	Action
If no replicate sample, no MSD, and no LCS/LCSD was analyzed for each matrix or for each data package,	qualify all associated detects of the same matrix as "J" and all associated non-detects as "UJ."
If an FB or EB was used for the replicate analysis and no MSD or LCSD was run,	qualify all associated detects of the same matrix as "J" and all associated non- detects as "UJ."
If the original result and replicate result are both >5X the PQL, and the RPD falls outside of appropriate control limits,	qualify all associated detects of the same matrix as "J" and all associated non- detects of the same matrix as "UJ."
If the original and/or replicate result is <5X the PQL (including non-detects) and the difference between the original result and replicate result is > the PQL,	qualify all associated detects of the same matrix as "J" and all associated non- detects of the same matrix as "UJ."

4.3.8 LCS

Data for LCSs are generated to provide information on the accuracy of the analytical method and on laboratory performance, including sample preparation.

Criteria: An LCS should be analyzed for all methods at a frequency of once per data package, once per matrix, or once per 20 analytical samples, whichever is most frequent.

> The LCS must have recovery calculated for all single-component compounds or at least one multi-component compound, if applicable. For very large analyte lists or for known poor performers, the laboratory may have received an exemption for one or more analytes. Analytes with exemptions will be identified in the case narrative.

The LCS must meet all sample acceptance criteria. If the MS/MSD and the samples meet all QC acceptance criteria, but the surrogate and/or internal standard recovery acceptance criteria are not met in the LCS analysis, the LCS must be reanalyzed.

The LCS should meet all method-specific LCS requirements and acceptance criteria. If the recovery acceptance criteria are not reported, the criteria in Appendix E or 70% to 130% should be used for evaluation.

If the laboratory analyzed an LCS/LCSD as a measure of precision, both the LCS and LCSD must meet the acceptance criteria.

General laboratory precision and accuracy can be evaluated using the LCS acceptance criteria and the interlaboratory comparison data given in Appendix E. Individual LCS recoveries may be evaluated against the criteria in Appendix E if the laboratory's criteria are significantly different from those in the tables.

Evaluation	Action
If, based on professional judgment, the laboratory's internal acceptance criteria are excessively wide or significantly biased,	notify the program manager.
If the frequency of the LCS did not meet the specified criteria,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If results are reported for target compounds that are not in the LCS,	may qualify detects for these compounds as "J" and non-detects as "UJ" based on professional judgment. Compounds missing under an exemption may be qualified based on professional judgment.

If the LCS criteria are not met and reanalysis was not performed, then the laboratory performance and method accuracy are in question. Professional judgment should be used to determine whether data should be qualified for all target compounds or just those compounds associated with the failed LCS compound. The following may be used as guidance in qualifying data.

IMPORTANT NOTICE:

Evaluation	Action
If the LCS %R is > the upper acceptance limit,	qualify all associated detects as "J+."
If the LCS %R is < the lower acceptance limit,	qualify all associated detects for that compound as "J-" and all associated non-detects as "UJ" if the %R is ≥20% and as "R" if the %R is <20%.
If %Rs for more than half of the compounds in the LCS analysis are below the acceptance range,	qualify all associated detects as "J-", and all associated non-detects as "UJ" if the failures are marginally low and as "R" if %Rs are significantly below acceptance limits.
	Note: If %Rs for more than half of the compounds in the LCS analysis are below the acceptance range, the laboratory has not shown that it can actually meet program required DLs.
If %Rs for more than half of the compounds in the LCS analysis are above the acceptance range,	qualify all associated detects as "J+."
If %Rs for more than half of the compounds in the LCS analysis are outside the acceptance range, both above and below,	qualify all associated detects as "J" and all associated non-detects as "UJ."

4.3.9 **TAL Compound Identification**

These criteria are established to ensure that adequate chromatographic resolution and instrument sensitivity is achieved by the chromatographic system.

Criteria: The laboratory must report RT window data for each GC column used to analyze samples. The RT of the ICV (or the first CCV of the day) should fall within the RT window established by the ICAL. RTs of subsequent CCVs should fall within the RT window established by the ICV or the first CCV of the day.

If RT windows were not reported,

If RT windows are not available, or if an RT for a standard exceeds the associated windows,

If RT windows are not available, or if an all associated detects as "NJ" and all associated non-detects as "R."

Emphasize the possibility of either false negatives or false positives, as appropriate, in the data validation report.

Procedure No: AOP 00-03

Issue No: 05

4.3.10 Sample Carry-over

Sample carry-over may occur when a high-concentration sample is analyzed immediately prior to another field sample. Steps must be taken to avoid introduction of false positive results in the second sample analysis due to instrument contamination.

Criteria: The absence of sample carry-over must be determined and verified. If examination of the run logs indicates that any samples in the analytical run of interest required dilution, and there is no documentation of a rinse or blank analysis immediately following the original undiluted analysis then sample carry-over may be suspected in the subsequent sample.

Evaluation	Action
If any target compound found in the sample requiring dilution exceeded the high calibration standard and was also found in the following sample at a concentration <5X the PQL,	qualify the results for that compound in the second sample as "R" or "NJ", based on professional judgment.
If no data are available for the sample that required dilution and the laboratory has not documented that carry-over was evaluated, and the compound(s) was (were) also found in the following sample at concentrations <5X the PQL,	qualify the results for that compound in the second sample as "N."

Data Validation Procedure for Chemical and Radiochemical Data Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

4.3.11 Dilutions

Criteria: The PQLs must be adjusted to reflect all sample dilutions, concentrations, splits,

clean-up activities, and dry weight factors that are not accounted for by the

method.

Samples must be diluted and reanalyzed when any analytes exceed the calibration range. Data from original sample runs should be included when any sample requires dilution due to one or more compounds exceeding the calibration range.

The original undiluted results document the actual MDLs for non-detects.

Evaluation	Action
If the PQLs have not been properly adjusted,	request an amended report from the laboratory.
If an initial dilution was required because of expected high concentrations of nontarget analytes or because one or more target analyte were expected to greatly exceed the instrument working range and the laboratory was not able to analyze the undiluted sample,	note the dilution and elevated MDLs in the data validation report.
If any target compound exceeded the calibration range and	
the original undiluted sample result was reported,	qualify all detects which exceeded the calibration range as "J."
the original undiluted sample run was not provided,	request this information from the laboratory.
the sample was diluted and reanalyzed and the diluted sample data were reported,	qualify all non-detects from the diluted analysis as "UJ."
If data from the original sample run are unavailable,	refer to Section 4.3.4 for assessment of initially diluted samples with low surrogate recovery.

Criteria: The laboratory shall strive to make dilutions in such a way that the final concentration is measured in the mid-range of the calibration curve, and that results are not reported from measurements below the lowest concentration standard.

Evaluation	Action
If the concentration (reported result / dilution factor) of any detect from diluted samples is < that of the lowest	qualify all associated detects from the diluted analysis as "J."
concentration standard,	

Criteria: The extraction efficiency for extremely high concentrations of analytes has generally not been determined for most methods. If the analysis requires an extraction and dilutions of > 100,000:1 the efficiency of the extraction may be suspect.

Evaluation	Action
If dilutions of > 100,000:1 was required,	qualify all associated detects as "J"

4.3.12 Quantification and Confirmation

Criteria: Detected compound results must be confirmed using a second GC/HPLC column. For 8330A/B HPLC HE analysis, the primary (C-18) and secondary (CN) columns recommended in the method should be used for analysis. However, if a different confirmation column is used, it must produce a separation that is substantially different from that obtained on the primary column. This is demonstrated by providing a separation in which the confirmation-column retention order is different from that of the primary column for all or most the compounds.

> The laboratory shall report RPDs between the results obtained from the two GC/HPLC columns. RPDs are not evaluated if the analyte is not detected on the primary column. (see note below)

Evaluation	Action
If the results from the second column confirmation are not reported,	qualify all detects as "NJ."
If the method-specified columns were not used for HPLC HE analysis, and the elution on the secondary column was not reversed relative to the primary column,	notify the program manager.
If the RPD between detects for a particular analyte from two analytical columns is >40% and ≤ 75%	
for PCB, pesticide, and herbicide analyses,	qualify the reported result as "J."
for high explosive (HE) analysis,	report the result from the primary column and qualify it as "J."

An RPD between results for a particular analyte from two analytical columns that is >75% may indicate that there is a significant coelution or interference problem. As applied here, a coelution is two target analytes, or one target and one non-target analyte, that have peaks at the same RT, and an interference is a non-target analyte with a peak at a target analyte RT. That is, a coelution is a quantity that cannot be verified, and an interference is a result that is a false positive.

A general review of the actual chromatograms may be required to determine the best qualification. If the chromatogram includes a significant number of extraneous peaks outside of the target analyte RT windows, interferences are likely on one or both of the columns. Asymmetrical peak shape is indicative of coelution, and shifts in RTs may indicate either coelution or interference. A review of the beginning and ending CCV RTs will give the reviewer an indication of instrument stability during the analysis.

If one of the results is < the PQL and the other is much > the PQL, suspect interference or a false positive. Values around the PQL should be evaluated using both RPD and absolute differences. For example, results of $1\mu g/L$ and $5\mu g/L$ have an RPD of 133% but would not be significantly different from each other for analyses with a PQL of $5\mu g/L$. An attempt should be made to determine whether the peak is primarily due to interference or if the peak has a significant contribution from the target analyte.

Note: It is not uncommon to find MDLs for GC/HPLC methods as determined using 40 CFR 136 to be artificially low, which may result in false positives due to random instrument noise for concentrations below the PQL.

In general, rejection of data with results much > the PQL will require additional supporting analytical information such as GC/MS or diode array spectral matching (see Appendix F).

IMPORTANT NOTICE:

Evaluation	Action
If the RPD is >75% and	
one result is <5X the PQL and the other result is > the PQL and >10X the first result,	qualify the reported result as "R."
both results are <5X the PQL	
for PCB, pesticide, and herbicide analyses,	qualify the reported result as "NJ."
for HE analysis,	report the result from the primary column and qualify it as "NJ."
both results are much > the PQL, one or both peaks may have contribution due to coelution, and	and quarry it as 143.
it is apparent that the peak is primarily due to the target analyte,	qualify the reported result as "J+."
it is not apparent that the peak is primarily due to the target analyte,	qualify the reported result as "NJ+."
If rejecting data where both results are much > the PQL,	include a complete description of the justification and supporting data used in the data validation report.

Procedure No: AOP 00-03

Issue No: 05

In the analysis of waste samples, the separation techniques may not completely isolate the analytes of concern from other compounds and the chromatograms may contain multiple extraneous peaks. The more peaks there are in the chromatograms, the more likely it is that false positives will be reported.

Evaluation	Action
If a large number of unidentified peaks are seen in the chromatograms or if several	results may be qualified as "N" using professional judgment.
additional peaks are located near a reported analyte RT in both spectra,	

Criteria: Although confirmation is not required for non-detects it is a common laboratory practice to use a dual column system and perform the confirmation analysis on all samples. Occasionally, there may be QC failures that occur on one of the columns that are acceptable on the other column. Laboratories may choose to report the analytes with acceptable results from one column and the remaining analytes from the other column. The following guidelines should be used when this occurs.

This practice may only be used for reporting non-detects from both columns.

All OC elements must be reported for both columns.

This can only be used when no primary column is specified, such as in SW-846 Method 8082A. When a primary column is specified, such as in SW-846 Method 8330B, all OC for the primary column must be acceptable.

The QC must be completely acceptable for each analyte on one or the other column. That is, the laboratory cannot use an acceptable LCS for an analyte on one column and an acceptable CCV for that analyte on the other column to justify acceptable performance.

Evaluation	Action
If both results are reported and qualification is required,	use the results from the column with the best performance with the caveat that results shall be reported from only the primary column for HPLC HE analysis.

4.3.13 Method-specific Analytical Requirements-Organic GC and HPLC

The additional analytical requirements given below are organized by SW-846 method. These requirements should be checked if the level of deliverable (level III or level IV) allows.

4.3.13.1 Method 8081A, Organochlorine Pesticide by GC

If discussion of water sample clean-up procedures was not included in the data package, it can be assumed that clean-up was not necessary and no discussion is required in the data validation report. For soil analysis, Florisil clean-up is required for all sample extracts.

Criteria: The laboratory must include a discussion of any clean-up procedures performed on the samples.

> An instrument blank consisting of clean solvent containing only the surrogate compounds shall be analyzed at the beginning and end of each analytical run, and once every 20 analytical samples.

Evaluation	Action
If discussion of sample clean-up procedures is missing or incorrect,	notify the laboratory and note the discrepancy in the data validation report.
If clean-up procedures were documented in the data package,	discuss the clean-up procedures used in the data validation report.
If no instrument blank was run, or if frequency criteria were not met,	may qualify detects <5X the MDL as "J" based on professional judgment.

Criteria: The total % breakdown for both dichlorodiphenyltrichloroethane (DDT) and endrin must each be $\leq 15\%$.

Evaluation	Action
If DDT breakdown is > 15%,	beginning with the samples following the last <i>in-control</i> standard, qualify all detects for DDT as "J" and all detects for dichlorodiphenyldichloroethane (DDD) and dichlorodiphenyldichloroethylene (DDE) as "NJ."
If DDT breakdown is >15% and DDT was not detected in any sample analyzed after the last in-control standard but DDD and DDE were detected in any of those samples,	qualify the result for DDT in the sample with DDD and DDE detects as "R."
If endrin breakdown is >15%,	beginning with the samples following the last <i>in-control</i> standard, qualify all detects for endrin as "J" and detects for endrin ketone and endrin aldehyde as "NJ."
If endrin breakdown is >15% and endrin was not detected in any sample analyzed after the last in-control standard, but endrin aldehyde and endrin ketone were detected in any of those samples,	qualify the result for endrin in the sample with the endrin aldehyde and endrin ketone detects as "R."

Note: A closing CCV is not required for toxaphene or chlordane if these compounds are non-detect in all samples.

4.3.13.2 Method 8082A, PCB Aroclors by GC

Criteria: PCB analysis shall be performed according to the requirements listed in SW-846 Method 8082A.

The laboratory must include a discussion of any clean-up procedures performed on the samples. If discussion of sample clean-up procedures was not included in the data package, it can be assumed that clean-up was not necessary and no discussion is required in the data validation report. The laboratory case narrative shall include a thorough discussion of any problems encountered regarding target compound recognition and/or quantitation and especially addressing suspected environmental degradation of compounds. Reported results shall be justified with such discussion and supporting documentation.

PCBs reported as total PCBs or as individual congeners are qualified in accordance with the Section 4.2.14.6.

Evaluation	Action
If clean-up procedures were documented in the data package,	discuss the clean-up procedures used in the data validation report.
If the discussion does not appear to justify the results reported by the laboratory,	notify the laboratory; more supporting documentation may be required from the laboratory.
If the laboratory identifies any aroclors as degraded,	qualify all associated detects as "J."

4.3.13.3 Method 8151A, Chlorinated Herbicides by GC

Criteria: Chlorinated herbicide analysis shall be performed according to the requirements listed in the SW-846 Method 8151A.

The LCS shall contain each of the specified target chlorinated herbicides at concentrations near the midpoint of the calibration range.

Evaluation	Action
If results are reported for target compounds that are not in the LCS,	may qualify all associated detects as "J" based on professional judgment.
If LCS analytes are not at concentrations near the midpoint of the calibration range,	note the finding in the data validation report and notify the laboratory.

4.3.13.4 Confirmation of Polyaromatic Hydrocarbon (PAH) by Method 8310

The primary analysis should be done by HPLC on a C18 column using a diode array detector. Confirmation is done qualitatively using spectral matching and/or quantitatively using a fluorescence detector. This method presupposes a high expectation of detecting the compounds. When used as a screening tool both confirmation methods should be employed. If a co-eluting compound is present that is detected by both the diode array detector and the fluorescence detector, the primary method of determining if interference is present is the spectral match from the diode array detector.

An effort to determine whether the peak is primarily from the target compound or due to interference should be made. This is determined by comparison of the sample diode array spectra to the reference spectra in accordance with Appendix F.

Evaluation	Action
If the diode array spectra were used for confirmation and no diode array spectra were included in the data package,	qualify the result as "NJ."
If the sample absorption spectra does not match the standard absorption spectra or the percent difference spectra does not exhibit a relatively straight line,	qualify the result as "R."
If the analyte was misidentified by the laboratory,	request an amended report from the laboratory.
If identification of the analyte was hampered by interferences such that it is not certain that a positive identification could be made or that the quantification may be biased high,	qualify all associated results as "N" or "NJ" based on professional judgment or request additional data from the laboratory.

The second evaluation compares the calculated values from the two detectors when a two-detector system is used. When one of the results is < the PQL and the other is much > the PQL (i.e., near or above the mid-point in the calibration curve), suspect interference or a false positive. Values around the PQL should be evaluated using both the RPDs and absolute differences. For example, results of $1\mu g/L$ and $5\mu g/L$ have an RPD of 133% but would not be significantly different from each other for analyses with a PQL of $5\mu g/L$.

Evaluation	Action
If results from the second detector confirmation were not reported,	qualify all detects as "NJ."
If the RPD between detects for a particular analyte from two analytical detectors is >40% and ≤75%,	report the result from the diode array detector and qualify it as "J."

Evaluation (concluded)	Action (concluded)
If the RPD is > 75% and	Note : If the RPD is >75%, one or both peaks may be due to coelution.
one result is <5X the PQL and the other result is > the first result,	qualify the result as "R."
both results are <5X the PQL,	qualify the result as "NJ."
both results are >5X the PQL and it appears that the peak is primarily due to the target analyte (spectral match),	qualify the result as "J+."
both results are >5X the PQL and it is not apparent that the peak is primarily due to the target analyte,	qualify the result as "NJ+."
If rejecting data where both results are much > the PQL,	include a complete description of the justification for the rejection and supporting data used in the data validation report.
	Note: In general, rejection of data with results >5X the PQL will require additional supporting analytical information such as GC/MS spectral matching.

4.3.13.5 Method 8015C or 8015D, Non-halogenated Organics Using GC/Flame Ionization Detector (Gasoline Range Organics/Diesel Range Organics)

Confirmation on a second column is not typically required for gasoline range organics (GRO) and diesel range organics (DRO) reported by this method.

GRO and DRO results represent all peaks detected over a designated RT range on the chromatogram. The RT assessment is performed as described in Section 4.3.8 for all reported RT markers.

Evaluation	Action
If RT windows are exceeded,	qualify all associated detects as "J" and all associated non-detects as "UJ."

4.4 Procedure for Liquid Chromatography/Mass Spectrometry/Mass Spectrometry (LC/MS/MS) Validation

The requirements addressed within this section are applicable to all LC/MS/MS analytical techniques. LC/MS/MS is a highly selective analysis that utilizes four means of compound discrimination: chromatographic separation, negative ion generations (where applicable), mass selection, and daughter fragmentation. It is theoretically possible that two different compounds could have the same RT and generate the same ion, but it is highly unlikely that these two compounds would fragment to the same daughter ion.

4.4.1 Instrument Calibration for LC/MS/MS

Initial Calibration

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing acceptable qualitative and quantitative data for compounds on the TAL. Initial calibration demonstrates that the instrument is capable of acceptable performance at the beginning of the analytical run and of producing a linear curve. In the absence of or in addition to, method-specific calibration acceptance criteria, the following general calibration acceptance criteria should be applied.

If an IS is used to calculate analytical results, the slope or RRF values are evaluated as directed below. If the analysis does not use an IS to quantitate analytical results, the value of the slope or CF is not evaluated.

The laboratory may establish a calibration curve using either the linear regression (linear curve) approach or the RRF approach. If both approaches are used to quantify and report target analytes within the same data package, calibration is to be assessed on an analyte-by-analyte basis.

Criteria: LC/MS/MS instrument calibration shall be performed using a minimum of five calibration standards. The lowest point of the curve must be at or below the PQL.

> If calibration curves are used, five standards are required for a linear (first-order) calibration model, six standards are required for quadratic (second-order) model, and seven standards are required for third-order polynomial. Higher order curves should not normally be used. If the laboratory uses a higher-order equation to establish a calibration curve, it should be evaluated for appropriate application.

Daily instrument calibration is required for perchlorate analysis.

Evaluation	Action
If an insufficient number of calibration standards were used, the PQLs were incorrect or all points were not analyzed within a 24-hour period:	qualify all associated detects as "J" and all associated non-detects as "UJ."

Evaluation (concluded)	Action (concluded)
If the instrument for perchlorate analysis was not calibrated daily,	qualify all associated detects as "J" and all associated non-detects as "UJ."

RRFs

Criteria: The %RSD for the RRFs obtained from the five ICAL standards must be $\leq 20\%$.

Evaluation	Action
For analyses using an IS for analyte quantitation, if the average RRF for any target analyte is < the specified minimum RRF, or <0.050 if no minimum is specified,	qualify all associated detects as "J" and all associated non-detects as "UJ" if the RRF is ≥0.010 and as "R" if the RRF is <0.010.
If any target compound has a %RSD	
>20% but ≤40%,	qualify all associated detects as "J" and, may qualify all associated non-detects as "UJ" if any other calibration criteria have been exceeded for that compound.
>40% but ≤60%,	qualify all associated detects as "J" and all associated non-detects as "UJ."
> 60%,	qualify all associated detects as "J" and all associated non-detects as "R."

Linear Curves

Criteria: The r^2 of the ICAL curve shall be ≥ 0.990 and have a slope ≥ 0.050 for each compound. The absolute value of the intercept shall be $\leq 3X$ the MDL.

> For perchlorate, forcing the calibration curve through a zero intercept is an acceptable practice and usually results in more accurate quantitation for low level results.

Evaluation Action qualify all associated detects as "J" all For analyses using an IS for analyte quantitation, if the slope for any target associated non-detects as "UJ" if the analyte is < the specified minimum RRF, or slope is ≥ 0.010 and as "R" if the slope is < 0.050 if no minimum RRF is specified, < 0.010. If any target compound has a r^2 : qualify all associated detects as "J" and <0.990 but ≥ 0.90 , may qualify all associated non-detects as "UJ" if any other calibration criteria have been exceeded for that compound. qualify all associated detects as "J" and <0.90 but ≥ 0.80 . all associated non-detects as "UJ." qualify all associated detects as "J" and < 0.80,all associated non-detects as "R." If the intercept for any target analyte is qualify all associated detects <3X the positive and > the MDL, intercept as "J+." When results are reported at the MDL: If the intercept for any target compound is negative with an absolute value... qualify all associated detects <3X the > the MDL but $\leq 3X$ the MDL, absolute value of the intercept as "J-" and all associated non-detects as "UJ." qualify all associated detects <3X the >3X the MDL, absolute value of the intercept as "J-" and all associated non-detects as "R."

Procedure No: AOP 00-03

Issue No: 05

4.4.2 Calibration Verification

Compliance requirements for satisfactory initial and continuing instrument calibration verification are established to ensure that the instrument is capable of producing acceptable qualitative and quantitative data for target compounds. The ICV independently verifies the calibration, and CCV establishes the 12-hour relative RRFs on which the quantitations are based and checks satisfactory performance of the instrument on a day-to-day basis.

The evaluation of CCV data applies to all CCVs that bracket samples of interest.

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Criteria: An ICV standard is analyzed immediately following an ICAL. For perchlorate analysis, the ICV is always evaluated for %D criteria. For HE analysis, the ICV standard analysis results are not required to be reported in the data package unless the samples in the SDG were analyzed after the ICAL but before a CCV standard analysis was performed. In this case, the ICV %D is assessed according to the calibration verification criteria described below for the associated samples. If a CCV is analyzed prior to samples and ICV data are also reported in the package, both the ICV %D and the appropriate CCV %D are to be assessed as described below. If both ICV %D and CCV %D infractions occur, the worst infraction should be evaluated for result qualification.

A CCV standard must be analyzed:

- 1) at the beginning of each analytical run;
- 2) at least once every 10 samples; and
- 3) and at the end of each analytical run.

If multiple CCVs were analyzed to obtain a passing CCV, the calibration is not verified and the calibration frequency is not met.

Evaluation	Action
If the ICV/CCV standards were not analyzed at the proper frequency, or if either a required ICV or CCV was not analyzed, or if all target compounds were not present in any ICV or CCV standard,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If all required ICVs and CCVs were not analyzed,	qualify all associated detects as "J" and all associated non-detects as "R."

RRFs

Criteria: The %D between the ICV and/or CCV RRFs and the average RRF obtained from the ICAL shall be calculated according to the formula in Section 6.3 and must be $\leq 20\%$ for HE and $\leq 15\%$ for perchlorate. The evaluation of CCV data applies to all CCVs that bracket samples of interest.

Evaluation	Action
If the %D was reported with the wrong sign (e.g., +%D for a negative bias),	document the occurrence in the data validation report and assess any infractions using the correct sign.

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Evaluation (concluded)	Action (concluded)
If the %D between an ICAL RRF or CF and an ICV or CCV RRF or CF for any target analyte is	
>20% for HE or >15% for perchlorate and positive (high bias),	qualify all associated detects as "J+."
>20% for HE or >15% for perchlorate but ≤40% and negative (low bias),	qualify all associated detects as "J-" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>40% but ≤60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
>60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "R."

Linear Curves

Criteria: The %D between the ICV or CCV standard concentrations and their true values shall be calculated according to the formula in Section 6.3 and must be ≤20%

shall be calculated according to the formula in Section 6.3 and must be $\leq 20\%$ for HE and $\leq 15\%$ for perchlorate. The evaluation of CCV data applies to all

CCVs that bracket samples of interest.

Evaluation	Action
If the %D was reported with the wrong sign (e.g., +%D for negative bias),	document the occurrence in the data validation report and assess any infractions using the correct sign.

Evaluation (concluded)	Action (concluded)
If the %D between a measured ICV and/or CCV concentration and its true value for any analyte is	
>20% for HE or >15% for perchlorate and positive (high bias),	qualify all associated detects as "J+."
>20% for HE or >15% for perchlorate but ≤40% and negative (low bias),	qualify all associated detects as "J-" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>40% but ≤60% and negative,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
>60% and is negative,	qualify all associated detects as "J-" and all associated non-detects as "R."

4.4.3 RLV

A RLV standard (i.e., CRI), of the same origin as the calibration standard must be analyzed at the beginning and end of each perchlorate analysis run and at the beginning only if each HE analytical run as a measure of accuracy near the PQL. This analysis may be referred to as a PS. Analysis of a CRI is required for both HE and perchlorate methods. CRI standard concentrations are at 2X the MDLs for perchlorate analysis, and at no more than 2X the PQL for HE analysis.

The laboratory may run more than the required CRIs in a batch. In this case, the bracketing CRIs for perchlorate and the CRI immediately preceding the samples for HE will be used for the CRI evaluation.

With the adoption of Mehod 8330B for HE analysis, the laboratory has agreed to perform a reanalysis of the low-level calibration standard called the IRA. Acceptance criteria for the IRA are 80 to 120%.

Criteria: The advisory recovery acceptance criteria for perchlorate analysis are 70% to 130%. For HE analysis, recoveries must be within limits specified by the laboratory. If recovery acceptance criteria are not reported, the recovery acceptance range shall be 70% to 130%.

Evaluation Action qualify all detects <5X the PQL as "J" If frequency criteria are not met, and all non-detects as "UJ." qualify all associated detects <5X the PQL as "J+." If the R is >130% for the CRI or >120% for the IRA. If the %R is <70% but >30% for the CRI qualify all associated detects <5X the or <80% but >30% for the IRA, PQL as "J-" and all associated nondetects as "UJ." qualify all associated detects <5X the If the %R is <30%, PQL as "J-" and all associated nondetects as "R."

Procedure No: AOP 00-03

Issue No: 05

4.4.4 Blanks

For perchlorate analysis, refer to Section 4.5.3 for assessment of blanks.

The following applies for HE analysis.

The preparation batch consists of a group of no more than 20 samples of the same matrix processed on the same day. All samples in a batch must be initiated on the same day. Each batch must contain a MB.

An initial calibration blank (ICB) must be analyzed to verify the baseline immediately following calibration and prior to analytical sample analysis. A continuing calibration blank (CCB) must be analyzed after each CCV and at the end of every analytical sequence in order to bracket all sample analyses. All CCBs that bracket samples of interest shall be reported and assessed. If a bracket has an ICB and no CCB, then the ICB should be treated as a CCB for validation purposes.

The purpose of laboratory (or field) blank analysis is to determine the essence and magnitude of contamination resulting from laboratory (or field) activities.

The criteria for evaluation of blanks apply to any blank associated with the samples and include MBs and, if submitted, EBs and FBs. Action in the case of unsuitable blank results depends on the circumstances and origin of the blank. In instances where more than one blank is associated with a given sample, qualification should be based upon a comparison with the associated blank having the highest concentration of a contaminant. For purposes of evaluating multiple blanks, each preparation batch may be considered an independent event in evaluating preparation blanks, and each 12-hour run sequence may be considered an independent event for evaluating FBs and EBs.

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Criteria: The concentration of each target analyte found in the blank must be < the associated MDL. The sample results must not be corrected by subtracting any blank value. If QC problems exist with any blank, all data associated with the. case must be carefully evaluated to determine whether there is an inherent bias in the data, or if the problem is an isolated occurrence not affecting other data

Evaluation	Action
If a compound found in a blank is also found in the field sample,	qualify the sample result for that compound in accordance with the scenarios given below.
If gross contamination exists,	qualify results for all compounds affected as "R" due to interference.
If inordinate numbers of other target compounds are found at low levels in the blank(s)	discuss the presence of those compounds in the data validation report as it may be indicative of a problem at the laboratory.

Blank Type	Blank Result	Sample Result	Action
Method, Storage,	Detect	Non-detect	No qualification.
Field, Equipment, Instrument	≤PQL	≤PQL	Qualify as non-detect "U" at PQL.
1100		>PQL but ≤5X the blank value	Qualify "J+."
	>PQL	≤PQL	Qualify as non-detect "U" at PQL.
		>PQL but ≤2X blank value	Qualify as non-detect "U" at sample result and request corrective action from laboratory.
		>PQL and >2X blank value but ≤5X blank value	Qualify "J+" and request corrective action from laboratory.
	Gross Contamination	Detect	Report at sample result and qualify "J+" or "R," based on professional judgment and request corrective action from laboratory.

Gross contamination is not specifically defined but will be evaluated using professional judgment on a case by case basis. The infraction may be a high concentration of a single analyte or lowlevel contamination involving several analytes in the blank.

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Procedure No: AOP 00-03 Issue No: 05 Effective Date: 06/19/2017

4.4.5 Surrogate Recovery – HE analysis only

Laboratory performance on individual samples is established by means of surrogate spikes. All samples are spiked with surrogate compounds prior to sample preparation. The evaluation of the results of these surrogate spikes is not necessarily straightforward. The sample itself may produce matrix effects due to such factors as interference and high concentrations of analytes. Because the effects of the sample matrix are frequently outside the control of the laboratory and may present relatively unique problems, the review and validation of data based on specific sample results is frequently subjective and demands analytical experience and professional judgment. The evaluation of surrogate recoveries and ISs should be performed concurrently. Accordingly, this section consists primarily of guidelines, in some cases with several optional approaches suggested.

Criteria: Sample and blank surrogate recoveries must be within limits specified by the laboratory. Surrogate compound recoveries shall be calculated using the procedure described in SW-846 Method 8000C. Reported recoveries shall be accompanied by the applicable acceptance limits.

> Results from spiked or replicate QC samples that have surrogate %Rs < 10% cannot be used to evaluate associated sample results. Associated samples should be qualified for lack of accuracy and/or precision data as applicable.

Evaluation	Action
If the surrogate recovery acceptance criteria were not reported in the data package, If, based on professional judgment, the laboratory's internal acceptance criteria are excessively wide or biased,	request amended data from the laboratory. notify the program manager.
If an initial dilution was performed on any sample and at least one surrogate recovery is < the lower acceptance limit but ≥10%, or all surrogate recoveries are <10% and the results for one or more compounds are > the POL,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
If an initial dilution was performed on any sample, any surrogate %R is <10%, and all results are non-detect,	qualify all sample results as "R."

Evaluation (concluded)	Action (concluded)	
If there are two or more analyses for a particular fraction at the same dilution,	determine which contains the best data to report using the considerations below, qualify all data from the rejected analysis as "R" and document the reason for rejecting data from one analysis in the data validation report.	
	Considerations should include:	
For surrogate spike recoveries out of specification, the following approaches are suggested based on a review of all data	 surrogate recovery (marginal vs. gross deviation); holding times; comparison of the values of the target analytes reported in each fraction; and performance of ISs. 	
from the batch, especially considering the		
apparent complexity of the sample matrix.		
If the surrogate is out of specification low,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the recovery is ≥10% and as "R" if the recovery is <10%.	
If the surrogate is out of specification high,	qualify all detects as "J+."	

Criteria: In the case of a blank analysis with surrogate out of specification, special consideration must be given to the validity of associated sample data. The basic

> concern is whether the blank problems represent an isolated problem with the blank alone or whether there is a fundamental problem with the analytical process.

If one or more samples in the batch show acceptable surrogate recovery, r the blank problem may be considered an isolated occurrence. However, even if this judgment allows some use of the affected data, analytical problems remain that must be corrected by the laboratory.

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation	Action
If surrogate recovery in the blank does not meet acceptance criteria,	may qualify all detects < the PQL in all samples associated with the blank as "J" and all non-detects in all samples associated with the blank as "UJ."

4.4.6 IS Performance

IS criteria ensure that instrument sensitivity and response are stable and acceptable during each analysis.

Criteria: The laboratory may use an IS to calculate the result, or it may use the IS as a RT check only (perchlorate). If the IS is used for quantification, the IS area counts must not vary by more than 70% to 130% and the RT of the IS must not vary by more than ± 30 seconds from the average of those obtained from the calibration standards or from the mid-level calibration standard.

> If the IS is only used as an RT check, the RRT of the IS must fall within the acceptance range of 0.98 to 1.02, and the IS recovery should be evaluated using the surrogate criteria. If recovery acceptance limits are not reported in the data package, recovery should be evaluated based on reported MS acceptance limits.

Evaluation	Action
If there are two analyses for a particular sample,	determine which contains the best data to report based on the considerations below, qualify all data from the rejected analysis as "R" and document the reason for rejecting data from one analysis in the data validation report. Considerations should include: 1. magnitude of the RT shift; 2. holding times; 3. comparison of the values of the target compounds reported in each sample; and. 4. surrogate recovery.
· · · · · · · · · · · · · · · · · · ·	i. Sairogate recovery.

Evaluation (concluded)	Action (concluded)
If the IS was used for quantification and	
its area count is >130% of the average of that obtained from the calibration standards,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
its area count is <70% of the average of that obtained from the calibration standards,	qualify all associated detects as "J+" and all associated non-detects as "UJ" if the area count is ≥25% and as "R" if the area count is <25%.
	Note: If extremely low area counts are reported, or if performance exhibits a major abrupt drop-off, then a severe loss of sensitivity is indicated.
its RT varies by more than ±30 seconds,	qualify all associated detects as "N" or "R" and all associated non-detects as "R."
If the IS was used as an RT check and the RRT does not fall within the acceptance range,	qualify all associated detects as "N" or "R" and all associated non-detects as "R."
If the IS was used as an RT check,	evaluate the IS area counts according to Section 4.4.5.

4.4.7 MS/MSD

Data for MS/MSD pairs are generated to determine long-term precision and accuracy of the analytical method on samples various matrices and to demonstrate acceptable compound recovery by the laboratory at the time of sample analysis.

Criteria: The MS/MSD data shall not be used to evaluate associated field sample results unless the MS/MSD sample was from the same client and of similar matrix.

An MS and MSD sample shall be analyzed at a frequency of once per data package, once per 20 samples of similar matrix, or once per sample matrix, whichever is more frequent.

The laboratory shall not use FBs or EBs to satisfy these requirements if the laboratory can identify these blanks.

For HE, the MS and MSD accuracy and precision acceptance criteria shall be those calculated by the laboratory using the procedure given in SW-846 Method 8000C. If the recovery acceptance criteria are not given, recovery limits of 70% to 130% and 30% RPD should be used as the criteria. For solid and waste samples, it may be appropriate to accept up to a 40% RPD, based on the professional judgment of the reviewer.

For perchlorate, the MS/MSD recovery acceptance criteria are 75% to 125% and 20% RPD. For solid and waste samples, it may be appropriate to accept up to a 30% RPD, based on the professional judgment.

The MS and MSD %R must be within the limits, unless the sample concentration is >4X the spike concentration (see Section 4.1.20).

The MS and MSD analyses must meet all sample analysis acceptance criteria. The MS and MSD may be used results in conjunction with other QC results to determine the need for qualification of the data. An effort to determine to what extent the results of the MS/MSD affect the associated data should first be made. This determination should be made considering the MS/MSD sample matrix, the surrogate and IS %Rs, and the LCS results. Professional judgment should be used to determine whether MS/MSD failure warrants qualification of only the results for the failed compounds or if results for all the compounds associated with the failed MS compound and its associated IS are affected. Generally, unless evidence exists to warrant qualification of other compounds, only the compounds in the MS spiking mixture shall be qualified.

For programs that require application of one final qualifier to sample results, if a recovery (accuracy) infraction is identified in one or both of the MS samples along with an RPD (precision) infraction between the MS and MSD, the sample is qualified for the accuracy infraction. For example, if an analyte has low MS recovery and the RPD is not within criteria, the data are qualified as "J-."

Evaluation	Action
If the MS/MSD sample was from another client or of a dissimilar matrix; the frequency of the MS/MSD did not meet specified criteria; no MS/MSD was analyzed; an FB or EBs was used for MS/MSD purposes.	qualify all detects as "J" and all non- detects as "UJ."
If no other measure of precision (i.e., LCSD or replicate) is available,	qualify all detects as "J" and all non- detects as "UJ."
If the surrogate, IS and LCS %Rs are within the required acceptance criteria and	

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Evaluation (concluded)	Action (concluded)
either %R for any target compound is > the upper acceptance limit,	qualify all associated detects as "J+."
either %R for any target compound is < the lower acceptance limit, and $\geq 10\%$,	qualify all associated detects as "J-" and all associated non-detects as UJ" if the recovery is ≥20% and as "R" if the recovery is <20%.
If the RPD for any target compound does not meet the acceptance criteria or %Rs fail both high and low,	qualify all associated detects as "J" and all associated non-detects as "UJ."

4.4.8 Replicate

Replicate analyses are indicators of laboratory precision based on each sample matrix. If a replicate was performed instead of a MSD, the following criteria are applied. If insufficient sample was submitted to analyze a MS/MSD or replicate, the laboratory may run an LCSD to measure precision.

Criteria: A replicate sample shall be analyzed at a frequency of once per data package, once per 20 samples of similar matrix, or once per sample matrix, whichever is more frequent. All sample acceptance criteria must be met in the replicate analysis.

> Samples identified as FBs or EBs should not be used for replicate sample analysis.

For HE values ≥5X the PQL, the replicate precision acceptance criteria shall be that calculated by the laboratory using the procedure given in SW-846 Method

8000C. If the acceptance criteria are not given, an RPD of ≤30% should be used as the acceptance criteria. For solid and waste samples, it may be appropriate to accept up to a 40% RPD, based on the professional judgment.

For perchlorate, the replicate precision acceptance criterion is 20% RPD for sample values >5X the PQL. For solid and waste samples, it may be appropriate to accept up to a 30% RPD, based on the professional judgment.

For both HE and perchlorate analysis, a control limit of \pm the PQL shall be used for sample values > the PQL but <5X the PQL, including the case when only one of the replicate sample values is > the PQL but <5X the PQL.

No precision criteria apply when both replicate sample values are < PQL.

Evaluation	Action
If a replicate sample, MSD, and LCS/LCSD were not analyzed for each matrix or for each	qualify all detects of the same matrix as "J" and all non-detects of the same
data package,	matrix as "UJ."

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Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation (concluded)	Action (concluded)
If an FB or EB was used for the replicate analysis and no MSD or LCSD was run,	qualify all detects of the same matrix as "J" and all non-detects of the same matrix as "UJ."
If the original result and replicate result for any target compound are both ≥5X the PQL, and the RPD exceeds the appropriate control limit,	qualify all associated detects of the same matrix as "J" and all associated non- detects all associated as "UJ."
If the original and/or replicate result for any target compound is > the PQL but <5X the PQL (including non-detects) and the difference between the original result and replicate result is > the PQL,	qualify all associated detects of the same matrix as "J" and all associated non- detects of the same matrix as "UJ."

4.4.9 LCS

Data for LCSs are generated to provide information on the accuracy of the analytical method and on laboratory performance including sample preparation.

Criteria: An LCS should be analyzed for all methods at a frequency of once per data package, once per matrix, or once per 20 analytical samples, whichever is most frequent.

> The LCS must meet all sample acceptance criteria and all method-specific LCS requirements. The LCS for HE must meet laboratory-derived acceptance criteria. If the MS/MSD and the samples meet all QC acceptance criteria, but the surrogate and/or IS recovery acceptance criteria are not met for the LCS analysis, the LCS must be reanalyzed. If the recovery acceptance criteria are not reported, the criteria in Appendix E or 70% to 130% should be used for evaluation.

The recovery acceptance limits for perchlorate are 85% to 115%.

If the laboratory analyzed an LCS/LCSD as a measure of precision, both the LCS and LCSD must meet the acceptance criteria.

Evaluation	Action
If, based on professional judgment, the laboratory's internal acceptance criteria are excessively wide or significantly biased,	notify the program manager.
If the frequency of the LCS did not meet the specified criteria,	qualify all detects as "J" and all non- detects as "UJ."

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Evaluation (concluded)	Action (concluded)
If results are reported for target compounds that are not in the LCS,	may qualify detects for those compounds as "J" and non-detects for those compounds may be qualified as "UJ" based on professional judgment.

If the LCS criteria are not met and batch reanalysis was not performed, then the laboratory performance and method accuracy are in question. The following may be used as guidance in qualifying data.

Evaluation	Action
If the LCS %R is > the upper acceptance limit,	qualify all associated detects as "J+."
If the LCS %R is < the lower acceptance limit,	qualify all associated detects as "J-" and all associated non-detects as "UJ" if the R is $\geq 20\%$ and as "R" if R is $\leq 20\%$.
For HE analysis, if %Rs for more than half of the compounds in the LCS analysis are below the acceptance range,	qualify all detects as "J-" and all non- detects as "UJ" if the %Rs are marginally low and as "R" if %Rs are significantly below acceptance limits.
For HE analysis, if %Rs for more than half of the compounds in the LCS analysis are above the acceptance range,	qualify all detects as "J+."
For HE analysis, if %Rs for more than half of the compounds in the LCS analysis are outside the acceptance range, both above and below, or if an LCS/LCSD pair was analyzed and the recoveries of any target analyte are both above and below acceptance criteria,	qualify all detects as "J" and non-detects as "UJ."

4.4.10 Sample Carry-over

Sample carry-over may occur when a high-concentration sample is analyzed immediately prior to another field sample. Steps must be taken to avoid introduction of false positive results in the second sample analysis due to instrument contamination.

Criteria: The absence of sample carry-over must be determined and verified. If examination of the run logs indicates that any samples in the analytical run of interest required dilution and there is no documentation of a rinse or blank analysis immediately following the original undiluted analysis, then sample carry-over may be suspected.

Evaluation	Action
If any compound found in the sample requiring dilution exceeds the high calibration standard and was also found in the following sample at concentration <5X the PQL,	qualify the results for that compound in the second sample as "R."
If <u>no data</u> are available for the sample that required dilution and the laboratory has not documented that carry-over was evaluated, and any compound was also found in the following sample at concentration <5X the PQL,	qualify the results for that compound in the second sample as "N."

4.4.11 Dilutions

Criteria: The PQLs must be adjusted to reflect all sample dilutions, concentrations, splits, clean-up activities, and dry weight factors that are not accounted for by the method.

> Samples must be diluted and reanalyzed when any analyte exceeds the calibration range.

Original sample runs should be included when any sample requires dilution due to one or more compounds exceeding the calibration range.

The original undiluted results document the actual MDLs for non-detects.

Evaluation	Action
If the PQLs have not been properly adjusted,	request an amended report from the laboratory.
If an initial dilution was required because of expected high concentrations of non-target analytes or because one or more target analytes were expected to greatly exceed the instrument working range, and the laboratory was not able to analyze the undiluted sample.	note the dilution and evaluated MDLs in the data validation report.

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Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation (concluded)	Action (concluded)
If any target compound exceeds the calibration range and	
the original undiluted sample result was reported,	qualify all detects from the undiluted analysis that exceed the calibration range as "J."
the sample is diluted and reanalyzed, and the diluted sample data were reported,	qualify all non-detects from the diluted analysis as "UJ."
the original undiluted sample data was not provided,	request this information from the laboratory.
If data from the original sample run are unavailable,	refer to Section 4.2.5 for assessment of initially diluted samples with low surrogate recovery.

Criteria: The laboratory shall strive to make dilutions in such a way that the final concentration is measured in the mid-range of the calibration curve, and that results are not reported from measurements below the lowest concentration standard.

Evaluation	Action
If the concentration (reported result /	qualify all detects from the diluted
dilution factor) from diluted sample is <	analysis as "J."
that of the lowest concentration standard,	

4.4.12 Perchlorate Chlorine Ratios

Criteria: The natural isotopic abundances for the chlorine isotopes give a 35Cl/37Cl ratio of approximately 3.08. Laboratories must statistically derive isotope ratio acceptance criteria to be used as an additional confirmation of analyte identity.

> When the laboratory does not specify acceptance isotope ratio acceptance criteria for results between the MDL and the PQL, the individual sample isotope ratio acceptance limits shall be 3.08 ±20% (approximately 3 sigma of a typical isotope ratio population). For results above the PQL, the individual sample isotope ratio acceptance limits shall be $3.08 \pm 15\%$ (approximately 2 sigma of a typical population).

When isotope ratio acceptance criteria are not met, the laboratory must provide supporting data and explanatory case narrative comments in the data package.

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Evaluation	Action
If the isotope ratios were not reported,	calculate the ratio if the raw data were supplied or request an amended report from the laboratory if the raw data were not supplied.
If the isotope ratios are outside acceptance limits,	qualify detects as "NJ" or "R" based on professional judgment.
If supporting data and explanation were not provided,	request an amended report from the laboratory.

4.4.13 Perchlorate Interference Check Standard

Criteria: The laboratory shall analyze an interference check standard (ICS) from a matrix containing 500 ppm each of chloride, sulfate, carbonate, and bicarbonate in every batch. The concentration of this standard will be at the PQL. To demonstrate that perchlorate is adequately isolated and recovered under the specific conditions used, this standard should recover within $\pm 20\%$ of the known value.

Evaluation	Action
If frequency criteria were not met,	note the deficiency in the data validation report.
If the recovery is not within ±20% of the known value,	note the deficiency in the data validation report.

4.4.14 Method-specific Analytical Requirements – Organic LC/MS/MS

The additional analytical requirements addressed below are organized by method. These requirements should be checked if the level of deliverable allows.

Method 1694: Pharmaceuticals and Personal Care Products (PPCP) in Water, Soil, Sediment, and Biosolids by HPLC/MS/MS

Validation is to be performed as data deliverables allow.

Note: MS/MSD and replicate analyses are not required for Method 1694.

Initial Calibration

Criteria: If isotope dilution calibration is used, the %RSD must be <20%. If IS calibration is used, the %RSD must be <35%. Absolute RTs for last-eluting compounds in each of the four calibration groups must be \geq the reference RT listed in Tables 3, 5, 7, and 9 of Method 1694.

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Evaluation	Action
If the %RSD is:	
>20% for any compound calibrated using isotope dilution method or >35% for any compound calibrated using IS calibration, but ≤ 40%,	qualify all associated detects as "J" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."
>40% but <60% for either a labeled or unlabeled calibration standard,	qualify all associated detects as "J" and all associated non-detects as "UJ."
>60% for either a labeled or unlabeled calibration standard,	qualify all associated detects as "J" and all associated non-detects as "R."
If the RT of the last eluting compound in any of the four analysis groups is < the reference RT,	qualify all associated detects as "J" and all associated non-detects as "UJ."

Calibration Verification

Criteria: A QC Check Sample (from a second source vendor, similar to an ICV) shall be analyzed at least quarterly. The CCV %D must be $\pm 30\%$ of the ICAL response factor for all labeled and unlabeled compounds. The LC peaks for all native and labeled compounds must be present with a signal to noise ratio of at least 10. The retention times of the native and labeled compounds must be within 15 seconds (0.25 minutes) of the respective RTs in the most recent calibration verification standard.

Evaluation	Action
If there is no evidence in the data package of a QC Check sample (ICV) analysis,	note in the data validation report without qualification.
If the %D between an ICAL RRF and an ICV or CCV RRF for any labeled or unlabeled compound is	
>30% and positive (high bias),	qualify all associated detects as "J+."
>30% and negative (low bias),	qualify all associated detects as "J-" and, if any other calibration criteria have been exceeded for that compound, qualify all associated non-detects as "UJ."

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 Evaluation (concluded)
 Action (concluded)

 >40% but ≤60% and negative,
 qualify all associated detects as "J-" and all associated non-detects as "UJ."

 >60% and negative,
 qualify all associated detects as "J-" and

all associated non-detects as "R."

Procedure No: AOP 00-03

Issue No: 05

Evaluation	Action
If the LC peaks for all native and labeled compounds do not have a signal to noise ratio of \geq 10,	qualify detects as "J_" and non-detects as "UJ."
If the LC peaks for all native and labeled compounds do not have a signal to noise ratio of ≥ 3 ,	qualify detects as "J-" and non-detects as "R."

Evaluation	Action
If any native or labeled compound RT has shifted more than 15 seconds (0.25 minutes) from the RT of the most recent calibration verification standard,	qualify detects as "J_" and non-detects as "R".

Labeled Compound Recovery

Criteria: Labeled Compound recovery must meet criteria given in Table 12 of the method for LCS (OPR) samples, and field samples.

Evaluation	Action
If the labeled compound %R is < method acceptance criteria lower acceptance limit but >2.5%,	qualify all associated detects for that sample fraction as "J+" and all non-detects for that sample fraction as "UJ."
If the labeled compound %R for is <2.5%,	qualify all results for the associated result as "R."
If the %R for any sample fortification solution compound is > the upper limit of the method acceptance criteria	qualify all detects for that sample fraction as "J-" and all non-detects for that sample fraction as "UJ."

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Surrogate Recovery

Follow criteria listed in Section 4.4.5.

IS Performance

An IS performance criterion is not given in Method 1694.

Target Analyte Identification

Criteria: The signal to noise ratio for each native compound identified above the MDL in a

sample, MB, or LCS must be ≥ 2.5 , and ≥ 10 in associated calibration standards, except for the lowest ICAL standard, which must have a signal to noise ratio of 3. The RT of the peak for all identified compounds must be ± 15 seconds (0.25)

minutes) of the most recent CCV.

Evaluation	Action
If signal to noise ratio and/or RT criteria are not met for any reported compound,	notify the program manager and request an amended report.

4.5 Validation Guidelines for Confirmation by LC/MS/MS

These guidelines are for qualification of the original data based on the confirmation data obtained by LC/MS/MS and apply to HE by SW-846 Method 8330B and perchlorate by EPA Method 314. It should be noted that no confirmation LC/MS/MS results are qualified. If the original sample result is a detect, and the corresponding LC/MS/MS result is a detect, the original result is considered to be confirmed, although confirmation qualifiers may be applied. As with all validation guidelines, professional judgment is the final criteria.

4.5.1 Required LC/MS/MS Data

The laboratory is expected to include all calibration and QC data normally supplied in a level 4 data package; however, only the following information is necessary to evaluate the confirmation.

- 1) Form I Sample Results
- 2) CRI Summary
- 3) IRA Summary
- 4) CCV %D Summary
- 5) IS Recovery
- 6) MB results

4.5.2 LC/MS/MS QC are Acceptable

The laboratory is expected to meet all QC acceptance criteria when performing confirmation by LC/MS/MS. Note: Multiple QC failures should result in "NJ" or no qualification to original results.

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If all LC/MS/MS QC acceptance criteria are met:

Evaluation	Action
If the original result is ≥ the PQL and the LC/MS/MS result is non-detect,	qualify the original result as "R."
If the original result is < the PQL and the LC/MS/MS result is non-detect,	qualify the original result as "R."
If the original result and the LC/MS/MS result are > the PQL, and	
the %D is >40%,	qualify the original result as "J."
If the original result is \geq the PQL and the LC/MS/MS result is $<$ the PQL,	qualify the original result as "NJ+."
If the original result is $<$ the PQL and the LC/MS/MS result is \ge the PQL,	qualify the original result as "J"

4.5.3 Method Blank

The MB performed on LC/MS/MS should be < the MDL.

Evaluation	Action
If the LC/MS/MS MB concentration is ≥ the MDL and	
the original result is > the PQL, but <5X the MB concentration,	qualify the original result as "R."
the original result is < the PQL and <5X the MB concentration,	qualify the original result as "NJ+."

4.5.4 Continuing Calibration

If the original result is a detect and the LC/MS/MS result is a non-detect and the LC/MS/MS CCV is outside acceptance criteria:

Evaluation	Action
If the CCV %D is positive (high bias), >20% and	
the original result is \geq the PQL,	qualify the original result as "R."
the original result is < the PQL,	qualify the original result as "R."
If the CCV %D is negative (low bias), >20% and	
the original result is \geq the PQL,	qualify the original result as "R."
the original result is < the PQL,	qualify the original result as "NJ."

4.5.5 PS/CRI

If the original result is a detect, the LC/MS/MS result is a non-detect, and the LC/MS/MS PS/CRI are outside acceptance criteria:

Evaluation	Action
If the spike %R is > the upper limit and	
the original result is \geq PQL,	qualify the original result as "R."
the original result is < PQL,	qualify the original result as "R."
If the spike %R is < the lower limit and	
the original result is \geq the PQL,	qualify the original result as "R."
the original result is < the PQL,	qualify the original result as "NJ."

IS Performance 4.5.6

In the case of confirmation analysis, IS performance is assessed only as it reflects instrument sensitivity, not calculated bias; and only applies if the IS is used for quantification.

If the original result is a detect; the LC/MS/MS result is a non-detect; and the IS is outside acceptance criteria:

Evaluation	Action
If the IS %R is > the upper limit and	
the original result is ≥ the PQL,	qualify the original result as "R."
the original result is < the PQL,	qualify the original result as "R."
If the IS %R is < the lower limit and	
the original result is \geq the PQL,	qualify the original result as "R."
the original result is < the PQL,	qualify the original result as "NJ."

4.6 Procedure for Inorganic Data Validation

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing acceptable qualitative and quantitative data for target analytes. Initial calibration demonstrates that the instrument is capable of acceptable performance at the beginning of the analytical run, and CCV documents that the ICAL is still valid.

4.6.1 Initial Calibration

Criteria: Instruments used for all analyses other than ion chromatography (IC) must be calibrated daily and each time the instrument is set up as noted below.

> Inductively coupled plasma-atomic emission spectroscopy (ICP)-AES) radial-viewing analysis: A blank and at least one standard must be used in establishing the analytical curve. ICP-AES axial-viewing analysis: A blank and at least three standards must be used in establishing the analytical curve.

Inductively coupled plasma-mass spectrometry (ICP-MS) analysis: A blank and at least one standard must be used in establishing the analytical curve.

Mercury analysis by cold vapor atomic absorption: A blank and at least four standards must be used in establishing the analytical curve.

Cyanide analysis: A blank and at least three standards, one of which must be at the PQL, must be used in establishing the analytical curve.

IC analysis: A blank and at least three standards, one of which must be at the PQL, must be used in establishing the analytical curve. Daily calibration is not required if acceptable calibration verification is performed prior to the analytical run.

Flow Injection and Colorimetric analysis: A blank and at least three standards, one of which must be at the PQL, must be used in establishing the analytical curve.

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Evaluation	Action
If the minimum number of standards was not used for ICAL,	qualify all detects as "J" and all non- detects as "UJ."
If the instrument was not calibrated at the proper frequency,	qualify all sample results as "R."
If only one standard was used for trace axial view ICP-AES,	notify the laboratory and the program that the laboratory was not compliant with the <u>contract SOW</u> .

Criteria: The correlation coefficient (r) of the ICAL curve shall be ≥ 0.995 , and the absolute value of the intercept shall be $\leq 3X$ the MDL.

> Note: The sample results may be reported with non-detects at the MDL or at the PQL value. See below for appropriate evaluation.

The r assessment need only be performed on those curves established using at least three standards and a blank (four-point curve).

The intercept shall be assessed for all inorganic calibration curves, with the following exception: The laboratory may report both a low-level and a highlevel calibration curve for some inorganic analytes. In this case, the intercepts for the higher level curve should not be evaluated. Qualifiers should still be applied if the linearity criteria are not met for the calibration curve used to quantitate the samples.

Evaluation	Action
If any compound has a r:	
<0.995 but ≥0.90,	qualify all associated detects as "J" and all associated non-detects may be qualified as "UJ" if any other calibration criteria have been exceeded for that analyte.
$< 0.90 \text{ but } \ge 0.80,$	qualify all associated detects as "J" and all associated non-detects as "UJ."
<0.80,	qualify all associated detects as "J" and all associated non-detects as "R."

Evaluation (concluded)	Action (concluded)
When results are reported at the MDL:	
If the intercept for any target compound is negative with an absolute value	
> the MDL but ≤3X the MDL,	qualify all associated detects <3X the absolute value of the intercept as "J-" and all associated non-detects as "UJ."
>3X the MDL,	qualify all associated detects <3X the absolute value of the intercept as "J-" and all associated non-detects as "R."
If any compound has an intercept that is positive and > the MDL,	qualify all associated detects <3X the intercept as "J+."
When results are reported at the PQL:	
If the intercept for any target compound is negative with an absolute value	
> the MDL but ≤2X the PQL,	qualify all associated detects <3X the absolute value of the intercept as "J-" and all associated non-detects as "UJ."
>2X the PQL,	qualify all associated detects <3X the absolute value of the intercept as "J-" and all associated non-detects as "R."

4.6.2 CCV

Criteria: ICV and CCV: An ICV standard must be analyzed after instrument calibration and prior to sample analysis. A CCV standard must be analyzed once every 10 injections or every two hours, whichever is more frequent. The evaluation of CCV data applies to all CCVs that bracket samples of interest.

> ICV and CCV analysis results must be within the recovery acceptance criteria of 90% to 110% of the true value for all analytes.

Evaluation	Action
If the ICV and CCV standards were not analyzed at the proper frequency, or if either a required ICV or CCV was not analyzed, or if not all target compounds were present in any ICV or CCV standard,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If not all required ICVs and CCVs were analyzed,	qualify all associated detects as "J" and all associated non-detects as "R."
If the ICV or CCV %R is	
<90% but ≥75%,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
$>110\%$ but $\le 125\%$,	qualify all associated detects as "J+."
<75%,	qualify all associated detects "J-" and all associated non-detects as "R."
>125%,	qualify all associated detects as "R."

4.6.3 Blanks

Blank analysis results are assessed to determine the existence of contamination problems. The criteria for evaluation of blanks apply to any laboratory blank associated with the samples. See Section 4.1.2 for general chemistry QC exemptions.

Criteria: An ICB must be analyzed to verify the baseline immediately following calibration and prior to analytical sample analysis. A CCB must be analyzed after each CCV and at the end of every analytical sequence in order to bracket all sample runs. All CCBs that bracket samples of interest shall be reported and assessed.

A minimum of one MB (or preparation blank) should be analyzed for every 20 samples. The same reagents used for the sample digestion must be used to prepare the MB. In those cases for which reagents are automatically added to all samples by an autoanalyzer, the ICB is equivalent to a preparation blank. FBs and EBs are treated as preparation blanks.

If any QC problems exist with any blank, all data associated with the batch must be evaluated to determine whether there is an inherent bias in the data for the batch, or if the problem is an anomaly not affecting other data.

If the absolute value of the ICB or CCB result is > the PQL, the analysis should have been terminated and the problem corrected by the laboratory. If any analyte concentration in the blank is > the PQL, the lowest reported concentration in the associated samples must be $\geq 10X$ the concentration in the blank. Samples having analyte concentrations <10X that of the blank but > the PQL shall be re-digested and/or reanalyzed.

No contaminants \geq MDL should be present in the blanks.

When there is blank contamination and reanalysis is not possible, the data may need to be qualified. Use the blank (ICB, CCB, MB, FB, or EB) with the highest concentration associated with the samples of interest to qualify data. If a CCB is used to qualify data, it must bracket the sample of interest.

The effect of MB values versus ICB/CCB values on sample results is not straightforward and will vary depending on analytical method. Professional judgment is required to properly assess the effect of blank data on sample results. As a general guideline, in the case of conflicting positive and negative MB and ICB/CCB values, the MB values will take precedence over ICB/CCB values when applying qualifications to associated sample results.

Blank Type	Blank Result	Sample Result	Action
ICB/CCB, MB	If blank frequency		Note the deficiency in
	criteria were not met,		the data validation
			report and request a
	1		corrective action from
			the laboratory.
ICB/CCB, MB, EB, FB	If any associated blank	non-detect	No qualification
	value is positive and is	detect ≤PQL	Qualify "U" at PQL or
	\geq the MDL but \leq the		optionally "J+."
]	PQL,	detect > PQL but $\leq 5X$	Qualify "J+."
		blank value	
		detect >PQL and >5X	May be qualified "J+"
		blank value	based on professional
			judgment.
ICB/CCB, MB, EB, FB	If any associated blank	non-detect	No qualification.
į	value is positive and >	detect ≤ the PQL	Qualify "U" at PQL.
	the PQL,	detect > PQL but $\leq 5X$	Qualify "J+."
		blank value	
		detect >PQL and >5X	May be qualified "J+"
		blank value	based on professional
			judgment.

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Effective Date: 06/19/2017 Issue No: 05

Procedure No: AOP 00-03

Blank Type	Blank Result	Sample Result	Action
ICB/CCB, MB, EB, FB	If the absolute value of	non-detect	Qualify "UJ" at PQL.
	negative blank is >MDL but ≤PQL,	detect <5X the absolute value of the blank	Qualify "J-" or "J."
ICB/CCB, MB	If the absolute value of negative blank is >PQL and the analysis was not	non-detect	Qualify "R" and request a corrective action from the laboratory.
	terminated by the laboratory,	detect ≤10X the absolute value of the blank	Qualify "J-" and request a corrective action from the laboratory.
EB, FB	If the absolute value of negative blank is >PQL,	non-detect detect ≤10X the absolute value of the blank	Qualify "R." Qualify "J"

4.6.4 MS

The MS sample analysis is performed as a measure of the ability to recover analytes in a particular matrix.

Criteria: The MS data shall not be used to evaluate data unless the MS sample was from the same client and of similar matrix.

An MS sample shall be analyzed at a frequency of once per data package, once per 20 samples of similar matrix, or once per sample matrix, whichever is more frequent.

Samples identified as FBs and EBs cannot be used for MS analysis.

Spiking levels shall be approximately at the mid-point of the calibration range.

The MS recovery acceptance criteria are 75% to 125%, unless the sample concentration is >4X the spike concentration (see Section 4.1.20).

MS analysis shall be performed for all analytes other than sodium, potassium, magnesium, and calcium.

For methods, which require a digestion, PSs are occasionally performed. The recovery acceptance criteria on a PS are 75% to 125%. For methods which do not require digestion (e.g. IC, ion-specific electrode, and colorimetric techniques), MSs shall be analyzed. These spikes may be referred to as a post spikes or analytical spikes. These should be evaluated using the recovery acceptance criteria of a MS, 75% to 125%.

Evaluation	Action
If the MS sample was from another client or of a dissimilar matrix, the frequency of the MS did not meet specified criteria, an MS was not analyzed, or an FB or EB was used for MS analysis,	qualify all detects as "J" and all non- detects as "UJ."
If an MS %R is	
>125%,	qualify all associated detects as "J+."
<75% but ≥30%,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
<30%,	qualify all associated detects as "J-" and all associated non-detects as "R."
If an MS/MSD pair was analyzed and recoveries of any target analyte are both above and below acceptance criteria,	qualify all detects as "J" and all non- detects as "UJ."
If the MS %R is >125% and a PS was analyzed (if required when sample concentration is <4x spike added),	if the PS %R is >125%, qualify detects as "J+." If the PS %R is ≤125%, qualify detects as "J."
If the MS %R is <75% but ≥30% and a PS was analyzed (if required when sample concentration is <4x spike added),	if the PS %R is <75%, qualify detects as "J-" and non-detects as "UJ." If the PS %R is ≥75%, qualify detects as "J" and non-detects as "UJ." if the PS %R is <75%, qualify detects as "J"
If the MS %R is <30% and a PS was analyzed,	if the PS %R is <75%, qualify detects as "J-" and non-detects as "R." If the PS %R is ≥75%, qualify detects as "J" and non-detects as "UJ."

4.6.5 Replicate

Replicate analyses are indicators of laboratory precision based on each sample matrix.

Criteria: One replicate must be analyzed for each matrix or each batch, with a minimum frequency of one per 20 samples.

> Samples identified as FBs or EBs should not be used for replicate or MSD analysis.

An acceptance limit of 20% for the RPD shall be used for sample values $\geq 5X$ the PQL. For solid and waste samples, it may be appropriate to accept an RPD of up to 35% based on professional judgment.

A control limit of \pm the PQL shall be used for sample values > the PQL but <5Xthe PQL, including the case when only one of the replicate sample values is > the PQL but <5X the PQL.

No precision criteria apply when both replicate sample values are < the PQL.

When a replicate was not performed but an MSD was analyzed, the MS/MSD RPDs are evaluated as specified in Section 4.3.5. If neither a replicate nor an MS/MSD were analyzed, the laboratory may run an LCSD to measure precision. LCS/LCSD RPDs are evaluated as specified in Section 4.3.5.

Evaluation	Action
If no replicate sample, MSD, or LCSD were analyzed for each matrix or for each data package, or if an FB or EB was used for the replicate analysis,	qualify all detects of the same matrix as "J" and all non-detects of the same matrix as "UJ."
If the original result and replicate result are both ≥5X the PQL and the RPD exceeds the appropriate control limit,	qualify all associated detects of the same matrix as "J" and all associated non-detects of the same matrix as "UJ."
If the original and/or replicate result is > the PQL but <5X the PQL (including non-detects) and the difference between the original result and replicate result is > the PQL,	qualify all associated detects of the same matrix as "J" and all associated non-detects of the same matrix as "UJ."
If the RPD between the original sample and duplicate sample result is >100%,	use professional judgment.

4.6.6 LCS

Data for LCSs are generated to provide information on the accuracy of the analytical method and the overall laboratory performance, including sample preparation.

Criteria: LCSs shall be analyzed using the same sample preparation and analysis methods used for samples, with one LCS analyzed with each batch of up to 20 samples. Multiple LCS analyses may not be used to meet acceptance criteria; that is, if multiple LCSs are analyzed for a batch and any failures occur, the failed LCS will be used to qualify the data. For all aqueous LCS results, the recovery acceptance criteria are 80% to 120%, except antimony and silver. The recovery acceptance criteria for silver and antimony are laboratory-specified. LCS failures for silver and antimony shall be discussed in the data validation report but shall not be subject to the reanalysis requirement.

For all solid LCS results, the recovery acceptance criteria are established by the agency that prepared the reference material or statistically-derived criteria developed by the laboratory. The laboratory should report these acceptance

criteria on the LCS reporting form. If solid LCS acceptance criteria are not provided, then 30% to 150% should be used to assess soil results. If this situation occurs, it should be noted in the data validation report. A solid LCS should be analyzed for mercury in solid analyses.

An aqueous LCS is not required for mercury or cyanide analyses. Since the ICV is always digested/distilled for these analyses, it is equivalent to an LCS.

Evaluation	Action
If an LCS was not analyzed,	qualify all detects as "J" and all non- detects as "UJ."
Aqueous LCS If the LCS %R is	
>120%,	qualify all associated detects as "J+."
<80% but ≥50%,	qualify all associated detects as "J-" and all associated non-detects "UJ."
<50%,	qualify all associated detects as "J-" and all associated non-detects as "R."
Solid LCS If the LCS %R is	
> the upper control limit,	qualify all associated detects as "J+."
≥30% but < the lower control limit,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
<30%,	qualify all associated detects as "J-" and all associated non-detects as "R."

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Evaluation (concluded)	Action (concluded)
If an LCS/LCSD pair was analyzed and recoveries for any target analyte are both above and below acceptance criteria,	qualify all detects as "J" and all non- detects as "UJ."
If an aqueous LCS was analyzed for soil matrices,	qualify all detects as "J" and all non- detects as "UJ."

4.6.7 RLV

Criteria: RLV standards (LLCCV for ICP-AES, ICP-MS and AA) standards are analyzed

at the beginning of each analytical run as a measure of accuracy near the reporting limit. The standards are prepared with concentrations at the PQLs.

The recovery acceptance criteria for these analyses is 80% - 120%.

Evaluation	Action
If the RLV recovery is	
>120%,	qualify all associated detects <5X the PQL as "J+."
<80% but ≥30%,	qualify all associated detects <5X the PQL as "J-" and all associated non-detects as "UJ."
<30%,	qualify all associated detects <5X the PQL as "J-" and all associated non-detects as "R."

4.6.8 Method-specific analytical requirements (inorganic)

4.6.8.1 ICP-AES and ICP-MS Methods

ICS

The ICP-AES and ICP-MS ICSs (interference check sample solution A (ICS A) and interference check sample solution AB (ICS AB)) verify the instrument's interelement and background correction factors.

Criteria: An ICS A must be analyzed at the beginning of each sample analysis run.

Absolute values for all ICS A target analytes, except those in the ICS A solution, must be \leq the MDL.

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Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation	Action
If the ICS A sample was not analyzed at the required frequency,	note the deficiency in the data validation report.
If the sample concentrations of aluminum, calcium, iron, and/or magnesium are < their respective concentrations in the ICS A solution,	accept the sample results without qualification.
If the sample concentrations of aluminum, calcium, iron, and/or magnesium are comparable to or > their respective concentrations in the ICS A solution, and the ICS A result for a non-spiked analyte is	
positive and \geq the MDL,	qualify all associated sample detects < 50X the ICS A result as "J+."
negative and the absolute value of the result is > the MDL but ≤2X the MDL,	qualify all associated detects <50X the absolute value of the ICS A result as "J-" and all associated non-detects as "UJ."
negative and the absolute value of the result is >2X the MDL,	qualify all associated detects <50X the absolute value of the ICS A result as "J-" and all associated non-detects as "R."
positive result ≥MDL, but element is not present in the ICS AB solution,	qualify all associated sample detects <50X the ICS AB result as "J+."
negative and the absolute value of the result is > the MDL but ≤2X the MDL, but element is not present in the ICS AB solution,	qualify all associated detects <50X the absolute value of the ICS AB result as "J-" and all associated non-detects as "UJ."

Criteria: An ICS AB must be analyzed at the beginning of each sample analytical run.

ICS AB results for the target analytes in the ICS AB solution must be within 80% to 120% of the true value.

If the recovery criteria are not met, the analyst may either terminate the analysis or continue and re-analyze the failed constituents at a later time.

Evaluation	Action
If the ICS AB was not analyzed at the required frequency,	note the deficiency in the data validation report.
If the concentrations of aluminum, calcium, iron, and magnesium in the sample are < their respective concentrations in the ICS AB solution,	accept the sample results without qualification.
If the sample concentrations of aluminum, calcium, iron, and magnesium are comparable to or > their respective concentrations in the ICS AB solution and the ICS AB recovery for an analyte is	
>120%,	qualify all associated detects as "J+."
<80% but ≥50%, <50%,	qualify all associated detects as "J-" and all associated non-detects as "UJ." qualify all associated detects as "J-" and all associated non-detects as "R."

ICP Serial Dilution (SD)

The ICP SD monitors physical or chemical interferences that may exist in each sample matrix.

Criteria: A SD must be analyzed for each matrix in an analytical run. If the undiluted results for the sample used for SD are ≥50X the MDL, then the %D between a 5X dilution result and the original result must agree within 10%.

> Samples with elevated concentrations that require dilutions >50X or that require multiple dilutions must also meet these requirements. However, care should be used in evaluating the result from the undiluted sample since it may be above the linear range of the instrument and would not apply.

No acceptance criterion applies when the undiluted sample result is <50X the MDL.

Evaluation	Action
If frequency requirements are not met,	qualify all detects ≥50X the MDL as "J."
If the result for any analyte in the sample used for SD analysis is ≥50X the MDL and the %D is >10%,	qualify detects for all samples of the same matrix in the batch as "J" and non-detects all samples of the same matrix in the batch as "UJ."

Instrument Tuning for ICP-MS

Tuning and performance criteria are established to ensure mass resolution and identification. These criteria are not sample specific. Conformance is determined using standards. Therefore, these criteria should be met in all circumstances.

Criteria: The ICP-MS tune shall be evaluated daily. The tuning solution must contain elements representing all of the regions of interest. The mass calibration must be within 0.1 atomic mass units (AMU) of the true value. The resolution must be verified to be <0.9 AMU full width at 10% peak height.

Evaluation	Action
If tunes were not run daily or if all mass calibration and resolution criteria were not met,	use professional judgment to determine which data should be used. It is suggested that all associated detects should be qualified as "J" and all associated non-detects should be qualified as "UJ."
If multiple QC failures also occurred,	qualify all results as "R."

IS Performance for ICP-MS

IS criteria ensure that ICP-MS sensitivity and response are stable and acceptable during each analysis. They also allow for monitoring of indigenous quantities of the ISs.

Criteria: The intensity of the IS in the samples must fall within 60% to 125% of the intensity of the IS in the ICAL standard. The intensity of the IS in the bracketing CCVs and CCBs must fall within 80% to 120% of the intensity of the IS in the ICAL standard.

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Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation	Action
If no IS was used,	qualify all results as "R."
If the IS intensity for a sample is	
≥30% but <60% of the intensity of the IS in the calibration standard,	qualify all associated detects as "J+" and all associated non-detects as "UJ."
>125% but <160% of the intensity of the IS in the calibration standard,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
<30% or >160 % of the intensity of the IS in the calibration standard,	qualify all associated results as "R."
If both the CCV and CCB have IS intensities outside of the recovery limits,	all associated sample results may be qualified as "J/UJ" due to instrument drift based on professional judgment.

4.6.8.2 Total Organic Carbon (TOC) by SW-846 Method 9060

Criteria: Quadruplicate analyses are required. The average is to be reported.

Evaluation	Action
If quadruplicate analyses were not run,	qualify all detects as "J" and all non- detects as "R."

4.6.8.3 Total Cyanide and Cyanide Amenable to Chlorination

After evaluation of total cyanide data (and application of appropriate qualifiers) using Sections 4.6.1 through 4.6.7, proceed to the following guidance for further evaluation of analytical data for total and amenable cyanide,

Criteria: Sample preparation includes distillation of the samples. In addition to the field samples, the QC samples and one standard and the ICV must be distilled. The LCS meets the requirement for distillation of one standard if the concentrations of the LCS and ICV are different.

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samples and/or standards were not distilled,

Evaluation

If the samples, appropriate QC samples, and appropriate standards were not distilled,

If the field samples were distilled but the QC qualify all associated detects as "J" and

Procedure No: AOP 00-03

all associated non-detects as "UJ."

Issue No: 05

The remainder of this section is provided as guidance for the assessment of data for cyanide amenable to chlorination. Total cyanide data are to be reviewed according to the guidance in Sections 4.6.1 through 4.6.6.

Cyanide amenable to chlorination (decomposed by chlorination) is derived by measuring the total cyanide and the cyanide remaining after chlorination (referred to as chlorinated cyanide below). The amenable cyanide is calculated as the difference between the total cyanide and the chlorinated cyanide. Biases in the cyanide after chlorination will result in a bias in the opposite direction for the calculated amenable cyanide result.

The actual analysis of the sample is the same for the total and the chlorinated analysis; only the sample preparation is different. The laboratory will generally run the total and the chlorinated samples together in the same batch and will use the same calibration and calibration checks for both the total and chlorinated cyanide samples. The laboratory should identify which sample is the total sample and which sample is the chlorinated sample.

LCS and MS solutions are generally of the form that will not decompose with chlorination giving a %R of zero for chlorinated cyanide. When the chlorinated cyanide recovery is 0%, the reported amenable cyanide recovery is equal to the total cyanide recovery. Alternatively, the LCS and MS solutions may not decompose with chlorination and will give a recovery of 0% for amenable cyanide. The laboratory should discuss in the case narrative the form of solution that was used for LCS and MS analyses. The chlorinated cyanide LCS and MS recoveries may not be reported. In this case, the chlorinated cyanide LCS and MS recoveries can be determined from the raw data. The amenable cyanide LCS and MS data should be evaluated using the criteria given for total cyanide.

When total and amenable cyanide are analyzed together in the same run, all initial and continuing calibration qualification applied to the total cyanide results should also be applied to the amenable cyanide results. When total and chlorinated cyanide are not analyzed in the same run, qualifications applied to the total and chlorinated cyanide results should also be applied to the amenable cyanide results. Signed qualification for chlorinated cyanide results should be reversed for amenable cyanide results; that is, a "J+" for chlorinated cyanide results would be a "J-" for the associated amenable cyanide.

Major differences in total and chlorinated results are generally attributed to incomplete destruction of cyanide complexes such as thiocyanide. Non-detects for total cyanide with significant cyanide results for chlorinated cyanide (negative amenable cyanide) may indicate a significant presence of thiocyanide or other cyanide complexes in the sample.

Criteria: A CCV standard must be analyzed once every 10 injections or every two hours, whichever is more frequent. The evaluation of CCV data applies to all CCVs that

bracket samples of interest.

The recovery acceptance criteria for CCV analysis results must be within 90% to

Evaluation	Action
If a chlorinated cyanide CCV %R is	
<90% but ≥75%,	qualify all associated detects for amenable cyanide as "J+."
>110% but <125% and the amenable cyanide result is < the total cyanide result,	qualify all associated detects for amenable cyanide as "J-" and all associated non-detects for amenable cyanide as "UJ."
<75%,	qualify all associated detects for amenable cyanide as "R."
>130% and the amenable cyanide result is < the total cyanide result,	qualify both detects and non-detects for amenable cyanide as "R."

Criteria: A minimum of one MB should be analyzed for every 20 samples. The same reagents used for the sample must be used to prepare the MB. A CCB must be analyzed after each CCV and at the end of every analytical sequence. All CCBs that bracket samples of interest shall be reported and assessed.

> If cyanide is detected in an MB or CCB, the chlorinated sample results must be assessed to determine the impact on amenable cyanide results.

No contaminants \geq the MDL should be present in the blanks.

Evaluation	Action
If a chlorinated cyanide MB or CCB value is positive and ≥ the MDL and the chlorinated sample result is a detect <5X the MB/CCB value,	qualify all associated detects for amenable cyanide as "J-"and, if the total cyanide result is > the MDL, qualify all associated non-detects for amenable cyanide as "UJ."
If the absolute value of the negative chlorinated cyanide MB or CCB value is > the MDL and the chlorinated sample result is < 5X the MDL or non-detect,	qualify all associated detects for amenable cyanide <5X the absolute value of the blank as "J+."

IMPORTANT NOTICE:

Criteria: The absolute value of a negative amenable cyanide result must be <3X the MDL.

Note: Laboratories will generally report the amenable cyanide as non-detect when the chlorinated result is > the total cyanide result. The raw data may need to be reviewed to determine the actual negative amenable cyanide result.

Evaluation	Action
If the absolute value of a negative amenable cyanide result is >3X the MDL and the total cyanide is a non-detect,	note in the data validation report but do not qualify any results.
If the absolute value of a negative amenable cyanide result is >3X the MDL and the total cyanide is a detect,	qualify the amenable cyanide result as "UJ" if the absolute value of the amenable cyanide result is >3X but ≤10X the absolute value of the blank and as "R" if the absolute value of the amenable cyanide result is >10X the absolute value of the blank.

4.6.8.4 Total/Partial Inorganic Analyte Results

Several inorganic analytes are analyzed and reported as total and partial results, (i.e. total chromium/hexavalent chromium, total kjeldahl nitrogen (TKN)/ ammonia, hardness/calcium and magnesium, total alkalinity/carbonate and bicarbonate, total cyanide/amenable cyanide, and total phosphorus/phosphate. In these cases, it is expected that the partial value will be \leq the total value. These reported values may or may not be obtained from the same analytical method. When the reported result for the partial analyte is > the result for the total analyte, one or both results are suspect. The extent to which the quality of the data is affected must be determined. The following criteria should be used for guidance.

Note: Comparisons are made at the elemental level, that is, total nitrogen should be > the nitrogen in ammonia not > the ammonia.

Criteria: When both a partial and a total result are reported, the result for the partial analyte must be \leq the result for the total analyte.

If the partial result is > the total result, the laboratory should be contacted for further information. If the laboratory cannot be contacted or cannot provide sufficient explanation, the following criteria apply.

If the partial result is > the total result and both results are $\geq 5X$ the PQL, then the RPD between the two values should be $\leq 20\%$.

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home page.

If the partial result is > the total result and one or both results are <5X the PQL, then the difference between the two values should be \le the partial analyte's PQL.

Evaluation	Action
If the partial result is > the total result, and	
both the total and partial results are $\geq 5X$ the PQL and the RPD is $\geq 20\%$,	may qualify one or both results as "R" or "J" based on professional judgment.
one or both results are <5X the PQL and the difference between the two results is > the partial analyte's PQL,	may qualify one or both results as "R" or may qualify all associated detects as "J" (with or without bias) or may qualify all associated non-detects as "UJ" based on professional judgment.

Partial Analyte Conversions

To Convert	Co :	Multiply By
O-phosphate	Phosphorus	0.326
Ammonia	Nitrogen	0.824
Ca	Hardness	2.497 (2.5, if titrated)
Mg	Hardness	4.118 (4.12, if titrated)
	(total hardness is the sum of the	
	calculated Ca and Mg hardness results)	

Alkalinity Relationships

The results obtained from the phenolphthalein and total alkalinity determinations offer a mean for stoichiometric classification of the three principal forms of alkalinity present in many waters.

- Carbonate alkalinity is present when the phenolphthalein alkalinity is not zero and is not < the total alkalinity.
- Hydroxide alkalinity is present if the phenolphthalein alkalinity is > half the total alkalinity.
- Bicarbonate alkalinity is present if the phenolphthalein alkalinity is < half the total alkalinity.

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Result of Titration	Alkalinity 86	Carbonate Alkalinity as CaCO ₃	Bicarbonate Alkalinity as CaCO ₃
P=0	0	0	Т
P< 1/2 T	0	2P	T-2P
P= 1/2 T	0	2P	0
P> ½ T	2P-T	2(T-P)	0
P=T	T	0	0

P- phenolphthalein alkalinity; T- total alkalinity

Phenolphthalein alkalinity is the term traditionally used for the quantity measured by titration to pH 8.3. It is not routinely reported by the laboratories but could be calculated using the amount of titrant used to reach pH 8.3. There is usually a column on the alkalinity worksheet that contains this information. Since total alkalinity is reported, it should be verified that the carbonate, bicarbonate, and hydroxide values do not exceed the total.

No conversion is required to compare hexavalent chromium to total chromium.

The calculation for amenable cyanide is detailed in Section 4.6.8.3.

4.6.8.5 Data Validation for Analyses by NIOSH Method 7300

This procedure is for the analysis of elements capable of detection by ICP-AES analysis, including Be, in air samples. The samples are collected onto filters at a flow rate of 1 to 4 L/min. The working range of this method is 0.005 to 2.0 milligrams (mg) per cubic meter (2.5-1000 μg/sample) for each element in a 500-L air sample.

The laboratory must follow the requirements specified in NIOSH 7300 as well as any requirements specified in the AHIA Accreditation Requirements. Compliance requirements for satisfactory data reporting include:

- Case narrative
- Initial calibration data
- Continuing calibration data
- Media blanks, FBs, and preparation blank data
- LCS/LCSDs
- Sample results
- Instrument run logs

Initial Calibration

Compliance requirements for satisfactory instrument calibration are established to ensure that the instrument is capable of producing acceptable quantitative data. Initial calibration demonstrates that the instrument is capable of acceptable performance at the beginning of the analysis run, and CCV documents that the ICAL is still valid.

Criteria: Calibrate the spectrometer according to the manufacturer's recommendations.

Typically, an acid blank and a $10\mu\text{g/mL}$ multi-element* working standard is

used.

*refer to method for chemically compatible combinations of elements

Evaluation	Action
If the minimum number of standards as defined in the criteria section was not used for ICAL,	qualify all associated detects as "J" and all associated non-detects as "UJ."
If the instrument was not calibrated as required,	qualify all associated detects and all associated non-detects as "R."

Calibration Verification

Criteria: A CCV standard must be analyzed for every 10 samples. The recovery

acceptance criteria for analysis results must be within the 90% to 110% of the

true value for all analytes.

Evaluation	Action
If the CCV frequency criteria were not met	qualify all associated detects as "J" and all associated non-detects as "R."
If the ICV or CCV %R is	
≥75% but <90%,	qualify all associated detects as "J-" and all associated non-detects as "UJ."
≥110% but <125%,	qualify all associated detects as "J+."
If the ICV or CCV %R is	
<75%,	qualify all associated detects as "J-" and all associated non-detects as "R."
>125%,	qualify all associated detects as "R."

Blanks

Blank analysis results are assessed to determine the existence of contamination. During sampling, two to ten FBs are collected per sample set. During sample preparation, a reagent blank (MB) and media blanks are included with the samples during the digestion process. The average media blank result (ug/mL) is subtracted from the sample result (ug/mL) in the final

IMPORTANT NOTICE:

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

calculation. In instances where more than one blank (FB or MB) associated with a given sample is ≥ the MDL, qualification is to be performed using the associated blank with the highest concentration of contaminant.

Criteria: The same reagents used for the sample digestion must be used to prepare the MB. No contaminants > the MDL may be present in the blanks.

Evaluation	Action
If the frequency criteria were not met,	note the deficiency in the data validation report.
If problems with any blank exist,	all data associated with the batch must be evaluated to determine whether there is an inherent variability in the data for the batch or if the problem is an anomaly not affecting other data.
If any analyte concentration in the blank is in excess of the PQL,	the lowest reported concentration in the associated samples must be > 10X the concentration in the blank or results must be qualified or rejected.
If a blank value is ≥ the MDL,	qualify all associated detects <5X the blank concentration as "J."

LCS/LCSD

The LCS/LCSD serves as a measure of the overall performance of all steps in the analysis, including sample preparation.

Criteria: The LCS/LCSD must be analyzed using the same preparation and analysis methods used for samples, with one LCS/LCSD analyzed for each batch of up to 10 samples.

> All LCS/LCSD results for air filters must fall within the recovery and RPD control limits established by the agency that prepared the reference material or by statistically-derived limits developed by the laboratory. The laboratory is to include these limits on the LCS/LCSD reporting form.

Professional judgment may be used to determine the need for qualification of sample results based on whether or not the LCS, LCSD, or both meet QC acceptance criteria.

Evaluation Action If the LCS/LCSD was not analyzed, qualify all associated detects as "J" and all associated non-detects as "UJ." If the %R is... qualify all associated detects as "J+." > upper control limit, >30% and < the lower control limit, qualify all associated detects as "J-" and all associated non-detects as "UJ." qualify all associated detects as "J-" and all <30%, associated non-detects as "R." qualify detects for associated compounds as If RPD criteria were not met, "J" and non-detects as "UJ.

Procedure No: AOP 00-03

Issue No: 05

Dilutions

The PQLs must be adjusted to reflect all sample dilutions. Original undiluted results document the actual MDLs for non-detected compounds.

Criteria:

It must be determined that the laboratory strove to make dilutions in such a way that the final concentration was measured in the mid-range of the calibration curve, and that the laboratory did not report results from measurements above the highest concentration standard.

Evaluation	Action
If any samples required dilution because one or more analytes exceeded the calibration range, and	
the original undiluted results were reported,	qualify all associated detects > the high standard as "J."
only the diluted results were reported,	qualify all associated non-detects as "UJ."

4.6.8.6 Method 300.0/9056A, Determination of Inorganic Anions by Ion Chromatography and Method 314.0/9058, Determination of Perchlorate

Criteria: RTs for ICV and CCV analyses must be within 30 seconds (0.5 minutes) of those in the ICAL midpoint.

Evaluation	Action
If any target analyte RT in the CCV varies by more than ±30 seconds from the midpoint retention time obtained during ICAL,	qualify all associated detects as "J" and all associated non-detects as "UJ."

Procedure No: AOP 00-03

Issue No: 05

4.7 Procedure for Radiochemical Analyses Validation

4.7.1 Quantification

Criteria: Radiochemical analytical results shall be reported as measured and shall include the TPU at the 95% confidence level $(2-\sigma)$.

Note: Some programs may request the result be reported with the $1-\sigma$ uncertainty. When this is the case the reported uncertainty must be multiplied by two for evaluation of quantitation and replicate error ratio (RER).

The laboratory shall report all results regardless of concentration or sign and shall not report any result as "less than."

The laboratory shall include a sample minimum detectable activity (MDA) calculated using sample-specific parameters.

For programs that require application of one final qualifier to sample results, if the "BD" qualifier is applied to a sample result, the result shall not be further qualified as "J" due to other QC failures.

Note: Some programs may request the "U" qualifier instead of the "BD" qualifier.

Extremely large errors/uncertainties may indicate inappropriate error calculation. If large errors/uncertainties are reported with the results, the laboratory should verify the calculations.

Evaluation	Action
If the sample result is < the 2-σ TPU,	qualify the result as "BD."
If the sample result is < the MDA,	qualify the result as "BD."
If the sample result is \geq the MDA but $<3X$ the MDA,	qualify the result as "J."
If the absolute value of a negative result (excluding gamma spectroscopy, addressed below) is > the MDA,	qualify the result as "R."

Evaluation (concluded)	Action (concluded)
Gamma Spectroscopy: If the absolute	qualify the result as "R."
value of a negative sample result is >2X	
the MDA,	

If the "BD" qualifier is applied to a sample result, the result shall not be further qualified as "J" due to other QC failures.

4.7.2 Blanks

Blank analysis results are assessed to identify contamination. The criteria for evaluation of blanks apply to all blanks associated with the samples.

Criteria: One MB (or preparation blank) must be analyzed for each matrix and each batch, or for every 20 samples, whichever is most frequent.

MB analysis is required for all analyses requiring sample preparation.

Samples associated with any preparation blank result that is \geq the MDA shall be redigested and reanalyzed. Exceptions to this requirement are samples for which the measured sample concentration is $\geq 10X$ the preparation blank value.

Evaluation	Action
If the prep blank was not analyzed at the proper frequency and there are sample concentrations ≥ the MDA but <5X the MDA,	qualify those results as "J."
If the blank result is positive and is statistically >0.0 (i.e., > the 2- σ TPU and \geq the MDA),	qualify all associated sample results ≥ the MDA but <5X the blank value as "NJ+."
Evaluation (concluded)	Action (concluded)
If the absolute value of a negative blank result is > the MDA,	qualify all associated detects ≥ the MDA but <5X the MDA as "NJ"
	The program may require results < the MDA to be qualified "UJ."
If the absolute value of a negative blank result is >5X the MDA,	notify the laboratory and qualify all associated sample results as "R."

IMPORTANT NOTICE:

Evaluation (concluded)

If the absolute value of the negative blank result is > 5X the MDA for liquid scintillation analyses such as tritium, where the calibration blank is subtracted from the result,

Action (concluded)

notify the laboratory and qualify all sample results <5X the MDA as "R."

Procedure No: AOP 00-03

Issue No: 05

4.7.3 Sample-Specific Chemical/Tracer Recovery

An addition of a known quantity of radioactive or chemically similar material to a sample prior to chemical separation is used to determine the amount of the analyte recovered.

Criteria: Recovery guidelines for tracer and carrier results shall be 50% to 105%.

Optionally, low tracer recoveries may be evaluated from the total area counts.

Samples with low recoveries but with tracer area counts >400 counts may or may not be qualified based on professional judgment.

The quantity of tracer material used should be adequate to provide a maximum of 10% uncertainty at the 95% confidence level in the measured recovery.

Evaluation	Action
If a recovery for a chemical carrier or tracer isotope is	
>105% but ≤125%,	qualify all associated results ≥ the MDA as "J" The program may require results < the MDA to be qualified "UJ."
If a recovery for a chemical carrier or tracer isotope is	
>125%,	qualify all associated results as "R."
If a recovery for a chemical carrier is	
<50% but ≥20%,	qualify all associated results ≥ the MDA as "J+."
<20%,	qualify all associated results ≥ the MDA as "J+" and all associated results < the MDA as "R."

Evaluation (concluded)	Action (concluded)
If a recovery for a tracer isotope is	
≥10% but <50%,	qualify all associated results ≥ the MDA as "J+."
<10%,	qualify all associated results ≥ the MDA as "J+" and all associated results < the MDA as "R."

4.7.4 MS

MS analyses are performed on field samples, except as noted below, as a measure of the ability to recover the analyte from a particular matrix.

Criteria: The MS data shall not be used to evaluate sample data unless the MS sample was from the same client and of similar matrix.

> The recovery acceptance criteria for MS results must be within 75% to 125% unless the sample result is >4X the spike added (see Section 4.1.20).

One MS sample shall be analyzed from each batch, with a minimum frequency of one per 20 samples.

Samples identified as FBs or EBs shall not be used to satisfy the spike analysis requirement.

If an MS result fails to meet recovery criteria, all associated samples shall be redigested and reanalyzed. Unfiltered water samples and unprepared solid samples are exempt from the reanalysis requirement. Results for unfiltered water samples and unprepared solid samples for which the MS failed the acceptance criteria may be reported and qualified without reanalysis.

MSs are not required for gamma spectroscopy, radon-222, or any analyses utilizing standard addition spike or a tracer or carrier that is chemically identical to the analyte. In addition, radium-226 analyses that employ a barium-133 tracer are exempt from the MS requirements. For radium-228 analysis, an MS is required if the final actinium separation, which is not traced by barium-133, does not incorporate a carrier recovery.

Evaluation	Action
If the MS sample was from another client or of a dissimilar matrix, the frequency criteria of the MS was not met, no MS was analyzed, or an FB or EB was used for the MS,	qualify all results ≥ the MDA as "J." The program may require results < the MDA to be qualified "UJ."

Evaluation (concluded)	Action (concluded)
If an MS %R is	
<25%,	qualify all associated results ≥ the MDA as "J-" and all associated results < the MDA as "R."
<75% but ≥25%,	qualify all associated results ≥ the MDA as "J" The program may require results < the MDA to be qualified "UJ."
>125% but ≤150%,	qualify all associated results ≥ the MDA as "J+."
>150%,	qualify all associated results as "R."

4.7.5 Replicate

Replicate analyses indicate laboratory precision based on each sample matrix.

If an MS/MSD was analyzed in place of a replicate, the following criteria are applied to the MS/MSD results. If insufficient sample was submitted to analyze a replicate or MS/MSD, the laboratory may analyze an LCS/LCSD to measure precision using the following criteria for evaluation.

Criteria: One replicate sample shall be analyzed from each batch with a minimum frequency of one per 20 samples. The replicate data shall not be used to evaluate associated sample data unless the replicate sample was from the same client and of similar matrix.

The RER calculated using the $2-\sigma$ TPU is used to determine replicate precision for radiochemical results.

The radiochemical replicate determinations shall agree when the 95% confidence level uncertainties are considered. That is, the RER shall be <1.0.

Samples identified as FBs or EBs shall not be used to satisfy the replicate analysis requirement.

No precision criteria applies to samples with activities < the MDA, including those where one result is > the MDA and one result is < the MDA.

Replicate analyses may not be possible in tritium analyses when the moisture content is too low or the sample size is too small. A discussion of this problem shall be included in the laboratory case narrative, with no qualifiers applied.

Evaluation	Action
If no replicate sample, no MSD, and no LCSD was analyzed for each matrix or for each data package,	qualify all results ≥ the MDA of the same matrix as "J." The program may require results < the MDA to be qualified "UJ."
If frequency criteria are not met, or if an FB or EB was used for the replicate,	qualify all results ≥ the MDA as "J." The program may require results < the MDA to be qualified "UJ."
	Note: Some programs may not require replicate evaluation on non-client samples. For these programs, note this in the data validation report, with no qualifications applied.
If the RER is >1.0 and ≤ 3.0 ,	qualify all associated results ≥ the MDA as "J." The program may require results < the MDA to be qualified "UJ."
If the RER is >3.0,	qualify all associated results for that analyte as "R." Note: Tritium in soils are not qualified "R" when the RER is >3.0.

4.7.6 LCS

The LCS serves as a measure of the overall performance of all steps in the analysis, including sample preparation.

Criteria: One LCS shall be analyzed for each batch up to 20 samples.

For aqueous LCS analytical results, the recovery acceptance criteria shall be within 80% to 120% of the true value. For solid LCS results, the recovery acceptance criteria shall be within the control limits specified by the agency

that prepared the reference material or statistically-derived limits developed by the laboratory. The laboratory shall report the control limits in the QC portion of the deliverable. Multiple LCS analyses may not be used to meet acceptance criteria; that is, if multiple LCSs are analyzed for a batch and any failures occur, the failed LCS will be used to qualify the data.

Evaluation Action If LCS frequency criteria are not met, note the deficiency in the data validation report. If the LCS %R for any analyte is... <30% or >150%, qualify all associated sample results as "R." < the lower control limit but $\geq 30\%$, qualify all associated sample results \geq the MDA as "J-." The program may require results < the MDA to be qualified "UJ." > the upper control limit but $\le 150\%$, qualify all associated sample results \geq the MDA as "J+." If an LCS/LCSD pair was analyzed and qualify all results ≥ the MDA as "J." recoveries of any target analyte were both above and below acceptance criteria, If an aqueous LCS was used for solid qualify all associated results≥ the MDA matrices. as "J." The program may require results < the MDA to be qualified "UJ."

Procedure No: AOP 00-03

Issue No: 05

4.7.7 Instrument Control Charts

In general, there are four types of control charts used to monitor radiochemistry instrumentation performance: efficiency, resolution, centroid, and background.

• Efficiency Control Charts: Used for all instrumentation. A radioactive control source (that does not have to match the counting geometry of the samples) is counted and decay-corrected counts, count rate, activity or efficiency of the source is plotted. Displayed on the same plot are the average, ± 2 - σ control limits, and ± 3 - σ control limits, or just simply upper and lower control limits. Since the frequency of instrument calibration typically ranges from monthly to annually, a control source is counted to show that the instrument response is stable and that the efficiency calibration is valid for the sample count.

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Resolution Control Charts: Used for instrumentation that utilize multi-channel analyzers to create spectra (with the exception of liquid scintillation counters). The full-width halfmaximum (FWHM) of one or multiple peaks of the control source are plotted. Displayed on the same plot are the upper and lower FWHM control limits. This plot shows that there is no increase in instrument noise to negatively impact spectral resolution for the sample count.

- Centroid Control Charts: Used for instrumentation that use multi-channel analyzers to create spectra (with the exception of liquid scintillation counters). The centroid of one or multiple peaks of the control source is plotted. Displayed on the same plot are the upper and lower centroid control limits. This plot shows that the instrument gain is stable and that drift that could lead to poor peak integration or misidentification of peaks has not occurred.
- Background Control Charts: Used for most instrumentation, though has limited value for data validation. An instrument background is performed and the counts or count rate is plotted. Displayed on the same plot are the average, $\pm 2-\sigma$ limit, and $\pm 3-\sigma$ control limits, or just simply upper and lower control limits. This plot shows that the detector has not become contaminated with radioactivity or that the instrument noise has not increased to the point to cause unwanted counts.

For evaluation, a data point outside the control limit means outside the $\pm 3-\sigma$ control limit when the laboratory provides σ -type control charts.

Instead of control charts, the laboratory may provide control summaries that provide control data of multiple detectors and/or types of charts. This is acceptable as long as all the information needed to evaluate instrument control is provided in the summary.

All Radiochemistry Instrumentation

Criteria: The instrument raw data will clearly contain the detector ID and count start date and time for all samples. The control charts will list the detector IDs and list the range of dates plotted. The date range must be current up to the count start date of the sample. For controls that are counted daily or before use, the charts must be updated to the actual date of sample count. For controls that are counted weekly/monthly, the charts must be updated to within a week/month of the sample count. See the specific instrumentation criteria below for control count frequency requirements.

> In general, only the control chart data point appropriate for the sample count start date is evaluated. If the next control chart data point is plotted and shows an extreme outlier, the stability of the counter during the time of the sample count may need to be investigated.

The laboratory should have made an attempt to recount the sample (if possible) and verify the original count results. Professional judgment is needed in this situation, especially if the sample result looks suspect.

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation	Action
If the control chart is missing from the data package or not updated,	request an amended report from the laboratory.
If instrument control frequency was not met for the sample count,	note the deficiency in the data validation report.

Gas Proportional Instrumentation

Criteria: Alpha and beta control sources will be counted daily or before counter use and plotted using efficiency control charts. A background will be counted daily or before counter use and plotted using alpha and beta background control charts.

> If the sample analyte is an alpha-emitter, only the alpha control charts are evaluated. If the sample analyte is a beta-emitter, only the beta control charts are evaluated.

Evaluation	Action
If the efficiency control point is	
above the upper control limit,	qualify all associated results ≥ the MDA as "J+"
below the lower control limit,	qualify all associated results ≥ the MDA as "J" The program may require results < the MDA to be qualified "UJ."
If the background control point is	
above the upper control limit,	qualify all associated results ≥ the MDA but <3X the MDA as "J+."
below the lower control limit,	qualify all associated results ≥ the MDA but <3X the MDA as "J" The program may require results < the MDA to be qualified "UJ."

Liquid Scintillation Instrumentation

Criteria: Vendor supplied unquenched H-3, C-14, and blank control sources will be counted daily or before counter use and plotted using efficiency and background control charts.

For low-energy beta analysis, (such as H-3, Ni-63 or Pu-241) only the H-3 efficiency and background control charts are evaluated. For mid- to high-energy beta analysis, (such as C-14, Cl-36, Sr-90 or Tc-99) only the C-14 efficiency and background control charts are evaluated.

Evaluation	Action
If the efficiency control point is	
above the upper control limit,	qualify all associated results ≥ the MDA as "J+."
below the lower control limit,	qualify all associated results ≥ the MDA as "J" The program may require results < the MDA to be qualified "UJ."
If the background control point is	
above the upper control limit,	qualify all associated results ≥ the MDA but < 3X the MDA as "J+."
below the lower control limit,	qualify all associated results ≥ the MDA but <3X the MDA as "J" The program may require results < the MDA to be qualified "UJ."

Lucas Cell Instrumentation

Criteria: A control source will be counted daily or before counter use and plotted using efficiency control charts.

Evaluation	Action
If the efficiency control point is	
above the upper control limit,	qualify all associated results ≥ the MDA as "J+."
below the lower control limit,	qualify all associated results ≥ the MDA as "J" The program may require results < the MDA to be qualified "UJ."

IMPORTANT NOTICE:

Alpha Spectrometer Instrumentation

Criteria: If calibrated monthly, the calibration standard can also be used as the control source for generating efficiency control chart data. If a semi-annual or annual calibration is performed, a control source will be counted at least monthly and plotted using efficiency control charts. Only one peak is necessary to control chart. The laboratory should perform pulser control checks at least weekly (preferably daily or before use). The pulser checks will confirm that the instrument gain and resolution are stable. Backgrounds will be counted at least weekly and monitored by the laboratory for contamination. Pulser check results and background control charts do not have to be included or evaluated in the data package.

Evaluation	Action
If the efficiency control point is outside of the control limits and the tracer is measured simultaneously with the analyte,	note the deficiency in the data validation report. Note: Although the analyte results will not be biased, the reported tracer yield may be biased.
If an efficiency control point is above the upper control limit and a tracer is not measured simultaneously with the analyte (Ba-133, Np-239 or Th-234),	qualify all associated results ≥ the MDA as "J+."
If an efficiency control point is below the lower control limit and a tracer is not measured simultaneously with the analyte (barium-133, neptunium-239, or thorium-234),	qualify all associated results ≥ the MDA as "J" The program may require results < the MDA to be qualified "UJ."
If the detector background was not counted within one week of the sample count start date,	note the deficiency in the data validation report.

Gamma Spectroscopy Instrumentation

Criteria: A source will be counted daily or before counter use and plotted using efficiency control charts. At a minimum, two peaks need to be control charted for efficiency, resolution (FWHM), and centroid. These peaks are a lowenergy peak (< 100 kilo electron volts [kev]) and a high-energy peak (> 1000 kev). Backgrounds will be counted at least weekly and monitored by the

laboratory for contamination; however, background control charts do not have to be included or evaluated in the data package.

If the low-energy efficiency control point is outside control limits and the highenergy control point is within limits, technically only the low-energy gamma emitting target analytes need qualification. Since the determination of what energy range requires qualification is not straightforward and requires professional judgment, it is acceptable to qualify all target analytes in this situation.

Evaluation	Action
If the efficiency control point is	
above the upper control limit,	qualify all associated results≥ the MDA as "J+."
below the lower control limit,	qualify all associated results ≥ the MDA as "J" The program may require results < the MDA to be qualified "UJ."
If the resolution control point is outside (above or below) of the control limits,	notify the laboratory and qualify all associated sample results as "R."
If the centroid control point is outside (above or below) of the control limits,	note the deficiency in the data validation report.
If the detector background was not counted within one week of the sample count start date,	note the deficiency in the data validation report.

Method-Specific Analytical Requirements - Radiochemical 4.7.8

Gamma Spectroscopy

The laboratory may rejection of a specific gamma spectroscopy analyte result due to various analytical quality issues (e.g., interference, low abundance, no valid peak, or uncertain identification). These data shall be assessed based on professional judgment.

Criteria: The laboratory qualifiers shall be reviewed for "X"- qualified data.

Evaluation	Action
If the result is recommended for rejection by the laboratory,	may qualify the result as "R" based on professional judgment.

Gross Alpha Beta

The flaming of planchets in the gross alpha beta method may result in the loss of beta emitters. The omission of the flaming step may result in diminished alpha particle transmission.

Criteria: The sample preparation documentation shall be examined to determine whether the planchets were flamed. The preferred approach is to count beta, flame the planchets, and count alpha.

Evaluation	Action	
If the planchets were flamed prior to counting for gross beta,	qualify all beta results ≥ the MDA as "J" The program may require results < the MDA to be qualified "UJ."	
If the planchets were not flamed,	qualify all alpha results ≥ the MDA as "J" The program may require results < the MDA to be qualified "UJ."	

Total/Partial Radiochemical Results

Occasionally radiochemical analytes are analyzed and reported as total and partial results, for example, total radium by gross alpha and radium-226 by radon emanation. These reported values are necessarily not obtained from the same analytical method. The following criteria should be used for guidance.

Criteria: When both a partial and a total result are reported, the result for the partial analyte must be \leq the result for the total analyte. If the reported result for the partial analyte is > the result for the total analyte, one or both results are suspect.

The extent to which quality of the data is affected must be determined.

Evaluation	Action	
If the partial result is > the total result,	the laboratory should be contacted for further information.	
If the laboratory cannot be contacted or cannot provide sufficient explanation, the following criteria apply:		
If the total result is \leq the MDA and the partial result is $>$ MDA but $<$ 2X MDA,	no qualification (other than "BD" or "J" due to quantification) of either result is warranted as the results are statistically similar enough.	

Data Validation Procedure for Chemical and Radiochemical Data Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Evaluation (concluded)	Action (concluded)
If the total result is \leq the MDA and the partial result is $> 2X$ MDA,	qualify the total result as "J"
If both results are > MDA and the total result is < the partial, and the RER between the two results is >1.0,	qualify the total result as "NJ-" due to a suspected false negative.

Data Validation Reports

A data validation report shall be produced to discuss the data review and validation; and to document, based on instrumentation and methodology, the OC elements examined. The SMO or Program Manager uses the data validation report to evaluate and determine if nonconformance, corrective actions, or penalties should be pursued. If this procedure is modified based on the professional judgment of the data validator, the data validation report must document the adjustments. Any method-specific QC requirements not addressed in this document must be documented by the data validator in the data validation report or included as an addendum to the procedure. The database administrator submits the data validation report to the SNL/NM Customer Funded Record Center for archiving. The data validation report shall include the following (as appropriate):

5.1 Sample Findings Summary and Validation EDD Files

A data table or spreadsheet summarizing flagged data resulting from the data review and validation. The sample findings summary is to be used by database personnel to facilitate the data entry of data validation qualifiers to the electronic database. However, when laboratory EDD files are available the SNL/NM Validation EDD Generator is used to produce a sample findings summary and validation EDD file. The validation EDD file is subsequently used for electronic data entry of data validation qualifiers to the database. The sample findings summary and validation EDD file shall include the following:

- The site name
- ARCOC number
- Sample number(s)
- Analysis or individual analytes
- Data validation qualifiers
- Any relevant comments

5.2 Data Validation Narrative (format may vary by project)

A summary of samples and all qualifiers applied to the data as a result of the validation process. The narrative shall include the following:

- The date issued
- The names of those to whom the report is issued

- The validator's name
- The laboratory name and SDG identifier
- ARCOC number
- Type of analysis addressed in the report
- Sample/analyte qualification and a general description of why qualification was applied
- Data validation procedure and revision used
- Any relevant comments

5.3 Data Qualification Summary

A summary of the process used for review and validation. The data qualification summary includes the following sections (as appropriate):

- Sample Shipping/Receiving (refer to Section 4.1.1). Are all shipping/receiving and ARCOC issues that could affect data quality and defensibility discussed, and qualifications properly applied?
- Holding Times and Preservation (refer to Section 4.1.1). Are all holding time and preservation issues that could affect data quality discussed, and qualifications properly applied?
- Calibration. Are all calibration (initial and/or continuing/verification) issues that could affect data quality discussed, and qualifications properly applied?
- Tuning. Are all tuning issues that could affect data quality discussed, and qualifications properly applied?
- IS. Are all IS issues that could affect data quality discussed, and qualifications properly applied?
- Isotope Ratios. Are all isotope abundance issues that could affect data quality discussed, and qualifications properly applied?
- Surrogates. Are all surrogate issues that could affect data quality discussed, and qualifications properly applied?
- TICs. If required, are the identification and qualification of TICs discussed?
- Confirmation. Was second-column analysis discussed, and were qualifications properly applied?
- RLV (CRI/LLCCV). Are all CRI/LLCCV issues that could affect data quality discussed, and qualifications properly applied?
- ICP ICS. Are all ICS issues that could affect data quality discussed, and qualifications properly applied?
- ICP SD. Are all SD issues that could affect data quality discussed, and qualifications properly applied?
- Tracer/Carrier. Are all tracer and/or carrier issues that could affect data quality discussed, and qualifications properly applied?
- Blanks. Are all detections of target analytes in all applicable blanks discussed, and qualifications properly applied?
- LCS. Are all LCS issues that could affect data quality discussed, and qualifications properly applied?
- MS. Are all MS issues that could affect data quality discussed, and qualifications properly applied?

- Laboratory Replicates. Are all the laboratory replicate issues that could affect data quality discussed, and qualifications properly applied?
- DLs/Dilutions. Is the appropriateness of the reported DLs discussed? Are sample dilutions discussed?
- Other QC. Are all QC issues that could affect data quality, other than those previously addressed, discussed? Include a brief description of any laboratory nonconformance reports that directly impacted data quality.
- Corrective Action Reports. Discuss or attach laboratory correspondence covering any corrective
 action, clarification, or modification to the report that was required to complete the validation
 process.

5.4 Validation Notes/Worksheets (as appropriate)

The validation notes/worksheets document results of the review and data validation by methodology and show QC results that do not meet acceptance criteria (that is, failures). These notes/worksheets identify data for which holding time/preservation requirements and calibration acceptance criteria were not met; laboratory blanks, FBs, EBs, or TBs were contaminated; surrogate recovery criteria were exceeded; MS/MSDs exceeded limits; and LCS %Rs and replicate RPDs or RERs exceeded acceptance limits. In addition, the validation notes/worksheet identify the validator and the laboratory; include the validator's comments and notes; and include ARCOC numbers, SDG number, types and number of samples analyzed, and sample numbers.

5.5 CVR and ARCOC

The CVR and ARCOC records also are pertinent to data review and validation. These records are supplied by the SMO and the laboratory and are copied and attached to the data validation report.

6 DEFINITIONS

6.1 Data Qualifier Definitions

Data qualifiers are commonly used during the validation process to classify sample data as to their conformance to QC requirements. For the purposes of this procedure, the following code letters and associated definitions are provided:

- BD (below DL) Used in radiochemistry to identify results that are not statistically different from zero.
- J The associated value is an estimated quantity.
- J+ The associated numerical value is an estimated quantity with a suspected positive bias.
- J- The associated numerical value is an estimated quantity with a suspected negative bias.
- N Presumptive evidence of the presence of the material.
- NJ Presumptive evidence of the presence of the material at an estimated quantity.
- NJ+ Presumptive evidence of the presence of the material at an estimated quantity with a suspected positive bias.

- NJ- Presumptive evidence of the presence of the material at an estimated quantity with a suspected negative bias.
- R The data are unusable (compound may or may not be present). Resampling and reanalysis are necessary for verification.
- U The analyte was analyzed for but was not detected. The associated numerical value is the sample quantitation limit.
- UJ The analyte was analyzed for but was not detected. The associated value is an estimate and may be inaccurate or imprecise.

Datum is unqualified if the quality parameters indicate the method was appropriate and that the reported result reflects the true value within the expected analytical uncertainty.

Datum is qualified as estimated (J) if the reported result can be used to infer an estimate of the true value (with a suspected positive or negative bias, as may be indicated), but the quality parameters indicate an uncertainty in the result that is > the expected analytical uncertainty.

Datum is qualified as presumptive (N) if there is question as to whether the analyte is indigenous to the sample or if there is question regarding the identity of the analyte.

Datum is qualified as presumptive and estimated (NJ) when there is evidence of the presence of the material at an estimated quantity (with a suspected positive or negative bias, as may be indicated).

Datum is qualified as unusable (R) if the quality parameters do not support the reported result as a valid indicator of the true value.

Datum is qualified as estimated non-detects (UJ) for results reported as < the DL and for which some other quality concerns exist.

6.2 Sample Detection/Quantification Limits

For purposes of this procedure, the following definitions are provided:

- MDA Minimum detectable activity. A radiological DL. A sample with activity concentration at the minimum detectable concentration (MDC) has a 95% probability of being measured above the decision level, which is the lowest threshold used to distinguish a positive result (i.e., a detect). For the purposes of data validation, the MDC equals the minimum detectable activity.
- MDL Method detection limit. The minimum concentration of a substance that can be measured (quantified) and reported with 99% confidence that the analyte concentration is > zero. This measure of instrument sensitivity takes into account all solutions that have been subjected to all sample preparation steps for the method. In data packages the MDL may be referred to as the DL. For organic data, the MDL will be one-fifth the PQL, and the value associated with the "U" qualifier or the value of the low standard will be used as the POL.

PQL Practical quantitation limit. The lowest concentration of analytes in a sample that can be reliably determined and quantified within specified limits of precision and accuracy by the indicated methods under routine laboratory operating conditions. For the purposes of this procedure, the PQL is considered to be 5X the value of the MDL if not defined by the laboratory. In data packages the PQL may be referred to as the contract-required DL or RL. For inorganic data, the PQL will be 5X the MDL, and the value associated with the "U" qualifier will be used as the MDL.

6.3 Formulas

The %D for LCSs, standards, and SD is calculated as follows:

%D =	[M-T] x 100 T
where,	%D = percent difference M = measured value T = true value or sample value for SD

The RPD for replicate samples is calculated as follows:

$$RPD = \underbrace{\begin{bmatrix} S - R \end{bmatrix}}_{(S + R)/2} \times 100$$
where,
$$RPD = \text{relative percent difference}$$

$$S = \text{sample value (original)}$$

$$R = \text{replicate sample value}$$

The RPD for MS/MSD samples is calculated as follows:

RPD =		<u>/ISD</u> x /ISD)/2	100
where,	RPD MS R	= = =	relative percent difference MS value MSD value

The %R for spiked samples is calculated as follows:

%R =
$$\frac{SSR-SR}{SA}$$
 x 100
where, $\frac{SSR}{SA}$ = spiked sample result $\frac{SR}{SA}$ = sample result $\frac{SR}{SA}$ = spike added

The RER is used to determine replicate precision for radiochemical results. The RER is given by:

$$RER = \underbrace{ \begin{bmatrix} S - R \end{bmatrix}}_{F_{95S} + F_{95R}}$$
 where,
$$RER = \text{replicate error ratio}_{S = \text{sample value (original)}}$$

$$R = \text{replicate sample value}_{F_{95S}} = \text{sample uncertainty (95\% or 2-σ)}_{F_{95R}}$$

$$F_{95R} = \text{replicate uncertainty (95\% or 2-σ)}_{F_{95R}}$$

The linear curve equation is given by:

NOTE: Intercept calculation may vary depending on the laboratory instrumentation. The instrument software used for determining calibration curves often differs from the usual y = mx+b linear equation. When this occurs, it is up to the validator to determine the actual equation used and the corresponding slope and intercept.

The %RSD is calculated as follows:

GLOSSARY OF TERMS

2-σ error: The error reported at the 95% confidence interval.

Acceptance limits: Ranges of acceptable results for each type of QC measurement. They may be defined on a program-specific basis, or they may be derived internally at a laboratory from historic QC performance data. May also be referred to as control limits.

Accuracy: The closeness of agreement between an observed value and the true value. "Precision" is a measure of the reproducibility of a value, without knowledge of the true value. The classic example used to illustrate these terms is a dartboard example: The placement of four darts thrown at a dartboard is considered accurate if the darts are each close to the bull's-eye (regardless of their proximity to one another). Hence, to be both accurate and precise the four darts would need to be grouped closely together and be close to the bull's-eye.

Analyte: That which is analyzed for. This can be chemical (chromium, benzene), biological (fecal coliform bacteria), mineral (asbestos fibers), or radiological (alpha and beta emissions).

Analytical run: The interval (i.e., period of time or series of measurements) within which the accuracy and precision of the measuring system is expected to be stable. Within the analytical run, controls are often analyzed to confirm stability.

Batch: A group of samples which behave similarly with respect to the sampling or the testing procedures being employed and which are processed as a unit. For QC purposes, if the number of samples in a group is > 20, then each group of 20 samples or less will all be handled as a separate batch.

Bias: The difference between the reported result and the true result. Bias may be introduced through field or laboratory variability and error or due to substances in the sample that interfere with the analytical system's ability to provide an accurate measurement. Because the true concentration of an analyte in an environmental sample is generally never known, bias is estimated by using surrogates, MSs, LCSs, and other indicators of analytical accuracy.

Calibration: The process of correlating instrument signal response with analyte concentration. An instrument must be properly calibrated in order to produce accurate results.

Chemical carrier: An identical or similar carrier material used to infer the degree to which the separation processes were effective in separating the analyte from the matrix. Measured gravimetrically or chemically.

Congener: A congener refers to any one particular compound of the same chemical family. For example, there are 209 congeners of Chlorinated Biphenyls (CBs).

Contamination: A component of a sample from an extract that is not representative of the environmental source of the sample. Contamination may stem from other samples, sampling equipment, while in transit, from laboratory reagents, laboratory environment, or analytical instruments. Blanks (instrument blanks, MBs, preparation blanks, TBs, EBs, and FBs) may be used to assess contamination.

Effective Date: 06/19/2017 Issue No: 05

Procedure No: AOP 00-03

Control sample: A QC sample introduced into a process to monitor the performance of the system.

Correlation coefficient (r) or coefficient of determination (r²): A statistical evaluation of the linearity of a calibration curve, i.e. "goodness of fit."

Detect: Sample result ≥the MDL.

Duplicate: A second aliquot of a sample that is treated the same as the original aliquot of sample. (See definitions for field duplicate and replicate.)

Environmental sample: A sample taken unaltered (as much as possible) from the environment (as opposed to a blank, performance evaluation sample, MS sample, etc.). Environmental sample may be referred to as "field sample."

Equipment blank (EB): A sample of analyte-free media (for example, clean water poured over a bailer) that has been used to rinse the sampling equipment. The EB is collected after completion of decontamination and prior to collection of environmental samples. This blank is useful in documenting adequate decontamination of sampling equipment. An EB also may be referred to as a "rinsate blank."

Field blank (FB): A sample containing an analyte-free matrix that is collected and processed in exactly the same manner as an equivalent environmental sample (for example, clean water is poured into a sample container in the same physical location where the environmental sample is collected and is subsequently handled, processed, and analyzed exactly as an equivalent environmental sample). The FB is used to identify contamination resulting from field sample collection techniques.

Field duplicate: A duplicate sample generated in the field used to determine sampling and analytical precision.

Holding time: The period between collection of samples by the samplers and preparation and/or analysis of samples by the laboratory (see Appendix A for required hold times). If the method specifies a holding time to extraction and a holding time to analysis, then two holding times are evaluated. If no holding time to extraction is specified, then the listed holding time is the holding time to analysis. That is, the laboratory cannot extract a sample and store the extract in order to meet holding time. However, professional judgment may be applied here. If the sample preparation includes MS and LCS samples and both of these pass it may be inferred that the stability of the extract has been verified.

Instrument blank: A blank designed to determine the level of contamination associated with the analytical instruments.

Internal standard (IS): A chemical compound added to every blank, sample, and standard extract at a known concentration that is used to (1) compensate for analyte concentration changes that might occur during storage of the extract, and (2) compensate for quantification variations that can occur during analysis. ISs are used as the basis for quantifying target analytes.

Isomer: A chemical species with the same number and types of atoms as another chemical species, but possessing different properties. For example, 2,3,7,8-TCDD refers to only one of the 22 possible TCDD isomers; that isomer which is chlorinated in the 2,3,7,8-position of the dibenzo-p-dioxin ring structure.

Effective Date: 06/19/2017

Procedure No: AOP 00-03 Issue No: 05

Isotope dilution: A means of determining a naturally occurring (native) compound by reference to the same compound in which one or more atoms has been isotopically enriched.

Laboratory control sample (LCS): A known matrix that is spiked with compounds representative of the target analytes at known concentrations. The spiking occurs prior to sample preparation and analysis. An LCS is used to document laboratory overall performance.

Matrix: The substrate that contains the analyte of interest (for example, surface water, drinking water, air, soil, tissue, etc.).

Matrix interference: Bias introduced because something in the sample interferes with the analytical system's ability to provide an accurate measurement. The interference may be physical (turbidity in stormwater runoff may block light transmission in an analysis based on ultraviolet absorbance) or chemical (a chemical similar to the analyte of interest may increase the response of the instrument, resulting in a positive bias).

Matrix spike (MS): A measured amount of sample spiked with a known concentration of target analyte(s). The spiking occurs prior to sample preparation and analysis. An MS is used to assess the bias of a method in a given sample matrix.

Matrix spike duplicate (MSD): Intralaboratory (within the same laboratory) split-samples spiked with identical concentrations of target analyte(s). The spiking occurs prior to sample preparation and analysis. MSDs are used to assess the precision and bias of a method in a given sample matrix.

Method blank (MB): An analyte-free matrix that is prepared and processed at the laboratory in exactly the same manner as an equivalent environmental sample (that is, all reagents are added in the same volumes or proportions as used in sample processing). The MB is used to document contamination resulting from the analytical process.

Non-detect: Sample result < the MDL.

Ongoing Precision and Recovery (OPR): A MB spiked with known quantities of analytes and analyzed as a sample. Its purpose is to assure that results produced by the laboratory remain within the limits specified in EPA Method 1668A for precision and recovery.

Precision: The proximity to one another of the results for multiple measurements on the same sample (i.e., a measure of the repeatability of a measurement process). This does not address proximity to a true value; it is possible for multiple results to show very high precision and yet be completely incorrect by comparison with a true value. Precision is quantified, for example, by calculating the RPD between the result obtained for a sample and that obtained from an associated duplicate or replicate sample. As with accuracy, there is an assumed correlation between quantitative precision as determined via QC analyses and the inferred precision in measurements of unknowns.

Quality control (QC): The system of routine technical activities whose purpose is to measure and control the quality of a product or service so that it meets the needs of users. In other words, QC activities are the tactics used to measure and control quality.

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Radioactive tracer: A radioactive isotope of the analyte that is added to the sample to correct for any losses of the analyte during the chemical separations or other processes employed in the analysis.

Relative dilution factor: The dilution factor ratio (value ≥1) of two samples. For example, if one sample has a dilution factor of 2 and another sample has a dilution factor of 10, the relative dilution factor is 5.

Relative response factor (RRF): A measure of the relative mass spectral response of an analyte compared to its IS. RRFs are determined by analysis of standards and are used in the calculation of concentrations of analytes in samples.

Relative retention time (RRT): The ratio of the RT of a compound to that of a standard (such as an IS).

Replicate (also may be called "sample duplicate"): A duplicate sample generated in the laboratory used to determine analytical precision.

Reporting limit verification (RLV): A low-level verification standard of the same origin as the calibration standard run as a measure of accuracy near the PQL. The RLV is known as the CRI for ICP-AES, ICP-MS, and LC/MS/MS methods, CRA for AA methods, and CRDL for cyanide methods.

Response (also may be called "instrument response"): The signal output of an analytical instrument in which the intensity of the signal is proportionate to the concentration detected. Response is measured by peak area or peak height.

Retention time (RT): The time a target analyte is retained on a chromatography column before elution. The identification of a target analyte is dependent on a target compound's RT falling within the specified RT window established for that compound. RT is dependent on the nature of the column's stationary phase, column diameter, temperature, flow rate, and other parameters.

Sample delivery group (SDG): A group of samples that are processed together by the laboratory. Ideally, all the samples in a batch will be similar enough that matrix QC measurements performed with the batch will be representative of all the samples in the batch.

Spike: A known amount of analyte that is introduced purposely into a sample (either an environmental sample or a blank) for the purpose of determining whether the analytical system can accurately measure the analyte.

Surrogate: A chemical that is similar to the target analyte(s) in chemical composition and behavior in the analytical process but that is not expected to be present in the sample. Surrogates are added to all the environmental samples, blanks, and QC samples in the analytical batch during the preparation stage of the analysis. Surrogates are used to monitor the performance of the analytical process. An example would be the use of fluorinated organic compounds in an analysis that looks for chlorinated and brominated organic compounds. Surrogates also may be called "system monitoring compounds" (SMCs).

Target analyte: A chemical that is being looked for in an analysis.

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Tentatively identified compound (TIC): A compound that is outside the standard list of analytes in a GC/MS method but that is reported based on a tentative match between the instrument response and the instrument's computer library. The identification and quantitation of these compounds is tentative.

Trip blank (TB): A sample of analyte-free media (such as distilled/deionized water) taken from the laboratory to the sampling site and returned to the laboratory unopened. A TB is used to document contamination attributable to shipping and field handling procedures. This type of blank is useful in documenting contamination of volatile organic samples.

Reason Codes

Programs may require that general validation codes be included in their computer databases or EDDs. The following codes are the default codes used when this is required.

- H1 Holding time exceeded for sample analysis
- H2 Holding time exceeded for sample extraction
- H3 Holding time exceeded by >2X the specified holding time
- TP1 Sample improperly preserved
- TP2 Sample not preserved
- TP3 Sample not maintained at required temperature
- TP4 Required sample or extract clean up not performed
- TP5 Sample received unpreserved and preserved at the laboratory
- I1 Initial calibration not reported
- 12 Initial calibration not independently verified
- 13 Slope r² or RRF %RSD criteria not met
- 14 Minimum RRF / Slope not met
- 15 Intercept too large
- 16 Insufficient number of calibration standards used
- C1 Continuing calibration frequency not met
- C2 Continuing calibration %D failed high
- C3 Continuing calibration %D failed low
- B MB contamination at concentration > MDL
- B1 TB contamination at concentration > MDL
- B2 FB/EB contamination at concentration >MDL
- B3 Calibration blank contamination at concentration >MDL
- B4 Negative value for calibration blank absolute value >the MDL
- B5 Negative value for MB absolute value > the MDL
- B6 Negative value for FB/EB/TB absolute value > the MDL
- B7 MB contamination at activity ≥the MDA
- B8 MB frequency not met
- B9 Instrument or calibration blank frequency not met
- IS1 IS / tracer recovery failed high
- IS2 IS / tracer recovery failed low but ≥10%
- IS3 IS / tracer recovery failed low and <10%

- S1 Surrogate(s) failed high
- S2 Surrogate(s) failed low
- S3 Multiple random surrogate failures
- FR1 Result exceeds calibration range
- FR2 No result reported sample lost or damaged
- FR3 Result is less than the MDA / MDL or \leq the 2- σ TPU
- FR4 Negative result absolute value >2X the MDA/MDL
- FR5 RT criteria not met
- FR6 Ion mass ratio criteria not met
- FR7 Result is \geq the MDA and \leq 3X the MDA
- MS1 MS not analyzed or not applicable
- MS2 MS analyte(s) recovery failed high
- MS3 MS analyte(s) recovery failed low
- MS4 MS analytes recovery failed both high and low
- MS5 MS/MSD RPD failed
- RP1 Replicate not analyzed or not applicable
- RP2 Replicate RPD failed
- L1 LCS frequency not met
- L2 LCS analyte(s) recovery failed high
- L3 LCS analyte(s) recovery failed low
- L4 LCS analytes recovery failed both high and low
- L5 LCS/LCSD RPD failed
- DL1 RLV frequency not met1
- DL2 RLV percent recovery failed high1
- DL3 RLV percent recovery failed low1
- CK1 ICS frequency not met
- CK2 ICS analyte(s) failed high
- CK3 ICS analyte(s) failed low
- D1 SD failed %D
- D2 Inappropriate initial dilution
- V1 Conformation analysis not done
- V2 Conformation RPD exceeds criteria
- V3 Confirmation analysis by second method did not confirm original result (LCMSMS)
- X1 Non-specified data quality concern see validation report
- X2 Analysis failed to meet method requirements see validation report
- X3 Required QC documentation missing
- Z1 Spectral identification criteria not met²
- Z2 Minimum peak criteria not met³

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SNL/NM Statement of Work for Analytical Laboratories, Current Revision, Sample Management Office, Sandia National Laboratories/New Mexico, Albuquerque, New Mexico.

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U.S. Department of Energy NNSA Service Center, Model Data Validation Procedure, Current Revision

DoD DOE Consolidated Quality Systems Manual (QSM) for Environmental Laboratories, Version 5.1, January 2017

USEPA Contract Laboratory Program National Functional Guidelines for Organic Superfund Methods Data Review, January 2017.

USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Superfund Methods Data Review, January 2017.

40 CFR 136, Protection of Environment: Guidelines Establishing Test Procedures for the Analysis of **Pollutants**

EPA Method 314, Determination of Perchlorate in Drinking Water by Ion Chromatography

EPA Method 1613B, Tetra-thru-Octa (CDDs) Chlorinated Dioxins and Furans (CDFs)

EPA Method 1668A, Chlorinated Biphenyl Congeners in Water, Soil, Sediment, and Tissue by High-Resolution Gas Chromatography/High-Resolution Mass Spectrometry (HRGC/HRMS)

^{1:} refers to OC for LLCCV/CRI analyses.

^{2:} used when rejecting results that have been X qualified by the lab for interference or short half-life, and also for failed organic spectral matching (GC/MS, diode array, HPLC, etc.).

^{3:} used when rejecting results that have been X qualified by the lab for low abundance, no valid peak, or peak not meeting identification criteria.

EPA Method 1694, Pharmaceuticals and Personal Care Products in Water, Soil, Sediment, and Biosolids by HPLC/MS/MS

EPA Method TO-15 (Rev. 1), Determination of Volatile Organic Compounds (VOCs) in Air Collected in Specially Prepared Canisters and Analyzed by Gas Chromatography/Mass Spectrometry (GC/MS)

SW-846 Method 428, Determination of Polychlorinated Dibenzo-p-dioxin (PCDD), Polychlorinated dibenzofuran (PCDF), and Polychlorinated Biphenyl Emissions from Stationary Sources

SW-846 Method 8000C, Determinative Chromatographic Separations

SW-846 Method 8015C, Nonhalogenated Organics Using GC/FID

SW-846 Method 8015D, Nonhalogenated Organics Using GC/FID

SW-846 Method 8081A, Organochlorine Pesticides by Gas Chromatography

SW-846 Method 8082A, Polychlorinated Biphenyls (PCBs) by Gas Chromatography

SW-846 Method 8151A, Chlorinated Herbicides by GC Using Methylation or Pentafluorobenzylation Derivatization

SW-846 Method 8260B, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry (GC/MS)

SW-846 Method 8260C, Volatile Organic Compounds by Gas Chromatography/Mass Spectrometry (GC/MS)

SW-846 Method 8270C, Semivolatile Organic Compounds by Gas Chromatography/Mass Spectrometry (GC/MS)

SW-846 Method 8270D, Semivolatile Organic Compounds by Gas Chromatography/Mass Spectrometry (GC/MS)

SW-846 Method 8280B, The Analysis of Polychlorinated Dibenzo-p-Dioxins and Polychlorinated Dibenzofurans by High Resolution Gas Chromatography/Low Resolution Mass Spectrometry (HRGC/LRMS)

SW-846 Method 8290A, *Polychlorinated Dibenzodioxins (PCDDs) and Polychlorinated Dibenzofurans (PCDFs) by High-Resolution Gas Chromatography/High-Resolution Mass Spectrometry (HRGC/HRMS)*

SW-846 Method 8310, Polynuclear Aromatic Hydrocarbons

SW-846 Method 8330B, Nitroaromatics and Nitramines by High Performance Liquid Chromatography (HPLC)

NIOSH 7300, Elements by ICP (Nitric/Perchloric Acid Ashing)

IMPORTANT NOTICE:

Appendix A

Sample Preservation and Holding Times

Sample Preservation Techniques and Holding Times

Method	<u>Parameters</u>	<u>Matrix</u>	Volume/Container	<u>Preservation</u>	Holding 1 <u>Sample</u>	imes <u>Extract</u>
2310B, 2320B	Acidity, Alkalinity	Water	500 mL Plastic or Glass	≤6 °C	14 Days	NA
300.0, 9056A	Bromide, Chloride, Fluoride, Sulfate	Water	1 L Plastic	≤6 °C	28 Days	NA
5210B	BOD	Water	1 L Plastic	≤6 °C	48 Hours	NA
9012, 335.4	Total Cyanide Amenable Cyanide	Water Solid/Other	1 L Plastic 125 mL Glass Jar	≤6°C; NaOH; pH > 12 ≤6°C	14 Days 14 Days	NA NA
9060	TOC	Water Solid/Other	250 mL Amber Glass 125 mL Glass Jar	≤6 °C; H ₂ SO ₄ ; pH < 2 ≤6 °C	28 Days 28 Days	NA NA
200.7, 200.8, 6010B, 6020A	All metals except Cr(VI) and Hg	Water Solid/Other	500 mL Plastic 250 mL Glass Jar	HNO₃; pH < 2	180 Days 180 Days	NA NA
7197, 7196A 218.6 3060A	Cr(VI)	Water Water Solid/Other	500 mL Plastic 500 mL Plastic 250 mL Glass Jar	≤6 °C ≤6°C, pH 9-9.5 ≤6 °C	24 Hours 28 Days 30 Days	NA NA 7 Days
245.2, 7470A, 7471A	Hg	Water Solid/Other	500 mL Plastic 250 mL Glass Jar	HNO₃; pH < 2 ≤6 °C	28 Days 28 Days	NA NA
2340C, 130.1	Hardness	Water		HNO₃ ; pH < 2 ≤6 °C	180 Days	NA
350.1, 353.2 365.4, 351.2	Ammonium, Nitrate + Nitrite, Total Phosphorus, TKN	Water	1 L Plastic	≤6 °C; H₂SO₄; pH < 2 ≤6°C; not acidified	28 Days 24 Hours	NA NA

Note: 40 mL vials require caps with Teflon lined septum. All other containers require Teflon lined screw cap lids to minimize container contamination and loss of analyte. *If samples are shipped to the laboratory in EnCoreTM samplers, samples must be extruded and placed in sample containers within 48 hours of sample collection.

IMPORTANT NOTICE:

Sample Preservation Techniques and Holding Times

Method	Parameters	<u>Matrix</u>	Volume/Container	<u>Preservation</u>	Holding Ti <u>Sample</u>	imes <u>Extract</u>
300.0, 354.1	Nitrate, Nitrite,	Water	500 mL Plastic	≤6 °C	48 Hours	NA
365.1, 365.3	Ortho Phosphorus	Water	500 mL Plastic	≤6 °C; H ₂ SO ₄ ; pH < 2	48 Hours	NA
314.0, 9058	Perchlorate by IC	Water	250 mL Plastic or Glass	≤6 °C	28 Days	NA
6850, 331.0 6860, 330.0	Perchlorate by HPLC/MS/MS Perchlorate by IC/ESI/MS/MS	Water	250 mL Plastic or Glass	≤6 °C	28 Days	60 days
0000, 330.0	reichlorate by 10/23//W3/W3	Solid	4 oz. Wide-mouth jar	≤6 °C	28 Days	60 days
410.4	Chemical Oxygen Demand (COD)	Water	250 mL Glass	≤6 °C; H ₂ SO ₄ ; pH < 2	28 Days	NA
1664A	Total Recoverable Oil and Grease	Water Solid/Other	1 L Glass r 125 mL Glass Jar	≤6 °C; H ₂ SO ₄ or HCl; pH < 2 ≤6 °C	28 Days 28 Days	NA NA
9070A, 9071B	Total Recoverable Oil and Grease	Water Solid/Other	1 L Glass r 125 mL Glass Jar	≤6 °C; HCl; pH < 2 ≤6 °C	28 Days 28 Days	NA NA
ASTM D-854	Specific Gravity	Water	500 mL Plastic or Glass	None	None	
9030B, 9034 4500S ² -D	Sulfide	Water Solid/Other	1 L Glass r 125 mL Glass Jar	≤6 °C; NaOH; Zinc acetate; pH > 9 ≤6 °C	7 Days 7 Days	NA NA
2540C, D	TDS, TSS	Water	1 L Plastic	≤6 °C	7 Days	NA

Note: 40 mL vials require caps with Teflon lined septum. All other containers require Teflon lined screw cap lids to minimize container contamination and loss of analyte. *If samples are shipped to the laboratory in EnCoreTM samplers, samples must be extruded and placed in sample containers within 48 hours of sample collection.

IMPORTANT NOTICE:

Sample Preservation Techniques and Holding Times

<u>Method</u>	<u>Parameters</u>	<u>Matrix</u>	Volume/Container	<u>Preservation</u>	Holding Tir Sample	mes <u>Extract</u>
160.4	Volatile solids (volatile residue)	Water	Plastic or glass	≤6 °C	7 Day	NA
9020B	TOX	Water Solid/Other	1 L Amber Glass 125 mL Glass Jar	≤6 °C; H ₂ SO ₄ ; pH < 2 ≤6 °C	28 Days 28 Days	NA NA
418.1	Total Petroleum Hydrocarbon (TPH)	Water	1 L Amber Glass	≤6 °C; HCl; pH < 2	28 Days	NA
1664A	TPH	Water	1 L Amber Glass	≤6°C; H ₂ SO ₄ or HCl; pH < 2	28 Days	NA
8440	TPH	Solid/Other	125 mL Glass Jar	≤6 °C	28 Days	NA
9066 420.4	Total Recoverable Phenols	Water Solid	1 L Glass 125 mL Glass Jar	≤6 °C; H ₂ SO ₄ ; pH < 4 ≤6 °C	28 Days 28 Days	NA NA
9040C 4500H⁺ B	рH	Water	125 mL Plastic	≤6 °C	ASAP	NA
120.1, 9050A	Specific Conductance	Water	125 mL Plastic	≤6 °C	ASAP	NA
All radiochemic except Rn-222	•	Water Solid/Other	1 L Plastic (2 x 2 L Preferred) 250 mL Glass Jar	HNO₃; pH < 2	180 Days 180 Days	NA NA
7500-Rn B	Radon 222	Water	125 mL Glass	None	96 Hours	NA
906.0	Tritium	Water Solid/Other	1 L Glass Sample size will vary with mo	isture content	180 Days 180 Days	NA NA

Note: 40 mL vials require caps with Teflon lined septum. All other containers require Teflon lined screw cap lids to minimize container contamination and loss of analyte. *If samples are shipped to the laboratory in EnCoreTM samplers, samples must be extruded and placed in sample containers within 48 hours of sample collection.

IMPORTANT NOTICE:

Sample Preservation Techniques and Holding Times

Method	<u>Parameters</u>	<u>Matrix</u>	Volume/Container	Preservation	Holding Ti Sample	mes <u>Extract</u>
8015C/D	Petroleum Hydrocarbons (Diesel Range Organics)	Water³ Soil/Other	2 x 1 L Amber Glass Bottle 250 mL Glass Jar	≤6 °C ≤6 °C	7 Days 14 Days	40 Days 40 Days
	Petroleum Hydrocarbons (Gasoline Range Organics)	Water⁴ Soil/Other	3 x 40 mL Glass Vial 125 mL Glass Jar	≤6 °C; HCl; pH < 2 ≤6 °C	14 Days 14 Days	NA NA
5035A/ 8015C/D	Petroleum Hydrocarbons (Gasoline Range Organics)	Soil	4 x 40 mL Glass Vial	≤6 °C, 2 Vials NaHSO₄ 1 Vial CH₃OH, 1 Vial No Preservative	*14 days	NA
8021B	Halogenated Volatile Organics	Water ⁴ Soil/Other	3 x 40 mL Glass Vial 125 mL Glass Jar	≤6 °C; HCl; pH < 2 ≤6°C	14 Days 14 Days	NA NA
5035A/8021B	Halogenated Volatile Organics	Soil	4 x 40 mL Glass Vial	≤6°C, 2 Vials NaHSO₄ 1 Vial CH₃OH, 1 Vial No Preservative	*14 days	NA
8081A	Organochlorine Pesticides	Water ³ Soil/Other	4 L Amber Glass Bottle 250 Glass Jar	≤6 °C ≤6 °C	7 Days 14 Days	40 Days 40 Days
8082A	PCBs	Water ³ Soil/Other	4 L Amber Glass Bottle 250 Glass Jar	≤6 °C ≤6 °C	1 Year 1 Year	1 Year 1 Year
8141B	Organophosphorous Compounds	Water ³ Soil/Other	4 L Amber Glass Bottle 250 Glass Jar	≤6 °C; NaOH or H₂SO₄; pH 5-8 ≤6 °C	7 Days 14 Days	40 Days 40 Days
8151A	Chlorinated Herbicides	Water ³ Soil/Other	4 L Amber Glass Bottle 250 Glass Jar	≤6 °C ≤6 °C	7 Days 14 Days	40 Days 40 Days

Note: 40 mL vials require caps with Teflon lined septum. All other containers require Teflon lined screw cap lids to minimize container contamination and loss of analyte. *If samples are shipped to the laboratory in EnCoreTM samplers, samples must be extruded and placed in sample containers within 48 hours of sample collection.

IMPORTANT NOTICE:

Sample Preservation Techniques and Holding Times

Method	<u>Parameters</u>	<u>Matrix</u>	Volume/Container	Preservation	Holding T <u>Sample</u>	imes <u>Extract</u>
8260B, 624	Volatile Organics by GC-MS	Water ^{1,2,4}	3 x 40 mL Glass Vial	≤6 °C; HCl; pH < 2 ≤6 °C; not acidified	14 Days 7 Days	NA NA
		Soil/Other	125 mL Glass Jar	≤6 °C	14 Days	NA
5035A/ 8260B	Volatile Organics by GC-MS	Soil	4 x 40 mL Glass Vial	≤6 °C, 2 Vials NaHSO₄ 1 Vial CH₃OH, 1 Vial No Preservative	*14 days	NA
8270C, 625	Semivolatile Organics by GC-MS	Water ³ Soil/Other	4 L Amber Glass Bottle 250 mL Glass Jar	≤6 °C ≤6°C	7 Days 14 Days	40 Days 40 Days
8280B	Polychlorinated Dioxins and Furans by HRGC/LRMS	Water ³ Soil/Other	4 L Amber Glass Bottle 250 mL Glass Jar	≤6 °C ≤6 °C	30 Days 30 Days	45 Days 45 Days
8290A	Dioxins and Furans By HRGC/HRMS	Water ³ Soil/Other	4 L Amber Glass Bottle 250 mL Glass Jar	≤6 °C ≤6 °C	30 Days 30 Days	45 Days 45 Days
1613B	Dioxins and Furans by Isotope Dilution HRGC/HRMS	Water ³ Solid	Amber Glass Amber Glass Jar	≤6 °C ≤6 °C	1 Year 1 Year	1 Year 1 Year
1668A/C	PCB Congeners by HRGC/HRMS	Water ³ Solid	Amber Glass Amber Glass Jar	≤6 °C, ≤6 °C	1 Year 1 Year	1 Year 1 Year
1694	PPCP	Water ³ Solid	Amber Glass Amber Glass Jar	≤6 °C ≤6 °C	7 Days 7 Days	40 Days 40 Days
8318A	N-Methylcarbamate Pesticides by HPLC	Water Soil/Other	4 L Amber Glass Bottle 250 mL Glass Jar	≤6 °C; 0.1 <u>N</u> CICH ₂ CO ₂ H, pH 4 - 5 ≤6 °C	7 Days 7 Days	40 Days 40 Days

Note: 40 mL vials require caps with Teflon lined septum. All other containers require Teflon lined screw cap lids to minimize container contamination and loss of analyte. *If samples are shipped to the laboratory in EnCoreTM samplers, samples must be extruded and placed in sample containers within 48 hours of sample collection.

IMPORTANT NOTICE:

Effective Date: 06/19/2017

Procedure No: AOP 00-03 Issue No: 05

Sample Preservation Techniques and Holding Times

Method	<u>Parameters</u>	Matrix	Volume/Container	Preservation	Holding Tir Sample	nes <u>Extract</u>
8330B	Nitroaromatics and Nitramines by HPLC	Water Soil/Other	4 L Amber Glass Bottle 250 mL Glass Jar	≤6 °C ≤6 °C	7 Days 14 Days	40 Days 40 Days
610, 8310	PAHs by HPLC	Water ³ Soil/Other	Amber Glass/Teflon lined cap 250 mL Glass Jar	≤6 °C <6 °C	7 Days 14 Days	40 Days 40 Days
TO-15	VOC in Air	SUMMA® C	anister		30 Days	
8330A	High Explosives by HPLC	Water Solid	Amber Glass/Teflon lined cap Amber Glass/Teflon lined cap		7 Days 14 Days	40 Days 40 Days
8330B (modified)	High Explosives by LC/MS/MS	Water Solid	Amber Glass/Teflon lined cap Amber Glass/Teflon lined cap		7 Days 14 Days	40 Days 40 Days
	Gene-Trac-DHC (Dehalococcoides)	Water	I L Plastic, ≤6 °C		10 Days	NA
	Gene-Trac-VC (Vinyl Chloride reductase)	Water	I L Plastic, ≤6 °C		10 Days	NA

¹ If vinyl chloride, styrene, or 2-chloroethylvinylether are analytes of interest, collect a second set of samples without acid preservative and analyze within 7 days.

Note: 40 mL vials require caps with Teflon lined septum. All other containers require Teflon lined screw cap lids to minimize container contamination and loss of analyte. *If samples are shipped to the laboratory in EnCoreTM samplers, samples must be extruded and placed in sample containers within 48 hours of sample collection.

IMPORTANT NOTICE:

² If acrolein and acrylonitrile are analytes of interest, adjust to pH 4-5.

³ If residual chlorine is present, preserve with 80 mg of sodium thiosulfate per liter of water.

⁴ If residual chlorine is present, preserve with 10 mg of sodium thiosulfate per 125 mL of water.

Appendix B

Data Reporting Requirements

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

If any quality control (QC) samples are analyzed using a different initial calibration (ICAL) than that of the field samples, the laboratory must include a calibration report for the calibration affecting the QC samples. This calibration data shall only be used to evaluate the QC samples, and only if the QC samples fail to meet recovery or relative percent difference (RPD) acceptance criteria. The laboratory is not required to report calibration data associated with QC samples from another sample delivery group.

If required data is not present, contact the laboratory to request an amended report. Documentation may include the following, as appropriate.

Gas Chromatography/Mass Spectrometry (GC/MS)

- Case narrative
- Instrument tuning data
- ICAL data
- Applicable calibration verification data
- Continuing calibration check data
- Instrument and preparation blank data
- Surrogate data
- Internal standard (IS) performance data
- Matrix spike/matrix spike duplicate (MS/MSD) data
- Laboratory control sample (LCS) data
- Sample results and analytical data for the requested target analytes, including results from dilutions, if analyzed
- Identification and data for any sample tentatively identified compounds
- Instrument run logs
- ARCOC and shipping documents
- Login worksheet
- Laboratory replicate data, if analyzed

Dioxins and Furans by High Resolution Gas Chromatography/High Resolution Mass Spectrometry (HRGC/HRMS)

- Case narrative
- Column performance check data
- ICAL data
- Applicable calibration verification data
- Continuing calibration check data
- Preparation blank data
- Labeled compound data
- Ongoing precision and recovery (OPR) data
- Ion abundance ratio data
- Sample results and analytical data for the requested target analytes, including results from dilutions, if analyzed
- Instrument run logs
- ARCOC and shipping documents
- Login worksheet
- Laboratory replicate data, if analyzed

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Gas Chromatography (GC) and High-Performance Liquid Chromatography (HPLC)

- Case narrative
- ICAL calibration data, including secondary column, if appropriate
- Applicable calibration verification data
- Continuing calibration check data
- Retention time windows
- Instrument and preparation blank data
- Surrogate data
- MS/MSD data
- LCS data
- Sample results and analytical data for the target analytes, including data from dilutions, if analyzed
- Confirmation data and RPD between the results
- Instrument run logs
- ARCOC and shipping documents
- Login worksheet
- Laboratory replicate data, if analyzed

Polychlorinated Biphenyl (PCB) Congeners, EPA Method 1668A

- Case Narrative
- ICAL data, including relative retention time (RRT) windows
- Calibration verification data including ion abundance ratios and RRTs
- Preparation blank data
- OPR data
- Clean-up standard data
- Labeled compound data
- Sample results and analytical data for the requested target analytes, including results from dilutions, if analyzed
- Ion abundance ratio for all detected sample results, labeled compounds, and clean-up standards
- RRTs for all detected sample results, labeled compounds, and clean-up standards
- Instrument run logs
- Sample preparation data
- Lipid data (tissue samples only)
- ARCOC and shipping documents
- Login worksheet

High Explosives (HE) and Perchlorate by Liquid Chromatography/Mass Spectrometry/Mass Spectrometry (LC/MS/MS)

- Case narrative
- ICAL data
- Applicable calibration verification data
- Initial calibration blank (ICB) data

- Continuing calibration check data
- · Continuing calibration blank (CCB) data
- Low level calibration verification (CRI) ¹ data
- Low level calibration standard (IRA for HE only)
- Instrument and preparation blank data
- Surrogate data (HE only)
- IS or Method of Standard Addition performance data
- MS/MSD data
- LCS data
- Retention time data
- Isotope Ratio data (perchlorate only)
- Sample results and analytical data for the requested target analytes, including results from dilutions, if analyzed
- Instrument run logs
- ARCOC and shipping documents
- · Login worksheet
- Laboratory replicate data, if analyzed

Inorganic

- Case narrative
- ICAL data
- Applicable calibration verification data
- ICB data
- Continuing calibration data
- CCB data
- Instrument tuning data
- Instrument and preparation blank data
- MS data
- LCS data
- Laboratory replicate data
- Inductively coupled plasma-atomic emission spectroscopy (ICP-AES) and inductively coupled plasma-mass spectrometry (ICP-MS) interference check sample data
- ICP SD data
- PQL verification (CRI/LLCCV)²
- Sample results and analytical data, including data from dilutions, if analyzed
- Instrument run logs
- ARCOC and shipping documents
- Login worksheet

¹ CRI = reporting limit verification (RLV) for LC/MS/MS method.

² CRI = RLV for LC/MS/MS method.

LLCCV = for AA, ICP-AES and ICP-MS methods

Radiochemistry

- Case narrative
- Instrument and preparation blank data
- Applicable calibration verification data
- MS data
- LCS data
- Laboratory replicate data
- · Sample results
- Carrier or chemical tracer data
- Instrument run logs
- ARCOC and shipping documents
- Login worksheet
- Control Charts

Appendix C

Surrogate Recovery Limits

home page.

Procedure No: AOP 00-03

Issue No: 05

Guidelines for Surrogate Recovery Limits

Volatile Organics – Water	Mean	Lower	Upper
_		Limit	Limit
1,2-Dichloroethane-d4	95	72	119
4-bromofluorobenzene	98	76	119
Dibromofluoromethane	100	85	115
Toluene-d8	102	83	120
Volatile Organics - Solid			
4-bromofluorobenzene	101	84	118
Toluene-d8	100	84	116
Semivolatile Organics – Water			
2-fluorobiphenyl	79	48	112
Terphenyl-d14	92	51	135
2,4,6-Tribromophenol	82	42	124
2-Fluorophenol	63	19	108
Nitrobenzene-d5	76	41	111
Semivolatile Organics - Solid			
2-fluorobiphenyl	72	43	103
Terphenyl-d14	78	32	125
2,4,6-Tribromophenol	80	36	126
2-Fluorophenol	70	37	104
Phenol-d5/d6	71	40	102
Nitrobenzene-d5	69	37	102
Pesticides – Water			
Decachlorobiphenyl	83	32	135
Tetrachlorometaxylene (TCMX)	81	25	138
Pesticides - Solid			
Decachlorobiphenyl	94	56	132
TCMX	97	69	124
Polychlorinated Biphenyl			
(PCB) – Water			
Decachlorobiphenyl	88	42	133
PCB - Solid			
Decachlorobiphenyl	91	58	125
High Explosives (HE) - Water			
3,4-Dinitrotoluene	86	33	139
2-Methyl-4-nitroaniline	86	33	139
1,4-Dintrobenzene	86	33	139
1,2-Dintrobenzene	86	33	139
HE - Solid			
3,4-Dinitrotoluene	98	56	140
2-Methyl-4-nitroaniline	98	56	140
1,4-Dintrobenzene	98	56	140
1,2-Dintrobenzene	98	56	140

IMPORTANT NOTICE:

Appendix D

Gas Chromatography/Mass Spectrometry (GC/MS)
Internal Standards

Procedure No: AOP 00-03 Issue No: 05 Effective Date: 06/19/2017

Laboratories may vary the compounds calculated off of any internal standard (IS) and should identify within the report which compounds were calculated from each IS. If this information is not readily available the following tables may be used as guidelines.

GC/MS Volatile Organic Analysis Internal Standard Tables

Fluorobenzene

1 IUOI ODCIIZCIIC			
Chloromethane	Vinyl Chloride	Bromomethane	Chloroethane
Acetone	1,1-Dichloroethene	Methylene Chloride	Carbon Disulfide
1,1-Dichloroethane	Trans-1,2-Dichloroethene	2-Butanone	2,2-Dichloropropane
Cis-1,2-Dichloroethene	Chloroform	Bromochloromethane	1,1,1-Trichloroethane
1,1-Dichloropropene	Carbon Tetrachloride	1,2-Dichloroethane	Benzene
Trichloroethene	1,2-Dichloropropane	Bromodichloromethane	Dibromomethane
4-Methyl-2-pentanone	Cis-1,3-Dichloropropene	Trichlorofluoromethane	Acetonitrile
Acrolein	Acrylonitrile	n-Butyl alcohol	2-Chloro-1,3-butadiene
Dichlorodifluoromethane	1,4-Dioxane	Ethyl acetate	Iodomethane
Isobutyl alcohol	Methacrylonitrile	Methyl methacryalate	Methyl isobutyl ketone
Propionitrile	Trichlorofluoromethane		

Chlorobenzene-d5

Citor obcuzenc-u3					
Toluene	Trans-1,3-	1,1,2-Trichloroethane	2-Hexanone		
	Dichloropropene				
1,3-Dichloropropane	Tetrachloroethene	Chlorodibromomethane	1,2-Dibromoethane		
Chlorobenzene	1,1,1,2-tetrachloroethane	Ethylbenzene	m,p-Xylenes		
o-Xylene	Styrene	Bromoform	1,1,2,2-Tetrachloroethane		
Ethyl methacrylate					

1,4-Dichlorobenzene-d4

1,2,3-Trichloropropane	Bromobenzene	1,2-Dibromo-3-	1,2-Dichlorobenzene
		chloropropane	
1,3-Dichlorobenzene	1,4-Dichlorobenzene	Hexachlorobutadiene	Napthalene
Pentachloroethane	1,2,4-Trichlorobenzene		

Effective Date: 06/19/2017 Issue No: 05

Procedure No: AOP 00-03

GC/MS Semivolatile Organic Analysis Internal Standard Tables

1,4-Dichlorobenzene-d4

2-Fluorophenol	Phenol-d5	2-Chlorophenol-d4	1,2-Dichlorobenzene-d4
Phenol	Bis (2-Chloroethyl) ether	2-Chlorophenol	1,3-Dichlorobenzene
1,4-Dichlorobenzene	1,2-Dichlorobenzene	2-Methylphenol	2,2'-oxybis(2- Chloropropane
4-Methylphenol	N-Nitroso-di-n- propylamine	Hexachloroethane	Pyridine
Acetophenone	Aniline	Methyl metanesulfonate	N-Nitrosodiethylamine
N-Nitrosodimethylamine	N- Nitrosomethylethylamine	N-Nitrosomorpholine	N-Nitrosopiperidine
N-Nitrosopyrrolidine			

Naphthalene-d8

Nitrobenzene-d5	Nitrobenzene	Isophorone	2-Nitrophenol
2,4-Dimethylphenol	Bis (2-Chloroethoxy)	2,4-Dichlorophenol	1,2,4-Trichlorobenzene
	methane		
Naphthalene	4-Chloroaniline	Hexachlorobutadiene	4-Chloro-3-methylphenol
2-Methylnaphthalene	Benzoic Acid	2.6-Dichlorophenol	Hexachloropropene
N-Nitrosodi-n-butylamine	Safrole		

Acenaphthene-d10

2,4,6-Tribromophenol	2-Fluorobiphenyl	Hexachlorocyclopentadiene	2,4,6-Trichlorophenol
2,4,5-Trichlorophenol	2-Chloronaphthalene	2-Nitroaniline	Dimethylphthalate
Acenaphthylene	2,6-Dinitrotoluene	3-Nitroaniline	Acenaphthene
2,4-Dinitrophenol	4-Nitrophenol	Dibenzofuran	2,4-Dinitrotoluene
Diethylphthalate	4-Chlorophenyl phenyl ether	Fluorene	4-Nitroaniline
2-sec-Butyl-2,6- dinitrophenol	Isosafrole	N-Nitro-o-toluidine	Pentachlorophenol
1,2,4,5- Tetrachlorobenzene	2,3,4,6-Tetrachlorophenol		

Phenanthrene-d10

2,4-Dinitro-2-	N-Nitrosodiphenylamine	4-Bromophenyl phenyl	Hexachlorobenzene
methylphenol		ether	
Pentachlorophenol	Phenanthrene	Anthracene	Carbazole
Di-n-butylphthalate	Fluoranthene	Methapyrilene	Pentachloronitrobenzene
Phenacetin	Pronamide		

Chrysene-d12

Pyrene	Butylbenzylphthalate	3,3'-Dichlorobenzidine	Benzo(a)anthracene
Chrysene	Bis(2-Ethylhexl)phthalate	2-Acethylaminofluorene	Chlorobenzilate

Perlene-d12

Di-n-octylphthalate	Benzo(b)fluoranthene	Benzo(k)fluoranthene	Benzo(a)pyrene
Indeno(1,2,3-cd)pyrene	Dibenz(a,h)anthracene	Benzo(g,h,i)perylene	Hexachlorophene
3-Methylcholanthrene			

IMPORTANT NOTICE:

Appendix E

Laboratory Control Limits

Organic Laboratory Control Sample (LCS) Criteria Guidelines (Volatile Compounds)

Volatile Compound	CAS #*		Water			Solid	
		Mean	Low	High	Mean	Low	High
Acetone	67-64-1	99.5	39	160	99.6	36	164
Benzene	71-43-2	99.4	79	120	99.2	77	121
Bromobenzene	108-86-1	99.7	80	120	99.3	78	121
Bromochloromethane	74-97-5	100.8	78	123	101.4	78	125
Bromodichloromethane	75-27-4	101.8	79	125	101	75	127
Bromoform	75-25-2	97.8	66	130	99.1	67	132
Bromomethane	74-83-9	97	53	141	98.3	53	143
2-Butanone	78-93-3	99.6	56	143	99.6	51	148
n-Butylbenzene	104-51-8	101.1	75	128	98.7	70	128
sec-Butylbenzene	135-98-8	101.1	77	126	99	73	126
tert-Butylbenzene	98-06-6	101	78	124	98.8	73	125
Carbon disulfide	75-15-0	98.8	64	133	97.9	63	132
Carbon tetrachloride	56-23-5	103.8	72	136	102.3	70	135
Chlorobenzene	108-90-7	100	82	118	99.7	79	120
Chloroethane	75-00-3	99	60	138	98.8	59	139
Chloroform	67-66-3	101.1	79	124	100.3	78	123
Chloromethane	74-87-3	94.5	50	139	93.3	50	136
2-Chlorotoluene	95-49-8	100	79	122	98.5	75	122
4-Chlorotoluene	106-43-4	99.9	78	122	98.3	72	124
Cyclohexane	110-82-7	100.4	71	130	98.9	67	131
Dibromochloromethane	124-48-1	100	74	126	100.2	74	126
1,2-Dibromo-3-chloropropane	96-12-8	94.9	62	128	96.6	61	132
1,2-Dibromoethane	106-93-4	99	77	121	100.1	78	122
Dibromomethane	74-95-3	101.1	79	123	101.1	78	125
1,2-Dichlorobenzene	95-50-1	99.4	80	119	99.1	78	121
1,3-Dichlorobenzene	541-73-1	99.7	80	119	98.9	77	121
1,4-Dichlorobenzene	106-46-7	98.3	79	118	97.5	75	120
Dichlorodifluoromethane	75-71-8	92	32	152	88.9	29	149
1,1-Dichloroethane	75-34-3	101.3	77	125	100.4	76	125
1,2-Dichloroethane	107-06-2	100.3	73	128	100.5	73	128
1,1-Dichloroethene	75-35-4	101	71	131	100.3	70	131
cis-1,2-Dichloroethene	156-59-2	100.1	78	123	99.9	77	123
trans -1,2-Dichloroethene	156-60-5	99.5	75	124	99.2	74	125
1,2-Dichloropropane	78-87-5	100.1	78	122	99.5	76	123
1,3-Dichloropropane	142-28-9	99.1	80	119	99.1	77	121
2,2-Dichloropropane	594-20-7	99.7	60	139	99.7	67	133
1,1-Dichloropropene	563-58-6	102	79	125	100.5	76	125
cis-1,3-Dichloropropene	10061-01-5	99.5	75	124	99.8	74	126
trans-1,3-Dichloropropene	10061-02-6	100	73	127	100.9	71	130
Ethylbenzene	100-41-4	99.8	79	121	99.1	76	122

Volatile Compound	CAS#*		Water			Solid	
		Mean	Low	High	Mean	Low	High
Hexachlorobutadiene	87-68-3	100.1	66	134	98.1	61	135

Organic LCS Criteria Guidelines (Volatile Compounds) (concluded)

Volatile Compound	CAS#*		Water			Solid	
•		Mean	Low	High	Mean	Low	High
2-Hexanone	591-78-6	97.9	57	139	99.1	53	145
Isopropylbenzene	98-82-8	101.5	72	131	100.8	68	134
4-Isopropyltoluene	99-87-6	102	77	127	100.3	73	127
Methyl acetate	79-20-9	96	56	136	98.7	53	144
Methylcyclohexane	108-87-2	101.8	72	132	99.4	66	133
Methylene chloride	75-09-2	99.4	74	124	98.9	70	128
4-Methyl-2-pentanone	108-10-1	98.5	67	130	99.6	65	135
Naphthalene	91-20-3	94.6	61	128	95.6	62	129
n-Propylbenzene	103-65-1	101	76	126	98.9	73	125
Styrene	100-42-5	100.5	78	123	100.2	76	124
tert-Butyl methyl ether	1634-04-4	97.3	71	124	98.9	73	125
1,1,1,2-Tetrachloroethane	630-20-6	101.1	78	124	101.1	78	125
1,1,2,2-Tetrachloroethane	79-34-5	96.4	71	121	97	70	124
Tetrachloroethene	127-18-4	101.3	74	129	100.5	73	128
Toluene	108-88-3	100.1	80	121	99.3	77	121
1,2,3-Trichlorobenzene	87-61-6	98.7	69	129	97.8	66	130
1,2,4-Trichlorobenzene	120-82-1	99.8	69	130	98	67	129
1,1,1-Trichloroethane	71-55-6	102.7	74	131	101.6	73	130
1,1,2-Trichloroethane	79-00-5	99.5	80	119	99.7	78	121
Trichloroethene	79-01-6	101.1	79	123	100.2	77	123
Trichlorofluoromethane	75-69-4	103	65	141	101	62	140
Trichlorotrifluoroethane	76-13-1	103	70	136	100.8	66	136
1,2,3-Trichloropropane	96-18-4	97.5	73	122	99.1	73	125
1,2,4-Trimethylbenzene	95-63-6	99.6	76	124	98.7	75	123
1,3,5-Trimethylbenzene	108-67-8	99.5	75	124	98.4	73	124
Vinyl acetate	108-05-4	100.2	54	146	100.3	50	151
Vinyl chloride	75-01-4	97.4	58	137	95.6	56	135
m,p-Xylenes	179601-23-	100.5	80	21	100.4	77	124
o-Xylene	95-47-6	100	78	122	100	77	123
Xylenes (total)	1330-20-7	100.1	79	121	100.7	78	124

^{*}chemical abstract service number

Organic LCS Criteria Guidelines (Semivolatile Compounds)

Semivolatile Compound	CAS#*		Water			Solid	
		Mean	Low	High	Mean	Low	High
Polynuclear Aromatics		1					
2-Methynaphthalene	91-57-6	80.7	40	121	80.1	38	122
Acenaphthene	83-32-9	84.5	47	122	81.3	40	123
Acenaphthylene	208-96-8	85.3	41	130	81.8	32	132
Anthracene	120-12-7	89.6	57	123	85.2	47	123
Benz(a)anthracene	56-55-3	91.6	58	125	87.4	49	126
Benzo(b)fluoranthene	205-99-2	92	53	131	88.3	45	132
Benzo(k)fluoranthene	207-08-9	93.2	57	129	89.6	47	132
Benzo(g,h,i)perylene	191- 24-2	92	50	134	88.5	43	134
Benzo(a)pyrene	50-32-8	90.8	54	128	86.9	45	129
Chrysene	218-01-9	91.3	59	123	87.1	50	124
Dibenz(a,h)anthracene	53-70-3	92.7	51	134	89.5	45	134
Fluoranthene	206-44-0	92.6	57	128	88.3	50	127
Fluorene	86-73-7	88.8	52	124	84.2	43	125
Indeno(1,2,3-cd)pyrene	193-39-5	92.6	52	134	89.3	45	133
Naphthalene	91-20-3	80	40	121	78.8	35	123
Phenanthrene	85-01-8	89.6	59	120	85.4	50	121
Pyrene	129-00-0	91.1	57	126	87.2	47	127
Phenolic/Acidic							
2,4-Dichlorophenol	120-83-2	84	47	121	80.9	40	122
2, 4-Dimethylphenol	105-67-9	77.5	31	124	78.4	30	127
2,4-Dinitrophenol	51-28-5	82.9	23	143	**		
2-Chlorophenol	95-57-8	77.5	38	117	77.3	34	121
2-Methylphenol (o-Cresol)	95-48-7	73	30	114	77	32	122
3/4-Methylphenol (m/p-Cresol)	65794-96-9	69.7	29	110	76.5	34	119
4-Methylphenol	106-44-5	72.5	25	120	84.1	42	126
2-Nitrophenol	88-75-5	84.6	47	123	79.6	36	123
4,6-Dinitro-2-methylphenol	534-52-1	90.1	44	137	80.7	29	132
4-Chloro-3-methylphenol	59-50-7	85.5	52	119	83.3	45	122
Pentachlorophenol	87-86-5	86.4	35	138	78.7	25	133
4-Nitrophenol	100-02-7	**			80.6	30	132
Phenol	108-95-2	**			77.3	34	121
2,4,5-Trichlorophenol	95-95-4	88.1	53	123	82.6	41	124
2,4,6-Trichlorophenol	88-06-2	87.2	50	125	82.1	39	126
Basic							
3,3'-Dichlorobenzidine	91-94-1	77.9	27	129	71.3	22	121
4-Chloroaniline	106-47-8	75.3	33	117	61.3	17	106
Phthalate Esters							
Bis(2-ethylhexyl) phthalate	117-81-7	95.2	55	135	91.9	51	133
Butyl benzyl phthalate	85-68-7	93.3	53	134	90.3	48	132
Di-n-butyl phthalate	84-74-2	93	59	127	89.4	51	128
Di-n-octyl phthalate	117-84-0	95.5	51	140	92.4	45	140

Organic LCS Criteria Guidelines (Semivolatile Compounds) (concluded)

Semivolatile Compound	CAS #*	,	Water			Solid	
		Mean	Low	High	Mean	Low	High
Diethyl phthalate	84-66-2	90.1	56	125	87.2	50	124
Dimethyl phthalate	131-11-3	86	45	127	85.9	48	124
Nitrosoamines							
N-Nitrosodimethylamine	62-75-9				71.6	23	120
N-Nitrosodiphenylamine	86-30-6	86.8	51	123	82.7	38	127
N-Nitroso-di-n-propylamine	621-64-7	84	49	119	78.2	36	120
Chlorinated Aliphatics							
Bis(2-chloroethoxy) methane	111-91-1	83.9	48	120	78.4	36	121
Bis(2-chloroethyl) ether	111-44-4	80.8	43	118	75.4	31	120
Bis(2-chloroisopropyl) ether	39638-32-	83.4	37	130	82	33	131
	9						
Hexachlorobutadiene	87-68-3	73.1	22	124	77.3	32	123
Hexachloroethane	67-72-1	68	21	115	72.2	28	117
Halogenated Aromatics							
1,2,4-Trichlorobenzene	120-82-1	72.6	29	116	75.7	34	118
1,2-Dichlorobenzene	95-50-1	71.4	32	111	74.6	33	117
1,3-Dichlorobenzene	541-73-1	68.6	28	110	72.6	30	115
1,4-Dichlorobenzene	106-46-7	70.4	29	112	73.1	31	115
2-Chloronaphthalene	91-58-7	78	40	116	77.5	41	114
4-Bromophenyl phenyl ether	101-55-3	89.1	55	124	85.1	46	124
4-Chlorophenyl phenyl ether	7005-72-3	86.7	53	121	83	45	121
Hexachlorobenzene	118-74-1	88.7	53	125	83.5	45	122
Nitroaromatics							
2,4-Dinitrotoluene	121-14-2	92.3	57	128	86.8	48	126
2,6-Dinitrotoluene	606-20-2	90.7	57	124	85	46	124
2-Nitroaniline	88-74-4	90.8	55	127	85.4	44	127
3-Nitroaniline	99-09-2	84.4	41	128	75.9	33	119
4-Nitroaniline	100-01-6	**			**		
Nitrobenzene	98-95-3	83	45	121	77.8	34	122
Neutral Aromatics							
Carbazole	86-74-8	91.1	60	122	86.3	50	123
Dibenzofuran	132-64-9	85.3	53	118	81.5	44	120
Others							
Benzyl alcohol	100-51-6	71.2	31	112	75.7	29	122
Isophorone	78-59-1	83.3	42	124	75.9	30	122

^{*}chemical abstract service number

^{**}not listed in QSM

Organic LCS Criteria Guidelines (Pesticides and Polychlorinated Biphenyl [PCB])

Pesticide	CAS#*		Water			Solid	
		Mean	Low	High	Mean	Low	High
Aldrin	309-00-2	89.5	45	134	90.5	45	136
α-benzene hexachloride (BHC)	319-84-6	95.8	54	138	90.9	45	137
β-ВНС	319-85-7	96.3	56	136	93.1	50	136
δ-ВНС	319-86-8	97.2	52	142	93.3	47	139
γ-BHC (Lindane)	58-89-9	96.4	59	134	92.1	49	135
α-Chlordane	5103-71-9	94.3	60	129	93.7	54	133
4,4'- Dichlorodiphenyldichloroethane (DDD)	72-54-8	99.6	56	143	97.7	56	139
4,4'- Dichlorodiphenyldichloroethylene (DDE)	72-55-9	96	57	135	95.3	56	134
4,4'- Dichlorodiphenyltrichloroethane (DDT)	50-29-3	97	51	143	95.8	50	141
Dieldrin	60-57-1	98	60	136	95.7	56	136
Endosulfan I	959-98-8	93.8	62	126	92.2	53	132
Endosulfan II	33213-65-9	93.4	52	135	93.1	53	134
Endosulfan sulfate	1031-07-8	97.2	62	133	95.9	55	136
Endrin	72-20-8	98.7	60	138	98.1	57	140
Endrin aldehyde	7421-93-4	91.1	51	132	86	35	137
Endrin ketone	53494-70-5	95.9	58	134	95.5	55	136
Heptachlor	76-44-8	91.9	54	130	92.6	47	136
Heptachlor epoxide	1024-57-3	96.9	61	133	93.9	52	136
4,4'-Methoxychlor	72-43-5	99	54	145	97.6	52	143
PCB							
Aroclor – 1016	12674-11-2	87.1	46	129	90.1	47	134
Aroclor – 1254	11097-69-1	80.1	34	127	101.2	67	135
Aroclor – 1260	11096-82-5	89.4	45	134	96.6	53	140

^{*}chemical abstract service number

Organic LCS Criteria Guidelines (Nitroaromatics and Nitramines) [8330B]

Nitroaromatics and Nitramines	CAS#*		Water			Solid	
		Mean	Low	High	Mean	Low	High
2-Amino-4,6-Dinitrotoluene	35572-78-2	99.4	79	120	96.5	71	123
(2-Am-DNT)							
4-Amino-2,6-Dinitrotoluene	19406-51-0	100.3	76	125	95.4	64	127
(4-Am-DNT)							
1,3-Dinitrobenzene (DNB)	99-65-0	98.7	78	120	96.3	73	119
2,4-Dinitrotoluene (24DNT)	121-14-2	98.9	78	120	98	75	121
2,6-Dinitrotoluene (26DNT)	606-20-2	102	77	127	98	79	117
Hexahydro-1,3,5-trinitro-1,3,5-triazine	121-82-4	99.1	68	130	97.9	67	129
(RDX)						L	
Methyl-2,4,6-trinitrophenylnitramine (Tetryl)	479-45-8	95.8	64	128	101.3	68	135
Nitrobenzene (NB)	98-95-3	99.3	65	134	97.9	67	129
2-Nitrotoluene (2NT)	88-72-2	98.4	70	127	96.8	70	124
3-Nitrotoluene (3NT)	99-08-1	98.8	73	125	97.7	67	129
4-Nitrotoluene (4NT)	99-99-0	99.1	71	127	97.3	71	124
Pentaerythritol tetranitrate (PETN)	78-11-5	100.2	73	127	100.1	72	128
Octahydro-1,3,5,7-tetranitro-1,3,5,7-	2691-41-0	100	65	135	99.1	74	124
tetrazocine (HMX)							
1,3,5-Trinitrobenzene (135TNB)	99-35-4	99	73	125	98	80	116
2,4,6-Trinitrotoluene (TNT)	118-96-7	97	71	123	95.8	71	120

^{*}chemical abstract service number

Organic LCS Criteria Guidelines (TO-15)

Analyte	CAS #*			
		Mean	Low	High
1,1,1-Trichloroethane	71-55-6	96.7	68	125
1,1,2,2-Tetrachloroethane	79-34-5	95.9	65	127
1,1,2-Trichloroethane	79-00-5	95.9	73	119
1,1,2-Trifluoro-1,2,2trichloroethane	76-13-1	96.1	66	126
[Freon-113]				
1,1-Dichloroethane	75-34-3	97	68	126
1,1-Dichloroethene	75-35-4	97.3	61	133
1,2,4-Trichlorobenzene	120-82-1	98.5	55	142
1,2,4-Trimethylbenzene	95-63-6	99.2	66	132
1,2-Dibromoethane	106-93-4	98.2	74	122
1,2-Dichloro-1,1,2,2tetrafluoroethane	76-14-2	92.4	63	121
1,2-Dichlorobenzene	95-50-1	95.7	63	129
1,2-Dichloroethane	107-06-2	96.8	65	128
1,2-Dichloropropane	78-87-5	95.7	69	123
1,3,5-Trimethylbenzene	108-67-8	98.3	67	130
1,3-Dichlorobenzene	541-73-1	97.1	65	130
1,4-Dichlorobenzene	106-46-7	95.8	60	131
2-Butanone [MEK]	78-93-3	98.4	67	130
2-Hexanone	591-78-6	95.4	62	128
4-Ethyltoluene	622-96-8	97.9	67	129
4-Methyl-2-pentanone [MIBK]	108-10-1	98.5	67	130
Acetone	67-64-1	92.7	58	128
Benzene	71-43-2	93.8	69	119
Benzyl chloride	100-44-7	98.7	50	147
Bromodichloromethane	75-27-4	99.9	72	128
Bromoform	75-25-2	102.3	66	139
Bromomethane	74-83-9	98.6	63	134
Carbon disulfide	75-15-0	95.6	57	134
Carbon tetrachloride	56-23-5	99.6	68	132
Chlorobenzene	108-90-7	94.5	70	119
Chlorodibromomethane	124-48-1	99.9	70	130
Chloroethane	75-00-3	94.7	63	127
Chloroform	67-66-3	95.3	68	123
Chloromethane	74-87-3	95.2	59	132
cis-1,2-Dichloroethene	156-59-2	95.6	70	121
cis-1,3-Dichloropropene	10061-01-5	98.8	70	128
Dichlorodifluoromethane [Freon-12]	75-71-8	93.6	59	128
Ethylbenzene	100-41-4	96.8	70	124
Hexachlorobutadiene	87-68-3	96.7	56	138
m/p-Xylene [3/4-Xylene]	179601-23-	97.3	61	134
Methylene chloride	75-09-2	88.8	62	115

o-Xylene	95-47-6	96.3	67	125
Styrene	100-42-5	100.1	73	127
Tetrachloroethene	127-18-4	95.2	66	124
Toluene	108-88-3	92.7	66	119
trans-1,2-Dichloroethene	156-60-5	95.5	67	124
trans-1,3-Dichloropropene	10061-02-6	104	75	133
Trichloroethene	79-01-6	96.7	71	123
Trichlorofluoromethane [Freon-11]	75-69-4	93.7	62	126
Vinyl acetate	108-05-4	97.4	56	139

593-60-2

75-01-4

98.4

95.1

Procedure No: AOP 00-03

126

127

64

Issue No: 05

Vinyl bromide

Vinyl chloride
*chemical abstract service number

Appendix F

Mass Spectra Acceptability

Procedure No: AOP 00-03 Effective Date: 06/19/2017 Issue No: 05

Mass Spectra Acceptability

Ideal spectral identification of a target analyte by a mass spectrometer data system is performed by comparing three characteristic ions (i.e., a primary or quantitation ion, a secondary ion, and a tertiary ion) from one mass spectrum to the same characteristic ions in the reference mass spectrum. The three characteristic ions from the mass spectrum are defined as the three ions of greatest relative intensity or any ions over 30% relative intensity if less than three such ions occur in the reference spectrum. In most cases, ions with the greatest abundance are used for evaluation; however, if target analytes suffer from coelution, interferences may dictate the use of less abundant ions for evaluation. Some analytes generate a mass spectrum that is of such a simple nature that a tertiary ion is not sufficiently abundant (e.g., low molecular weight analytes and analytes that do not fragment sufficiently upon electron impact ionization). In this case, two ions are used for identification.

For evaluation of analyte spectra, all of the following factors should be considered before an acceptability judgment is made.

Retention Times (RT)

The intensities of the primary, secondary, and, if applicable, tertiary ions at the established RT of the target analyte are shown in the extracted ion current profiles (EICP). The RTs for the secondary and tertiary ion profiles should be the same as the primary ion. Depending on peak shape and chromatographic interferences, the RTs could differ by a few hundredths of a minute; however, the RTs between primary and secondary ions should not vary by more than 0.03 minutes.

Relative Retention Times (RRT)

The RRT of the target analyte in the sample should agree to within 0.06 RRT units of the same analyte in the reference standard (either the midpoint standard of the initial calibration or the daily continuing calibration verification).

Ion Ratios

The most intense ion in a spectrum is assigned a relative abundance of 100 and is known as the base peak. The intensities of all other ions in the spectrum are compared to the intensity of the base peak to obtain an intensity ratio (or ion ratio). The ion ratios for the three characteristic ions from the sample spectrum are compared to the ion ratios for the same ions from the reference spectrum. Relative intensities should agree to within ±30%. For example, an ion that has an abundance of 50% when compared to the base peak in the reference spectrum will have a range of 20-80% as its acceptance criteria for that same ion in the sample spectrum.

Ion ratio evaluation is performed by the laboratory. Gas chromatography/mass spectrometry data systems automatically flag target analytes with a "Q" or a "#" on the quantitation report when ratio comparison criteria are exceeded. Because interferences and varying instrument conditions can affect relative abundances, the presence of a Q flag does not necessarily indicate an invalid identification; however, a Q flag in addition to other guideline failures may result in the need for additional data in order to make an acceptability judgment.

IMPORTANT NOTICE:

Effective Date: 06/19/2017

Procedure No: AOP 00-03 Issue No: 05

Visual Comparison of Mass Spectra

The sample spectrum should be visually compared to the reference spectrum for pattern similarity. When a peak elutes in a discrete manner and the reference spectrum was obtained under similar conditions as the sample spectrum, the mass spectral pattern from the sample will be similar to the mass spectral pattern of the reference spectrum. If coelution occurs or a high level of background is present, the partial or total spectral pattern from the reference spectrum should be visible within the spectral pattern of the sample spectrum, with sample ion ratios emulating the reference ion ratios, depending on the inherent complexity of the analyte spectrum.

Identification of Target Analytes Present in Samples at Low Levels

Spectra from target analytes that are detected at levels around the established method detection limit (MDL) should be examined carefully for the presence of secondary and, if applicable, tertiary ions. For a qualitative identification to be made, all ions used by the instrument method should be present. Because background noise, column bleed, and interferences can hamper identification, correct subtraction is important to identification. Supplemental data may be required from the laboratory in order to properly separate chromatographic interferences. These data may include a library search with listed spectral fit (or match) quality (i.e., a "Q report") or additional EICPs displaying interfering ions for RT comparison.

Interferences

Identification of target analytes is hampered when sample components are not resolved chromatographically (i.e., there is co-elution of non-target and/or target analytes) and produce mass spectra containing ions from more than one analyte. When gas chromatography (GC) peaks, EICPs, or spectra show evidence of interference (e.g., GC or EICP peak appears broadened with shoulders, obviously overlapping peaks are present, or extraneous ions are present in the spectrum), supplemental data may be required from the laboratory in order to properly evaluate analyte spectra. These data may include a Q report or additional EICPs displaying interfering ions for RT comparison.

Guidelines for Use of Supplemental Data

O Report

If a target analyte is detected by the mass spectrometer data system and its identification is questionable, the mass spectrum for that analyte may be subjected to a computer comparison against a library of established mass spectra (i.e., a "library search"). This search generates a Q report that shows the mass spectrum being searched, the compounds in the library whose spectra most closely matches the mass spectrum being searched, and a Chemical Abstract Services (CAS) number and a match quality rating from 1-100 (with 100 being a perfect match fit) for each of those compounds. A Q report obtained for a spectrum at the RT of the analyte in question can sometimes help identify that analyte. Ideally, if a data system identifies a target analyte using the identification parameters discussed (i.e., RT, RRT, and major ion intensity ratios), a library search of the analyte should yield concurrent results with a match rating of >75. The following variables affect match quality ratings:

IMPORTANT NOTICE:

Effective Date: 06/19/2017 Issue No: 05

Procedure No: AOP 00-03

The Analyte Concentration

For identification of an analyte of low concentration (i.e., detections at or just above the MDL), ions of >50% relative intensity in library spectrum should be present in sample spectrum. If minor ions in the analyte spectrum are absent due to low concentration, the match quality rating may be low.

The Nature of the Spectrum

If the spectrum of the analyte in question is relatively complex (i.e., the spectrum contains multiple ions of >50% relative intensity), match quality ratings will generally be higher. If the spectrum of the peak in question only yields two or three ions within scanning range, separation from interferences and background is sometimes not possible, making match quality ratings lower. Also, if the spectrum of the analyte in question has one or more ions common to known contaminants, misidentification will be more common (e.g., acetone with its primary ion of 43 is sometimes hard to distinguish from early eluting, low molecular weight hydrocarbons that have the same ion as their primary ions).

The presence of one or more interferences can affect match quality ratings. One dominant interference can yield spectral match quality ratings that are high but whose best match is primarily due to the interfering non-target analyte, not the analyte in question. Multiple interferences will usually yield match quality ratings that are low due to the inability of the software to make a match without a dominant pattern.

The Conditions Under Which the Library Spectrum is Obtained

If spectral quality matches are low, especially for the target analyte in question, the library spectra for the analyte in question should be considered suspect. Mass spectra in established libraries are normally generated under wider scan ranges than are dictated in methods. Low molecular weight analytes in the library may have one or more characteristic ions below the scanning range of the environmental analytical method, rendering the spectra in the low molecular weight range only partially comparable. If it is obvious that the library mass spectrum for an analyte was obtained under different scanning conditions as the sample spectrum, match quality ratings may be reduced.

Additional EICPs

The review of EICPs of all ions of >50% relative abundance, including the primary, secondary, and tertiary ions of the analyte in question as well as ions from interfering analytes, is one way to determine interference separability and abundance contribution. If some or all of the ions from an interfering analyte (i.e., those not contained in the target analyte in question) maximize at the same RT as the target analyte in question, it is possible that interfering ions are contributing abundance to the analyte in question. Also, if peak shape is variable, it is possible that two or more compounds are co-eluting and are contributing a range of ions and overlapping chromatographic peak shapes. If EICPs show that the characteristic ions from the analyte in question elute at the same RT and that RT differs from the EICPs of the ions from interfering compounds, then it is possible that the target analyte in question is present.

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