

U.S. Environmental Protection Agency, Region 2 Field Operations Quality Procedures

ADMINISTRATIVE STANDARD OPERATING PROCEDURE

Standard Operating Procedure for ICP-MS Data Validation				
	Effective Date	I	Number	
	3/21/2022	QA-HWSS-A-009		
	Autl	nor		
Name:	Agustin Aoanan			
Title:	SEEP Grantee			
Division/Bra	anch/Section: LSASD/HWSB/HWS	S		
Signature:			Date:	
	Review & A	Approvals		
Name:	Kim Brandon-Bazile			
Title:	Chemist, HWSS			
Signature:			Date:	
Name:	Donna Ringel			
Title:	Chief, HWSS			
Signature:			Date:	
Name:	Jon Gabry			
Title:	Chief, HWSB			
Signature:			Date:	
Name:				
Title:				
Signature:			Date:	

The table below identifies information about the reviews conducted of this Standard Operating Procedure (SOP).

REVIEW HISTORY			
Date	Reviewer Name	Changes Required (Y/N)	

The table below identifies changes to this controlled document and the respective effective date(s) over time.

	REVISION HISTORY			
Revision Number	Revision Description	Effective Date		
0	Original Issue (Note: Replaces SOP HW-3b, Rev. 1 ISM02.2 – ICP-MS Data Validation, September 2016)			

NOTICE

The policies and procedures set forth here are intended as guidance to the United States Environmental Protection Agency (USEPA) and other governmental employees. They do not constitute rule-making by the USEPA and may not be relied upon to create a substantive or procedural right enforceable by any other person. The Government may take action that is at a variance with the policies and procedures in this Standard Operating Procedure (SOP).

TABLE OF CONTENTS

1.0	PURPOSE AND APPLICABILITY	4
2.0	SUMMARY OF PROCESS OR METHODOLOGY	4
3.0	DEFINITIONS	4
4.0	RESPONSIBILITIES/QUALIFICATIONS	6
5.0	REFERENCES	6
6.0	PROCEDURAL STEPS	7
7.0	DATA AND RECORDS MANAGEMENT	9
8.0	QUALITY ASSURANCE AND QUALITY CONTROL	10
9.0	APPENDICES	10
	Appendix A - Data Validation Criteria and Actions	
	Appendix B - Data Assessment Report Template	
	Appendix C - Definitions/Glossary of Terms	
	Appendix D - SOP Change Request Form	

1.0 PURPOSE AND APPLICABILITY

This document is designed to promote uniformity of data review of analytical data generated through the US EPA Contract Laboratory Program (CLP) Statement of Work (SOW) for Superfund Analytical Methods SFAM01.1 and any future editorial revisions of SFAM01.1. It is applicable to the review of Contract Laboratory Program (CLP) data of various matrices (water, soil, sediment, waste, wipes, etc.) generated using Inductively Coupled Plasma – Mass Spectrometry (ICP-MS) for metal analyses.

The guidelines presented in this document will aid in establishing (a) if data meets the specific technical and quality control (QC) criteria established in the SOW, and (b) the validity and extent of bias of any data not meeting the specific technical and QC criteria established in the SOW. It must be understood by the user that acceptance of data not meeting technical requirements is based upon many factors, including, but not limited to, site-specific technical requirements, the need to facilitate the progress of specific projects, and the availability for resampling. The user should note that while this document is to be used as an aid in the formal data review process, the site-specific quality assurance project plan (QAPP), as well as professional judgement, should also be used to determine the ultimate validity of data, especially in those cases where all data does not meet specific technical criteria.

2.0 SUMMARY OF PROCESS OR METHODOLOGY

This document provides the criteria for performing technical quality assurance reviews of metal data generated by the CLP. Criteria are based on the quality assurance/quality control and technical requirements specified in Exhibit D of SOW SFAM01.1. This SOP incorporates much of the content of the National Functional Guidelines (NFG) and provides additional guidance specific to EPA Region 2.

Upon receipt by EPA Region 2, CLP data in the Sample Delivery Group (SDG) undergoes a technical quality assurance review based upon the criteria in this document. A report of this review is prepared by the data validator, reviewed by the EPA Task Order Contracting Officer Representative (TOCOR), and provided to the data user.

3.0 **DEFINITIONS**

3.1. See Appendix C – Definitions/Glossary of Terms

3.2. Acronyms and Abbreviations

The following acronyms and abbreviations are applicable to this document.

%D	Percent Difference
%R	Percent Recovery
%RI	Percent Relative Intensity

%SolidsPercent Solids, (also %S)ASBAnalytical Services BranchCCBContinuing Calibration BlankCCSContract Compliance ScreeningCCVContract Laboratory ProgramCLPSSContract Laboratory Program Support SystemDARData Assessment ReportDFDilution FactorDLDetection LimitDVData ValidationEDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESSElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration VerificationICSLaboratory Control SampleICSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNGGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality AssuranceQAPPQuality AssuranceQAPPQuality AssuranceQAPPSample Delivery GroupSEDD <td< th=""><th>%RSD</th><th>Percent Relative Standard Deviation</th></td<>	%RSD	Percent Relative Standard Deviation
ASBAnalytical Services BranchCCBContinuing Calibration BlankCCSContract Compliance ScreeningCCVContract Laboratory ProgramCLPSSContract Laboratory Program Support SystemDARData Assessment ReportDFDilution FactorDLDetection LimitDVData ValidationEDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality AssuranceQAPPQuality AssuranceQAPPQuality AssuranceQAPPQuality ControlQLQuantitation LimitRSDSampling and Analysis PlanSDGSample Delivery Group		
CCBContinuing Calibration BlankCCVContract Compliance ScreeningCCVContract Laboratory ProgramCLPSContract Laboratory Program Support SystemDARData Assessment ReportDFDilution FactorDLDetection LimitDVData ValidationEDDElectronic Data DeliverablesEDMExest Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSLaboratory Control SampleICVInitial Calibration BlankICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality AssuranceQAPPQuality ControlQLQuality AssuranceQAPPSamplie Control Center CoordinatorSAPSample Delivery Group		
CCSContract Compliance ScreeningCCVContinuing Calibration VerificationCLPContract Laboratory ProgramCLPSSContract Laboratory Program Support SystemDARData Assessment ReportDFDilution FactorDLDetection LimitDVData ValidationEDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality AssuranceQAPPQuality AssuranceQAPPQuality AssuranceQAPPQuality AssuranceQAPPQuality AssuranceQAPPQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center Coordinat		•
CCVContinuing Calibration VerificationCLPContract Laboratory ProgramCLPSSContract Laboratory Program Support SystemDARData Assessment ReportDFDilution FactorDLDetection LimitDVData ValidationEDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESSElectronic Data Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWVSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality Assurance Project PlanQCRegional Sample Control Center CoordinatorSSPSampling and Analysis PlanSDGSample Delivery Group		
CLPContract Laboratory ProgramCLPSSContract Laboratory Program Support SystemDARData Assessment ReportDFDilution FactorDLDetection LimitDVData ValidationEDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLBBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group		
CLPSSContract Laboratory Program Support SystemDARData Assessment ReportDFDilution FactorDLDetection LimitDVData ValidationEDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAFSampling and Analysis PlanSDGSample Delivery Group		
DARData Assessment ReportDFDilution FactorDLDetection LimitDVData ValidationEDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)EXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLESLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality AssuranceQAPPQuality AssuranceQAPPRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSampling and Analysis Plan		
DFDilution FactorDLDetection LimitDVData ValidationEDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group		
DLDetection LimitDVData ValidationEDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group		
DVData ValidationEDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality AssuranceQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group		
EDDElectronic Data DeliverablesEDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNatrix SpikeNFGNatrix SpikeOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality AssuranceQAPPQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAFSampling and Analysis PlanSDGSample Delivery Group		
EDMEXES Data ManagerEDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality ControlQLQuality ControlQLQuality ControlSAPSample Control Center CoordinatorSAPSample Delivery Group		
EDSEnvironmental Data ServicesEICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality ControlQLQuality ControlQLQuality ControlSAPSample Control Center CoordinatorSAPSample Delivery Group		
EICCElectronic Internal Chain of CustodyEPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality AssuranceQLQuality ControlQLQuality ControlQLRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSDGSampling and Analysis PlanSDGSampling and Analysis Plan		
EPAEnvironmental Protection Agency (see also USEPA)ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group		
ESATEnvironmental Services Assistance TeamEXESElectronic Data Exchange and Evaluation SystemHWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSDGSampling and Analysis PlanSDGSample Delivery Group	EPA	
HWSSHazardous Waste Support SectionICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	ESAT	
ICIon ChromatographyICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality ControlQLQuanitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSDGSample Delivery Group	EXES	Electronic Data Exchange and Evaluation System
ICBInitial Calibration BlankICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	HWSS	Hazardous Waste Support Section
ICP-MSInductively Coupled Plasma – Mass SpectrometryICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSDGSample Delivery Group	IC	Ion Chromatography
ICSInterference Check SampleICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSDGSample Delivery Group	ICB	Initial Calibration Blank
ICVInitial Calibration VerificationLCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSDGSample Delivery Group	ICP-MS	Inductively Coupled Plasma – Mass Spectrometry
LCSLaboratory Control SampleLEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSDGSample Delivery Group	ICS	Interference Check Sample
LEBLeachate Extraction BlankMDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	ICV	Initial Calibration Verification
MDLMethod Detection LimitMSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	LCS	Laboratory Control Sample
MSMatrix SpikeNFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	LEB	Leachate Extraction Blank
NFGNational Functional GuidelinesOSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	MDL	Method Detection Limit
OSRTIOffice of Superfund Remediation and Technology InnovationPDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	MS	Matrix Spike
PDFPortable Document FormatQAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	NFG	National Functional Guidelines
QAQuality AssuranceQAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	OSRTI	Office of Superfund Remediation and Technology Innovation
QAPPQuality Assurance Project PlanQCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	PDF	Portable Document Format
QCQuality ControlQLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group		Quality Assurance
QLQuantitation LimitRPDRelative Percent DifferenceRSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	QAPP	Quality Assurance Project Plan
RPD Relative Percent Difference RSCC Regional Sample Control Center Coordinator SAP Sampling and Analysis Plan SDG Sample Delivery Group		
RSCCRegional Sample Control Center CoordinatorSAPSampling and Analysis PlanSDGSample Delivery Group	QL	
SAP Sampling and Analysis Plan SDG Sample Delivery Group		Relative Percent Difference
SDG Sample Delivery Group	RSCC	
	SAP	
SEDD Staged Electronic Data Deliverable	SDG	
	SEDD	Staged Electronic Data Deliverable

SMO	Sample Management Office
SOP	Standard Operating Procedure
SOW	Statement of Work
SP	SharePoint
SPLP	Synthetic Precipitation Leaching Procedure
TCLP	Toxicity Characteristic Leaching Procedure
TDS	Total Dissolved Solids
тос	Total Organic Carbon
TOCOR	Task Order Contracting Officer Representative
TR/COC	Traffic Report/Chain of Custody
TSS	Total Suspended Solids
USEPA	United States Environmental Protection Agency

3.3. Data Qualifier Definitions

Data qualifier definitions are provided in the beginning of Appendix A.

4.0 **RESPONSIBILITIES/QUALIFICATIONS**

4.1. Qualifications

Data Validators must be familiar with the current CLP SOW, EDM and the documents referenced in Section 5.0 below.

4.2. Responsibilities

- 4.2.1. EPA TOCOR (when applicable) will review data assessments reports and other deliverables prepared by contract data validators. They will update the MS Planner DV Flowboard indicating the progress of SDGs, post final deliverables to the EDS SharePoint site and send notification to clients via the established workflow.
- 4.2.2. Data Validator will follow the criteria and actions provided in this document and prepare Data Assessment Reports (DAR)) and Summary Reports, as necessary. If the validator is an ESAT contractor employee, they will consult the EPA TOCOR when questions arise. They will update the DV Flowboard indicating progress of SDGs.

5.0 REFERENCES

National Functional Guidelines for Inorganic Superfund Methods Data Review, EPA 540-R-20-006, November 2020.

Contract Laboratory Program (CLP) Statement of Work (SOW) Superfund Analytical Method (SFAM) SFAM01.1

FA-0010.1, Standard Operating Procedure for Development and Use of Field SOPs, December 2015.

U.S. EPA, 2007. Guidance for the Preparation of Standard Operating Procedures (SOPs) for Quality-Related Documents. EPA QA/G-6, EPA/600/B-07/001. April 2007.

QA-HWSS-A-001, Document Control Room, Data Dissemination and Archive Operations. Revision 0, January 2021.

6.0 PROCEDURAL STEPS

6.1. EXES Processing

At the Sample Management Office (SMO) the data package and EDD are checked for compliance with the CLP SOW. A Contract Compliance Screening Report (CCS) is issued and posted on the SMO portal web site. The EDD is processed electronically to evaluate QC performance against the NFG and Region 2 criteria by EXES. An electronic report of the EXES review is also posted on the SMO portal web site.

6.2. Initial Notification

The EICC SharePoint web application is setup to send an e-mail alert notification to EPA and ESAT data validators when a new data package is received and available for review and validation. Entry of data into the EICC SharePoint site will automatically trigger an e-proxy card to populate on the DV Flowboard in MS Planner.

Alternate electronic systems may be applied in the future.

6.3. DV Flowboard Updates

Updates to the DV Flowboard will be performed as per SOP QA-HWSS-A-001, Document Control Room, Data Dissemination and Archive Operations (or most current version).

6.4. Data Package Inspection

The EXES Data Manager (EDM) is a useful tool in the data review process. EDM will identify any missing and/or incorrect information in the data package. When available, the EDM should be reviewed as part of the initial data package inspection. The CLP laboratory may submit a reconciliation package for any missing items or to correct the data. If there are any concerns regarding the data package, contact the TOCOR.

An initial review of the data package is to be performed, taking into consideration all information specific to the sample data package, (e.g., modified analysis requests, trip report/chain-of-custody documentation, SDG narratives, etc.). The reviewer should also have a copy of the Quality Assurance Project Plan (QAPP) or similar document for the project for which the samples were analyzed. The criteria for data validation outlined in the QAPP will supersede that in this SOP. The reviewer should access the HWSS SP Documents Dashboard to obtain a copy of the relevant documents.

The SDGs or cases routinely have unique samples that require special attention from the reviewer. These include field blank, equipment blanks, trip blanks, and field duplicates which must be identified in the sample records. The sampling records (i.e., trip reports or COC records) should identify:

- 1) The Region where the samples were taken,
- 2) The case number,
- 3) The complete list of samples with the following information as applicable:
 - a. Sample matrix,
 - b. Field blanks (i.e., equipment, rinsate and trip),
 - c. Field duplicates,
 - d. Field spikes,
 - e. Shipping dates,
 - f. Preservatives, and
 - g. Laboratories involved

6.5. Data Review/Validation

The EXES electronic validation will apply most of the criteria and actions provided in Appendix A. The data validator will examine the EXES report to identify any issues that warrant further investigation. All EXES rejected data will be manually evaluated. The data validator will use the criteria and actions in Appendix A, as well as their own professional judgement to manually assess this data.

To use this SOP effectively, the reviewer should understand the analytical method. The exact number of samples, their assigned numbers, type of matrix, and the number of laboratories involved in the analysis are essential information for the validator.

The TR/COC documentation includes sample descriptions and date(s) of sampling. The reviewer must consider lag times between sampling and start of analysis when assessing technical sample holding times.

The laboratory's SDG narrative is another source of general information. Notable problems with matrices, insufficient sample volume for analysis or reanalysis, samples received in broken containers, preservation and unusual events should be documented in the SDG narrative. The reviewer should also inspect any email, telephone or any communication logs

detailing any discussion of sample or analysis issues between the laboratory, the CLP Sample Management Office and USEPA Region 2.

All data is initially marked as "Reportable" (YES) in EDM before validation is begun. Sometimes, due to dilutions and/or re-analyses being performed, there will be multiple results for a single analyte from a sample. The following criteria and professional judgement are used to determine which result should be reported:

- 1) the analysis with the lower QL,
- 2) the analysis with the better QC results, and/or
- 3) the analysis with the higher result

Data validator will reconcile results from the multiple runs to provide results in one run and report. The analyte values and their respective QLs are then transferred into a single sample run. The runs and results that are not to be used are marked "not reportable" or entered "NO" in the "Reportable" fields of the EDM.

6.6. Data Assessment Report

The data validator will prepare a Data Assessment Report documenting the results of their data review. This report will be formatted in accordance with the template provided in Appendix B. Modifications to the template are allowed at the discretion of the user.

6.7. Summary Report

If requested by the client on the Analytical Request Form, the data validator will prepare a Summary Report using the HWSS Summary Report application.

7.0 DATA AND RECORDS MANAGEMENT

7.1. DATA MANAGEMENT

Posting data to the SP EDS site is done in accordance with QA-HWSS-A-001, "Document Control Room, Data Dissemination and Archive Operations".

7.2. RECORDS MANAGEMENT

The data files uploaded to the EDS SharePoint site include:

- 1) Data Assessment Report (Adobe PDF),
- 2) Edited/Validated Sample Summary Report from SMO portal (Adobe PDF),
- 3) Edited/Validated EQuIS EDD report from SMO portal (MS Excel),
- 4) Generated Summary Report (MS Excel), if applicable, and
- 5) Generated Summary Report with Hits Only (MS Excel), if applicable.

In addition to the above stated documents, data validators also forward the following EXES files, which are not uploaded to EDS SharePoint:

- 6) The CCS Report from the SMO Portal (Adobe PDF), .
- 7) Edit History Report from the SMO Portal (Adobe PDF)

All files stated above are saved to the Local Area Network (LAN) G: drive at DESADIV/HWSS/DATA VALIDATION/Site Name/Case #/SDG #. Files are renamed using the following naming convention, Case#_SDG#_Filetype.*, e.g., 12345_MBAB12_S3VEM.xlsx or 12345_BAB12-M_S3VEM.xlsx.

Note: The letter "M" in the beginning of the SDG name or appended as "-M" signifies that the analyses are inorganic. "M" in the file type signifies that the data has been manually validated by ESAT and/or EPA Staff.

Additional records management procedures are discussed in QA-HWSS-A-001, "Document Control Room, Data Dissemination and Archive Operations".

8.0 QUALITY ASSURANCE AND QUALITY CONTROL

- **8.1.** This SOP will be reviewed annually. Reviews will be documented on the Review History Table on page 2 of the SOP. The SOP shall be updated every 5 years, or more frequently, when necessary, due to significant changes.
- **8.2.** The "Request for SOP Change Form" is used to document changes and is appended to the final SOP until such time as the changes are incorporated into the body of the text of the SOP.

9.0 APPENDICES

- Appendix A Data Validation Criteria and Actions
- Appendix B Data Assessment Report Template
- Appendix C Definitions/Glossary of Terms
- Appendix D SOP Change Request Form (CRF)

Appendix A

Data Validation Criteria and Actions

TABLE OF CONTENTS

I.	DATA QUALIFIER DEFINITIONS	14
	ICP-MS TABLE 1. DATA QUALIFIER DEFINITIONS	14
н.	PRESERVATION AND HOLDING TIMES	15
	ICP-MS TABLE 2. PRESERVATION AND HOLDING TIME ACTIONS	15
III.	TUNE ANALYSIS	16
	ICP-MS TABLE 3. TUNE ACTIONS	16
IV.	CALIBRATION	16
	ICP-MS TABLE 4. CALIBRATION ACTIONS	17
v.	BLANKS	
	ICP-MS TABLE 5. BLANKS ACTIONS	19
VI.	INTERFERENCE CHECK SAMPLE	20
	ICP-MS TABLE 6. INTERFERENCE CHECK ACTIONS	21
VII.	LABORATORY CONTROL SAMPLE	22
	ICP-MS TABLE 7. LCS ACTIONS	22
VIII.	LABORATORY DUPLICATE SAMPLE ANALYSIS	22
	ICP-MS TABLE 8. LABORATORY DUPLICATE SAMPLE ACTIONS	23
IX.	SPIKE SAMPLE ANALYSIS	24
	ICP-MS TABLE 9. SPIKE SAMPLE ACTIONS	25
х.	SERIAL DILUTION	26
	ICP-MS TABLE 10. SERIAL DILUTION ACTIONS	26
XI.	INTERNAL STANDARDS	27
	ICP-MS TABLE 11. INTERNAL STANDARD ACTIONS	28
XII.	FIELD DUPLICATE SAMPLE ANALYSIS	28
	ICP-MS TABLE 12. FIELD DUPLICATE SAMPLE ACTIONS	29

XIII.	TARGET ANALYTE QUANTITATION	. 30
	ICP-MS TABLE 13. TARGET ANALYTE QUANTITATION - PERCENT SOLIDS OF SEDIMENT ACTION	30

I. Data Qualifier Definitions

The following table provides brief explanations of the qualifiers assigned to results during the data review process. The reviewer should use these qualifiers as applicable.

Data Qualifier	Definition
U	The analyte was analyzed for but was not detected above the level of the adjusted detection limit or quantitation limit, as appropriate.
J	The result is an estimated quantity. The associated numerical value is the approximate concentration of the analyte in the sample.
J+	The result is an estimated quantity, but the result may be biased high.
J-	The result is an estimated quantity, but the result may be biased low.
IJ	The analyte was analyzed for but was not detected. The reported quantitation limit is approximate and may be inaccurate or imprecise.
R	The data are unusable. The sample results are rejected due to serious deficiencies in meeting QC criteria. The analyte may or may not be present in the sample.

ICP-MS Table 1. Data Qualifier Definitions

NOTES:

- 1. Comments for sample results with data qualifiers other than "U" or no qualification based on professional judgement must be included in the DAR.
- 2. With familiarity of project data objectives and/or consultation with project staff, the reviewer should be able to refine the use of data qualifiers to avoid ambiguity. For example, if critical site decisions are to be made based on the data, the reviewer may decide to apply an "R" qualifier rather than a "UJ"
- 3. Although a "J+" or a "J-" may be seen as less ambiguous than a "J", the reviewer should reserve the application of directional bias indicators to those situations when there is an overwhelming influence in one direction. The exercise of professional judgment is critical, especially in situations where ambiguity exists due to opposing factors, to objectively interpret the effects of all factors.
- 4. Criteria, evaluation, quantitation limits (QLs), calculations, acceptable ranges and related parameters and definitions are detailed in the applicable Statement of Work (SOW) and/or National Functional Guidelines (NFG) documents referenced.

II. Preservation and Holding Times

A. Review Items

Laboratory Results Reports, sampling documentation [e.g., Chain of Custody (COC) Records], sample receipt forms, sample preparation logs, raw data, and narrative in the data package, checking for: pH, shipping container temperature, holding time, and other sample conditions.

B. Objective

The objective is to determine the validity of the analytical results based on the sample shipping and storage conditions and the holding time of the sample.

C. Action:

Refer to ICP-MS Table 2 below for the evaluation criteria and corresponding actions for detected and non-detected target analyte results in the deficient samples. Apply the actions to each field sample and field blank for which the preservation or holding time criteria was not met.

If a discrepancy is found between the sample analysis dates on the Laboratory Results Reports and in the raw data, perform a more comprehensive review to determine the correct date to be used to establish the holding time.

	Action	
Criteria	Detect	Non-detect
Aqueous/water samples received with $pH \ge 2$ and pH adjusted by laboratory	No qualification	No qualification
Aqueous/water samples received with $pH \ge 2$ and pH not adjusted	J-	R
Technical Holding Time: Aqueous/water samples > 180 days	J-	R
Technical Holding Time: Soil/sediment/waste samples > 180 days	J-	R
Samples properly preserved and analyzed within specified holding time	No qualification	No qualification

ICP-MS Table 2. Preservation and Holding Time Actions

III. Tune Analysis

A. Review Items

Laboratory instrument performance check (Tune) reports (if available), instrument printouts and raw data in the data packages.

B. Objective

The ICP-MS tune serves as an initial demonstration of instrument stability and precision.

C. Action:

Refer to ICP-MS Table 3 below for the evaluation criteria and corresponding actions for detected and non-detected target analyte results in the deficient ICP-MS Tunes. For ICP- MS tunes that do not meet the technical criteria, apply the actions to all samples reported from the analytical sequence.

	Action	
Criteria	Detect	Non-detect
Tune not performed	R	R
Tune not performed with required isotopes and/or number of scans	J	IJ
Resolution of mass calibration not within 0.1 u	J	UJ
%RSD > 5%	J	UJ
Tune properly analyzed with required isotopes, mass resolution and %RSD within specified limits	No qualification	No qualification

ICP-MS Table 3. Tune Actions

IV. Calibration

A. Review Items

Laboratory initial calibration and calibration verification reports (if available), preparation logs, calibration standard logs, instrument logs, instrument printouts, and raw data in the data package.

B. Objective

The objective is to determine the validity of the analytical results based on initial calibration and calibration verification.

C. Action:

Refer to ICP-MS Table 4 below for the evaluation criteria and corresponding actions for detected and non-detected target analyte results in the samples associated with deficient initial calibrations or calibration verification standard.

- 1. For initial calibrations or ICV standard analyses that do not meet the technical criteria, apply the actions to all associated samples reported from the analytical sequence.
- 2. For CCV standards analyses that do not meet the technical criteria, apply the actions to all samples analyzed between a previous technically acceptable analysis of the QC sample and a subsequent technically acceptable analysis of the QC sample in the analytical sequence.
- 3. If the instrument was not calibrated with a blank and at least 5 calibration standards, or if the calibration curve does not include standards at required concentrations (e.g., a blank and at least one at or below the QL but above the MDL), qualify detects as estimated (J) and non-detects as estimated (UJ).
- **NOTE:** For critical samples, a further in-depth evaluation of the calibration curve may be warranted to determine if additional qualification is necessary.

	Action	
Criteria	Detect	Non-detect
Calibration not performed or not performed at specified frequency	R	R
Calibration incomplete (insufficient number ofstandards or required concentrations missing)	J	UJ
Not at least one calibration standard at or below the QL for each analyte	J	UJ
For linear fits, the correlation coefficient < 0.995; %D outside ±30%; or other specified statistical test values outside limits	J	UJ
ICV/CCV not performed at specified frequency	J	UJ
ICV/CCV %R < 75%	J-	UJ
ICV/CCV %R 75-89%	J	UJ
ICV/CCV %R 90-110%	No qualification	No qualification
ICV/CCV %R 111-125%	J	No qualification
ICV/CCV %R > 125%	J+	No qualification

ICP-MS Table 4. Calibration Actions

ICV/CCV %RSD > 5%	J	UJ
Instrument blank analyzed prior to CCV	Use professional	Use professional
	judgment	judgment

V. <u>Blanks</u>

A. Review Items

Laboratory blanks reports (if available), preparation logs, calibration standard logs, instrument logs, and raw data in the data package, and sampling trip reports.

B. Objective

The objective is to determine the validity of the analytical results based on the blank responses by determining the existence and magnitude of contamination resulting from laboratory (or field) activities or baseline drift during analysis.

C. Action:

Refer to ICP-MS Table 5 below for the evaluation criteria and corresponding actions for detected and non-detected target analyte results in the samples associated with deficient blanks.

- 1. For ICB analyses that do not meet the technical criteria, apply the actions to all associated samples reported from the analytical sequence.
- 2. For CCB analyses that do not meet the technical criteria, apply the actions to all associated samples analyzed between a previous technically acceptable analysis of the CCB and a subsequent technically acceptable analysis of the CCB in the analytical sequence.
- 3. For Preparation Blank analyses that do not meet the technical criteria, apply the actions to all associated samples prepared in the same preparation batch.
- 4. Action regarding 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 contaminant.
- If the absolute value of an ICB or a CCB result is ≥ QL, the analysis should have been terminated and the affected samples re-analyzed. If samples were not re-analyzed, qualify as described in Table 5 below.
- 6. All samples associated with the Preparation Blank with concentrations < 10x the Preparation Blank concentration and ≥ QL should have been redigested and reanalyzed. If the associated samples were not redigested and reanalyzed, qualify as described in Table 4 below.

- If an analyte result in a diluted sample analysis is < QL, the final analyte result should be checked against a less dilute run and reported from that analysis. However, if no less-dilute analysis is reported, use professional judgment to decide whether to report from the dilution.
- 8. For blank results \leq (-MDL) but > (-QL), the possibility of false negative exists.
- **NOTE:** Do not qualify blanks with blank results. The blank analyses may not involve the same weights, volumes, or dilution factors as the associated samples. In particular, soil/sediment or waste sample results reported Laboratory Results Reports will not be on the same basis (units, dilution) as the calibration blank data. It may be easier to work with the raw data and/or convert the ICB or CCB results to the same units as the soil/sediment or waste samples for comparison purposes.

Blank Type	Blank Result	Sample Result	Action for Samples
	Not analyzed at	Non-detect	UJ
ICB/CCB	CCB the specified frequency Detect		J
		Non-detect	No qualification
ICB/CCB	Detect < QL	Detect < QL	Report at QL and qualify U
		≥QL	No qualification
	≤ (-MDL) but	Non-detect	UJ
ICB/CCB	ICB/CCB > (-QL) Detect		No qualification
		Non-detect	No qualification
		Detect < QL	Report at QL and qualify U
ICB/CCB	≥QL	≥ QL but < ICB/CCB Result	Report at ICB/CCB Result and qualify U
		≥ ICB/CCB Result	No qualification
		Non-detect	UJ
		Detect < QL	J-
ICB/CCB	≤ (-QL)	≥ QL but < 10x QL	J-
		≥ 10x QL	No qualification

ICP-MS Table 5. Blanks Actions

Preparation	Not analyzed at	Non-detect	UJ
Blank/LEB	the specified frequency	Detect	J
Preparation		Non-detect	No qualification
Blank/LEB/ Field Blank/	ank/LEB/ eld Blank/ Detect < QL Detect < QL		Report at QL and qualify U
Rinse Blank		≥QL	No qualification
Preparation Blank/LEB/	≤ (-MDL) but	Non-detect	IJ
Field Blank/ Rinse Blank	> (-QL)	Detect	No qualification
		Non-detect	No qualification
Preparation		Detect < QL	Report at QL and qualify U
		≥ QL but ≤ the PB/LEB/FB/RB Result	Report at Blank Result and qualify U
Blank/LEB/ Field Blank/ Rinse Blank	≥ QL > PB/LEB/FB/RB Result but < 10x the PB/LEB/FB/RB Result		Report at Sample Result and qualify J+
		≥ 10x the Preparation Blank/LEB/Field Blank/ Rinse Blank Result	No qualification
Droparation		Non-detect	UJ
Preparation Blank/LEB/	≤ (-QL)	Detect < QL	J-
Field Blank/ Rinse Blank	≤ (-QL)	≥ QL but < 10x QL	J-
		≥ 10x QL	No qualification

VI. Interference Check Sample

A. Review Items

Laboratory interference checks reports (if available), instrument printouts and raw data in the data package.

B. Objective

The objective is to determine the validity of the analytical results based on the instrument's ability to overcome interferences typical of those found in samples.

C. Action:

Refer to ICP-MS Table 6 below for the evaluation criteria and corresponding actions for detected and non-detected target analyte results in the samples associated with deficient ICSs.

- 1. For an ICS analysis that does not meet the technical criteria, apply the actions to all samples reported from the analytical sequence.
- **NOTE:** The same result units should be used when comparing analyte results in samples to those in the ICS. Unit conversion may be necessary when soil/sediment/waste samples are evaluated.

Apply action only if the concentration of interferents Aluminum (Al), Calcium (Ca), Iron (Fe), and Magnesium (Mg) in the sample are found to be greater than their respective concentrations in the ICS.

2. Actions regarding the interpretation and/or the subsequent qualification of ICP data due to the ICS analytical results can be complex. Use professional judgment to determine the need for the associated sample data to be qualified. Obtain additional information from the laboratory, if necessary. Record all interpretive situations in the Data Review Narrative.

	Action	
Criteria	Detect	Non-detect
ICS not analyzed	R	R
ICS not analyzed in specified sequence	J	UJ
ICSAB %R < 50%	J-	R
ICS %R 50-84% [or ICS found value is < (true value –2x QL), whichever is lower]	J-	UJ
ICS %R 85-115%	No qualification	No qualification
ICS %R 116-150% [or ICS true value is > (true value + 2x QL), whichever is greater]	J+	No qualification
ICS %R > 150%	J+	No qualification
ICSA results ≥ DLs or MDLs, but not present in ICS (potential false positives)	J+	No qualification
Negative ICSA results, but not present in ICS (potential false negatives)	J- for results < 10x (negative sampleresult)	IJ

ICP-MS Table 6. Interference Check Actions

VII. Laboratory Control Sample

A. Review Items

Laboratory LCS reports (if available), preparation logs, instrument printouts, and raw data in the data package.

B. Objective

The objective is to determine the validity of the analytical results based on the recovery of the digested Laboratory Control Sample (LCS).

C. Action

Refer to ICP-MS Table 7 for the evaluation criteria and corresponding actions for detected and non- detected target analytes in the samples associated with deficient LCSs. For an LCS analysis that does not meet the technical criteria, apply the actions to all samples in the same preparation batch.

Matrix spike data can be reviewed to determine batch quality if an LCS was not prepared and analyzed with the samples.

	Action	
Criteria	Detect	Non-detect
LCS not prepared with sample	J	UJ
LCS not prepared at specified concentration	J	UJ
Aqueous/water and soil/sediment/waste %R < 40%	J-	R
Aqueous/water and soil/sediment/waste %R 40-69%	J-	UJ
Aqueous/water and soil/sediment/waste %R 70-130%	No qualification	No qualification
Aqueous/water and soil/sediment/waste %R 131-150%	J+	No qualification
Aqueous/water and soil/sediment/waste %R > 150%	R	No qualification

ICP-MS Table 7. LCS Actions

VIII. Laboratory Duplicate Sample Analysis

A. Review Items

Data Package Cover Page, laboratory duplicate reports (if available), preparation logs, instrument printouts, and raw data in the data package.

B. Objective

The objective of the duplicate sample analysis is to demonstrate acceptable method precision by the laboratory at the time of analysis.

C. Action:

Refer to ICP-MS Table 8 below for the evaluation criteria and corresponding actions for detected and non-detected target analyte results in the samples associated with deficient duplicates.

- 1. For a laboratory duplicate sample analysis that does not meet the technical criteria, apply the actions to the field sample used to prepare the duplicate sample only. Exercise professional judgment in determining sample similarity when making use of all available data, including: site and sampling documentation (e.g., location and type of sample, descriptive data, soil classification); field test data (e.g., pH, E_h, conductivity, chlorine); and laboratory data for other parameters [e.g., Total Suspended Solids (TSS), Total Dissolved Solids (TDS), Total Organic Carbon (TOC), alkalinity or buffering capacity, reactive sulfide, anions]. Additionally, use the sample data (e.g., similar concentrations of mercury) in determining similarity between samples in the data package. Two possible determinations are: 1) all of the samples are considered sufficiently similar to the duplicate sample and all of the samples should be qualified; or 2) only some of the samples in the data package are similar to the duplicate sample, and that only these samples should be qualified.
- 2. Note the potential effects on the data due to out-of-control duplicate sample results in the Data Review Narrative.
- 3. For high RPDs (i.e., > 100%), use professional judgment to qualify the data as this may be indicative of a sampling problem.

	Action	
Criteria	Detect	Non-detect
Duplicate analysis not performed at the specified frequency	J	UJ
<i>Aqueous</i> : Both original sample and duplicate sample results are ≥ 5x QL and 20% < RPD ≤ 100%	J	NA
Soil/Sediment: Both original sample and duplicate sample results are ≥ 5x QL and 35% < RPD ≤ 100%	J	NA
<i>Aqueous</i> : Both original sample and duplicate sample results are ≥ 5x QL and RPD ≤ 20%	No qualification	No qualification
Soil/Sediment: Both original sample and duplicate sample results are ≥ 5x QL and RPD ≤ 35%	No qualification	No qualification

ICP-MS Table 8. Laboratory Duplicate Sample Actions

Both original sample and duplicate sample results are ≥ 5x QL and RPD > 100%	R	NA
Original sample or duplicate sample result < 5x QL and absolute difference between sample and duplicate > QL	J	IJ
Original sample or duplicate sample result < 5x QL (including non-detects) and absolute difference between sample and duplicate ≤ QL	No qualification	No qualification

IX. Spike Sample Analysis

A. Review Items

Data Package Cover Page, laboratory matrix spike reports (if available), preparation logs, instrument printouts, and raw data in the data package.

B. Objective

The objective of the spiked sample analysis is to evaluate the effect of each sample matrix on the sample preparation procedures and the measurement methodology.

C. Action:

Refer to ICP-MS Table 9 below for the evaluation criteria and corresponding actions for detected and non-detected target and spike analyte results in the samples associated with deficient matrix spikes.

- 1. For a matrix spike sample analysis that does not meet the technical criteria, apply the actions only to the field sample used to prepare the Matrix Spike sample. Exercise professional judgment in determining sample similarity when making use of all available data, including site and sampling documentation (e.g., location and type of sample, descriptive data, soil classification); field test data (e.g., pH, E_h, conductivity, chlorine); and laboratory data for other parameters [e.g., Total Suspended Solids (TSS), Total Dissolved Solids (TDS), Total Organic Carbon (TOC), alkalinity or buffering capacity, reactive sulfide, anions]. Additionally, use the sample data (e.g., similar concentrations of analytes) in determining similarity between samples in the data package. Two possible determinations are: 1) all of the samples are considered sufficiently similar, and all samples should be qualified; or 2) only some of the samples in the data package are similar to the Matrix Spike sample, and that only these samples should be qualified.
- 2. Note the potential effects on the data due to out-of-control spiked sample results in the Data Review Narrative.
- **NOTE:** Matrix spike analysis is not required for SDG that contains only field blank samples.

When the Sample Result is reported as a non-detect, use SR = 0 only for calculating the %R.

Matrix spike analysis is not required for Calcium (Ca), Magnesium (Mg), Potassium (K), and Sodium (Na) for both matrices; Aluminum (Al) and Iron (Fe) for soil only.

Disregard the out-of-control spike recoveries for analytes whose unspiked concentrations are $\ge 4x$ the spike added.

Criteria	Action		
	Detect	Non-detect	
Matrix Spike analysis not performed at the specified frequency (qualify all samples associated with the matrix spike)	J	IJ	
Matrix Spike not prepared from field sample (qualify all samples associated with the matrix spike)	J	UJ	
Matrix Spike %R < 30% Post-digestion spike %R < 75%	J-	R	
Matrix Spike %R < 30% Post-digestion spike %R ≥ 75%	J	UJ	
Matrix Spike %R 30-74% Post-digestion spike %R < 75%	J-	IJ	
Matrix Spike %R 30-74% Post-digestion spike %R ≥ 75%	J	UJ	
Matrix Spike %R > 125% Post-digestion spike %R > 125%	J+	No qualification	
Matrix Spike %R > 125% Post-digestion spike %R ≤ 125%	J	No qualification	
Matrix Spike %R < 30% No post-digestion spike performed	J-	R	
Matrix Spike %R 30-74% No post-digestion spike performed	J-	UJ	
Matrix Spike %R 75-125% No post-digestion spike is required	No qualification	No qualification	
Matrix Spike %R > 125% No post-digestion spike performed	J+	No qualification	

ICP-MS Table 9. Spike Sample Actions

X. Serial Dilution

A. Review Items

Laboratory serial dilution reports (if available), instrument printouts, and raw data in the data package.

B. Objective

The objective of the serial dilution analysis is to determine if significant physical or chemical interferences exist due to sample matrix.

C. Action:

Refer to ICP-MS Table 10 below for the evaluation criteria and corresponding actions for detected and non-detected target analyte results in the samples associated with deficient serial dilution analyses.

- 1. For a serial dilution sample analysis that does not meet the technical criteria, apply the actions to all samples of the same matrix if the samples are considered sufficiently similar. Exercise professional judgment in determining sample similarity when making use of all available data, including: site and sampling documentation (e.g., location and type of sample, descriptive data, soil classification); field test data (e.g., pH, E_h, conductivity, chlorine); and laboratory data for other parameters [e.g., Total Suspended Solids (TSS), Total Dissolved Solids (TDS), Total Organic Carbon (TOC), alkalinity or buffering capacity, reactive sulfide, anions]. Additionally, use the sample data (e.g., similar concentrations of analytes) in determining similarity between samples in the data package. Two possible determinations are: 1) only some of the samples in the data package are similar to the serial dilution sample, and that only these samples should be qualified; or 2) no samples are sufficiently similar to the sample used for serial dilution, and thus only the field sample used to prepare the serial dilution sample should be qualified.
- 2. Note the potential effects on the reported data in the Data Review Narrative.
- **NOTE:** Serial dilution analysis is not required for SDG that contains only field blank samples.

Qualifiers based on serial dilutions are to be applied to the parent sample only.

Criteria	Action	
	Detect	Non-detect
Serial Dilution analysis not performed at the specified frequency	J	IJ
Aqueous: Sample concentration > 50x MDL, serial dilution sample concentration \geq QL, and %D \geq 100%	R	NA
Soil/Sediment: Sample concentration > 50x MDL, serial dilution sample concentration \ge QL, and %D \ge 120%	R	NA

ICP-MS Table 10. Serial Dilution Actions

Aqueous: Sample concentration > 50x MDL, serial dilution sample concentration ≥ QL, and 10% < %D < 100%	J	No qualification
Soil/Sediments: Sample concentration > 50x MDL, serial dilution sample concentration ≥ QL, and 15% < %D < 120%	J	No qualification
Aqueous: Sample concentration > 50x MDL and serial dilution sample concentration ≥ QL, and %D < 10%	No qualification	NA
Soil/Sediment: Sample concentration > 50x MDL and serial dilution sample concentration ≥ QL, and %D < 15%	No qualification	NA
Sample concentration > 5x QL and serial dilution sample concentration < QL	No qualification	No qualification
	Use	Use
Interferences present	professional	professional
	judgment	judgment

XI. Internal Standards

A. Review Items

Laboratory internal standard reports (if available), instrument printouts and raw data in the data package.

B. Objective

The objective of internal standard analysis is to determine the existence and magnitude of instrument drift and physical interferences.

C. Action:

Refer to ICP-MS Table 11 below for the evaluation criteria and corresponding actions for detected and non-detected target analyte results in the samples with deficient internal standards. Apply the actions to the affected analytes for each sample that does not meet the internal standard criteria.

If the Internal Standard %RI grossly exceeds the limits in both the original analysis and the diluted re-analysis, qualify the data based on the following considerations:

- If the %RI is greater than 200%, high recoveries are generally due to the natural presence of the internal standard isotope in the sample(s). This occurrence may have been detected in earlier sampling of the site. Apply another appropriate internal standard to the affected analytes, do not qualify the analytes based on the high internal standard.
- 2. If the Internal Standard %RI is less than 30%, it is possible that some form of signal suppression is taking place.

When two separate qualifiers are listed as actions, use professional judgment to qualify detects and non-detects based on the extent to which the criteria are not met.

	Action	
Criteria	Detect	Non-detect
Internal standards not analyzed	R	R
Less than the required number of internal standards analyzed	R	R
Target analyte not associated with internal standards	R	R
%RI 60 - 125%	No qualification	No qualification
%RI < 60% or > 125% and original sample reanalyzed at specified dilution with %RI 60-125%	No qualification	No qualification
%RI < 60% or > 125% and original sample reanalyzed at specified dilution with %RI < 60% or > 125%	J	UJ
Original sample not reanalyzed at specified dilution	Use professional judgment to qualify J or R	Use professional judgment to qualify UJ or R

ICP-MS Table 11. Internal Standard Actions

XII. Field Duplicate Sample Analysis

A. Review Items

Data Package Cover Page, sampling chain of custody page, laboratory duplicate reports (if available), preparation logs, instrument printouts, raw data in the data package, and sampling trip reports.

B. Objective

The objective of the field duplicate sample analysis is to demonstrate and guarantee the acceptability of the sampling method through the homogeneity of the sample group test data results.

C. Action:

Refer to ICP-MS Table 12 below for the evaluation criteria and corresponding actions for detected and non-detected target analyte results in the samples associated with deficient field duplicates.

- If a field duplicate samples pair was collected and analyzed, calculate, and report the RPD when the sample and its field duplicate values are both greater than or equal to (≥) 5x QL. Calculate and report the absolute difference when at least one value (sample or its field duplicate) is less than (<) 5x QL.
- 2. If one value is greater than (>) QL and the other value is non-detect, calculate the absolute difference between the value greater than (>) QL and the MDL, and use this criterion to qualify the results.

- 3. For a field duplicate sample analysis that does not meet the technical criteria, apply the actions to only the field samples and its duplicate. Exercise professional judgment in determining sample similarity when making use of all available data, including site and sampling documentation (e.g., location and type of sample, descriptive data, soil classification); field test data (e.g., pH, E_h, conductivity, chlorine); and laboratory data for other parameters [e.g., Total Suspended Solids (TSS), Total Dissolved Solids (TDS), Total Organic Carbon (TOC), alkalinity or buffering capacity, reactive sulfide, anions]. Additionally, use the sample data (e.g., similar concentrations of analytes) in determining similarity between samples in the data package. Two possible determinations are: 1) all of the samples are considered sufficiently similar, and all samples should be qualified; or 2) only some of the samples in the data package are similar to the field duplicate sample, and that only these samples should be qualified.
- 4. Any action should be in accordance with the project specifications and the criteria for acceptable field duplicate sample results.
- 5. In general, for QA/QC performance not within QAPP specification, qualify detects as estimated (J) and non-detects as estimated (UJ).

NOTE: Do not calculate RPD when either value is non-detect.

	Action		
Criteria	Detect	Non-detect	
<i>Aqueous</i> : Both original sample and its field duplicate sample results are ≥ 5x QL and RPD > 20% but < 100%	J	NA	
Soil/Sediment: Both original sample and its field duplicate sample results are ≥ 5x QL and RPD > 50% but < 100%	J	NA	
Both original sample and its field duplicate sample results are ≥ 5x QL and RPD ≤ 20% (<i>Aqueous</i>) / RPD ≤ 50% (<i>Soil/Sediment</i>)	No qualification	No qualification	
Aqueous: Original sample and/or its field duplicate sample result < 5x QL and absolute difference between sample and duplicate > QL	J	UJ	
Soil/Sediment: Original sample and/or its field duplicate sample result < 5x QL (including non-detects) and absolute difference between sample and duplicate > 2x QL	J	UJ	
Original sample and its field duplicate ≥ 5x QL and RPD ≥ 100%	Use professional judgment to qualify other than R	NA	

ICP-MS Table 12. Field Duplicate Sample Actions

XIII. Target Analyte Quantitation

A. Review Items

Laboratory result reports, sample preparation sheet, data package narrative, instrument printouts and raw data.

B. Objective

The objective is to ensure that the reported results and quantitation limits for target analytes reported by the laboratory are accurate and sufficient to meet requirements.

C. Action:

Refer to ICP-MS Table 13 below for the evaluation criteria and corresponding actions for detected and non-detected target analyte results in the samples with deficient quantitation limits. Apply the actions to the affected analytes for each sample that does not meet the quantitation criteria.

- 1. If sample results are < QLs and \geq MDLs or limits in the QAPP, qualify as estimated (J).
- 2. If any sample result was greater than the linear range for ICP-MS and the sample was not diluted to obtain the result reported on Form I, qualify the affected results as estimated (J).
- 3. If the percent solids of sediment for a sample are < 50% but ≥ 30%, qualify the affected results ≥ MDL as estimated (J), and the non-detects as estimated (UJ).
- 4. If the sample's percent solids of sediment are < 30%, check if the sample was prepared at greater mass to maintain the QLs. Use professional judgment when this was not completed.

ICP-MS Table 13. Target Analyte Quantitation - Percent Solids of Sediment	Actions
---	---------

	Action		
Criteria	Detect	Non-detect	
Sample result < QLs and \geq MDLs or limits in the QAPP	J	NA	
Sample result > the linear range for ICP-MS and the sample was not diluted to obtain the result reported on Form I, qualify the affected results as estimated, J	J	NA	
Percent solids of sediment sample < 50% but \ge 30%	J	UJ	
Percent solids of sediment sample < 30%, and was not prepared at greater mass to maintain QLs	Use professional judgment to qualify J or R	Use professional judgment to qualify UJ or R	

Appendix B

Data Assessment Report Template



UNITED STATES ENVIRONMENTAL PROTECTION AGENCY REGION 2 LSASD/HWSB/HWSS 2890 Woodbridge Avenue, Edison, NJ 08837

EXECUTIVE NARRATIVE

Case No.: Site: Number of Samples: Analysis: SDG No.: Laboratory: Sampling dates: Validation SOP:

QAPP: Contractor: Reference: DCN Number

SUMMARY OF DEFINITIONS:

Critical: Results have an unacceptable level of uncertainty and should not be used for making decisions. Data have been qualified "R" rejected.

Major: A level of uncertainty exists that may not meet the data quality objectives for the project. A bias is likely to be present in the results. Data has been qualified "J" estimated. "J+" and "J-" represent likely direction of the bias.

Minor: The level of uncertainty is acceptable. No significant bias in the data was observed.

Critical Findings:

Major Findings:

Minor Findings:

COMMENTS:

Reviewer Name(s):

Approver's Signature:

Name:

Date:

Affiliation: USEPA/R2/LSASD/HWSB/HWSS

Appendix C

Definitions/Glossary of Terms

Definitions/Glossary of Terms

Action Limit – A result for a Performance Evaluation (PE) sample that is outside the 99% ($\pm 3\sigma$) control limits. The laboratory may be required to apply and document corrective actions to bring the analytical results back into control.

Analyte – The element or ion an analysis seeks to determine, the element of interest.

Analytical Services Branch (ASB) – Directs the Contract Laboratory Program (CLP) from within the Office of Superfund Remediation and Technical Innovation (OSRTI) in the Office of Solid Waste and Emergency Response (OSWER).

Analytical Sample – Any prepared field sample or extract thereof that is introduced into an instrument for the purpose of measuring any target analyte. This definition excludes any instrument quality control samples (e.g., standards associated with initial calibration, Initial Calibration Verification (ICV), Initial Calibration Blank (ICB), Continuing Calibration Verification (CCV), Continuing Calibration Blank (CCB), and tune verifications). The following are also defined as analytical samples: diluted samples; matrix spike and matrix spike duplicate samples; duplicate samples; serial dilution samples, postdigestion/post-distillation spike samples; Laboratory Control Samples (LCSs); Performance Evaluation (PE) samples; Preparation/Method Blanks; Field Blanks (FBs); and Leachate Extraction Blanks (LEBs).

Associated Samples – Any sample related to a particular Quality Control (QC) analysis. For example, for Initial Calibration Verification (ICV), all samples analyzed under the same calibration curve. For duplicates, all Sample Delivery Group (SDG) samples digested/distilled of the same matrix.

Blank – An analytical sample that has negligible or unmeasurable amounts of a substance of interest. The blank is designed to assess specific sources of contamination. Types of blanks may include calibration blanks, preparation blanks, and field blanks. See the individual definitions for types of blanks.

Calibration – A set of operations that establish under specific conditions, the relationship between values indicated by a measuring instrument and the corresponding known values. The calibration standards should be prepared using the same type of reagents or concentration of acids as used in the sample preparation.

Calibration Blank – A blank solution containing all reagents and in the same concentration as those used in the analytical sample preparation. This blank is digested/distilled for mercury and cyanide. Calibration blanks are used to verify that the instrument baseline is stable, and the instrument is free of contamination.

Calibration Curve – A plot of instrument response versus concentration of standards.

Calibration Standards – A series of known standard solutions used by the analyst for calibration of the instrument (i.e., preparation of the calibration curve). The solutions may or may not be subjected to the preparation method, but contain the same matrix (i.e., the same amount of reagents and/or preservatives) as the sample preparations to be analyzed.

Case – A finite, usually predetermined number of samples collected over a given period from a particular project site. A case numbers is assigned by the Sample Management Office (SMO) and consists of one or more Sample Delivery Groups (SDGs).

Chain of Custody (COC) Record – A sample identification form completed by the sampler, which accompanies the sample during shipment to the laboratory and is used to document sample identity, sample chain of custody, sample condition, and sample receipt by the laboratory.

Contamination – A component of a sample or an extract that is not representative of the environmental source of the sample. Contamination may result from other samples, sampling equipment, or from introduction while in transit, from laboratory reagents, from the laboratory environment, or from analytical instruments.

Continuing Calibration Blank (CCB) – A reagent water sample that is run at specified interval and designed to detect any carryover contamination.

Contract Compliance Screening (CCS) – A screening of electronic and hardcopy data deliverables for completeness and compliance with the contract. This screening is performed under EPA direction by the Contract Laboratory Program (CLP) Sample Management Office (SMO) contractor.

Continuing Calibration Verification (CCV) – A single parameter or multi-parameter standard solution prepared from the same source as the initial calibration standards by the analyst and used to periodically verify the stability of the instrument calibration during analysis of samples. The CCV can be one of the calibration standards with the concentration near the middle of the calibration range. However, all parameters being measured by the particular system must be represented in this standard and the standard must have the same matrix (i.e., the same amount of reagents and/or preservatives) as the samples.

Control Limits – A range within which specified measurement results should fall to be compliant. Control limits may be mandatory, requiring corrective action if exceeded, or advisory, requiring that noncompliant data be flagged.

Contract Laboratory Program (CLP) – Supports the EPA's Superfund effort by providing a range of state-of-the-art chemical analytical services of known quality. This program is directed by the Analytical Services Branch (ASB) of the Office of Superfund Remediation and Technical Innovation (OSRTI) of USEPA.

Contract Required Quantitation Limit (CRQL) – Minimum level of quantitation acceptable under the contract Statement of Work (SOW).

Data Package Narrative – Portion of the data package which includes laboratory information, sample identification, and descriptive documentation of any problems encountered in processing the samples, along with corrective action taken and problem resolution.

Detection Limit (DL) - A generic term for the minimum measured concentration of a substance that can be reported with a specified confidence that the measured concentration is distinguishable from blank results. Includes Method Detection Limit (MDL), Limit of Detection (LOD), and other means of establishing this limit.

Duplicate – A second aliquot of a sample that is treated the same as the original sample in order to evaluate the precision.

Field Blank (FB) –A blank used to provide information about contaminants that may be introduced during sample collection, shipment, storage, and/or preparation and analysis in the laboratory. Examples of field blanks include trip blanks, rinse blanks, bottle blanks, equipment blanks, preservative blanks, decontamination blanks, etc.

Field Duplicate (FD) – A duplicate sample generated in the field, not in the laboratory.

Field Quality Control (FQC) – Any QC samples submitted from the field to the laboratory. Examples include, but are not limited to, field blanks, and field duplicates.

Field Sample – A portion of material received from the field to be analyzed for analytes of interest.

Holding Time – The maximum amount of time samples may be held before they are processed.

Holding Time (Contractual) – The maximum amount of time that the Contract Laboratory Program (CLP) laboratory may hold the samples from the sample receipt date until analysis and still be in compliance with the terms of the contract, as specified in the CLP Analytical Services Statement of Work (SOW). These times are the same or less than technical holding times to allow for sample packaging and shipping.

Holding Time (Technical) – The maximum amount of time that samples may be held from the collection date until analysis.

Initial Calibration – Analysis of analytical standards at a series of different specified concentrations; used to define the quantitative response, linearity, and dynamic range of the instrument to target analytes.

Initial Calibration Blank (ICB) – The first blank standard analysis to confirm the calibration curve.

Initial Calibration Verification (ICV) – The analysis of solution(s) prepared from stock standard solutions, metals, or salts obtained from a source separate from that utilized to prepare the calibration standards. The ICV is used to verify the concentration of the calibration standards and the adequacy of the instrument calibration. The ICV solution(s) should be traceable to National Institute of Standards and Technology (NIST) or other certified standard sources.

Interference Check Sample (ICS) – A solution containing both interfering and analyte elements of known concentration that can be used to verify background and interelement correction factors.

Internal Standard – A non-target element added to a sample at a known concentration after preparation but prior to analysis. Instrument responses to internal standards are monitored as a means of assessing overall instrument performance.

Laboratory – The place where the samples are processed and tested.

Laboratory Control Sample (LCS) – A reference matrix spiked with target analytes at a known concentration. LCSs are analyzed using the same sample preparation, reagents, and analytical methods employed for the samples received.

Leachate Extraction Blank (LEB) – A blank carried through the entire Toxicity Characteristic Leaching Procedure (TCLP) or Synthetic Precipitation Leaching Procedure (SPLP) extraction with the resulting leachate extracted, digested, or distilled by an appropriate aqueous method from the analytical method.

Matrix – The predominant material of which the sample to be analyzed is composed. For the purposes ofthis document, the matrices are aqueous/water, soil/sediment, and wipe. Matrix is <u>not</u> synonymous with phase (liquid or solid).

Matrix Spike – Aliquot of a sample (aqueous/water or soil/sediment) fortified (spiked) with known quantities of specific analytes and subjected to the entire analytical procedure to estimate recovery.

Method Detection Limit (MDL) – The minimum measured concentration of a substance that can be reported with 99% confidence such that the measured concentration is distinguishable from method blank results. Additional information about the procedure is provided in Title 40 of the Code of Federal Regulations (CFR), Chapter 1, Subchapter D, part 136, Appendix B, Definition and Procedure for the Determination of the Method Detection Limit, Revision 2.

Narrative (SDG Narrative) – Portion of the data package which includes laboratory, contract, Case, Sample Number identification, and descriptive documentation of any problems encountered in processing the samples, along with corrective action taken and problem resolution.

Office of Solid Waste and Emergency Response (OSWER) – The EPA office that provides policy, guidance, and direction for the EPA's solid waste and emergency response programs, including Superfund.

Percent Difference (%D) – The relative difference between two values (e.g., a measured and expected value) expressed as a percentage of one of the values (e.g., expected value).

Percent Solids (%Solids) – The proportion of solid in a soil/sediment sample determined by drying an aliquot of the sample.

Performance Evaluation (PE) Sample – A sample prepared by a third party at known concentrations that are unknown to the analytical laboratory and is provided to test whether the laboratory can produce analytical results within specified performance limits.

Post-Digestion Spike/Post-Distillation Spike – The addition of a known amount of standard after digestion or distillation (also identified as an analytical spike).

Preparation Blank – An analytical control that contains reagent water and reagents, which is carried through the entire preparation and analytical procedure.

Preparation Log – A record of sample preparation (e.g., digestion, extraction, distillation) at the laboratory.

Quality Assurance Project Plan (QAPP) – A formal document describing the management policies, objectives, principles, organizational authority, responsibilities, accountability, and implementation plan of an agency, organization, or laboratory for ensuring quality in its products and utility to its users.

Quantitation Limit (QL) – The minimum level of acceptable quantitation that is supported by the analysis of standards.

Raw Data – The originally recorded and unprocessed measurements from any measuring device such as analytical instruments, balances, pipettes, thermometers, etc. Reported data are processed raw measurement values that may have been reformatted from the original measurement to meet specific reporting requirements such as significant figures and decimal precision.

Relative Percent Difference (RPD) – The absolute of the relative difference between two values normalized to the mean of the two values expressed as a percentage.

Regional Sample Control Center Coordinator (RSCC) – In EPA Regions, coordinates sampling efforts and serves as the central point-of-contact for sampling questions and problems. Also assists in coordinating the level of Regional sampling activities to correspond with the monthly projected demand for analytical services.

Relative Standard Deviation (RSD) – As used in this document and the Statement of Work (SOW), the mean divided by the standard deviation, expressed as a percentage.

Sample – A portion of material to be analyzed that is contained in single or multiple containers and identified by a unique sample number.

Sample Delivery Group (SDG) – A unit within a sample Case that is used to identify a group of samples for delivery. An SDG is defined by the following, whichever is most frequent:

- a. Each 20 field samples [excluding Performance Evaluation (PE) samples] within a Case; or
- b. Each 7-calendar day period (3-calendar day period for 7-day turnaround) during which field samples in a Case are received (said period beginning with the receipt of the first sample in the SDG).
- c. Scheduled at the same level of deliverable.

In addition, all samples and/or sample fractions assigned to an SDG must be scheduled under the same contractual turnaround time. Preliminary Results have **no impact** on defining the SDG. Samples may be assigned to SDGs by matrix (i.e., all soil/sediment samples in one SDG, all aqueous/water samples in another) at the discretion of the laboratory.

Sample Identifier – A unique identification number that appears on the Chain of Custody (COC) Records or sampling forms which documents information for a sample.

Sample Management Office (SMO) – A contractor-operated facility operated under the SMO contract, awarded, and administered by the EPA. Provides necessary management, operations, and administrative support to the Contract Laboratory Program (CLP).

Sampling and Analysis Plan (SAP) – A document which specifies the procedural and analytical requirements for one-time, or time-limited, projects involving the collection of water, soil, sediment, or other samples taken to characterize areas of potential environmental contamination.

Serial Dilution (SD) – The dilution of a sample by a factor of five. When corrected by the Dilution Factor (DF), the diluted sample should agree with the original undiluted sample within specified limits. Serial dilution may reflect the influence of interferents [Inductively Coupled Plasma (ICP) only].

Soil – Synonymous with soil/sediment and sediment as used herein.

Statement of Work (SOW) – A document which specifies how laboratories analyze samples under a contract, such as the Contract Laboratory Program (CLP) analytical program.

Tune – A solution containing a range of isotope masses analyzed to serve as an initial demonstration of Inductively Coupled Plasma – Mass Spectrometry (ICP-MS) accuracy, resolution, and precision prior to calibration. May also be called Instrument Performance Check sample (IPC)

Warning Limit – A result for a Performance Evaluation (PE) sample that is outside the 95% ($\pm 2\sigma$) control limits. The laboratory should apply and document corrective actions to bring the analytical resultsback into control

Appendix D

SOP Change Request Form (CRF)

REQUEST FOR SOP CHANGE

Requestor Name:			Date of Initiation:			
Dept.:		SOP #:	Revisi	on #:	Date:	
SOP Title:						
Please Check One MINO		OR REVISION	. MA	MAJOR REVISION		
CHANGE(S) (Use attachment if necessary):						
CHANGE FROM:		• *				
CHANGE TO:						
REASON(S) FOR CHANGE(S):						
APPROVAI		NAME:		Signa	ture/Date	
EPA Branch Ch Section Chief/T Leader						
EPA TOCO	R					
REQUESTO	R					
Effective Dat	te		1			