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Innovative Technology Verification Report

XRF Technologies for Measuring Trace Elements in Soil and Sediment

Niton XLt 700 Series XRF Analyzer



Innovative Technology Verification Report

Niton XLt 700 Series XRF Analyzer

Prepared by

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National Exposure Research Laboratory Office of Research and Development U.S. Environmental Protection Agency

Notice

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Foreword

The U.S. Environmental Protection Agency (EPA) is charged by Congress with protecting the nation's natural resources. Under the mandate of national environmental laws, the Agency strives to formulate and implement actions leading to a compatible balance between human activities and the ability of natural systems to support and nurture life. To meet this mandate, EPA's Office of Research and Development (ORD) provides data and scientific support that can be used to solve environmental problems, build the scientific knowledge base needed to manage ecological resources wisely, understand how pollutants affect public health, and prevent or reduce environmental risks.

The National Exposure Research Laboratory is the Agency's center for investigation of technical and management approaches for identifying and quantifying risks to human health and the environment. Goals of the laboratory's research program are to (1) develop and evaluate methods and technologies for characterizing and monitoring air, soil, and water; (2) support regulatory and policy decisions; and (3) provide the scientific support needed to ensure effective implementation of environmental regulations and strategies.

EPA's Superfund Innovative Technology Evaluation (SITE) Program evaluates technologies designed for characterization and remediation of contaminated Superfund and Resource Conservation and Recovery Act (RCRA) sites. The SITE Program was created to provide reliable cost and performance data to speed acceptance and use of innovative remediation, characterization, and monitoring technologies by the regulatory and user community.

Effective monitoring and measurement technologies are needed to assess the degree of contamination at a site, provide data that can be used to determine the risk to public health or the environment, and monitor the success or failure of a remediation process. One component of the EPA SITE Program, the Monitoring and Measurement Technology (MMT) Program, demonstrates and evaluates innovative technologies to meet these needs.

Candidate technologies can originate within the federal government or the private sector. Through the SITE Program, developers are given an opportunity to conduct a rigorous demonstration of their technologies under actual field conditions. By completing the demonstration and distributing the results, the Agency establishes a baseline for acceptance and use of these technologies. The MMT Program is managed by ORD's Environmental Sciences Division in Las Vegas, Nevada.

Gary Foley, Ph.D. Director National Exposure Research Laboratory Office of Research and Development

Abstract

The Niton XLt 700 Series (XLt) XRF Services x-ray fluorescence (XRF) analyzer was demonstrated under the U.S. Environmental Protection Agency (EPA) Superfund Innovative Technology Evaluation (SITE) Program. The field portion of the demonstration was conducted in January 2005 at the Kennedy Athletic, Recreational and Social Park (KARS) at Kennedy Space Center on Merritt Island, Florida. The demonstration was designed to collect reliable performance and cost data for the XLt analyzer and seven other commercially available XRF instruments for measuring trace elements in soil and sediment. The performance and cost data were evaluated to document the relative performance of each XRF instrument.

This innovative technology verification report describes the objectives and the results of that evaluation and serves to verify the performance and cost of the XLt analyzer. Separate reports have been prepared for the other XRF instruments that were evaluated as part of the demonstration.

The objectives of the evaluation included determining each XRF instrument's accuracy, precision, sample throughput, and tendency for matrix effects. To fulfill these objectives, the field demonstration incorporated the analysis of 326 prepared samples of soil and sediment that contained 13 target elements. The prepared samples included blends of environmental samples from nine different sample collection sites as well as spiked samples with certified element concentrations. Accuracy was assessed by comparing the XRF instrument's results with data generated by a fixed laboratory (the reference laboratory). The reference laboratory performed element analysis using acid digestion and inductively coupled plasma – atomic emission spectrometry (ICP-AES), in accordance with EPA Method 3050B/6010B, and using cold vapor atomic absorption (CVAA) spectroscopy for mercury only, in accordance with EPA Method 7471A.

The Niton XLt is a small, field-portable instrument designed for chemical characterization of soils, sediment, and other thick homogeneous samples (plastics and metals). The analyzer features a miniaturized x-ray tube for the excitation source and a Peltier-cooled Si-PiN x-ray detector. The analyzer's standard software is programmed to analyze and automatically report 25 elements. The Niton XLt analyzer is designed to be used as either a hand-held instrument for in situ analysis or as a bench-top instrument, in a test stand with a sample drawer below the instrument, for ex situ analysis. The Niton XLt analyzer can be used to analyze elements under three primary scenarios: (1) bulk sample mode (includes soils, sediments, and metal alloys); (2) thin film mode (includes dust wipes and filters); and (3) plastics mode. XRF analyses using the Niton XLt analyzer comply with EPA Method 6200, "Field Portable XRF Spectrometry for the Determination of Elemental Concentrations in Soil and Sediment."

This report describes the results of the evaluation of the XLt analyzer based on the data obtained during the demonstration. The method detection limits, accuracy, and precision of the instrument for each of the 13 target analytes are presented and discussed. The cost of element analysis using the XLt analyzer is compiled and compared to both fixed laboratory costs and average XRF instrument costs.

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Acronyms, Abbreviations, and Symbols

μg	Micrograms
μA	Micro-amps
AC	Alternating current
ADC	Analog to digital converter
Ag	Silver
Am	Americium
ARDL	Applied Research and Development Laboratory, Inc.
As	Arsenic
ASARCO	American Smelting and Refining Company
BN	Burlington Northern
C	Celsius
Cd	Cadmium
CFR	Code of Federal Regulations
cps	Counts per second
CPU	Central processing unit
Cr	Chromium
CSV	Comma-separated value
Cu	Copper
CVAA	Cold vapor atomic absorption
EDXRF	Energy dispersive XRF
EDD	Electronic data deliverable
EPA	U.S. Environmental Protection Agency
ERA	Environmental Research Associates
ESA	Environmental site assessment
ESD	Environmental Sciences Division
ETV	Environmental Technology Verification (Program)
eV	Electron volts
Fe	Iron
FPT	Fundamental Parameters Technique
FWHM	Full width of peak at half maximum height
GB	Gigabyte
Hg	Mercury
Hz	Hertz

Acronyms, Abbreviations, and Symbols (Continued)

ICP-AES	Inductively coupled plasma-atomic emission spectrometry
ICP-MS	Inductively coupled plasma-mass spectrometry
IR	Infrared
ITVR	Innovative Technology Verification Report
KARS	Kennedy Athletic, Recreational and Social (Park)
keV	Kiloelectron volts
kg	Kilograms
KSC	Kennedy Space Center
kV	Kilovolts
LEAP	Light Element Analysis Program
LiF	Lithium fluoride
LIMS	Laboratory information management system
LOD	Limit of detection
mA	Milli-amps
MB	Megabyte
MBq	Mega Becquerels
MCA	Multi-channel analyzer
mCi	Millicuries
MDL	Method detection limit
mg/kg	Milligrams per kilogram
MHz	Megahertz
mm	Millimeters
MMT	Monitoring and Measurement Technology (Program)
Mo	Molybdenum
MS	Matrix spike
MSD	Matrix spike duplicate
NASA	National Aeronautics and Space Administration
NELAC	National Environmental Laboratory Accreditation Conference
NERL	National Exposure Research Laboratory
Ni	Nickel
NIOSH	National Institute for Occupational Safety and Health
NIST	National Institute for Standards and Technology
NRC	Nuclear Regulatory Commission
NSWC	Naval Surface Warfare Center
ORD	Office of Research and Development
OSWER	Office of Solid Waste and Emergency Response

Acronyms, Abbreviations, and Symbols (Continued)

P	Phosphorus
Pb	Lead
PC	Personal computer
PDA	Personal digital assistant
PCB	Polychlorinated biphenyls
Pd	Palladium
PE	Performance evaluation
PeT	Pentaerythritol
ppb	Parts per billion
ppm	Parts per million
Pu	Plutonium
QA	Quality assurance
QAPP	Quality assurance project plan
QC	Quality control
r ²	Correlation coefficient
RCRA	Resource Conservation and Recovery Act
Rh	Rhodium
RPD	Relative percent difference
RSD	Relative standard deviation
%RSD	Percent relative standard deviation
SAP	Sampling and analysis plan
SBMM	Sulphur Bank Mercury Mine
Sb	Antimony
Se	Selenium
Si	Silicon
SITE	Superfund Innovative Technology Evaluation
SOP	Standard operating procedure
SRM	Standard reference material
SVOC	Semivolatile organic compound
TAP	Thallium acid phthalate
Tetra Tech	Tetra Tech EM Inc.
Ti	Titanium
TSA	Technical systems audit
TSP	Total suspended particulates
TXRF	Total reflection x-ray fluorescence spectroscopy
U	Uranium
USFWS	U.S. Fish and Wildlife Service

Acronyms, Abbreviations, and Symbols (Continued)

V	Vanadium
V	Volts
VOC	Volatile organic compound
W	Watts
WDXRF	Wavelength-dispersive XRF
WRS	Wilcoxon Rank Sum
XRF	X-ray fluorescence
Zn	Zinc

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Chapter 1 Introduction

The U.S. Environmental Protection Agency (EPA), Office of Research and Development (ORD) conducted a demonstration to evaluate the performance of innovative x-ray fluorescence (XRF) technologies for measuring trace elements in soil and sediment. The demonstration was conducted as part of the EPA Superfund Innovative Technology Evaluation (SITE) Program.

Eight field-portable XRF instruments, which were provided and operated by six XRF technology developers, were evaluated as part of the demonstration. Each of these technology developers and their instruments are listed in Table 1-1. The technology developers brought each of these instruments to the demonstration site during the field

portion of the demonstration. The instruments were used to analyze a total of 326 prepared soil and sediment samples that contained 13 target elements. The same sample set was analyzed by a fixed laboratory (the reference laboratory) using established EPA reference methods. The results obtained using each XRF instrument in the field were compared with the results obtained by the reference laboratory to assess instrument accuracy. The results of replicate sample analysis were utilized to assess the precision and the detection limits that each XRF instrument could achieve. The results of these evaluations, as well as technical observations and cost information, were then documented in an Innovative Technology Verification Report (ITVR) for each instrument.

This ITVR documents EPA's evaluation of the Niton XLt 700 Series XRF analyzer based on the results of the demonstration.

1.1 Organization of this Report

This report is organized to first present general information pertinent to the demonstration. This information is common to all eight ITVRs that were developed from the XRF demonstration. Specifically, this information includes an introduction (Chapter 1), the locations where the field samples were collected (Chapter 2), the field demonstration (Chapter 3), the evaluation design (Chapter 4), and the reference laboratory results (Chapter 5).

The second part of this report provides information relevant to the specific instrument that is the subject of this ITVR. This information includes a description of the instrument (Chapter 6), a performance evaluation (Chapter 7), a cost analysis (Chapter 8), and a summary of the demonstration results (Chapter 9).

Developer Full Name	Distributor in the	Developer Short	Instrument Full	Instrument Short
	United States	Name	Name	Name
Elvatech, Ltd.	Xcalibur XRF Services	Xcalibur	ElvaX	ElvaX
Innov-X Systems	Innov-X Systems	Innov-X	XT400 Series	XT400
NITON Analyzers, A	NITON Analyzers, A	Niton	XLi 700 Series	XLi
Division of Thermo	Division of Thermo		XLt 700 Series	XLt
Electron Corporation	Electron Corporation			
Oxford Instruments	Oxford Instruments	Oxford	X-Met 3000 TX	X-Met
Analytical, Ltd.	Analytical, Ltd.		ED2000	ED2000
Rigaku, Inc.	Rigaku, Inc.	Rigaku	ZSX Mini II	ZSX Mini II
RÖNTEC AG	RÖNTEC USA	Rontec	PicoTAX	PicoTAX
(acquired by Bruker				
AXS, 11/2005)				

References are provided in Chapter 10. A verification statement for the instrument is provided as Appendix A. Comments from the instrument developer on the demonstration and any exceptions to EPA's evaluation are presented in Appendix B. Appendices C, D, and E contain the data validation summary report for the reference laboratory data and detailed evaluations of instrument versus reference laboratory results.

1.2 Description of the SITE Program

Performance verification of innovative environmental technologies is an integral part of EPA's regulatory and research mission. The SITE Program was established by the EPA Office of Solid Waste and Emergency Response and ORD under the Superfund Amendments and Reauthorization Act of 1986. The overall goal of the SITE Program is to conduct performance verification studies and to promote acceptance of innovative technologies that may be used to achieve long-term protection of human health and the environment. The program is designed to meet three primary objectives: (1) identify and remove obstacles to development and commercial use of innovative technologies; (2) demonstrate promising innovative technologies and gather reliable information on performance and cost to support site characterization and cleanup; and (3) maintain an outreach program to operate existing technologies and identify new opportunities for their use. Additional information on the SITE Program is available on the EPA ORD web site (www.epa.gov/ord/SITE).

The intent of a SITE demonstration is to obtain representative, high-quality data on the performance and cost of one or more innovative technologies so that potential users can assess a technology's suitability for a specific application. The SITE Program includes the following program elements:

 Monitoring and Measurement Technology (MMT) Program – Evaluates technologies that sample, detect, monitor, or measure hazardous and toxic substances. These technologies are expected to provide better, faster, or more costeffective methods for producing real-time data during site characterization and remediation studies than can conventional technologies.

- Remediation Technology Program Demonstrates innovative treatment technologies to provide reliable data on performance, cost, and applicability for site cleanups.
- Technology Transfer Program Provides and disseminates technical information in the form of updates, brochures, and other publications that promote the SITE Program and the participating technologies.

The demonstration of XRF instruments was conducted as part of the MMT Program, which is administered by the Environmental Sciences Division (ESD) of the National Exposure Research Laboratory (NERL) in Las Vegas, Nevada. Additional information on the NERL ESD is available on the EPA web site (www.epa.gov/nerlesd1/). Tetra Tech EM Inc. (Tetra Tech), an EPA contractor, provided comprehensive technical support to the demonstration.

1.3 Scope of the Demonstration

Conventional analytical methods for measuring the concentrations of inorganic elements in soil and sediment are time-consuming and costly. For this reason, field-portable XRF instruments have been proposed as an alternative approach, particularly where rapid and cost-effective assessment of a site is a goal. The use of a field XRF instrument for elemental analysis allows field personnel to quickly assess the extent of contamination by target elements at a site. Furthermore, the near instantaneous data provided by field-portable XRF instruments can be used to quickly identify areas where there may be increased risks and allow development of a more focused and cost-effective sampling strategy for conventional laboratory analysis.

EPA-sponsored demonstrations of XRF technologies have been under way for more than a decade. The first SITE MMT demonstration of XRF occurred in 1995, when six instruments were evaluated for their ability to analyze 10 target elements. The results of this demonstration were published in individual reports for each instrument (EPA 1996a, 1996b, 1998a, 1998b, 1998c, and 1998d). In 2003, two XRF instruments were included in a demonstration of field methods for analysis of mercury in soil and sediment. Individual ITVRs were also prepared for each of these two instruments (EPA 2004a, 2004b).

Although XRF spectrometry is now considered a mature technology for elemental analysis, fieldportable XRF instruments have evolved considerably over the past 10 years, and many of the instruments that were evaluated in the original demonstration are no longer manufactured. Advances in electronics and data processing, coupled with new x-ray tube source technology, have produced substantial improvements in the precision and speed of XRF analysis. The current demonstration of XRF instruments was intended to evaluate these new technologies, with an expanded set of target elements, to provide information to potential users on current state-of-theart instrumentation and its associated capabilities.

During the demonstration, performance data regarding each field-portable XRF instrument were collected through analysis of a sample set that included a broad range of soil/sediment types and target element concentrations. To develop this sample set, soil and sediment samples that contain the target elements of concern were collected in bulk quantities at nine sites from across the U.S. These bulk samples of soil and sediment were homogenized, characterized, and packaged into demonstration samples for the evaluation. Some of the batches of soil and sediment were spiked with selected target elements to ensure that representative concentration ranges were included for all target elements and that the sample design was robust. Replicate samples of the material in each batch were included in the final set of demonstration samples to assess instrument precision and detection limits. The final demonstration sample set therefore included 326 samples.

Each developer analyzed all 326 samples during the field demonstration using its XRF instrument and in accordance with its standard operating procedure. The field demonstration was conducted during the week of January 24, 2005, at the Kennedy Athletic, Recreational and Social (KARS) Park, which is part of the Kennedy Space Center on Merritt Island, Florida. Observers were assigned to each XRF instrument during the field demonstration to collect detailed information on the instrument and operating procedures, including sample processing times, for subsequent evaluation. The reference laboratory also analyzed a complete set of the demonstration samples for the target elements using acid digestion and inductively coupled plasma-atomic emission spectrometry (ICP-AES), in accordance with EPA Method 3050B/6010B, and using cold vapor atomic absorption (CVAA) spectroscopy (for mercury only) in accordance with EPA Method 7471A. By assuming that the results from the reference laboratory were essentially "true" values, instrument accuracy was assessed by comparing the results obtained using the XRF instrument with the results from the reference laboratory. The data obtained using the XRF instrument were also assessed in other ways, in accordance with the objectives of the demonstration, to provide information on instrument precision, detection limits, and interferences.

1.4 General Description of XRF Technology

XRF spectroscopy is an analytical technique that exposes a solid sample to an x-ray source. The xrays from the source have the appropriate excitation energy that causes elements in the sample to emit characteristic x-rays. A qualitative elemental analysis is possible from the characteristic energy, or wavelength, of the fluorescent x-rays emitted. A quantitative elemental analysis is possible by counting the number (intensity) of x-rays at a given wavelength.

Three electron shells are generally involved in emissions of x-rays during XRF analysis of samples: the K, L, and M shells. Multiple-intensity peaks are generated from the K, L, or M shell electrons in a typical emission pattern, also called an emission spectrum, for a given element. Most XRF analysis focuses on the x-ray emissions from the K and L shells because they are the most energetic lines. K lines are typically used for elements with atomic numbers from 11 to 46 (sodium to palladium), and L lines are used for elements above atomic number 47 (silver). M-shell emissions are measurable only for metals with an atomic number greater than 57 (lanthanum).

As illustrated in Figure 1-1, characteristic radiation arises when the energy from the x-ray source exceeds the absorption edge energy of inner-shell electrons, ejecting one or more electrons. The vacancies are filled by electrons that cascade in from the outer shells. The energy states of the electrons in the outer shells are higher than those of the inner-shell electrons, and the outer-shell electrons emit energy in the form of x-rays as they cascade down. The energy of this x-ray radiation is unique for each element.

An XRF analyzer consists of three major components: (1) a source that generates x-rays (a radioisotope or x-ray tube); (2) a detector that converts x-rays emitted from the sample into measurable electronic signals; and (3) a data processing unit that records the emission or fluorescence energy signals and calculates the elemental concentrations in the sample.

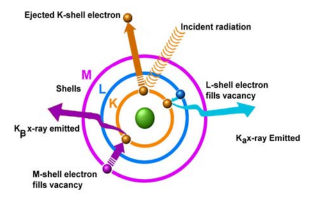


Figure 1-1. The XRF process.

Measurement times vary (typically ranging from 30 to 600 seconds), based primarily on data quality objectives. Shorter analytical measurement times (30 seconds) are generally used for initial screening, element identification, and hot-spot delineation, while longer measurement times (300 seconds or more) are typically used to meet higher goals for precision and accuracy. The length of the measuring time will also affect the detection limit; generally, the longer the measuring time, the lower the detection limit. However, detection limits for individual elements may become worse because of sample heterogeneity or the presence of other elements in the sample that fluoresce with similar x-ray energies.

The main variables that affect precision and accuracy for XRF analysis are:

- 1. Physical matrix effects (variations in the physical character of the sample).
- 2. Chemical matrix effects (absorption and enhancement phenomena)Spectral interferences (peak overlaps).
- 3. Moisture content above 10 percent, which affects x-ray transmission.

Because of these variables, it is important that each field XRF characterization effort be guided by a wellconsidered sampling and analysis plan. Sample preparation and homogenization, instrument calibration, and laboratory confirmation analysis are all important aspects of an XRF sampling and analysis plan. EPA SW-846 Method 6200 provides additional guidance on sampling and analytical methodology for XRF analysis.

1.5 Properties of the Target Elements

This section describes the target elements selected for the technology demonstration and the typical characteristics of each. Key criteria used in selecting the target elements included:

- The frequency that the element is determined in environmental applications of XRF instruments.
- The extent that the element poses an environmental consequence, such as a potential risk to human or environmental receptors.
- The ability of XRF technology to achieve detection limits below typical remediation goals and risk assessment criteria.
- The extent that the element may interfere with the analysis of other target elements.

In considering these criteria, the critical target elements selected for this study were antimony, arsenic, cadmium, chromium, copper, iron, lead, mercury, nickel, selenium, silver, vanadium, and zinc. These 13 target elements are of significant concern for site cleanups and human health risk assessments because most are highly toxic or interfere with the analysis of other elements. The demonstration therefore focused on the analysis of these 13 elements in evaluating the various XRF instruments.

1.5.1 Antimony

Naturally occurring antimony in surface soils is typically found at less than 1 to 4 milligrams per kilogram (mg/kg). Antimony is mobile in the environment and is bioavailable for uptake by plants; concentrations greater than 5 mg/kg are potentially phytotoxic, and concentrations above 31 mg/kg in soil may be hazardous to humans. Antimony may be found along with arsenic in mine wastes, at shooting ranges, and at industrial facilities. Typical detection limits for field-portable XRF instruments range from 10 to 40 mg/kg. Antimony is typically analyzed with success by ICP-AES; however, recovery of antimony in soil matrix spikes is often below quality control (QC) limits (50 percent or less) as a result of loss through volatilization during acid digestion. Therefore, results using ICP-AES may be lower than are obtained by XRF.

1.5.2 Arsenic

Naturally occurring arsenic in surface soils typically ranges from 1 to 50 mg/kg; concentrations above 10 mg/kg are potentially phytotoxic. Concentrations of arsenic greater than 0.39 mg/kg may cause carcinogenic effects in humans, and concentrations above 22 mg/kg may result in adverse noncarcinogenic effects. Typical detection limits for field-portable XRF instruments range from 10 to 20 mg/kg arsenic. Elevated concentrations of arsenic are associated with mine wastes and industrial facilities. Arsenic is successfully analyzed by ICP-AES; however, spectral interferences between peaks for arsenic and lead can affect detection limits and accuracy for XRF analysis when the ratio of lead to arsenic in 10 to 1 or more. Risk-based screening levels and soil screening levels for arsenic may be lower than the detection limits of field-portable XRF instruments.

1.5.3 Cadmium

Naturally occurring cadmium in surface soils typically ranges from 0.6 to 1.1 mg/kg; concentrations greater than 4 mg/kg are potentially phytotoxic. Concentrations of cadmium that exceed 37 mg/kg may result in adverse effects in humans. Typical detection limits for field-portable XRF instruments range from 10 to 50 mg/kg. Elevated concentrations of cadmium are associated with mine wastes and industrial facilities. Cadmium is successfully analyzed by both ICP-AES and fieldportable XRF; however, action levels for cadmium may be lower than the detection limits of fieldportable XRF instruments.

1.5.4 Chromium

Naturally occurring chromium in surface soils typically ranges from 1 to 1,000 mg/kg; concentrations greater than 1 mg/kg are potentially phytotoxic, although specific phytotoxicity levels for naturally occurring chromium have not been documented. The variable oxidation states of chromium affect its behavior and toxicity. Concentrations of hexavalent chromium above 30 mg/kg and of trivalent chromium above 10,000 mg/kg may cause adverse health effects in humans. Typical detection limits for field-portable XRF instruments range from 10 to 50 mg/kg. Hexavalent chromium is typically associated with metal plating or other industrial facilities. Trivalent chromium may be found in mine waste and at industrial facilities. Neither ICP-AES nor field-portable XRF can distinguish between oxidation states for chromium (or any other element).

1.5.5 Copper

Naturally occurring copper in surface soils typically ranges from 2 to 100 mg/kg; concentrations greater than 100 mg/kg are potentially phytotoxic. Concentrations greater than 3,100 mg/kg may result in adverse health effects in humans. Typical detection limits for field-portable XRF instruments range from 10 to 50 mg/kg. Copper is mobile and is a common contaminant in soil and sediments. Elevated concentrations of copper are associated with mine wastes and industrial facilities. Copper is successfully analyzed by ICP-AES and XRF; however, spectral interferences between peaks for copper and zinc may affect the detection limits and accuracy of the XRF analysis.

1.5.6 Iron

Although iron is not considered an element that poses a significant environmental consequence, it interferes with measurement of other elements and was; therefore, included in the study. Furthermore, iron is often used as a target reference element in XRF analysis.

Naturally occurring iron in surface soils typically ranges from 7,000 to 550,000 mg/kg, with the iron content originating primarily from parent rock. Typical detection limits for field-portable XRF instruments are in the range of 10 to 60 mg/kg. Iron is easily analyzed by both ICP-AES and XRF; however, neither technique can distinguish among iron species in soil. Although iron in soil may pose few environmental consequences, high levels of iron may interfere with analyses of other elements in both techniques (ICP-AES and XRF). Spectral interference from iron is mitigated in ICP-AES analysis by applying inter-element correction factors, as required by the analytical method. Differences in analytical results between ICP-AES and XRF for other target elements are expected when concentrations of iron are high in the soil matrix.

1.5.7 Lead

Naturally occurring lead in surface soils typically ranges from 2 to 200 mg/kg; concentrations greater than 50 mg/kg are potentially phytotoxic. Concentrations greater than 400 mg/kg may result in adverse effects in humans. Typical detection limits for field-portable XRF instruments range from 10 to 20 mg/kg. Lead is a common contaminant at many sites, and human and environmental exposure can occur through many routes. Lead is frequently found in mine waste, at lead-acid battery recycling facilities, at oil refineries, and in lead-based paint. Lead is successfully analyzed by ICP-AES and XRF; however, spectral interferences between peaks for lead and arsenic in XRF analysis can affect detection limits and accuracy when the ratio of arsenic to lead is 10 to 1 or more. Differences between ICP-AES and XRF results are expected in the presence of high concentrations of arsenic, especially when the ratio of lead to arsenic is low

1.5.8 Mercury

Naturally occurring mercury in surface soils typically ranges from 0.01 to 0.3 mg/kg; concentrations greater than 0.3 mg/kg are potentially phytotoxic.

Concentrations of mercury greater than 23 mg/kg and

concentrations of methyl mercury above 6.1 mg/kg may result in adverse health effects in humans. Typical detection limits for field-portable XRF instruments range from 10 to 20 mg/kg. Elevated concentrations of mercury are associated with amalgamation of gold and with mine waste and industrial facilities. Native surface soils are commonly enriched by anthropogenic sources of mercury. Anthropogenic sources include coal-fired power plants and metal smelters. Mercury is too volatile to withstand both the vigorous digestion and extreme temperature involved with ICP-AES analysis; therefore, the EPA-approved technique for laboratory analysis of mercury is CVAA spectroscopy. Mercury is successfully measured by XRF, but differences between results obtained by CVAA and XRF are expected when mercury levels are high.

1.5.9 Nickel

Naturally occurring nickel in surface soils typically ranges from 5 to 500 mg/kg; a concentration of 30 mg/kg is potentially phytotoxic. Concentrations greater than 1,600 mg/kg may result in adverse health effects in humans. Typical detection limits for fieldportable XRF instruments range from 10 to 60 mg/kg. Elevated concentrations of nickel are associated with mine wastes and industrial facilities. Nickel is a common environmental contaminant at metal processing sites. It is successfully analyzed by both ICP-AES and XRF with little interference; therefore, a strong correlation between the methods is expected.

1.5.10 Selenium

Naturally occurring selenium in surface soils typically ranges from 0.1 to 2 mg/kg; concentrations greater than 1 mg/kg are potentially phytotoxic. Its toxicities are well documented for plants and livestock; however, it is also considered a trace nutrient. Concentrations above 390 mg/kg may result in adverse health effects in humans. Typical detection limits for field-portable XRF instruments range from 10 to 20 mg/kg. Most selenium is associated with sulfur or sulfide minerals, where concentrations can exceed 200 mg/kg. Selenium can be measured by both ICP-AES and XRF; however, detection limits using XRF usually exceed the ecological risk-based screening levels for soil. Analytical results for selenium using ICP-AES and XRF are expected to be comparable.

1.5.11 Silver

Naturally occurring silver in surface soils typically ranges from 0.01 to 5 mg/kg; concentrations greater than 2 mg/kg are potentially phytotoxic. In addition, concentrations that exceed 390 mg/kg may result in adverse effects in humans. Typical detection limits for field-portable XRF instruments range from 10 to 45 mg/kg. Silver is a common contaminant in mine waste, in photographic film processing wastes, and at metal processing sites. It is successfully analyzed by ICP-AES and XRF; however, recovery may be reduced in ICP-AES analysis because insoluble silver chloride may form during acid digestion. Detection limits using XRF may exceed the risk-based screening levels for silver in soil.

1.5.12 Vanadium

Naturally occurring vanadium in surface soils typically ranges from 20 to 500 mg/kg; concentrations greater than 2 mg/kg are potentially phytotoxic, although specific phytotoxicity levels for naturally occurring vanadium have not been documented. Concentrations above 550 mg/kg may result in adverse health effects in humans. Typical detection limits for field-portable XRF instruments range from 10 to 50 mg/kg. Vanadium can be associated with manganese, potassium, and organic matter and is typically concentrated in organic shales, coal, and crude oil. It is successfully analyzed by both ICP-AES and XRF with little interference.

1.5.13 Zinc

Naturally occurring zinc in surface soils typically ranges from 10 to 300 mg/kg; concentrations greater than 50 mg/kg are potentially phytotoxic. Zinc at concentrations above 23,000 mg/kg may result in adverse health effects in humans. Typical detection limits for field-portable XRF instruments range from 10 to 30 mg/kg. Zinc is a common contaminant in mine waste and at metal processing sites. In addition, it is highly soluble, which is a common concern for aquatic receptors. Zinc is successfully analyzed by ICP-AES; however, spectral interferences between peaks for copper and zinc may influence detection limits and the accuracy of the XRF analysis. This page was left blank intentionally.

Chapter 2 Field Sample Collection Locations

Although the field demonstration took place at KARS Park on Merritt Island, Florida, environmental samples were collected at other sites around the country to develop a demonstration sample that incorporated a variety of soil/sediment types and target element concentrations. This chapter describes these sample collection sites, as well as the rationale for the selection of each.

Several criteria were used to assess potential sample collection sites, including:

- The ability to provide a variety of target elements and soil/sediment matrices.
- The convenience and accessibility of the location to the sampling team.
- Program support and the cooperation of the site owner.

Nine sample collection sites were ultimately selected for the demonstration; one was the KARS Park site itself. These nine sites were selected to represent variable soil textures (sand, silt, and clay) and iron content, two factors that significantly affect instrument performance.

Historical operations at these sites included mining, smelting, steel manufacturing, and open burn pits; one, KARS Park, was a gun range. Thus, these sites incorporated a wide variety of metal contaminants in soils and sediments. Both contaminated and uncontaminated (background) samples were collected at each site.

A summary of the sample collection sites is presented in Table 2-1, which describes the types of metalcontaminated soils or sediments that were found at each site. This information is based on the historical data that were provided by the site owners or by the EPA remedial project managers.

2.1 Alton Steel Mill Site

The Alton Steel Mill site (formerly the Laclede Steel site) is located at 5 Cut Street in Alton. Illinois. This 400-acre site is located in Alton's industrial corridor. The Alton site was operated by Laclede Steel Company from 1911 until it went bankrupt in July 2001. The site was purchased by Alton Steel, Inc., from the bankruptcy estate of Laclede Steel in May 2003. The Alton site is heir to numerous environmental concerns from more than 90 years of steel production; site contaminants include polychlorinated biphenyls (PCBs) and heavy metals. Laclede Steel was cited during its operating years for improper management and disposal of PCB wastes and electric arc furnace dust that contained heavy metals such as lead and cadmium. A Phase I environmental site assessment (ESA) was conducted at the Alton site in May 2002, which identified volatile organic compounds (VOCs), semivolatile organic compounds (SVOCs), total priority pollutant metals, and PCBs as potential contaminants of concern at the site.

Based on the data gathered during the Phase I ESA and on discussions with Alton personnel, several soil samples were collected for the demonstration from two areas at the Alton site, including the Rod Patenting Building and the Tube Mill Building. The soil in the areas around these two buildings had not been remediated and was known to contain elevated concentrations of arsenic, cadmium, chromium, lead, nickel, zinc, and iron. The matrix of the contaminated soil samples was a fine to medium sand; the background soil sample was a sand loam.

Table 2-2 presents historical analytical data (the maximum concentrations) for some of the target elements detected at the Alton site.

			Site-Specific Metals of Concern for XRF Demonstration											
Sample Collection Site	Source of Contamination	Matrix	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg	Ni	Se	Ag	Zn
	Steel manufacturing facility with metal arc													
	furnace dust. The site also includes a metal													
Alton Steel, Alton, IL	scrap yard and a slag recovery facility.	Soil		Х	Х	Х		Х	Х		Х			Χ
Burlington Northern-	Railroad yard staging area for smelter ores.													
ASARCO Smelter Site,	Contaminated soils resulted from dumping and													
East Helena, MT	spilling concentrated ores.	Soil		Х	Х				Х					
KARS Park – Kennedy														
Space Center, Merritt	Impacts to soil from historical facility													
Island, FL	operations and a former gun range.	Soil	Х	Х		Х	Х		Х					Х
	Abandoned open-pit sulfur and copper mine													
Leviathan Mine	that has contaminated a 9-mile stretch of													
Site/Aspen Creek, Alpine	mountain creeks, including Aspen Creek, with	Soil and												
County, CA	heavy metals.	Sediment		Х	Х	Х	Х	Х			Х			
Naval Surface Warfare	Open disposal and burning of general refuse													
Center, Crane Division,	and waste associated with aircraft													
Crane, IN	maintenance.	Soil	Х	Х	Х	Х	Х	Х	Х	Х	Х		Х	Х
	Silver Bow Creek was used as a conduit for													
Ramsay Flats–Silver Bow	mining, smelting, industrial, and municipal	Soil and												
Creek, Butte, MT	wastes.	Sediment		Х	Х		Х	Х	Х					Х
	Inactive mercury mine. Waste rock, tailings,													
Sulphur Bank Mercury	and ore are distributed in piles throughout the													
Mine	property.	Soil	Х	Х					Х	Х				
	Copper mining produced mill tailings that were													
Torch Lake Site (Great	dumped directly into Torch Lake,													
Lakes Area of Concern),	contaminating the lake sediments and													
Houghton County, MI	shoreline.	Sediment		Х		Х	Х		Х	Х		Х	Х	Х
	Abandoned smelter complex with	Seamont												
	contaminated soils and mineral-processing													
	wastes, including remnant ore piles,													
Wickes Smelter Site, decomposed roaster brick, slag piles and fines,														
Jefferson City, MT	and amalgamation sediments.	Soil	Х	Х	Х	Х	Х	Х	Х		X			Х
Notes (in order of appear		2011								1		l	1	
			DL.	т.	ad			0	1	Selei				
Sb: Antimony As: Arsenic	Cr: Chromium Cu: Copper		Pb:	Le	ad ercury				Se:	Selei				
As: Arsenic	Cu: Copper		Hg:		ercury				Ag:	Silve	1			

Table 2-1. Nature of Contamination in Soil and Sediment at Sample Collection Sites

Iron Note: Vanadium was not a chemical of concern at any of the sites and so does not appear on the table.

Fe:

Cd:

Cadmium

Ni:

Nickel

Zn:

Zinc

Metal	Maximum Concentration (mg/kg)
Arsenic	80.3
Cadmium	97
Chromium	1,551
Lead	3,556

Table 2-2. Historical Analytical Data, AltonSteel Mill Site

2.2 Burlington Northern-ASARCO Smelter Site

The Burlington Northern (BN)-ASARCO Smelter site is located in the southwestern part of East Helena, Montana. The site was an active smelter for more than 100 years and closed in 2002. Most of the ore processed at the smelter was delivered on railroad cars. An area west of the plant site (the BN property) was used for temporary staging of ore cars and consists of numerous side tracks to the primary railroad line into the smelter. This site was selected to be included in the demonstration because it had not been remediated and contained several target elements in soil.

At the request of EPA, the site owner collected samples of surface soil in this area in November 1997 and April 1998 and analyzed them for arsenic, cadmium, and lead; elevated concentrations were reported for all three metals. The site owner collected 24 samples of surface soil (16 in November 1997 and 8 in April 1998). The soils were found to contain up to 2,018 parts per million (ppm) arsenic, 876 ppm cadmium, and 43,907 ppm lead. One sample of contaminated soil and one sample of background soil were collected. The contaminated soil was a light brown sandy loam with low organic carbon content. The background soil was a medium brown sandy loam with slightly more organic material than the contaminated soil sample. Table 2-3 presents the site owner's data for arsenic, cadmium, and lead (the maximum concentrations) from the 1997 and 1998 sampling events.

Table 2-3. Historical Analytical Data, BN-
ASARCO Smelter Site

Metal	Maximum Concentration (ppm)
Arsenic	2,018
Cadmium	876
Lead	43,907

2.3 Kennedy Athletic, Recreational and Social Park Site

Soil and sediment at the KARS Park site were contaminated from former gun range operations and contain several target elements for the demonstration. The specific elements of concern for the KARS Park site include antimony, arsenic, chromium, copper, lead, and zinc.

The KARS Park site is located at the Kennedy Space Center on Merritt Island, Florida. KARS Park was purchased in 1962 and has been used by employees of the National Aeronautics and Space Administration (NASA), other civil servants, and guests as a recreational park since 1963. KARS Park occupies an area of Kennedy Space Center just outside the Cape Canaveral base. Contaminants in the park resulted from historical facility operations and impacts from the former gun range. The land north of KARS is owned by NASA and is managed by the U.S. Fish and Wildlife Service (USFWS) as part of the Merritt Island National Wildlife Refuge.

Two soil and two sediment samples were collected from various locations at the KARS Park site for the XRF demonstration. The contaminated soil sample was collected from an impact berm at the small arms range. The background soil sample was collected from a forested area near the gun range. The matrix of the contaminated and background soil samples consisted of fine to medium quartz sand. The sediment samples were collected from intermittently saturated areas within the skeet range. These samples were organic rich sandy loams. Table 2-4 presents historical analytical data (the maximum concentrations) for soil and sediment at KARS Park.

Table 2-4. Historical Analytical Data, KARS Park Site

Metal	Maximum Concentration (mg/kg)
Antimony	8,500
Arsenic	1,600
Chromium	40.2
Copper	290,000
Lead	99,000
Zinc	16,200

2.4 Leviathan Mine Site

The Leviathan Mine site is an abandoned copper and sulfur mine located high on the eastern slopes of the Sierra Nevada Mountain range near the California-Nevada border. Development of the Leviathan Mine began in 1863, when copper sulfate was mined for use in the silver refineries of the Comstock Lode. Later, the underground mine was operated as a copper mine until a mass of sulfur was encountered. Mining stopped until about 1935, when sulfur was extracted for use in refining copper ore. In the 1950s, the mine was converted to an open-pit sulfur mine. Placement of excavated overburden and waste rock in nearby streams created acid mine drainage and environmental impacts in the 1950s. Environmental impacts noted at that time included large fish kills.

Historical mining distributed waste rock around the mine site and created an open pit, adits, and solution cavities through mineralized rock. Oxygen in contact with the waste rock and mineralized rock in the adits oxidizes sulfur and sulfide minerals, generating acid. Water contacting the waste rock and flowing through the mineralized rock mobilizes the acid into the environment. The acid dissolves metals, including arsenic, copper, iron, and nickel, which creates conditions toxic to insects and fish in Leviathan. Aspen, and Bryant Creeks, downstream of the Leviathan Mine. Table 2-5 presents historical analytical data (the maximum concentrations) for the target elements detected at elevated concentrations in sediment samples collected along the three creeks. Four sediment and one soil sample were collected. One of the sediment samples was collected from the iron precipitate terraces formed from the acid mine drainage. The matrix of this sample appeared to be an orange silty clay loam. A second sediment sample was collected from the settling pond at the wastewater treatment system. The matrix of this sample was orange clay. A third sample was collected from the salt crust at the settling pond. This sample incorporated white crystalline material. One

background sediment and one background soil sample were collected upstream of the mine. These samples consisted of light brown sandy loam.

Table 2-5. Historical Analytical Data,Leviathan Mine Site

Metal	Maximum Concentration (mg/kg)			
Arsenic	2,510			
Cadmium	25.7			
Chromium	279			
Copper	837			
Nickel	2,670			

2.5 Navy Surface Warfare Center, Crane Division Site

The Old Burn Pit at the Naval Surface Warfare Center (NSWC), Crane Division, was selected to be included in the demonstration because 6 of the 13 target elements were detected at significant concentration in samples of surface soil previously collected at the site.

The NSWC, Crane Division, site is located near the City of Crane in south-central Indiana. The Old Burn Pit is located in the northwestern portion of NSWC and was used daily from 1942 to 1971 to burn refuse. Residue from the pit was buried along with noncombustible metallic items in a gully north of the pit. The burn pit was covered with gravel and currently serves as a parking lot for delivery trailers. The gully north of the former burn pit has been revegetated. Several soil samples were collected from the revegetated area for the demonstration because the highest concentrations of the target elements were detected in soil samples collected previously from this area. The matrix of the contaminated and background soil samples was a sandy loam. The maximum concentrations of the target elements detected in surface soil during previous investigations are summarized in Table 2-6.

Metal	Maximum Concentration (mg/kg)
Antimony	301
Arsenic	26.8
Cadmium	31.1
Chromium	112
Copper	1,520
Iron	105,000
Lead	16,900
Mercury	0.43
Nickel	62.6
Silver	7.5
Zinc	5,110

Table 2-6. Historical Analytical Data,NSWC Crane Division-Old Burn Pit

2.6 Ramsay Flats-Silver Bow Creek Site

The Ramsay Flats-Silver Bow Creek site was selected to be included in the demonstration because 6 of the 13 target elements were detected in samples of surface sediment collected previously at the site. Silver Bow Creek originates north of Butte, Montana, and is a tributary to the upper Clark Fork River.

More than 100 years of nearly continuous mining have altered the natural environment surrounding the upper Clark Fork River. Early wastes from mining, milling, and smelting were dumped directly into Silver Bow Creek and were subsequently transported downstream. EPA listed Silver Bow Creek and a contiguous portion of the upper Clark Fork River as a Superfund site in 1983.

A large volume of tailings was deposited in a lowgradient reach of Silver Bow Creek in the Ramsay Flats area. Tailings at Ramsay Flats extend several hundred feet north of the Silver Bow Creek channel. About 18 inches of silty tailings overlie texturally stratified natural sediments that consist of lowpermeability silt, silty clay, organic layers, and stringers of fine sand.

Two sediment samples were collected from the Ramsay Flats tailings area and were analyzed for a suite of metals using a field-portable XRF. The contaminated sediment sample was collected in Silver Bow Creek adjacent to the mine tailings. The matrix of this sediment sample was orange-brown silty fine sand with interlayered black organic material. The background sediment sample was collected upstream of Butte, Montana. The matrix of this sample was organic rich clayey silt with approximately 25 percent fine sand. The maximum concentrations of the target elements in the samples are summarized in Table 2-7.

Metal	Maximum Concentration (mg/kg)
Arsenic	176
Cadmium	141
Copper	1,110
Iron	20,891
Lead	394
Zinc	1,459

Table 2-7. Historical Analytical Data, Ramsay Flats-Silver Bow Creek Site

2.7 Sulphur Bank Mercury Mine Site

The Sulphur Bank Mercury Mine (SBMM) is a 160acre inactive mercury mine located on the eastern shore of the Oaks Arm of Clear Lake in Lake County, California, 100 miles north of San Francisco. Between 1864 and 1957, SBMM was the site of underground and open-pit mining at the hydrothermal vents and hot springs. Mining disturbed about 160 acres of land at SBMM and generated large quantities of waste rock (rock that did not contain economic concentrations of mercury and was removed to gain access to ore), tailings (the waste material from processes that removed the mercury from ore), and ore (rock that contained economic concentrations of mercury that was mined and stockpiled for mercury extraction). The waste rock, tailings, and ore are distributed in piles throughout the property.

Table 2-8 presents historical analytical data (the maximum concentrations) for the target elements detected at elevated concentrations in surface samples collected at SBMM. Two contaminated soil samples and one background soil sample were collected at various locations for the demonstration project. The mercury sample was collected from the ore stockpile and consisted of medium to coarse sand. The second contaminated soil sample was collected from the waste rock pile and consisted of coarse sand and gravel with trace silt. The matrix of the background soil sample was brown sandy loam.

Metal	Maximum Concentration (mg/kg)
Antimony	3,724
Arsenic	532
Lead	900
Mercury	4,296

Table 2-8. Historical Analytical Data, SulphurBank Mercury Mine Site

2.8 Torch Lake Superfund Site

The Torch Lake Superfund site was selected because native and contaminated sediment from copper mining, milling, and smelting contained the elements targeted for the demonstration. The specific metals of concern for the Torch Lake Superfund site included arsenic, chromium, copper, lead, mercury, selenium, silver, and zinc.

The Torch Lake Superfund site is located on the Keweenaw Peninsula in Houghton County, Michigan. Wastes were generated at the site from the 1890s until 1969. The site was included on the National Priorities List in June 1986. Approximately 200 million tons of mining wastes were dumped into Torch Lake and reportedly filled about 20 percent of the lake's original volume. Contaminated sediments are believed to be up to 70 feet thick in some locations. Wastes occur both on the uplands and in the lake and are found in four forms, including poor rock piles, slag and slag-enriched sediments, stamp sands, and abandoned settling ponds for mine slurry.

EPA initiated long-term monitoring of Torch Lake in 1999; the first monitoring event (the baseline study) was completed in August 2001. Table 2-9 presents analytical data (the maximum concentrations) for eight target elements in sediment samples collected from Torch Lake during the baseline study. Sediment samples were collected from the Torch Lake site at various locations for the demonstration. The matrix of the sediment samples was orange silt and clay.

Table 2-9. Historical Analytical Data, TorchLake Superfund Site

Metal	Maximum Concentration'(mg/kg)		
Arsenic	40		
Chromium	90		
Copper	5,850		
Lead	325		
Mercury	1.2		
Selenium	0.7		
Silver	6.2		
Zinc	630		

2.9 Wickes Smelter Site

The roaster slag pile at the Wickes Smelter site was selected to be included in the demonstration because 12 of the 13 target elements were detected in soil samples collected previously at the site.

The Wickes Smelter site is located in the unincorporated town of Wickes in Jefferson County, Montana. Wastes at the Wickes Smelter site include waste rock, slag, flue bricks, and amalgamation waste. The wastes are found in discrete piles and are mixed with soil. The contaminated soil sample was collected from a pile of roaster slag at the site. The slag was black, medium to coarse sand and gravel. The matrix of the background soil sample was a light brown sandy loam. Table 2-10 presents historical analytical data (maximum concentrations) for the roaster slag pile.

Table 2-10. Historical Analytical Data, WickesSmelter Site-Roaster Slag Pile

Metal	Maximum Concentration (mg/kg)
Antimony	79
Arsenic	3,182
Cadmium	70
Chromium	13
Copper	948
Iron	24,780
Lead	33,500
Nickel	7.3
Silver	83
Zinc	5,299

Chapter 3 Field Demonstration

The field demonstration required a sample set and a single location (the demonstration site) where all the technology developers could assemble to analyze the sample set under the oversight of the EPA/Tetra Tech field team. This chapter describes how the sample set was created, how the demonstration site was selected, and how the field demonstration was conducted. Additional detail regarding these topics is available in the *Demonstration and Quality Assurance Project Plan* (Tetra Tech 2005).

3.1 Bulk Sample Processing

A set of samples that incorporated a variety of soil and sediment types and target element concentrations was needed to conduct a robust evaluation. The demonstration sample set was generated from the bulk soil and sediment samples that were collected from the nine sample collection sites described in Chapter 2. Both contaminated (environmental) and uncontaminated (background) bulk samples of soil and sediment were collected at each sample collection site. The background sample was used as source material for a spiked sample when the contaminated sample did not contain the required levels of target elements. By incorporating a spiked background sample into the sample set, the general characteristics of the soil and sediment sample matrix could be maintained. At the same time, this spiked sample assured that all target elements were present at the highest concentration levels needed for a robust evaluation.

3.1.1 Bulk Sample Collection and Shipping

Large quantities of soil and sediment were needed for processing into well-characterized samples for this demonstration. As a result, 14 soil samples and 11 sediment samples were collected in bulk quantity from the nine sample collection sites across the U.S. A total of approximately 1,500 kilograms of unprocessed soil and sediment was collected, which yielded more than 1,000 kilograms of soil and sediment after the bulk samples had been dried.

Each bulk soil sample was excavated using clean

shovels and trowels and then placed into clean, plastic 5-gallon (19-liter) buckets at the sample collection site. The mass of soil and sediment in each bucket varied, but averaged about 25 kilograms per bucket. As a result, multiple buckets were needed to contain the entire quantity of each bulk sample.

Once it had been filled, a plastic lid was placed on each bucket, the lid was secured with tape, and the bucket was labeled with a unique bulk sample number. Sediment samples were collected in a similar method at all sites except at Torch Lake, where sediments were collected using a Vibracore or Ponar sediment sampler operated from a boat. Each 5-gallon bucket was overpacked in a plastic cooler and was shipped under chain of custody via overnight delivery to the characterization laboratory, Applied Research and Development Laboratory (ARDL).

3.1.2 Bulk Sample Preparation and Homogenization

Each bulk soil or sediment sample was removed from the multiple shipping buckets and then mixed and homogenized to create a uniform batch. Each bulk sample was then spread on a large tray at ARDL's laboratory to promote uniform air drying. Some bulk samples of sediment required more than 2 weeks to dry because of the high moisture content.

The air-dried bulk samples of soil and sediment were sieved through a custom-made screen to remove coarse material larger than about 1 inch. Next, each bulk sample was mechanically crushed using a hardened stainless-steel hammer mill until the particle size was sub-60-mesh sieve (less than 0.2 millimeters). The particle size of the processed bulk soil and sediment was measured after each round of crushing using standard sieve technology, and the particles that were still larger than 60-mesh were returned to the crushing process. The duration of the crushing process for each bulk sample varied based on soil type and volume of coarse fragments.

After each bulk sample had been sieved and crushed, the sample was mixed and homogenized using a

Model T 50A Turbula shaker-mixer. This shaker was capable of handling up to 50 gallons (190 liters) of sample material; thus, this shaker could handle the complete volume of each bulk sample. Bulk samples of smaller volume were mixed and homogenized using a Model T 10B Turbula shakermixer that was capable of handling up to 10 gallons (38 liters). Aliquots from each homogenized bulk sample were then sampled and analyzed in triplicate for the 13 target elements using ICP-AES and CVAA. If the relative percent difference between the highest and lowest result exceeded 10 percent for any element, the entire batch was returned to the shaker-mixer for additional homogenization. The entire processing scheme for the bulk samples is shown in Figure 3-1.

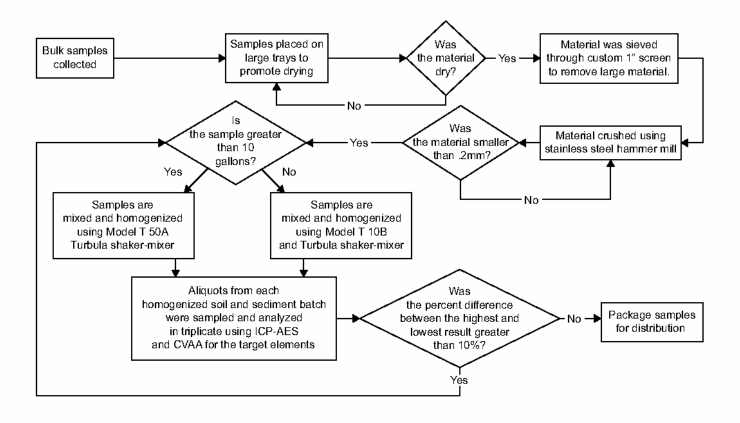


Figure 3-1. Bulk sample processing diagram.

3.2 Demonstration Samples

After the bulk soil and sediment sample material had been processed into homogenized bulk samples for the demonstration, the next consideration was the concentrations of target elements. The goal was to create a demonstration sample set that would cover the concentration range of each target element that may be reasonably found in the environment. Three concentration levels were identified as a basis for assessing both the coverage of the environmental samples and the need to generate spiked samples. These three levels were: (1) near the detection limit, (2) at intermediate concentrations, and (3) at high concentrations. A fourth concentration level (very high) was added for lead, iron, and zinc in soil and for iron in sediment. Table 3-1 lists the numerical ranges of the target elements for each of these levels (1 through 4).

3.2.1 Environmental Samples

A total of 25 separate environmental samples were collected from the nine sample collection sites described in Chapter 2. This bulk environmental sample set included 14 soil and 11 sediment samples. The concentrations of the target elements in some of these samples, however, were too high or too low to be used for the demonstration. Therefore, the initial analytical results for each bulk sample were used to establish different sample blends for each sampling location that would better cover the desired concentration ranges.

The 14 bulk soil samples were used to create 26 separate sample blends and the 11 bulk sediment samples were used to create 19 separate sample blends. Thus, there were 45 environmental sample blends in the final demonstration sample set. Either five or seven replicate samples of each sample blend were included in the sample set for analysis during the demonstration. Table 3-2 lists the number of sample blends and the number of demonstration samples (including replicates) that were derived from the bulk environmental samples for each sampling location.

3.2.2 Spiked Samples

Spiked samples that incorporated a soil and sediment matrix native to the sampling locations were created by adding known concentrations of target elements to the background samples. The spiked concentrations were selected to ensure that a minimum of three samples was available for all concentration levels for each target element.

After initial characterization at ARDL's laboratory, all bulk background soil and sediment samples were shipped to Environmental Research Associates (ERA) to create the spiked samples. The spiked elements were applied to the bulk sample in an aqueous solution, and then each bulk spiked sample was blended for uniformity and dried before it was repackaged in sample bottles.

Six bulk background soil samples were used at ERA's laboratory to create 12 separate spiked sample blends, and four bulk sediment samples were used to create 13 separate spiked sample blends. Thus, a total of 10 bulk background samples were used to create 25 spiked sample blends. Three or seven replicate samples of each spiked sample blend were included in the demonstration sample set. Table 3-3 lists the number of sample blends and the number of demonstration samples (including replicates) that were derived from the bulk background samples for each sampling location.

3.2.3 Demonstration Sample Set

In total, 70 separate blends of environmental and spiked samples were created and a set of 326 samples was developed for the demonstration by including three, five, or seven replicates of each blend in the final demonstration sample set. Thirteen sets of the demonstration samples, consisting of 326 individual samples in 250-milliliter clean plastic sample bottles, were prepared for shipment to the demonstration site and reference laboratory.

Analyte	Level 1 Target Range	Level 2 Target Range	Level 3 Target Range	Level 4 Target Range			
	(mg/kg)	(mg/kg)	(mg/kg)	(mg/kg)			
SOIL							
Antimony	40 - 400	400 - 2,000	>2,000				
Arsenic	20 - 400	400 - 2,000	>2,000				
Cadmium	50 - 500	500 - 2,500	>2,500				
Chromium	50 - 500	500 - 2,500	>2,500				
Copper	50 - 500	500 - 2,500	>2,500				
Iron	60 - 5,000	5,000 - 25,000	25,000 - 40,000	>40,000			
Lead	20 - 1,000	1,000 - 2,000	2,000 - 10,000	>10,000			
Mercury	20 - 200	200 - 1,000	>1,000				
Nickel	50 - 250	250 - 1,000	>1,000				
Selenium	20 - 100	100 - 200	>200				
Silver	45 - 90	90 - 180	>180				
Vanadium	50 - 100	100 - 200	>200				
Zinc	30 - 1,000	1,000 - 3,500	3,500 - 8,000	>8,000			
		SEDIMENT					
Antimony	40 - 250	250 - 750	>750				
Arsenic	20 - 250	250 - 750	>750				
Cadmium	50 - 250	250 - 750	>750				
Chromium	50 - 250	250 - 750	>750				
Copper	50 - 500	500 - 1,500	>1,500				
Iron	60 - 5,000	5,000 - 25,000	25,000 - 40,000	>40,000			
Lead	20 - 500	500 - 1,500	>1,500				
Mercury	20 - 200	200 - 500	>500				
Nickel	50 - 200	200 - 500	>500				
Selenium	20 - 100	100 - 200	>200				
Silver	45 - 90	90 - 180	>180				
Vanadium	50 - 100	100 - 200	>200				
Zinc	30 - 500	500 - 1,500	>1,500				

Sampling Location	Number of Sample Blends	Number of Demonstration Samples
Alton Steel Mill Site	2	10
Burlington Northern-ASARCO East Helena Site	5	29
Kennedy Athletic, Recreational and Social Park Site	6	32
Leviathan Mine Site	7	37
Naval Surface Warfare Center, Crane Division Site	1	5
Ramsay Flats—Silver Bow Creek Superfund Site	7	37
Sulphur Bank Mercury Mine Site	9	47
Torch Lake Superfund Site	3	19
Wickes Smelter Site	5	31
TOTAL *	45	247

 Table 3-2.
 Number of Environmental Sample Blends and Demonstration Samples

* Note: The totals in this table add to those for the spiked blends and replicates as summarized in Table 3-3 to bring the total number of blends to 70 and the total number of samples to 326 for the demonstration.

Sampling Location	Number of Spiked Sample Blends	Number of Demonstration Samples
Alton Steel Mill Site	1	3
Burlington Northern-ASARCO East Helena Site	2	6
Leviathan Mine Site	5	15
Naval Surface Warfare Center, Crane Division Site	2	6
Ramsey Flats—Silver Bow Creek Superfund Site	6	22
Sulphur Bank Mercury Mine Site	3	9
Torch Lake Superfund Site	4	12
Wickes Smelter Site	2	6
TOTAL *	25	79

Table 3-3. Number of Spiked Sample Blends and Demonstration Samples

* Note: The totals in this table add to those for the unspiked blends and replicates as summarized in Table 3-2 to bring the total number of blends to 70 and the total number of samples to 326 for the demonstration.

3.3 Demonstration Site and Logistics

The field demonstration occurred during the week of January 24, 2005. This section describes the selection of the demonstration site and the logistics of the field demonstration, including sample management.

3.3.1 Demonstration Site Selection

The demonstration site was selected from among the list of sample collection sites to simulate a likely field deployment. The following criteria were used to assess which of the nine sample collection sites might best serve as the demonstration site:

- Convenience and accessibility to participants in the demonstration.
- Ease of access to the site, with a reasonably sized airport that can accommodate the travel schedules for the participants.
- Program support and cooperation of the site owner.
- Sufficient space and power to support developer testing.
- Adequate conference room space to support a visitors day.
- A temperate climate so that the demonstration could occur on schedule in January.

After an extensive search for candidates, the site selected for the field demonstration was KARS Park, which is part of the Kennedy Space Center on Merritt Island, Florida. KARS Park was selected as the demonstration site for the following reasons:

- Access and Site Owner Support Representatives from NASA were willing to support the field demonstration by providing access to the site, assisting in logistical support during the demonstration, and hosting a visitors day.
- Facilities Requirements and Feasibility The recreation building was available and was of sufficient size to accommodate all the demonstration participants. Furthermore, the recreation building had adequate power to operate

all the XRF instruments simultaneously and all the amenities to fully support the demonstration participants, as well as visitors, in reasonable comfort.

• Ease of Access to the Site — The park, located about 45 minutes away from Orlando International Airport, was selected because of its easy accessibility by direct flight from many airports in the country. In addition, many hotels are located within 10 minutes of the site along the coast at Cocoa Beach, in a popular tourist area. Weather in this area of central Florida in January is dry and sunny, with pleasant daytime temperatures into the 70s (F) and cool nights.

3.3.2 Demonstration Site Logistics

The field demonstration was held in the recreation building, which is just south of the gunnery range at KARS Park. Photographs of the KARS Park recreation building, where all the XRF instruments were set up and operated, are shown in Figures 3-2 and 3-3.

A visitors day was held on January 26, 2005 when about 25 guests came to the site to hear about the demonstration and to observe the XRF instruments in operation. Visitors day presentations were conducted in a conference building adjacent to the recreation building at KARS Park (see Figure 3-4). Presentations by NASA and EPA representatives were followed by a tour of the XRF instruments in the recreation building while demonstration samples were being analyzed.



Figure 3-2. KARS Park recreation building.



Figure 3-3. Work areas for the XRF instruments in the recreation building.



Figure 3-4. Visitors day presentation.

3.3.3 EPA Demonstration Team and Developer Field Team Responsibilities

Each technology developer sent its instrument and a field team to the demonstration site for the week of January 24, 2005. The developer's field team was responsible for unpacking, setting up, calibrating, and operating the instrument. The developer's field team was also responsible for any sample preparation for analysis using the XRF instrument.

The EPA/Tetra Tech demonstration team assigned an observer to each instrument. The observer sat beside the developer's field team, or was nearby, throughout the field demonstration and observed all activities involved in setup and operation of the instrument. The observer's specific responsibilities included:

- Guiding the developer's field team to the work area in the recreation building at KARS Park and assisting with any logistical issues involved in instrument shipping, unpacking, and setup.
- Providing the demonstration sample set to the developer's field team in accordance with the sample management plan.
- Ensuring that the developer was operating the instrument in accordance with standard procedures and questioning any unusual practices or procedures.
- Communications with the developer's field team regarding schedules and fulfilling the requirements of the demonstration.
- Recording information relating to the secondary objectives of the evaluation (see Chapter 4) and for obtaining any cost information that could be provided by the developer's field team.
- Receiving the data reported by the developer's field team for the demonstration samples, and loading these data into a temporary database on a laptop computer.

Overall, the observer was responsible for assisting the developer's field team throughout the field demonstration and for recording all pertinent information and data for the evaluation. However, the observer was not allowed to advise the developer's field team on sample processing or to provide any feedback based on preliminary inspection of the XRF instrument data set.

3.3.4 Sample Management during the Field Demonstration

The developer's field team analyzed the demonstration sample set with its XRF instrument during the field demonstration. Each demonstration sample set was shipped to the demonstration site with only a reference number on each bottle as an identifier. The reference number was tied to the source information in the EPA/Tetra Tech database, but no information was provided on the sample label that might provide the developer's field team any insight as to the nature or content of the sample. Spiked samples were integrated with the environmental samples in a random manner so that the spiked samples could not be distinguished.

The demonstration sample set was divided into 13 subsets, or batches, for tracking during the field demonstration. The samples provided to each developer's field team were randomly distributed in two fashions. First, the order of the jars within each batch was random, so that the sample order for a batch was different for each developer's field team. Second, the distribution of sample batches was random, so that each developer's field team received the sample batches in a different order.

The observer provided the developer's field team with one batch of samples at a time. When the developer's field team reported that analysis of a batch was complete, the observer would reclaim all the unused sample material from that batch and then provide the next batch of samples for analysis. Chain-of-custody forms were used to document all sample transfers. When the analysis of all batches was complete, the observer assisted the developer's field team in cleanup of the work area and repackaging the instrument and any associated equipment. The members of the developer's field team were not allowed to take any part of the demonstration samples with them when they left the demonstration site.

Samples that were not in the possession of the developer's field team during the demonstration were held in a secure storage room adjacent to the demonstration work area (see Figure 3-5). The storage room was closed and locked except when the observer retrieved samples from the room. Samples were stored at room temperature during the demonstration, in accordance with the quality assurance/quality control (QA/QC) requirements established for the project.





3.3.5 Data Management

Each of the developer's field teams was able to complete analysis of all 326 samples during the field demonstration (or during the subsequent week, in one case when the developer's field team arrived late at the demonstration site because of delays in international travel). The data produced by each developer's field team were submitted during or at the end of the field demonstration in a standard Microsoft Excel[®] spreadsheet. (The EPA/Tetra Tech field team had provided a template.) Since each instrument provided data in a different format, the developer's field team was responsible for reducing the data before they were submitted and for transferring the data into the Excel spreadsheet.

The observer reviewed each data submittal for completeness, and the data were then uploaded into a master Excel spreadsheet on a laptop computer for temporary storage. Only the EPA/Tetra Tech field team had access to the master Excel spreadsheet during the field demonstration.

Once the EPA/Tetra Tech field team returned to their offices, the demonstration data were transferred to an Microsoft Access[®] database for permanent storage. Each developer's data, as they existed in the Access database, were then provided to the developer for review. Any errors the developers identified were corrected, and the database was then finalized. All statistical analysis and data evaluation took place on this final database.

Chapter 4 Evaluation Design

This chapter presents the approach for evaluating the performance of the XRF instruments. Specifically, the sections below describe the objectives of the evaluation and the experimental design.

The *Demonstration and Quality Assurance Project Plan* (Tetra Tech 2005) provides additional details on the overall demonstration approach. However, some deviations from the plan, involving data evaluation and laboratory audits, occurred after the demonstration plan was written. For completeness, the primary changes to the written plan are documented in the final section of this chapter.

4.1 Evaluation Objectives

The overall purpose of the XRF technology demonstration was to evaluate the performance of various field XRF instruments in detecting and quantifying trace elements in soils and sediments from a variety of sites around the U.S. The performance of each XRF instrument was evaluated in accordance with primary and secondary objectives. Primary objectives are critical to the evaluation and require the use of quantitative results to draw conclusions about an instrument's performance. Secondary objectives pertain to information that is useful but that will not necessarily require use of quantitative results to draw conclusions about an instrument's performance.

The primary and secondary objectives for the evaluation are listed in Table 4-1. These objectives were based on:

- Input from MMT Program stakeholders, including developers and EPA staff.
- General expectations of users of field measurement instruments.
- The time available to complete the demonstration.

• The capabilities of the instruments that the developers participating in the demonstration intended to highlight.

4.2 Experimental Design

To address the first four primary objectives, each XRF instrument analyzed the demonstration sample set for the 13 target elements. The demonstration samples originated from multiple sampling locations across the country, as described in Chapter 2, to provide a diverse set of soil and sediment matrices. The demonstration sample set included both blended environmental samples and spiked background samples, as described in Chapter 3, to provide a wide range of concentrations and combinations of elements.

When the field demonstration was completed, the results obtained using the XRF instruments were compared with data from a reference laboratory to evaluate the performance of each instrument in terms of accuracy and comparability (Primary Objective 2). The results for replicate samples were used to evaluate precision in various concentration ranges (Primary Objective 3) and the method detection limits (MDL) (Primary Objective 1). Each of these quantitative evaluations of instrument performance was carried out for each target element. The effect of chemical and spectral interferences and of soil characteristics (Primary Objectives 4 and 5) were evaluated to help explain extreme deviations or outliers observed in the XRF results when compared with the reference laboratory results.

A second important comparison involved the average performance of all eight XRF instruments that participated in the demonstration. For the first three primary objectives (MDL, accuracy, precision), the performance of each individual instrument was compared to the overall average performance of all eight instruments. Where the result of the instrument under consideration was less than 10 percent different from the corresponding result for all eight instruments, the result was considered "equivalent." A similar comparison was conducted with respect to cost (Primary Objective 7). These comparisons were intended to illustrate the performance of each XRF instrument in relation to its peers.

The evaluation design for meeting each objective, including data analysis procedures, is discussed in more detail in the sections below. Where specific deviations from these procedures were necessary for the data set associated with specific instruments, these deviations are described as part of the performance evaluation in Chapter 7.

4.2.1 Primary Objective 1 — Method Detection Limits

The MDL for each target element was evaluated based on the analysis of sets of seven replicate

samples that contained the target element at concentrations near the detection limit. The MDL was calculated using the procedures found in Title 40 Code of Federal Regulations (CFR) Part 136, Appendix B, Revision 1.11. The following equation was used:

$$MDL = t_{(n-1,1-\alpha=0.99)}(s)$$

where

MDL	= method detection limit
t	= Student's t value for a 99 percent
	confidence level and a standard
	deviation estimate with <i>n</i> -1 degrees
	of freedom
n	= number of samples
S	= standard deviation.

Table 4-1. Evaluation Objectives

Objective	Description
Primary Objective 1	Determine the MDL for each target element.
Primary Objective 2	Evaluate the accuracy and comparability of the XRF measurement to the results of laboratory reference methods for a variety of contaminated soil and sediment samples.
Primary Objective 3	Evaluate the precision of XRF measurements for a variety of contaminated soil and sediment samples.
Primary Objective 4	Evaluate the effect of chemical and spectral interference on measurement of target elements.
Primary Objective 5	Evaluate the effect of soil characteristics on measurement of target elements.
Primary Objective 6	Measure sample throughput for the measurement of target elements under field conditions.
Primary Objective 7	Estimate the costs associated with XRF field measurements.
Secondary Objective 1	Document the skills and training required to properly operate the instrument.
Secondary Objective 2	Document health and safety concerns associated with operating the instrument.
Secondary Objective 3	Document the portability of the instrument.
Secondary Objective 4	Evaluate the instrument's durability based on its materials of construction and engineering design.
Secondary Objective 5	Document the availability of the instrument and of associated customer technical support.

Based on the data provided by the characterization laboratory before the demonstration, a total of 12 sample blends (seven for soil and five for sediment) were identified for use in the MDL determination.

The demonstration approach specified the analysis of seven replicates for each of these sample blends by both the developer and the reference laboratory. It was predicted that these blends would allow the determination of a minimum of one MDL for soil and one MDL for sediment for each element, with the exception of iron. This prediction was based on the number of sample blends that contained concentrations less than 50 percent lower or higher than the lower limit of the Level 1 concentration range (from 20 to 50 ppm, depending on the element), as presented in Table 3-1.

After the field demonstration, the data sets obtained by the developers and the reference laboratory for the MDL sample blends were reviewed to confirm that they were appropriate to use in calculating MDLs. The requirements of 40 CFR 136, Appendix B, were used as the basis for this evaluation. Specifically, the CFR states that samples to be used for MDL determinations should contain concentrations in the range of 1 to 5 times the predicted MDL. On this basis, and using a nominal predicted reporting limit of 50 ppm for the target elements based on past XRF performance and developer information, a concentration of 250 ppm (5 times the "predicted" nominal MDL) was used as a threshold in selecting samples to calculate the MDL. Thus, each of the 12 MDL blends that contained mean reference laboratory concentrations less than 250 ppm were used in calculating MDLs for a given target element. Blends with mean reference laboratory concentrations greater than 250 ppm were discarded for evaluating this objective.

For each target element, an MDL was calculated for each sample blend with a mean concentration within the prescribed range. If multiple MDLs could be calculated for an element from different sample blends, these results were averaged to arrive at an overall mean MDL for the demonstration. The mean MDL for each target element was then categorized as either low (MDL less than 20 ppm), medium (MDL between 20 and 100 ppm), or high (MDL exceeds 100 ppm). No blends were available to calculate a detection limit for iron because all the blends contained substantial native concentrations of iron.

4.2.2 Primary Objective 2 — Accuracy

Accuracy was assessed based on a comparison of the results obtained by the XRF instrument with the results from the reference laboratory for each of the 70 blends in the demonstration sample set. The results from the reference laboratory were essentially used as a benchmark in this comparison, and the accuracy of the XRF instrument results was judged against them. The limitations of this approach should be recognized, however, because the reference laboratory results were not actually "true values." Still, there was a high degree of confidence in the reference laboratory results for most elements, as described in Chapter 5.

The following data analysis procedure was followed for each of the 13 target elements to assess the accuracy of an XRF instrument:

- 1. The results for replicate samples within a blend were averaged for both the data from the XRF instrument and the reference laboratory. Since there were 70 sample blends, this step created a maximum of 70 paired results for the assessment.
- 2. A blend that exhibited one or more non-detect values in either the XRF instrument or the reference laboratory analysis was excluded from the evaluation.
- 3. A blend was excluded from the evaluation when the average result from the reference laboratory was below a minimum concentration. The minimum concentration for exclusion from the accuracy assessment was identified as the lower limit of the lowest concentration range (Level 1 in Table 3-1), which is about 50 ppm for most elements.
- 4. The mean result for a blend obtained with the XRF instrument was compared with the corresponding mean result from the reference laboratory by calculating a relative percent difference (RPD). This comparison was carried out for each of the paired XRF and reference laboratory results included in the evaluation (up to 70 pairs) as follows:

$$\frac{(M_{\rm R} - M_{\rm D})}{\text{average } (M_{\rm R}, M_{\rm D})}$$

where

 M_R = the mean reference laboratory measurement M_D = the mean XRF instrument measurement.

- 5. Steps 1 through 4 provided a set of up to 70 RPDs for each element (70 sample blends minus the number excluded in steps 1 and 2). The absolute value of each of the RPDs was taken and summary statistics (minimum, maximum, mean and median) were then calculated.
- The accuracy of the XRF instrument for each target element was then categorized, based on the median of the absolute values of the RPDs, as either excellent (RPD less than 10 percent), good (RPD between 10 percent and 25 percent), fair (RPD between 25 percent and 50 percent), or poor (RPD above 50 percent).
- The set of absolute values of the RPDs for each instrument and element was further evaluated to assess any trends in accuracy versus concentration. These evaluations involved grouping the RPDs by concentration range (Levels 1 through 3 and 4, as presented in Table 3-1), preparing summary statistics for each range, and assessing differences among the grouped RPDs.

The absolute value of the RPDs was taken in step 5 to provide a more sensitive indicator of the extent of differences between the results from the XRF instrument and the reference laboratory. However, the absolute value of the RPDs does not indicate the direction of the difference and; therefore, does not reflect bias.

The populations of mean XRF and mean reference laboratory results were assessed through linear correlation plots to evaluate bias. These plots depict the linear relationships between the results for the XRF instrument and reference laboratory for each target element using a linear regression calculation with an associated correlation coefficient (r^2). These plots were used to evaluate the existence of general bias between the data sets for the XRF instrument and the reference laboratory.

4.2.3 Primary Objective 3 — Precision

The precision of the XRF instrument analysis for each target element was evaluated by comparing the results for the replicate samples in each blend. All 70 blends in the demonstration sample set (including environmental and spiked samples) were included in at least triplicate so that precision could be evaluated across all concentration ranges and across different matrices.

The precision of the data for a target element was evaluated for each blend by calculating the mean relative standard deviation (RSD) with the following equation:

$$RSD = \left|\frac{SD}{\overline{C}}\right| x \ 100$$

where

 $\begin{array}{ll} \text{RSD} &= \text{Relative standard deviation} \\ \text{SD} &= \text{Standard deviation} \\ \overline{C} &= \text{Mean concentration.} \end{array}$

1

The standard deviation was calculated using the equation:

$$SD = \left[\frac{1}{n-1}\sum_{k=1}^{n} \left(C_k - \overline{C}\right)^2\right]^{\frac{1}{2}}$$

where

SD	= Standard deviation
n	= Number of replicate
	samples
$\mathbf{C}_{\mathbf{k}}$	= Concentration of sample K
\overline{C}	= Mean concentration.

The following specific procedure for data analysis was followed for each of the 13 target elements to assess XRF instrument precision:

1. The RSD for the replicate samples in a blend was calculated for both data from the XRF instrument and the reference laboratory. Since there were 70 sample blends, this step created a maximum of 70 paired RSDs for the assessment.

- 2. A blend that exhibited one or more non-detect values in either the XRF or the reference laboratory analysis was excluded from the evaluation.
- 3. A blend was excluded from the evaluation when the average result from the reference laboratory was below a minimum concentration. The minimum concentration for exclusion from the precision assessment was identified as the lower limit of the lowest concentration range (Level 1 in Table 3-1), which was about 50 ppm for most elements.
- 4. The RSDs for the various blends for both the XRF instrument and the reference laboratory were treated as a statistical population. Summary statistics (minimum, maximum, mean and median) were then calculated and compared for the data set as a whole and for the different concentration ranges (Levels 1 through 3 or 4).
- The precision of the XRF instrument for each target element was then categorized, based on the median RSDs, as either excellent (RSD less than 5 percent), good (RSD between 5 percent and 10 percent), fair (RSD between 10 percent and 20 percent), or poor (RSD above 20 percent).

One primary evaluation was a comparison of the mean RSD for each target element between the XRF instrument and the reference laboratory. Using this comparison, the precision of the XRF instrument could be evaluated against the precision of accepted fixed-laboratory methods. Another primary evaluation was a comparison of the mean RSD for each target element between the XRF instrument and the overall average of all XRF instruments. Using this comparison, the precision of the XRF instrument could be evaluated against its peers.

4.2.4 Primary Objective 4 — Impact of Chemical and Spectral Interferences

The potential in the XRF analysis for spectral interference between adjacent elements on the periodic table was evaluated for the following element pairs: lead/arsenic, nickel/copper, and copper/zinc. The demonstration sample set included multiple blends where the concentration of one of these elements was greater than 10 times the concentration of the other element in the pair to facilitate this evaluation. Interference effects were identified through evaluation of the RPDs for these sample blends, which were calculated according to the equation in Section 4.2.2, since spectral interferences would occur only in the XRF data and not in the reference laboratory data.

Summary statistics for RPDs (mean, median, minimum, and maximum) were calculated for each potentially affected element for the sample blends with high relative concentrations (greater than 10 times) of the potentially interfering element. These summary statistics were compared with the RPD statistics for sample blends with lower concentrations of the interfering element. It was reasoned that spectral interference should be directly reflected in increased RPDs for the interference samples when compared with the rest of the demonstration sample set.

In addition to spectral interferences (caused by overlap of neighboring spectral peaks), the data sets were assessed for indications of chemical interferences. Chemical interferences occur when the x-rays characteristic of an element are absorbed or emitted by another element within the sample, causing low or high bias. These interferences are common in samples that contain high levels of iron, where low biases for copper and high biases for chromium can result. The evaluations for Primary Objective 4 therefore included RPD comparisons between sample blends with high concentrations of iron (more than 50,000 ppm) and other sample blends. These RPD comparisons were performed for the specific target elements of interest (copper, chromium, and others) to assess chemical interferences from iron. Outliers and subpopulations in the RPD data sets for specific target elements, as identified through graphical means (probability plots and box plots), were also examined for potential interference effects.

The software that is included with many XRF instruments can correct for chemical interferences. The results of this evaluation were intended to differentiate the instruments that incorporated effective software for addressing chemical interferences.

4.2.5 Primary Objective 5 — Effects of Soil Characteristics

The demonstration sample set included soil and sediment samples from nine locations across the U.S. and a corresponding variety of soil types and lithologies. The accuracy and precision statistics (RPD and RSD) were grouped by soil type (sample location) and the groups were compared to assess the effects of soil characteristics. Outliers and subpopulations in the RPD data sets, as identified through graphical means (correlation plots and box plots), were also examined for matrix effects.

4.2.6 Primary Objective 6 — Sample Throughput

Sample throughput is a calculation of the total number of samples that can be analyzed in a specified time. The primary factors that affect sample throughput are the time required to prepare a sample for analysis, to conduct the analytical procedure for each sample, and to process and tabulate the resulting data. The time required to prepare and to analyze demonstration samples was recorded each day that demonstration samples were analyzed.

Sample throughput can also be affected by the time required to set up and calibrate the instrument as well as the time required for quality control. The time required to perform these activities was also recorded during the field demonstration.

An overall mean processing time per sample and an overall sample throughput rate was calculated based on the total time required to complete the analysis of the demonstration sample set from initial instrument setup through data reporting. The overall mean processing time per sample was then used as the primary basis for comparative evaluations.

4.2.7 Primary Objective 7 — Technology Costs

The costs for analysis are an important factor in the evaluation and include the cost for the instrument, analytical supplies, and labor. The observer collected information on each of these costs during the field demonstration.

Based on input from each technology developer and from distributors, the instrument cost was established for purchase of the equipment and for daily, weekly, and monthly rental. Some of the technologies are not yet widely available, and the developer has not established rental options. In these cases, an estimated weekly rental cost was derived for the summary cost evaluations based on the purchase price for the instrument and typical rental to purchase price ratios for similar instruments. The costs associated with leasing agreements were also specified in the report, if available.

Analytical supplies include sample cups, spoons, xray film, Mylar[®], reagents, and personal protective equipment. The rate that the supplies are consumed was monitored and recorded during the field demonstration. The cost of analytical supplies was estimated per sample from these consumption data and information on unit costs.

Labor includes the time required to prepare and analyze the samples and to set up and dismantle the equipment. The labor hours associated with preparing and analyzing samples and with setting up and dismantling the equipment were recorded during the demonstration. The labor costs were calculated based on this information and typical labor rates for a skilled technician or chemist.

In addition to the assessment of the above-described individual cost components, an overall cost for a field effort similar to the demonstration was compiled and compared to the cost of fixed laboratory analysis. The results of the cost evaluation are presented in Chapter 8.

4.2.8 Secondary Objective 1 — Training Requirements

Each XRF instrument requires that the operator be trained to safely set up and operate the instrument. The relative level of education and experience that is appropriate to operate the XRF instrument was assessed during the field demonstration.

The amount of specific training required depends on the complexity of the instrument and the associated software. Most developers have established training programs. The time required to complete the developer's training program was estimated and the content of the training was identified.

4.2.9 Secondary Objective 2 — Health and Safety

The health and safety requirements for operation of the instrument were identified, including any that are associated with potential exposure from radiation and to reagents. Not included in the evaluation were potential risks from exposure to site-specific hazardous materials or physical safety hazards associated with the demonstration site.

4.2.10 Secondary Objective 3 — Portability

The portability of the instrument depends on size, weight, number of components, power requirements, and reagents required. The size of the instrument, including physical dimensions and weight, was recorded (see Chapter 6). The number of components, power requirements, support structures, and reagent requirements were also recorded. A qualitative assessment of portability was conducted based on this information.

4.2.11 Secondary Objective 4 — Durability

The durability of the instrument was evaluated by gathering information on the warranty and expected lifespan of the radioactive source or x-ray tube. The ability to upgrade software or hardware also was evaluated. Weather resistance was evaluated if the instrument is intended for use outdoors by examining the instrument for exposed electrical connections and openings that may allow water to penetrate.

4.2.12 Secondary Objective 5 — Availability

The availability of the instrument from the developer, distributors, and rental agencies was documented. The availability of replacement parts and instrumentspecific supplies was also noted.

4.3 Deviations from the Demonstration Plan

Although the field demonstration and subsequent data evaluations generally followed the *Demonstration and Quality Assurance Project Plan* (Tetra Tech 2005), there were some deviations as new information was uncovered or as the procedures were reassessed while the plan was executed. These deviations are documented below for completeness and as a supplement to the demonstration plan:

- 1. An in-process audit of the reference laboratory was originally planned while the laboratory was analyzing the demonstration samples. However, the reference laboratory completed all analysis earlier than expected, during the week of the field demonstration, and thereby created a schedule conflict. Furthermore, it was decided that the original pre-award audit was adequate for assessing the laboratory's procedures and competence.
- 2. The plan suggested that each result for spiked samples from the reference laboratory would be replaced by the "certified analysis" result, which was quantitative based on the amount of each element spiked, whenever the RPD between these two results was greater than 10 percent. The project team agreed that 10 percent was too stringent for this evaluation, however, and decided to use 25 percent RPD as the criterion for assessing reference laboratory accuracy against the spiked samples. Furthermore, it was found during the data evaluations that replacing individual reference laboratory results using this criterion would result in a mixed data set. Therefore, the 25 percent criterion was applied to the overall mean RPD for each element, and the "certified analysis" data set for a specific target element was used as a supplement to the reference laboratory result when this criterion was exceeded.
- 3. Instrument accuracy and comparability in relation to the reference laboratory (Primary Objective 2) was originally planned to be assessed based on a combination of percent recovery (instrument result divided by reference laboratory result) and RPD. It was decided during the data analysis, however, that the RPD was a much better parameter for this assessment. Specifically, it was found that the mean or median of the absolute values of the RPD for each blend was a good discriminator of instrument performance for this objective.
- 4. Although this step was not described in the plan, some quantitative results for each instrument were compared with the overall average of all XRF instruments. Since there were eight instruments, it was believed that a comparison of

this type did not violate EPA's agreement with the technology developers that one instrument would not be compared with another. Furthermore, this comparison provides an easyto-understand basis for assessing instrument performance.

5. The plan proposed statistical testing in support of Primary Objectives 4 and 5. Specifically, the Wilcoxon Rank Sum (WRS) test was proposed to assist in evaluating interference effects, and the Rosner outlier test was proposed in evaluating other matrix effects on XRF data quality (EPA 2000; Gilbert 1987). However, these statistical tests were not able to offer any substantive performance information over and above the evaluations based on RPDs and regression plots because of the limited sample numbers and scatter in the data. On this basis, the use of these two statistical tests was not further explored or presented.

Chapter 5 Reference Laboratory

As described in Chapter 4, a critical part of the evaluation was the comparison of the results obtained for the demonstration sample set by the XRF instrument with the results obtained by a fixed laboratory (the reference laboratory) using conventional analytical methods. Therefore, a significant effort was undertaken to ensure that data of the highest quality were obtained as the reference data for this demonstration. This effort included three main activities:

- Selection of the most appropriate methods for obtaining reference data,
- Selection of a high-quality reference laboratory, and
- Validation of reference laboratory data and evaluation of QA/QC results.

This chapter describes the information that confirms the validity, reliability, and usability of the reference laboratory data based on each of the three activities listed above (Sections 5.1, 5.2, and 5.3). Finally, this chapter presents conclusions (Section 5.4) on the level of data quality and the usability of the data obtained by the reference laboratory.

5.1 Selection of Reference Methods

Methods for analysis of elements in environmental samples, including soils and sediments, are well established in the environmental laboratory industry. Furthermore, analytical methods appropriate for soil and sediment samples have been promulgated by EPA in the compendium of methods, *Test Methods for Evaluating Solid Waste, Physical/Chemical Methods* (SW-846) (EPA 1996c). Therefore, the methods selected as reference methods for the demonstration were the SW-846 methods most typically applied by environmental laboratories to soil and sediment samples, as follows:

• Inductively coupled plasma-atomic emission spectroscopy (ICP-AES), in accordance with

EPA SW-846 Method 3050B/6010B, for all target elements except mercury.

 Cold vapor atomic absorption (CVAA) spectroscopy, in accordance with EPA SW-846 Method 7471A, for mercury only.

Selection of these analytical methods for the demonstration was supported by the following additional considerations: (1) the methods are widely available and widely used in current site characterizations, remedial investigations, risk assessments, and remedial actions; (2) substantial historical data are available for these methods to document that their accuracy and precision are adequate to meet the objectives of the demonstration; (3) these methods have been used extensively in other EPA investigations where confirmatory data were compared with XRF data; and (4) highly sensitive alternative methods were less suitable given the broad range of concentrations that were inherent in the demonstration sample set. Specific details on the selection of each method are presented below.

<u>Element Analysis by ICP-AES</u>. Method 6010B (ICP-AES) was selected for 12 of the target elements because its demonstrated accuracy and precision meet the requirements of the XRF demonstration in the most cost-effective manner. The ICP-AES method is available at most environmental laboratories and substantial data exist to support the claim that the method is both accurate and precise enough to meet the objectives of the demonstration.

Inductively coupled plasma-mass spectrometry (ICP-MS) was considered as a possible analytical technique; however, fewer data were available to support the claims of accuracy and precision. Furthermore, it was available in less than one-third of the laboratories solicited for this project. Finally, ICP-MS is a technique for analysis of trace elements and often requires serial dilutions to mitigate the effect of high concentrations of interfering ions or other matrix interferences. These dilutions can introduce the possibility of error and contaminants that might bias the results. Since the matrices (soil and sediment) for this demonstration are designed to contain high concentrations of elements and interfering ions, ICP-AES was selected over ICP-MS as the instrumental method best suited to meet the project objectives. The cost per analysis is also higher for ICP-MS in most cases than for ICP-AES.

Soil/Sediment Sample Preparation by Acid Digestion. The elements in soil and sediment samples must be dissolved from the matrix into an aqueous solution by acid digestion before analysis by ICP-AES. Method 3050B was selected as the preparation method and involves digestion of the matrix using a combination of nitric and hydrochloric acids, with the addition of hydrogen peroxide to assist in degrading organic matter in the samples. Method 3050B was selected as the reference preparation method because extensive data are available that suggest it efficiently dissolves most elements, as required for good overall recoveries and method accuracy. Furthermore, this method was selected over other digestion procedures because it is the most widely used dissolution method. In addition, it has been used extensively as the digestion procedure in EPA investigations where confirmatory data were compared with XRF data.

The ideal preparation reference method would completely digest silicaceous minerals. However, total digestion is difficult and expensive and is therefore seldom used in environmental analysis. More common strong acid-based extractions, like that used by EPA Method 3050B, recover most of the heavy element content. In addition, stronger and more vigorous digestions may produce two possible drawbacks: (1) loss of elements through volatilization, and (2) increased dissolution of interfering species, which may result in inaccurate concentration values.

Method 3052 (microwave-assisted digestion) was considered as an alternative to Method 3050B, but was not selected because it is not as readily available in environmental laboratories.

<u>Soil/Sediment Sample Preparation for Analysis of</u> <u>Mercury by CVAA</u>. Method 7471A (CVAA) is the only method approved by EPA and promulgated for analysis of mercury. Method 7471A includes its own digestion procedure because more vigorous digestion of samples, like that incorporated in Method 3050B, would volatilize mercury and produce inaccurate results. This technique is widely available, and extensive data are available that support the ability of this method to meet the objectives of the demonstration.

5.2 Selection of Reference Laboratory

The second critical step in ensuring high-quality reference data was selection of a reference laboratory with proven credentials and quality systems. The reference laboratory was procured via a competitive bid process. The procurement process involved three stages of selection: (1) a technical proposal, (2) an analysis of performance audit samples, and (3) an onsite laboratory technical systems audit (TSA). Each stage was evaluated by the project chemist and a procurement specialist.

In Stage 1, 12 analytical laboratories from across the U.S. were invited to bid by submitting extensive technical proposals. The technical proposals included:

- A current statement of qualifications.
- The laboratory quality assurance manual.
- Standard operating procedures (SOP) (including sample receipt, laboratory information management, sample preparation, and analysis of elements).
- Current instrument lists.
- Results of recent analysis of performance evaluation samples and audits.
- Method detection limit studies for the target elements.
- Professional references, laboratory personnel experience, and unit prices.

Nine of the 12 laboratories submitted formal written proposals. The proposals were scored based on technical merit and price, and a short list of five laboratories was identified. The scoring was weighed heavier for technical merit than for price. The five laboratories that received the highest score were advanced to stage 2. In stage 2, each of the laboratories was provided with a set of six samples to analyze. The samples consisted of three certified reference materials (one soil and two sediment samples) at custom spiking concentrations, as well as three pre-demonstration soil samples. The results received from each laboratory were reviewed and assessed. Scoring at this stage was based on precision (reproducibility of results for the three pre-demonstration samples), accuracy (comparison of results to certified values for the certified reference materials), and completeness of the data package (including the hard copy and electronic data deliverables). The two laboratories that received the highest score were advanced to stage 3.

In stage 3, the two candidate laboratories were subjected to a thorough on-site TSA by the project chemist. The audit consisted of a direct comparison of the technical proposal to the actual laboratory procedures and conditions. The audit also tracked the pre-demonstration samples through the laboratory processes from sample receipt to results reporting. When the audit was conducted, the project chemist verified sample preparation and analysis for the three pre-demonstration samples. Each laboratory was scored on identical checklists.

The reference laboratory was selected based on the highest overall score. The weights of the final scoring selection were as follows:

Scoring Element	Relative Importance
Audits (on site)	40%
Performance evaluation samples, including data package and electronic data deliverable	50%
Price	10%

Based on the results of the evaluation process, Shealy Environmental Services, Inc. (Shealy), of Cayce, South Carolina, received the highest score and was therefore selected as the reference laboratory. Shealy is accredited by the National Environmental Laboratory Accreditation Conference (NELAC). Once selected, Shealy analyzed all demonstration samples (both environmental and spiked samples) concurrently with the developers' analysis during the field demonstration. Shealy analyzed the samples by ICP-AES using EPA SW-846 Method 3050B/6010B and by CVAA using EPA SW-846 Method 7471A.

5.3 QA/QC Results for Reference Laboratory

All data and QC results from the reference laboratory were reviewed in detail to determine that the reference laboratory data were of sufficiently high quality for the evaluation. Data validation of all reference laboratory results was the primary review tool that established the level of quality for the data set (Section 5.3.1). Additional reviews included the on-site TSA (Section 5.3.2) and other evaluations (Section 5.3.3).

5.3.1 Reference Laboratory Data Validation

After all demonstration samples had been analyzed, reference data from Shealy were fully validated according to the EPA validation document, USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review (EPA 2004c) as required by the Demonstration and Quality Assurance Project Plan (Tetra Tech 2005). The reference laboratory measured 13 target elements, including antimony, arsenic, cadmium, chromium, copper, iron, lead, mercury, nickel, selenium, silver, vanadium, and zinc. The reference laboratory reported results for 22 elements at the request of EPA; however, only the data for the 13 target elements were validated and included in data comparisons for meeting project objectives. A complete summary of the validation findings for the reference laboratory data is presented in Appendix C.

In the data validation process, results for QC samples were reviewed for conformance with the acceptance criteria established in the demonstration plan. Based on the validation criteria specified in the demonstration plan, all reference laboratory data were declared valid (were not rejected). Thus, the completeness of the data set was 100 percent. Accuracy and precision goals were met for most of the QC samples, as were the criteria for comparability, representativeness, and sensitivity. Thus, all reference laboratory data were deemed usable for comparison to the data obtained by the XRF instruments. Only a small percentage of the reference laboratory data set was qualified as undetected as a result of blank contamination (3.3 percent) and estimated because of matrix spike and matrix spike duplicate (MS/MSD) recoveries (8.7 percent) and serial dilutions results (2.5 percent). Table 5.1 summarizes the number of validation qualifiers applied to the reference laboratory data according to QC type. Of the three QC types, only the MS/MSD recoveries warranted additional evaluation. The MS/MSD recoveries for antimony were marginally low (average recovery of 70.8 percent) when compared with the QC criterion of 75 to 125 percent recovery. It was concluded that low recoveries for antimony are common in analysis of soil and sediment by the prescribed methods and likely result from volatilization during the vigorous acid digestion process or spectral interferences found in soil and sediments matrices (or both). In comparison to antimony, high or low recoveries were observed only on an isolated basis for the other target metals (for example, lead and mercury) such that the mean and median percent recoveries were well within the required range. Therefore, the project team decided to evaluate the XRF data against the reference laboratory data for all 13 target elements and to evaluate the XRF data a second time against the ERA certified spike values for antimony only. These comparisons are discussed in Section 7.1. However, based on the validation of the complete reference data set and the low occurrence of qualified data, the reference laboratory data set as a whole was declared of high quality and of sufficient quality to make valid comparisons to XRF data.

5.3.2 Reference Laboratory Technical Systems Audit

The TSA of the Shealy laboratory was conducted by the project chemist on October 19, 2004, as part of the selection process for the reference laboratory. The audit included the review of element analysis practices (including sample preparation) for 12 elements by EPA Methods 3050B and 6010B and for total mercury by EPA Method 7471A. All decisionmaking personnel for Shealy were present during the TSA, including the laboratory director, QA officer, director of inorganics analysis, and the inorganics laboratory supervisor. Project-specific requirements were reviewed with the Shealy project team as were all the QA criteria and reporting requirements in the demonstration plan. It was specifically noted that the demonstration samples would be dried, ground, and sieved before they were submitted to the laboratory, and that the samples would be received with no preservation required (specifically, no chemical preservation and no ice). The results of the performance audit were also reviewed.

No findings or nonconformances that would adversely affect data quality were noted. Only two minor observations were noted; these related to the revision dates of two SOPs. Both observations were discussed at the debriefing meeting held at the laboratory after the TSA. Written responses to each of the observations were not required; however, the laboratory resolved these issues before the project was awarded. The auditor concluded that Shealy complied with the demonstration plan and its own SOPs, and that data generated at the laboratory should be of sufficient and known quality to be used as a reference for the XRF demonstration.

5.3.3 Other Reference Laboratory Data Evaluations

The data validation indicated that all results from the reference laboratory were valid and usable for comparison to XRF data, and the pre-demonstration TSA indicated that the laboratory could fully comply with the requirements of the demonstration plan for producing data of high quality. However, the reference laboratory data were evaluated in other ways to support the claim that reference laboratory data are of high quality. These evaluations included the (1) assessment of accuracy based on ERAcertified spike values, (2) assessment of precision based on replicate measurements within the same sample blend, and (3) comparison of reference laboratory data to the initial characterization data that was obtained when the blends were prepared. Each of these evaluations is briefly discussed in the following paragraphs.

Blends 46 through 70 of the demonstration sample set consisted of certified spiked samples that were used to assess the accuracy of the reference laboratory data. The summary statistics from samples with the reference laboratory results are shown in Table 5-2. The target for percent recovery was 75 to 125 percent. The mean percent recoveries for 12 of the 13 target elements were well within this accuracy goal. Only the mean recovery for antimony was outside the goal (26.8 percent). The low mean percent recovery for antimony supported the recommendation made by the project team to conduct a secondary comparison of XRF data to ERAcertified spike values for antimony. This secondary evaluation was intended to better understand the impacts on the evaluation of the low bias for antimony in the reference laboratory data. All other recoveries were acceptable. Thus, this evaluation

further supports the conclusion that the reference data set is of high quality.

All blends (1 through 70) were prepared and delivered with multiple replicates. To assess precision, percent RSDs were calculated for the replicate sample results submitted by the reference laboratory for each of the 70 blends. Table 5-3 presents the summary statistics for the reference laboratory data for each of the 13 target elements. These summary statistics indicate good precision in that the median percent RSD was less than 10 percent for 11 out of 13 target elements (and the median RSD for the other two elements was just above 10

	Nur	Number and Percentage of Qualified Results per QC type ¹												
	Metho	d Blank	MS	/MSD	Serial Dilution									
Element	Number	Percent ²	Number	Percent ²	Number	Percent ²								
Antimony	5	1.5	199	61.0	8	2.4								
Arsenic	12	3.7	3	0.9	10	3.1								
Cadmium	13	4.0	0	0	6	1.8								
Chromium	0	0	0	0	10	3.1								
Copper	1	0.3	0	0	8	2.4								
Iron	0	0	0	0	10	3.1								
Lead	0	0	34	10.5	11	3.4								
Mercury	68	20.9	31	9.5	4	1.2								
Nickel	0	0	0	0	10	3.1								
Selenium	16	4.9	0	0	3	0.9								
Silver	22	6.7	102	31.3	7	2.1								
Vanadium	0	0	0	0	9	2.8								
Zinc	1	0.3	0	0	10	3.1								
Totals	138	3.3	369	8.7	106	2.5								

Table 5-1. Number of Validation Qualifiers.

Notes:

MS Matrix spike.

MSD Matrix spike duplicate.

QC Quality control.

² Percents for individual elements are calculated based on 326 results per element. Total percents at the bottom of the table are calculated based on the total number of results for all elements (4,238).

This table presents the number of "U" (undetected) and "J" (estimated) qualifiers added to the reference laboratory data during data validation. Though so qualified, these results are considered usable for the demonstration. As is apparent in the "Totals" row at the bottom of this table, the amount of data that required qualifiers for any specific QC type was invariably less than 10 percent. No reference laboratory data were rejected (that is, qualified "R") during the data validation.

percent). Thus, this evaluation further supports the conclusion that the reference data set is of high quality.

ARDL, in Mount Vernon, Illinois, was selected as the characterization laboratory to prepare environmental samples for the demonstration. As part of its work, ARDL analyzed several samples of each blend to evaluate whether the concentrations of the target elements and the homogeneity of the blends were suitable for the demonstration. ARDL analyzed the samples using the same methods as the reference laboratory; however, the data from the characterization laboratory were not validated and were not intended to be equivalent to the reference laboratory data. Rather, the intent was to use the results obtained by the characterization laboratory as an additional quality control check on the results from the reference laboratory.

A review of the ARDL characterization data in comparison to the reference laboratory data indicated that ARDL obtained lower recoveries of several elements. When expressed as a percent of the average reference laboratory result (percent recovery), the median ARDL result was below the lower QC limit of 75 percent recovery for three elements — chromium, nickel, and selenium. This discrepancy between data from the reference laboratory and ARDL was determined to have no significant impact on reference laboratory data quality for three reasons: (1) the ARDL data were obtained on a rapid turnaround basis to evaluate homogeneity — accuracy was not a specific goal, (2) the ARDL data were not validated, and (3) all other quality measurement for the reference laboratory data indicated a high level of quality.

5.4 Summary of Data Quality and Usability

A significant effort was undertaken to ensure that data of high quality were obtained as the reference data for this demonstration. The reference laboratory data set was deemed valid, usable, and of high quality based on the following:

- Comprehensive selection process for the reference laboratory, with multiple levels of evaluation.
- No data were rejected during data validation and few data qualifiers were added.
- The observations noted during the reference laboratory audit were only minor in nature; no major findings or non-conformances were documented.
- Acceptable accuracy (except for antimony, as discussed in Section 5.3.3) of reference laboratory results in comparison to spiked certified values.
- Acceptable precision for the replicate samples in the demonstration sample set.

Based on the quality indications listed above, the reference laboratory data were used in the evaluation of XRF demonstration data. A second comparison was made between XRF data and certified values for antimony (in Blends 46 through 70) to address the low bias exhibited for antimony in the reference laboratory data.

Statistic	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg	Ni	Se	Ag	V	Zn
Number of %R values	16	14	20	12	20	NC	12	15	16	23	20	15	10
Minimum %R	12.0	65.3	78.3	75.3	51.7	NC	1.4	81.1	77.0	2.2	32.4	58.5	0.0
Maximum %R	36.1	113.3	112.8	108.6	134.3	NC	97.2	243.8	116.2	114.2	100.0	103.7	95.2
Mean %R ¹	26.8	88.7	90.0	94.3	92.1	NC	81.1	117.3	93.8	89.9	78.1	90.4	90.6
Median %R ¹	28.3	90.1	87.3	97.3	91.3	NC	88.0	93.3	91.7	93.3	84.4	95.0	91.3

Table 5-2. Percent Recovery for Reference Laboratory Results in Comparison to ERA Certified Spike Values for Blends 46 through 70

¹Values shown in bold fall outside the 75 to 125 percent acceptance criterion for percent recovery.

ERA = Environmental Resource Associates, Inc.

NC = Not calculated.

%R = Percent recovery.

Source of certified values: Environmental Resource Associates, Inc.

- Sb Antimony
- As Arsenic
- Cd Cadmium
- Cr Chromium
- Cu Copper
- Fe Iron
- Pb Lead
- Hg Mercury
- Ni Nickel
- Se Selenium
- Ag Silver
- V Vanadium
- Zn Zinc

Statistic	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg	Ni	Se	Ag	V	Zn
Number of %RSDs	43	69	43	69	70	70	69	62	68	35	44	69	70
Minimum %RSD	1.90	0.00	0.91	1.43	0.00	1.55	0.00	0.00	0.00	0.00	1.02	0.00	0.99
Maximum %RSD	78.99	139.85	40.95	136.99	45.73	46.22	150.03	152.59	44.88	37.30	54.21	43.52	48.68
Mean %RSD ¹	17.29	13.79	12.13	11.87	10.62	10.56	14.52	16.93	10.28	13.24	12.87	9.80	10.94
Median %RSD ¹	11.99	10.01	9.36	8.29	8.66	8.55	9.17	7.74	8.12	9.93	8.89	8.34	7.54

 Table 5-3. Precision of Reference Laboratory Results for Blends 1 through 70

¹Values shown in bold fall outside precision criterion of less than or equal to 25 %RSD.

%RSD = Percent relative standard deviation.

Based on the three to seven replicate samples included in Blends 1 through 70.

- Sb Antimony
- As Arsenic
- Cd Cadmium
- Cr Chromium
- Cu Copper
- Fe Iron
- Pb Lead
- Hg Mercury
- Ni Nickel
- Se Selenium
- Ag Silver
- V Vanadium
- Zn Zinc

Chapter 6 Technology Description

The XLt 700 Series XRF analyzer is manufactured by NITON Analyzers, a division of Thermo Electron Corporation (Niton). This chapter provides a technical description of the XLt based on information obtained from Niton and observation of this analyzer during the field demonstration. This section also identifies a Niton company contact, where additional technical information may be obtained.

6.1 General Description

The Niton XLt 700 Series XRF analyzer is a small, field-portable instrument designed for chemical characterization of soils, sediment, and other thick, homogeneous samples (plastics and metals). The analyzer features a miniaturized x-ray tube for the excitation source and a Peltier-cooled Si-PiN x-ray detector. The analyzer's standard software is programmed to analyze and automatically report 25 elements. The optional proprietary software package called PERFECT (for **P**rogrammable **E**xcitation by **R**egulation of **F**ilters, **E**nergy, **C**urrent, and **T**ime) provides additional capabilities for analyzing and reporting light elements (including vanadium and chromium) and optimizes the detection limits for a suite of elements specific to an application. Other features include an integrated touch-screen display; completely sealed housing to protect the analyzer from moisture and dust: lithium-ion batteries: an integrated bar code reader and virtual keypad; remote operation and custom report generation capability from a Windows-based PC; a shielded bench-top test stand: and Bluetooth wireless communication to a laptop or personal data assistant (PDA).

The XLt is factory calibrated to simultaneously analyze up to 25 elements, including all eight Resource Conservation and Recovery Act (RCRA) metals. The analyzer does not require a site- or material-specific calibration; however, it is capable of handling user-generated empirical calibrations, if required for specific applications.

The XLt is designed to be used either as a hand-held instrument for in situ analysis (Figure 6-1) or as a

bench-top instrument, in a test stand with a sample drawer below the instrument, for ex situ analysis. To analyze soil samples in the *in situ* mode, the instrument x-ray window is placed directly on the ground or on soils in a plastic bag; for *ex situ* analysis, samples are prepared in x-ray sample cups and are placed in the sample drawer at the bottom of the test stand, directly beneath the instrument x-ray window. In situ testing with the XLt allows for a semi-quantitative assessment of element concentrations at multiple locations or over large areas in a short time. Quantitative ex situ testing involves properly preparing the samples, placing the samples in x-ray sample cups, and analyzing them in a controlled area, typically free from dust and weather extremes. Most field applications incorporate a combination of in situ and ex situ testing.



Figure 6-1. In situ testing with the Niton XLt 700 Series analyzer.

The XLt can be used to analyze elements under three primary scenarios: (1) bulk sample mode (includes soils, sediments, and metal alloys); (2) thin film mode (includes dust wipes and filters); and (3) plastics mode. Two standard calibrations are provided under the bulk sample mode and include one for standard soils and one for industrial bulk (metal alloys). Additional user-defined calibrations can be programmed and used under the bulk sample mode. In the thin film mode, the XLt can be used to analyze thin samples (dust wipes) and other filter media used to capture airborne particulate matter.

XRF analysis using the XLt can comply fully with EPA Method 6200, "Field Portable XRF Spectrometry for the Determination of Elemental Concentrations in Soil and Sediment." Since XRF analysis is nondestructive, samples analyzed by XRF can be sent to a fixed analytical laboratory for confirmation of results.

The technical specifications for the XLt are presented in Table 6-1.

Niton has not published formal standard operating procedures for the XLt analyzer operations, but

recommends that users follow EPA Method 6200 and the instrument user's manual to ensure that the appropriate protocol is followed. The recommended steps include the following:

- (1) Insert the battery, turn on the system, and allow it to warm up for 15 minutes.
- (2) Ensure the date and time is correct.
- (3) Analyze the standard check samples (National Institute of Standards and Technology [NIST] 2709; NIST 2710; and blank) to ensure proper precision. Repeat this step every 4 to 6 hours, or after a battery exchange and reboot.
- (4) Download and delete data after 3,000 readings have been taken.

Weight:	3.0 pounds (1.4 kg)
Dimensions:	Hand-held, approx. 9.75 by 10.5 by 3.75 inches (248 by 273 by 95 millimeter).
Excitation Source:	Miniature x-ray tube and power supply (40 kilovolts [kV]/50 microamps [µA]
	maximum) with optional PERFECT technology.
Detector:	High-performance Si-PiN detector, Peltier cooled.
Signal Processing:	Hitachi SH-4 CPU ASICS high-speed DSP 4096 channel MCA
Element Range:	• Up to 25 standard elements in the range from titanium (atomic number: 22)
	to plutonium (atomic number: 94)
	Some nonstandard in-range elements available at additional cost
Batteries:	(2) Rechargeable lithium-ion battery packs with quick-swap capability; 6 to 12
	hour life between recharges (maximum depends on platform and duty cycle),
	2-hour recharge cycle.
Display:	¹ / ₄ Backlit VGA touch-screen LCD.
Testing Modes:	Bulk sample mode
	• Thin sample mode, including dust wipe mode and 37-millimeter filter mode
Standard Accessories:	• Soil Sampling Kit/Thin Sample Kit (varies by model and configuration)
	 Lockable, shielded waterproof carrying case
	• Shielded belt holster
	• Spare lithium-ion battery pack with holster
	• 110/220 volt AC battery charger/adapter
	• PC interface cable
	• Niton Data Transfer (NDT) PC software
	Safety lanyard
	Check and verification standards
	• Integrated bar code scan engine and virtual keypad for rapid and reliable
	entry of sample information

Table 6-1. Niton XLt XRF Analyzer Technical Specifications

6.2 Instrument Operations during the Demonstration

The XLt can be shipped or transported as checked or carry-on baggage. For transport to the field demonstration, the XLt was packed in a Pelican case that was 8 inches tall by 20 inches wide and 16 inches deep. The Pelican case was inspected by airport transportation security staff without concerns or delays. One additional large box was needed to hold all the accessories and supplies for routine analysis. A laptop computer is not required for analysis, but was used during the field demonstration for data downloading and to serve as a larger screen for viewing results and for assigning sample numbers.

6.2.1 Set up and Calibration

During the field demonstration, the XLt was used in the ex situ, bulk analysis mode. Instrument set-up involved placing the unit in the environmental test stand, connecting the analyzer to the laptop computer, and powering up both the analyzer and the computer. As part of the standard set-up routine, the analyzer was initially calibrated using the silver and tungsten shielding on the inside of the shutter to finetune the known peaks for these elements. Even though a warm-up time is not required, about 5 minutes is recommended to allow the analyzer to equilibrate with ambient conditions.

Niton included five calibration and reference samples with the analyzer to be used for calibration. Included were three NIST standards, one RCRA metals reference sample, and one silica blank. The Niton field team also used additional standards and samples with known element concentrations (the predemonstration samples) to further evaluate the calibration of the XLt analyzer. Individual element results and the error for each value were evaluated to verify that the analyzer was calibrated. The pre-set factory standard calibration for soil was selected for all routine analysis of soil and sediment and included the simultaneous analysis of up to 25 elements. The XLt analyzer software allows for empirical calibrations and corrections for any of the 25 elements. However, no empirical calibrations or corrections were used during the field demonstration.

6.2.2 Demonstration Sample Processing

Niton sent a two person team to the demonstration site to operate the two Niton instruments that were participating in the demonstration. One field team member was assigned to each instrument and completed the sample preparation, analysis, and data reduction for that instrument. Thus, the XLt had a dedicated operator for the entire length of the field demonstration, which showed how a single person could efficiently prepare and analyze samples in the field using the XLt.

Before sample processing was initiated, each sample set was arranged in numerical order. Custody seals were broken, and the soil samples were placed in standard 32-millimeter sample cups using a small stainless steel spatula (Figure 6-2). The sample cups were filled approximately 1/2 to 2/3 full. Each sample cup was then fitted with a small paper disc, polyester batting material behind the soil, and an end cap. The polyester batting and paper disc were necessary to hold the soil firmly against the upper Mylar[®] film when the sample cup is inverted. A colored self-adhesive dot was used to label each sample cup with the proper number and was attached to the bottom of the prepared sample cups.



Figure 6-2. Niton technician using a stainless steel scoop to fill a sample cup.

Prepared samples were placed in a queue and analyzed in order (Figure 6-3). Each sample was placed in the environmental sample holder and the drawer closed. The sample analysis was started from the laptop computer that was directly connected to the XLt instrument. A 120-second analysis run time was selected as an appropriate length that would simulate the choice of a typical customer under normal field conditions. At the end of the 120second test, the sample number recording screen was viewed on the laptop computer, and then the sample number was entered and the results were saved. The Niton data transfer (NDT) software has the option to save the data simultaneously to the laptop computer and to the XLt. The data were written to the computer using a comma-separated value (CSV) format.



Figure 6-3. Instrument set-up with samples awaiting analysis.

6.3 General Demonstration Results

The Niton operator for the XLt analyzed all 326 soil and sediment samples in 4 days using the standard soils calibration in the bulk sample mode. All analyses were completed in the ex situ mode after the samples were placed in the sample cups for analysis. Samples with results for iron above 50,000 ppm (5 percent) and samples with results for lead above 10,000 ppm (1 percent) were set aside for an additional 30-second analysis using the industrial bulk calibration in the bulk sample mode.

The industrial bulk calibration is not considered as accurate as the standard calibration for elements in soils below approximately 1 percent. However, the industrial bulk calibration is considered more accurate than the standard calibration for most elements in soils at concentrations above 1 percent. Iron is a common element in soils and is typically at concentrations above 10,000 ppm; therefore, the standard soil calibration range for iron extends to 50,000 ppm.

6.4 Contact Information

Additional information on Niton's XLt 700 Series XRF analyzer is available from the following source:

Mr. Dave Mercuro Niton Analyzers 900 Middlesex Turnpike, Building #8 Billerica, Massachusetts 01821 Telephone: (800) 875-1578, Ext. 333 Fax: 978-670-7430 E-mail: <u>dmercuro@niton.com</u> Internet: www.Niton.com

Chapter 7 Performance Evaluation

As discussed in Chapter 6, Niton analyzed all 326 demonstration samples of soil and sediment at the field demonstration site between January 25 and 27, 2005. A complete set of electronic data for the XLt in Microsoft Excel[®] spreadsheet format was delivered to the EPA/Tetra Tech field team before Niton demobilized from the site on January 28, 2005. All the data provided by Niton at the close of the demonstration are tabulated and compared with the reference laboratory data and the ERA-certified spike concentrations in Appendix D.

The XLt data set was reviewed and evaluated in accordance with the primary and secondary objectives of the demonstration. The findings of the evaluation for each objective are presented below.

7.1 Primary Objective 1 — Method Detection Limits

Samples were selected to calculate MDLs for each target element from the 12 potential MDL sample blends, as described in Section 4.2.1. The evaluation and selection of data for MDL calculation also addressed results reported as "not detected" by Niton. For many of the MDL blend results, element concentrations were below the statistical limits of detection (LODs) calculated by the XLt's instrument algorithms. These LODs are sample-specific and are calculated based on blank measurements, calibration routines, and relative element concentrations in the samples analyzed. (Additional information on calculating LODs is available from the developer, and a technical bulletin is available at http://www.NITONiton.com/docs/LODs.pdf.) In selecting samples from among the 12 blends for the calculation of MDLs, blends where one or more of the seven replicates was reported as "<LOD" were generally not used. In essence, this meant that all seven replicates had to have detected concentrations, as reported by Niton, to be used in this evaluation. Iron was not included in the MDL evaluation, as was discussed in Section 4.2.1.

The MDLs calculated for the XLt are presented in Table 7-1. As shown, the fewest MDLs were calculated for antimony, cadmium, copper, selenium,

and silver, with between two and four MDLs each. The most MDLs were calculated for chromium, nickel, and vanadium, with between nine and 12 MDLs each. With the exception of antimony and cadmium, at least one soil and one sediment MDL could be calculated for each of the target elements. Due to the lack of appropriate sediment blends, however, only soil MDLs could be calculated for antimony and cadmium.

Also shown in Table 7-1 are the mean MDLs calculated for each target element, which are classified as follows:

- Very low (1 to 20 ppm): arsenic, lead, mercury, selenium, and vanadium.
- Low (20 to 50 ppm): copper, silver, and zinc.
- Medium (50 to 100 ppm): chromium and nickel.
- High (greater than 100 ppm): antimony and cadmium.

The highest MDL of 216 ppm was calculated for antimony. The relatively high MDLs observed for antimony and cadmium should be qualified as based on limited data. Each of these mean MDLs was based on only two soil MDLs. Furthermore, the mean XRF concentrations reported for these elements were well above the predicted MDL range (generally greater than 300 ppm). In addition, these MDLs were calculated from Blends 8 and 12, which contained high concentrations of other elements such as arsenic, copper, lead, and zinc. Thus, the data for these elements are limited and may be biased high, producing uncertainty in the calculated MDLs. As an additional qualification, review of Table 7-1 indicates that the MDL calculated for nickel in the complex Blend 8 sample matrix (171 ppm) may be biased high when compared with other nickel MDLs. Blend 8 consisted of roaster slag from the Wickes Smelter site, a matrix that is further discussed in Sections 7.2 and 7.5 below.

The mean MDLs calculated for the XLt are compared in Table 7-2 with the mean LODs reported by Niton

			Antimony	7		Arsenic			Cadmium			Chromium	
Matrix	Blend No.	XLt MDL ²	XLt Conc. ³	Ref. Lab Conc. ⁴	XLt MDL ²	XLt Conc. ³	Ref. Lab Conc. ⁴	XLt MDL ²	XLt Conc. ³	Ref. Lab Conc. ⁴	XLt MDL ²	XLt Conc. ³	Ref. Lab Conc. ⁴
Soil	2	NC	ND	17	NC	ND	1.5	NC	ND	ND	38	299	167
Soil	5	NC	ND	ND	15	49	47	NC	ND	1.9	48	98	121
Soil	6	NC	ND	8	NC	389	477	NC	ND	12	50	101	133
Soil	8	234	646	118	NC	12,258	3,943	142	173	91	NC	ND	55
Soil	10	NC	ND	ND	20	36	39	NC	ND	0.96	86	107	116
Soil	12	198	326	62	NC	680	559	69	301	263	26	78	101
Soil	18	NC	ND	ND	NC	ND	9	NC	ND	ND	70	182	150
Sediment	29	NC	ND	ND	NC	ND	10	NC	ND	ND	NC	ND	63
Sediment	31	NC	ND	ND	NC	ND	11	NC	ND	ND	47	159	133
Sediment	32	NC	ND	ND	22	34	31	NC	ND	ND	55	79	75
Sediment	39	NC	ND	ND	11	19 ⁵	14	NC	ND	ND	46	93	102
Sediment	65	NC	ND	11	23	291	250	NC	ND	44	NC	345	303
Mean XLt M	DL	216			18			105			52	_	
			Copper			Lead			Mercury		Nickel		
	Blend	XLt	XLt	Ref. Lab									
Matrix	No.	MDL ²	Conc. ³	Conc. ⁴	MDL ²	Conc. ³	Conc. ⁴	MDL ²	Conc. ³	Conc. ⁴	MDL ²	Conc. ³	Conc. ⁴
Soil	2	19	42	47	NC	974	1,200	NC	ND	ND	47	123	83
Soil	5	NC	ND	49	11	70	78	NC	ND	ND	50	78	60
Soil	6	67	143	160	NC	3,703	3,986	NC	ND	0.83	60	101	70
Soil	8	NC	2,900	1,243	NC	46,986	33,429	NC	ND	15	171	277	57
Soil	10	NC	ND	31	13	60	72	NC	ND	0.14	65	78	60
Soil	12	NC	845	747	NC	4,423	4,214	NC	ND	1.8	49	137	91
Soil	18	NC	ND	50	NC	ND	17	7	31	56	58	289	213
Sediment	29	NC	1,877	1,986	27	40	33	NC	ND	0.24	28	154	72
Sediment	31	NC	1,522	1,514	20	55	51	15	23	ND	135	377	196
Sediment	32	NC	ND	36	23	28	26	12	17	ND	100	194	174
Sediment	39	66	98	94	9	36	27	10	28	ND	76	263	202
Sediment	65	40	80	69	13	35	25	11	38	32	90	325	214
Mean XLt M	DL	48			17			11			77		

 Table 7-1. Evaluation of Sensitivity -- Method Detection Limits for the Niton XLt¹

			Seleniu	m		Silver			Vanadium			Zinc	
Matrix	Blend No.	XLt MDL ²	XLt Conc. ³	Ref. Lab Conc.⁴	XLt MDL ²	XLt Conc. ³	Ref. Lab Conc. ⁴	XLt MDL ²	XLt Conc. ³	Ref. Lab Conc. ⁴	XLt MDL ²	XLt Conc. ³	Ref. Lab Conc. ⁴
Soil	2	NC	ND	ND	NC	ND	ND	NC	ND	1.2	23	24	24
Soil	5	NC	ND	ND	NC	ND	0.93	9	47	55	25	202	229
Soil	6	NC	ND	ND	NC	ND	14	23	47	56	NC	749	886
Soil	8	NC	ND	ND	35	224	144	17	29	34	NC	13,050	5,657
Soil	10	NC	ND	ND	NC	ND	ND	16	42	51	15	77	92
Soil	12	27	19 ⁵	15	27	46 ⁵	38	11	40	45	NC	2,554	2,114
Soil	18	NC	ND	ND	NC	ND	ND	13	65	67	21	86	90
Sediment	29	NC	ND	ND	NC	ND	ND	31	62	96	45	152	160
Sediment	31	NC	ND	ND	NC	ND	6.2	5	50	76	33	142	137
Sediment	32	7	11	4.6	NC	ND	ND	17	68	57	33	79	69
Sediment	39	7	10	ND	NC	ND	ND	13	39	38	61	158	137
Sediment	65	4	27	22	27	41	41	10	36	31	NC	1,946	1,843
Mean XLt M	DL	11			30			15			32		

 Table 7-1. Evaluation of Sensitivity -- Method Detection Limits for the Niton XLt¹ (Continued)

¹ Detection limits and concentrations are milligrams per kilogram (mg/kg), or parts per million (ppm).

² MDLs calculated from the 12 MDL sample blends for the Niton XLt in this technology demonstration (in bold typeface for emphasis).

³ This column lists the mean concentration reported for this blend by the XLt.

⁴ This column lists the mean concentration reported for this blend by the reference laboratory.

⁵ To increase the number of calculated MDLs for this metal, this blend was included despite the fact that detections were reported by the developer for only six of the seven replicates. This mean concentration and the corresponding MDL were calculated using the six replicate detected concentrations.

Conc. Concentration.

MDL Method detection limit.

NC The MDL was not calculated because reference laboratory concentrations exceeded five times the expected MDL range (approximately 50 ppm, depending on the element) or an insufficient number of detected concentrations were reported.

ND One or more results for this blend were reported as "Not Detected." Except where otherwise noted, blends with one or more ND result as reported by the XRF were not used for calculating the MDL.

Ref. Lab.Reference laboratory.

Element	XLt Mean MDLs ²	XLt Mean LODs ³	All XRF Instrument Mean MDLs ⁴	EPA Method 6200 ⁵ Mean Detection Limits
Antimony	216	91	61	55 ⁶
Arsenic	18	26	26	92
Cadmium	105	36	70	NR
Chromium	52	46	83	376
Copper	48	44	23	171
Lead	17	11	40	78
Mercury	11	30	23	NR
Nickel	62	191	50	100^{6}
Selenium	11	37	8	NR
Silver	30	29	42	NR
Vanadium	15	5^{4}	28	NR
Zinc	32	NR	38	89

 Table 7-2. Comparison of XLt MDLs to XLt Instrument LODs and EPA Method 6200 Data¹

- ¹ Detection limits are in units of milligrams per kilogram (mg/kg), or parts per million (ppm).
- 2 The mean MDLs calculated for this technology demonstration, as presented in Table 7-1.
- 3 The mean LODs reported by Niton for the 12 MDL blends.
- ⁴ The mean MDLs calculated for all eight XRF instruments that participated in this EPA technology demonstration.
- ⁵ Mean values calculated from Table 4 of Method 6200 (EPA 1998e, <u>www.epa.gov/sw-846</u>).
- ⁶ Only one value reported.
- EPA U.S. Environmental Protection Agency
- LOD Limit of detection
- MDL Method detection limit
- NR Not reported; no MDLs or LODs reported for this element.

for the MDL blends, the mean MDLs for all instruments that participated in the demonstration, and the mean MDLs derived from performance data presented in EPA Method 6200 (EPA 1998e). As shown, the mean MDLs for the XLt are generally comparable to Niton's LODs and are generally lower than the mean MDLs calculated from EPA Method 6200 data. Exceptions include antimony and cadmium where, as noted above, limited data for the XLt appear to have produced mean MDLs that are higher than the LODs and available Method 6200 data. When compared with the results for the demonstration as a whole (encompassing all eight XRF instruments), the XLt again exhibited high relative mean MDLs for antimony and cadmium as well as for copper, nickel, and selenium. Mean MDLs for the XLt were well below the all-instrument means for chromium, lead, mercury, and vanadium.

7.2 Primary Objective 2 — Accuracy and Comparability

The number of demonstration sample blends that met the criteria for evaluation of accuracy, as described in Section 4.2.2, was generally in the range of 40 to 70 for most elements. However, low relative numbers of acceptable blends were noted for antimony (24), cadmium (22), mercury (26), and selenium (25). RPDs between the mean XLt and mean reference laboratory concentrations were calculated for each blend that met the criteria for an element.

Table 7-3 presents the median RPDs, along with the number or RPD results used to calculate the median, for each target element. These statistics are provided for the demonstration as a whole, as well as for subpopulations grouped by media type (soil versus sediment) and concentration level (Levels 1 through 4, as documented in Table 3-1). Additional summary statistics for the RPDs (minimum, maximum, and mean) are provided in Appendix E (Table E-1).

Accuracy was classified as follows for the target elements based on the overall median RPDs:

- Very good (median RPD less than 10 percent): copper and selenium.
- Good (median RPD between 10 percent and 25 percent): arsenic, cadmium, chromium, iron, lead, silver, and zinc.
- Fair (median RPD between 25 percent and 50 percent): nickel and vanadium.
- Poor (median RPD greater than 50 percent): antimony and mercury.

The median RPD was used for this evaluation because it is less affected by extreme values than is the mean. (The initial evaluation of the RPD populations showed that they were right-skewed or lognormal.) However, the classification of the elements based on accuracy generally stayed the same when the mean RPD rather than the median RPD was used for the evaluation (Table E-1). Moreover, these classifications did not vary with media type (soil versus sediment) for any of the elements. Neither did they vary with concentration range, with the following exceptions:

• High median RPDs were observed in the Level 3 samples for arsenic (with concentrations greater than 2,000 ppm) and Level 1 samples for cadmium (with concentrations between 50 and 500 ppm). At 79 percent for arsenic and 38 percent for cadmium, these median RPDs were much higher than were observed for other concentration ranges. These RPDs appeared to be skewed high by the results for sample blends 8 and 9 from the Wickes Smelter site, which contained high concentrations of other elements (lead, copper, zinc, and iron).

- The best accuracy for mercury was observed in the Level 1 samples (with concentrations between 20 and 200 ppm) in both the soil and sediment matrices, where median RPDs of less than 50 percent were calculated. These samples were generally characterized by very low concentrations of other elements, including elements adjacent to mercury in the periodic table, such as cadmium and lead.
- Median RPDs for silver in the Level 3 concentration samples (in the range of 50 percent for both soil and sediment) were higher than in the Level 1 and 2 samples (where median RPDs ranged only as high as 17 percent). This effect was also observed for vanadium, where accuracy declined from "good" or "fair" into the "poor" range as concentrations increased. These trends appeared to be generalized rather than caused by limited data or extreme results, and may be related to the overall increasing complexity of the sample matrices as concentrations of these elements increased.

Section 5.3.3 discussed how the reference laboratory data for antimony were consistently biased low when compared with the ERA-certified spike concentrations. This effect may be caused by volatilization of the antimony compounds used for spiking, resulting in loss of antimony during the sample digestion process at the reference laboratory. Therefore, Table 7-3 includes a second accuracy evaluation for antimony, comparing the XLt results to the ERA-certified values. As shown, this comparison indicates far better performance for antimony than does the comparison to the reference laboratory results, with median RPDs in the range of 5 to 15 percent ("very good" to "good") for all media and concentration levels. Furthermore, these results suggest that the XLt analyzer may measure some antimony compounds more accurately in soil and sediment than the fixed-laboratory reference methods.

	Sample		Ant	imony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury	Nickel	Selenium	Silver	Vanadium	Zinc
Matrix	Group	Statistic	Ref. Lab	ERA Spike							-					
Soil	Level 1	Number	6		12	4	23	14	5	14	7	23	4	2	13	19
		Median	133.3%		19.0%	37.6%	20.4%	10.3%	16.3%	19.1%	47.6%	38.7%	15.2%	17.0%	8.3%	4.5%
	Level 2	Number	5	1	4	7	4	8	13	4	7	5	5	3	4	6
		Median	129.5%	7.1%	22.5%	10.7%	13.2%	27.6%	16.1%	17.5%	116.4%	39.0%	5.4%	17.4%	73.5%	11.8%
	Level 3	Number	4	3	4	2	2	2	12	8	2	6	4	7	4	9
		Median	113.8%	8.9%	79.4%	10.3%	17.1%	13.9%	5.7%	9.2%	126.3%	29.4%	3.9%	43.4%	103.7%	18.2%
	Level 4	Number							8	5						
		Median							15.0%	12.4%						
	All Soil	Number	15	4	20	13	29	24	38	31	16	34	13	12	21	34
		Median	125.9%	8.7%	24.5%	11.0%	19.2%	12.8%	13.5%	17.1%	92.0%	37.0%	5.4%	24.8%	19.3%	8.3%
Sediment	Level 1	Number	2	2	14	2	10	9	3	16	3	18	5	5	6	17
		Median	144.4%	6.7%	9.1%	10.2%	15.4%	8.9%	5.2%	12.4%	44.7%	52.6%	9.1%	12.7%	40.5%	16.8%
	Level 2	Number	4	4	4	4	3	4	19	4	4	6	4	4	8	5
		Median	138.5%	15.5%	12.6%	14.9%	12.9%	1.2%	12.5%	3.3%	80.4%	37.8%	9.7%	12.4%	53.2%	3.3%
	Level 3	Number	3	3	2	3	3	10	5	3	3	4	3	3	3	4
		Median	107.7%	13.5%	20.5%	9.4%	9.0%	5.8%	8.5%	8.2%	91.6%	27.5%	5.4%	53.2%	104.0%	8.5%
	Level 4	Number							5							
		Median							26.0%							
	All Sediment	Number	9	9	20	9	16	23	32	23	10	28	12	12	17	26
		Median	122.3%	13.5%	12.7%	9.8%	12.3%	5.6%	13.5%	8.0%	81.6%	40.1%	9.1%	15.1%	44.5%	12.9%
All Samples	Niton XLt	Number	24	13	40	22	45	47	70	54	26	62	25	24	38	60
		Median	125.5%	11.0%	18.1%	10.8%	16.4%	9.3%	13.5%	12.0%	88.8%	38.9%	6.8%	19.0%	40.5%	11.5%
All Samples	All XRF	Number	206	110	320	209	338	363	558	392	192	403	195	177	218	471
	Instruments	Median	84.3%	70.6%	26.2%	16.7%	26.0%	16.2%	26.0%	21.5%	58.6%	25.4%	16.7%	28.7%	38.3%	19.4%

Table 7-3. Evaluation of Accuracy - Relative Percent Differences Versus Reference Laboratory Data for the Niton XLt

All RPDs presented in this table are absolute values.

-- No samples reported by the reference laboratory in this concentration range.

ERA Environmental Resource Associates, Inc.

NC Not calculated.

Number Number of samples meeting criteria for accuracy evaluation (Section (4.2.2).

Ref. Lab Reference laboratory (Shealy Environmental Services, Inc.).

RPD Relative percent difference.

As an additional comparison, Table 7-3 also presents the median XRF instrument RPDs for the demonstration as a whole (across all eight instruments). Complete summary statistics for the RPDs across the eight XRF instruments are included in Appendix E. Table 7-3 indicates that the median RPDs for the XLt were higher than the all-instrument medians for antimony, mercury, and nickel, but were equivalent to or below them for the remaining 10 target elements.

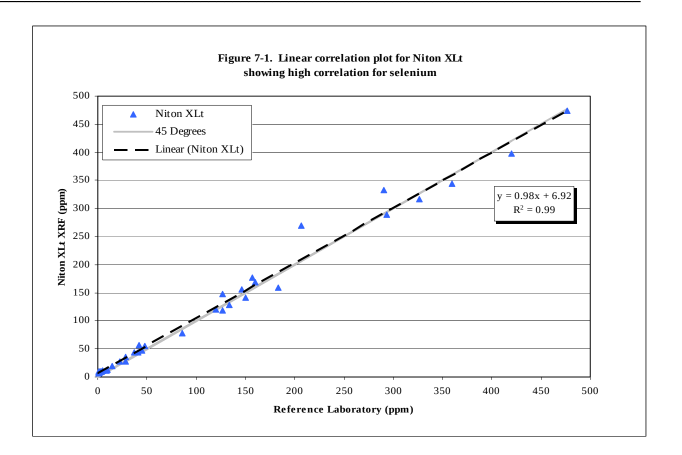
In addition to calculating RPDs, the evaluation of accuracy included preparing linear correlation plots of XLt concentration values against the reference laboratory values. These plots are presented for the individual target elements in Figures E-1 through E-13 of Appendix E. The plots include a 45-degree line showing the "ideal" relationship between the XLt data and the reference laboratory data, as well as a "best fit" linear equation (y = mx + b, where *m* is the slope of the line and *b* is the y-intercept of the line) and correlation coefficient (r^2) to help illustrate the "actual" relationship between the two methods. To be considered accurate, the correlation coefficient should be greater than 0.9, the slope (m) should be between 0.75 and 1.25, and the y-intercept (b) should be relatively close to zero (that is, plus or minus the mean MDL calculated in Table 7-1). Table 7-4 lists the results for these three correlation parameters and highlights in bold each target element that met all three accuracy criteria. This table shows that the results for antimony (correlated with the ERAcertified values only), cadmium, chromium, iron, and selenium met all three of these criteria. The correlation plot for selenium is displayed in Figure 7-1 as an example of the high correlations obtained for these elements.

Target Element	m	b	\mathbf{r}^2	Correlation	Bias	
Antimony (Ref. Lab) ¹	3.40	129	0.88	Moderate	High	
Antimony (Cert. Val.) ¹	1.12	18	0.99	High		
Arsenic	1.34	12	0.95	High	High	
Cadmium	1.20	28	0.98	High		
Chromium	1.00	26	0.95	High		
Copper	1.09	55	0.87	Moderate		
Iron	1.23	403 ²	0.94	High		
Lead	1.45	509 ²	0.95	High	High	
Mercury	0.24	25	0.98	High	Low	
Nickel	1.31	56	0.98	High	High	
Selenium	0.98	7	0.99	High		
Silver	1.24	18	0.71	Moderate		
Vanadium	0.22	40	0.51	Moderate	Low	
Zinc	1.77	456	0.76	Moderate	High	

 Table 7-4.
 Summary of Correlation Evaluation for the Niton XLt

¹ For antimony, correlation was assessed for the Niton XLt versus the reference laboratory data ("Ref. Lab") as well as versus the ERA-certified spike values ("Cert. Val.") for the spiked sample blends.

- For iron, no MDL was calculated and the high intercept value was the result of the extreme range of concentrations in the demonstration samples. The broad range of concentrations in the demonstration samples also produced a high intercept for lead.
- -- No bias observed.
- b Y-intercept of correlation line.
- m Slope of correlation line.
- r^2 Correlation coefficient of correlation line.



The elements with a high relative degree of correlation between the XLt and reference laboratory were generally the same elements for which accuracy was rated "Very Good" or "Good" through the evaluation of RPDs. Specifically, the elements with high correlation coefficients (in the range of 0.94 to 0.99) included arsenic, cadmium, chromium, iron, lead, nickel, selenium, and zinc. Mercury, which was rated "poor" by the RPD evaluation, also exhibited a high r^2 value. However, this correlation was affected by two extreme Level 4 concentrations (blends 21 and 22) that were more than three times higher than the next-highest concentrations in the mercury data set (see Figure E-8). Removing these extreme concentrations from the plots produced a much poorer correlation coefficient, in the range of 0.84, for the rest of the data set, consistent with the findings of the RPD evaluation. Other observations from the correlation plots are as follows:

• Slopes significantly greater than 1 in conjunction with moderately high correlation coefficients indicated a high bias in the XLt data for arsenic, lead, and zinc. However, further review of the

data indicated that removal of high outliers from complex blends 7, 8, and 9 (Wickes Smelter slag) improved the r^2 values and reduced or eliminated the positive bias for these elements. Though no bias was observed for copper, removal of a blend 7 outlier also improved r^2 for this element from 0.87 ("Moderate") to 0.95 ("High").

- Large deviations from zero were noted in the yintercepts for lead and iron. Examination of the plots for these elements (Figures E-6 and E-7) reveals that these deviations were small relative to the extreme range of concentrations in the demonstration samples.
- For antimony, the high bias in relation to the reference laboratory results was eliminated when the XLt results were compared to the ERA-certified values. Comparison to the ERA-certified values also produced a very high r² of 0.99. These findings agree with the RPD evaluation above in showing better performance for antimony by the XLt when ERA-certified spike values are used to assess accuracy.

• Low biases were observed for mercury and vanadium. Vanadium also had the lowest r² of all the target elements (0.51). These observations are consistent with the RPD evaluation, which found higher RPDs (lower accuracy) for these elements in the higher-concentration samples. The plot for vanadium is shown Figure 7-2. In the case of mercury, the low bias may reflect loss of mercury through volatilization during the high level of sample processing that preceded the demonstration.

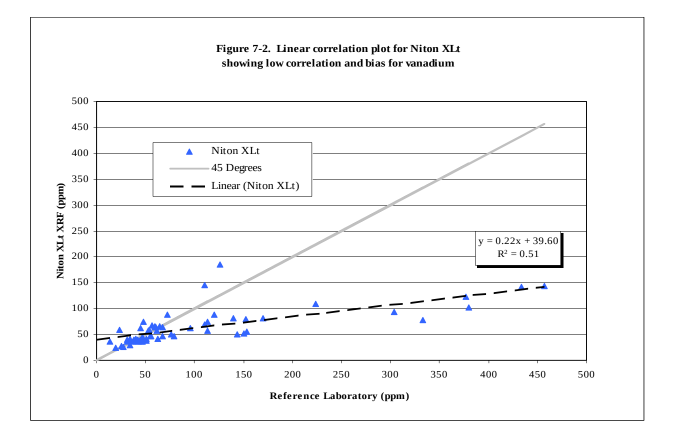
In conclusion, the evaluations of accuracy showed an acceptable overall level of performance by the XLt for the target elements. Correlations with the reference laboratory were generally high, and median RPDs were better for most elements in comparison to all eight XRF instruments as a whole. Niton's proven calibration and quantification algorithms for environmental media may have contributed to the high relative level of accuracy attained. However, the XLt demonstrated somewhat low relative performance for mercury and vanadium in the accuracy evaluation.

7.3 Primary Objective 3 — Precision

As outlined in Section 4.2.3, precision of the XLt data set was evaluated by calculating RSDs for the replicate measurements from each sample blend. Median RSDs for the various concentration levels and media (soil and sediment) are presented in Table 7-5. The table also presents the median RSDs for the demonstration data set as a whole for the XLt. Additional summary statistics for the RSDs (including minimum, maximum, and mean) are provided in Appendix E (Table E-2).

The RSD calculation revealed a high level of precision for the XLt in that the overall median RSDs were 10 percent or less for all target elements in both matrices. The ranges into which the median RSDs fell are further summarized below:

• RSD of 1 percent to 5 percent: cadmium, iron, lead, selenium, and zinc.



• RSD of 5 percent to 10 percent: antimony, arsenic, chromium, copper, mercury, nickel, silver, and vanadium.RSD of greater than 10 percent: none.

No differences were observed between the RSDs for soil and sediment. Use of the mean RSDs (Appendix E) as opposed to the median RSDs indicated a similarly high level of precision in the XLt results.

The high overall level of precision may have been facilitated by the high level of processing (homogenizing, sieving, crushing, and drying) performed on the sample blends before the demonstration (Chapter 3). This observation is consistent with the previous SITE MMT program demonstration of XRF technologies that occurred in 1995 (EPA 1996a, 1996b, 1998a, 1998b, 1998c, and 1998d). The high level of sample processing applied during both XRF technology demonstrations was necessary to minimize the effects of sample heterogeneity on the demonstration results and on comparability with the reference laboratories. During project design, site investigation teams that intend to compare XRF and laboratory data should similarly assess the need for sample processing steps to manage sample heterogeneity and improve data comparability.

Further review of the median RSDs in Table 7-5 based on concentration range reveals slightly higher RSDs (in other words, lower precision) for the target elements in Level 1 samples when compared with the rest of the data set. This observation indicates that analytical precision for the XRF results is somewhat concentration-dependent. Even for the Level 1 samples, however, the mean RSDs were relatively good, with the highest RSD being 26 percent (for cadmium in soil).

As an additional comparison, Table 7-5 also presents the median RSDs calculated for all XRF instruments that participated in the demonstration. Complete summary statistics for the RSDs across all XRF instruments are included in Appendix E. Table 7-5 indicates that the median RSDs for the XLt were equivalent to or below the all-instrument medians for all elements with the exception of antimony, copper, nickel, and silver, where slightly higher median RSDs were observed.

Table 7-6 presents median RSD statistics for the reference laboratory for comparison to the XLt data. These median RSD statistics were calculated using the same blends as those used to calculate the XLt RSD statistics. (Complete summary statistics are provided in Table E-3 of Appendix E.) Table 7-6 indicates that the XLt median RSDs were equivalent to or lower than the reference laboratory RSDs for 11 out of 13 target elements (only the chromium and nickel RSDs were slightly higher). Thus, the XLt exhibited slightly better precision overall than the reference laboratory. In comparison, the median RSDs for all XRF instruments were equivalent to or lower than the reference laboratory RSDs also for 11 of the 13 target elements (the exceptions were chromium and vanadium).

7.4 Primary Objective 4 — Impact of Chemical and Spectral Interferences

The mean RPD data from the accuracy evaluation were further processed to assess the effects of interferences. The RPD data for elements susceptible to interferences were grouped and compared based on the relative concentrations of potentially interfering elements. Of specific interest for the comparison were the potential effects of:

- High concentrations of lead on the RPDs for arsenic,
- High concentrations of nickel on the RPDs for copper (and vice versa), and
- High concentrations of zinc on RPDs for copper (and vice versa).

The rationale and approach for evaluation of these interferents are described in Section 4.2.4.

	Sample														
Matrix	Group	Statistic	Antimony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury	Nickel	Selenium	Silver	Vanadium	Zinc
Soil	Level 1	Number	6	12	4	23	14	5	14	7	23	4	2	13	19
		Median	12.8%	10.8%	26.4%	10.4%	17.1%	5.5%	8.7%	13.2%	12.8%	9.3%	19.8%	10.0%	6.3%
	Level 2	Number	5	4	7	4	8	13	4	7	5	5	3	4	6
		Median	6.9%	5.5%	2.6%	3.9%	4.2%	2.9%	2.6%	6.9%	8.1%	4.4%	6.6%	6.8%	2.4%
	Level 3	Number	4	4	2	2	2	13	8	2	6	4	7	4	9
		Median	3.1%	3.9%	2.2%	1.7%	1.5%	1.8%	2.0%	3.1%	1.8%	3.9%	3.9%	6.5%	2.8%
	Level 4	Number						7	5						
		Median						2.5%	3.9%						
	All Soil	Number	15	20	13	29	24	38	31	16	34	13	12	21	34
		Median	6.9%	7.3%	2.8%	9.9%	11.1%	2.4%	3.7%	7.3%	10.6%	4.4%	5.7%	8.5%	3.9%
Sediment	Level 1	Number	2	14	2	10	9	3	16	3	18	5	5	6	17
		Median	19.6%	10.4%	10.6%	20.2%	16.0%	4.4%	7.7%	8.6%	12.5%	6.9%	19.5%	15.0%	9.5%
	Level 2	Number	4	4	4	3	4	19	4	4	6	4	4	8	5
		Median	4.9%	3.3%	3.0%	2.6%	3.7%	1.8%	2.9%	4.8%	7.9%	4.3%	2.6%	4.0%	2.6%
	Level 3	Number	3	2	3	3	10	5	3	3	4	3	3	3	4
		Median	6.1%	5.1%	3.2%	1.8%	2.2%	2.0%	2.5%	3.2%	4.4%	4.7%	5.3%	4.5%	2.8%
	Level 4	Number						5							
		Median						2.1%							
	All Sediment	Number	9	20	9	16	23	32	23	10	28	12	12	17	26
		Median	6.1%	8.2%	3.2%	9.9%	3.5%	2.0%	4.9%	3.8%	9.2%	5.1%	5.9%	5.8%	7.1%
All Samples	Niton XLt	Number	24	40	22	45	47	70	54	26	62	25	24	38	60
		Median	6.7%	7.9%	3.0%	9.9%	5.7%	2.2%	4.1%	5.7%	10.0%	4.7%	5.9%	7.0%	4.8%
All Samples	All XRF	Number	206	320	209	338	363	558	392	192	403	195	177	218	471
	Instruments	Median	6.1%	8.2%	3.6%	12.1%	5.1%	2.2%	4.9%	6.8%	7.0%	4.5%	5.2%	8.5%	5.3%

Table 7-5. Evaluation of Precision - Relative Standard Deviations for the Niton XLt

-- No samples reported by the reference laboratory in this concentration range.

Number of samples meeting criteria for precision evaluation (Section (4.2.3).

RSD Relative standard deviation.

Matrix	Sample Group	Statistic	Antimony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury	Nickel	Selenium	Silver	Vanadium	Zinc
Soil	Ref. Lab	Number	17	23	15	34	26	38	33	16	35	13	13	21	35
		Median	9.8%	12.4%	9.0%	10.6%	9.1%	8.7%	13.2%	6.6%	10.0%	7.1%	7.5%	6.6%	9.1%
Sediment	Ref. Lab	Number	7	24	10	26	21	31	22	10	27	12	10	17	27
~~~~~		Median	9.1%	9.2%	8.2%	7.5%	8.9%	8.1%	7.4%	6.9%	7.3%	7.6%	6.6%	8.1%	6.9%
All	Ref. Lab	Number	24	47	25	60	47	69	55	26	62	25	23	38	62
Samples		Median	9.5%	9.5%	9.0%	8.4%	8.9%	8.5%	8.6%	6.6%	8.2%	7.4%	7.1%	7.2%	7.4%
All	Niton	Number	24	40	22	45	47	70	54	26	62	25	24	38	60
Samples	XLt	Median	6.7%	7.9%	3.0%	9.9%	5.7%	2.2%	4.1%	5.7%	10.0%	4.7%	5.9%	7.0%	4.8%
All	All XRF	Number	206	320	209	338	363	558	392	192	403	195	177	218	471
Samples	Instruments	Median	6.1%	8.2%	3.6%	12.1%	5.1%	2.2%	4.9%	6.8%	7.0%	4.5%	5.2%	8.5%	5.3%

# Table 7-6. Evaluation of Precision – Relative Standard Deviations for the Reference Laboratory versus the Niton XLt and All Demonstration Instruments

Notes:

Number Number of samples meeting criteria for precision evaluation (Section (4.2.3).

Ref. Lab Reference Laboratory.

RSD Relative standard deviation.

XRF X-ray fluorescence.

Interferent-to-element ratios were calculated using the mean concentrations reported for each blend by the reference laboratory, are classified as low (less than 5X), moderate (5 to 10X), or high (10X). Table 7-7 presents median RPD data for arsenic, nickel, copper, and zinc that are grouped based this classification scheme. Additional summary statistics are presented in Appendix E (Table E-4). The table indicates a clear increase in the median RPD for arsenic at the higher lead-to-arsenic ratios. Specifically, a median RPD of 13 percent at low interferent ratios increases to near 50 percent in the high interferent ratios. Using the criteria applied in Section 7.2, high concentrations of lead diminish the accuracy of the XLt from "good" to "fair" for arsenic. Similarly, Table 7-7 indicates that high concentrations of copper reduce instrument accuracy from "fair" to "poor" for nickel. In presenting statistics for unmodified RPDs as well as the absolute values of the RPDs, Table E-4 further shows that the interferences by lead and copper tend to increase the positive bias of the results for arsenic and nickel (as indicated by more negative unmodified RPDs).

Interestingly, Table 7-7 reveals no other trends in RPDs that would indicate significant potential interferences of nickel, copper, or zinc with each other. The only other significant difference in RPDs apparent in Table 7-7 is in the data for zinc and copper, where RPDs for blends with moderate zinc-to-copper ratios are much higher than for blends with either lower or higher ratios. This effect on RPDs for copper may be related to matrix effects other than zinc interference. The four samples included in this grouping include blends 7, 8, and 9, which contained high concentrations of elements other than zinc, such as arsenic, lead, and iron (see Section 7.5 below).

#### 7.5 Primary Objective 5 — Effects of Soil Characteristics

The population of RPDs between the XLt results and the reference laboratory results were further evaluated against sampling site and soil type. Separate sets of summary statistics were developed for the mean RPDs associated with each sampling site for comparison to the other sites and to the demonstration data set as a whole. The site-specific median RPDs are presented in Table 7-8, along with descriptions of soil/sediment type from observations during sampling at each site. Complete RPD summary statistics for each soil type (minimum, maximum, and mean) are presented in Table E-5 of Appendix E.

Another perspective on the effects of soil type was developed by graphically assessing outliers and extreme values in the mean RPD data sets for the target elements. This evaluation focused on correlating these values with sample types or locations for multiple elements across the data set. Outliers and extreme values are apparent in the correlation plots (Figures E-1 through E-13) and are further depicted for the target elements on box and whisker plots in Figure E-14.

Review of Table 7-8 and Figure E-14 indicates that high relative median RPDs are observed in the Wickes Smelter blends for a number of elements, including arsenic, cadmium, copper, silver, and vanadium. Evaluation of the outliers confirms this observation, indicating that two or more of the highest mean RPDs for these elements are from Wickes Smelter blends 7, 8, or 9. During the demonstration sample collection program (Chapter 2), the soil matrix from this site was described as roaster slag, consisting of a black, fairly coarse sand and gravel material. This slag is an intermediate product in processing ore, wherein volatile sulfide compounds are thermally removed, leaving concentrated heavy elements. In addition to arsenic, copper, and zinc, this matrix was found to contain high concentrations of lead and iron. These same Wickes Smelter blends were identified in the previous section as affecting instrument performance for some target elements. Thus, the data presented in Table 7-8 and Figure E-14 confirm that the complex slag matrixes of blends 7, 8, and 9 have significant effects on the accuracy of the XLt for some elements. RPDs for several target elements are presented for these blends in Table 7-9.

Parameter	Lead Effects on Arsenic		Copper Effects on Nickel			Nickel Effects on Copper			Zinc F	Effects on (	Copper	Copper Effects on Zinc			
Interferent/ Element Ratio	<5	5 – 10	>10	<5	5 - 10	>10	<5	5 - 10	>10	<5	5 - 10	>10	<5	5 – 10	>10
Number of Samples	29	7	5	44	6	14	39	1	7	32	4	11	47	3	11
Median RPD of Target Element ²	12.9%	24.5%	46.7%	34.4%	35.8%	71.2%	9.4%	7.9%	8.9%	8.2%	59.3%	9.4%	8.8%	16.8%	14.3%
Median Interferent Concentration	72	7460	2300	127	1076	2117	150	387	1767	163	4399	2300	147	742	2167
Median Target Element Concentration	95	1038	98	186	176	108	748	78	84	944	944	123	823	102	137

Table 7-7. Effects of Interferent Elements on the RPDs (Accuracy) of Target Elements¹

¹ Concentrations are reported in units of milligrams per kilogram (mg/kg), or parts per million (ppm).

² All median RPDs presented in this table are based on the population of absolute values of the individual RPDs.

< Less than.

> Greater than.

RPD Relative percent difference.

		Matrix		Antimony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead
Matrix	Site	Description	Statistic	-						
Soil	AS Fine to medium sand (steel		Number			2	2	3	3	3
		processing)	Median			14.9%	20.9%	9.4%	14.1%	9.9%
Soil	BN	Sandy loam, low organic (ore	Number	4	6	5	7	5	7	7
		residuals)	Median	134.6%	16.2%	9.3%	24.9%	12.3%	13.7%	8.6%
Soil	CN	Sandy loam (burn pit residue)	Number	1	1	1	1	3	3	3
			Median	100.5%	28.6%	11.0%	29.5%	12.4%	13.0%	22.3%
Soil &	KP	Soil: Fine to medium quartz sand.	Number	1			4	3	6	6
		Sed.: Sandy loam, high organic. (Gun and skeet ranges)	Median	6.0%			55.1%	11.4%	9.2%	20.4%
Sediment	Sediment LV	Clay/clay loam, salt crust (iron and other precipitate)	Number	4	9	5	7	5	12	8
			Median	116.8%	10.3%	9.4%	11.7%	9.3%	27.6%	21.2%
Sediment	RF	Silty fine sand (tailings)	Number	3	12	4	9	13	13	11
			Median	109.9%	15.3%	10.2%	10.5%	5.9%	5.5%	5.1%
Soil	SB	Coarse sand and gravel (ore and	Number	5	5	1	11	2	12	5
		waste rock)	Median	125.9%	13.5%	15.1%	11.2%	19.6%	6.4%	20.1%
Sediment	TL	Silt and clay (slag-enriched)	Number	3	1	2	1	7	7	4
			Median	154.8%	27.7%	15.5%	18.0%	4.6%	31.8%	8.5%
Soil	WS	Coarse sand and gravel (roaster	Number	3	6	2	3	6	7	7
		slag)	Median	138.2%	51.4%	73.5%	21.4%	28.5%	16.9%	12.4%
	All		Number	24	40	22	45	47	70	54
			Median	125.5%	18.1%	10.8%	16.4%	9.3%	13.5%	12.0%

## Table 7-8. Effect of Soil Type on the RPDs (Accuracy) for Target Elements

		Matrix		Mercury	Nickel	Selenium	Silver	Vanadium	Zinc
Matrix	Site	Description	Statistic	-					
Soil	AS	Fine to medium sand (steel	Number		3	1	1	2	3
		processing)	Median		39.0%	5.4%	22.8%	70.5%	3.3%
Soil	BN	Sandy loam, low organic (ore	Number		6	4	4	3	7
		residuals)	Median		36.7%	9.5%	19.6%	96.6%	17.9%
Soil	CN	Sandy loam (burn pit residue)	Number	2	3	2	2	1	3
			Median	79.4%	27.6%	14.8%	22.1%	93.9%	7.0%
Soil &	KP	Soil: Fine to medium quartz sand.	Number		3	3			2
		Sed.: Sandy loam, high organic. (Gun and skeet ranges)	Median		27.3%				10.8%
Sediment LV C		Clay/clay loam, salt crust (iron and other precipitate)	Number	4	11	5	4	9	8
	Median		69.1%	39.4%	9.1%	14.0%	38.3%	17.1%	
Sediment	RF	Silty fine sand (tailings)	Number	5	13	5	5	4	13
			Median	59.5%	35.4%	9.1%	11.6%	81.1%	11.5%
Soil	SB	Coarse sand and gravel (ore and	Number	12	11	3	1	10	10
		waste rock)	Median	107.5%	32.5%	4.5%	77.7%	7.6%	4.4%
Sediment	TL	Silt and clay (slag-enriched)	Number	3	6	4	4	7	7
			Median	104.9%	76.3%	10.8%	17.8%	42.0%	14.3%
Soil	WS	Coarse sand and gravel (roaster	Number		6	1	3	2	7
		slag)	Median		45.4%	3.4%	43.4%	16.4%	16.7%
	All		Number	26	62	25	24	38	60
			Median	88.8%	38.9%	6.8%	19.0%	40.5%	11.5%

#### Table 7-8. Evaluation of the Effect of Soil Type on RPDs (Accuracy) of Target Elements (Continued)

Notes:

AS Alton Steel Mill

- BN Burlington Northern railroad/ASARCO East
- CN Naval Surface Warfare Center, Crane Division
- KP KARS Park Kennedy Space Center
- LV Leviathan Mine/Aspen Creek
- RF Ramsey Flats Silver Bow Creek
- SB Sulfur Bank Mercury Mine
- TL Torch Lake Superfund Site
- WS Wickes Smelter Site

Other notes:

Number RPD

---

No samples reported by the reference laboratory in this concentration range. Number of demonstration samples evaluated. Relative percent difference (absolute value).

Blend	Arsenic	Cadmium	Copper	Iron	Lead	Nickel	Silver	Zinc
7	56.2%		38.6%	55.9%	12.4%	97.1%		36.9%
8	102.6%	61.7%	80.0%	92.1%	33.7%		43.4%	79.0%
9	121.5%	85.4%	108.3%	108.6%	56.3%	146.5%	61.7%	108.9%
Median for								
all blends	18.1%	10.8%	9.3%	13.5%	21.5%	38.9%	19.0%	11.5%

Table 7-9. RPDs Calculated for Wickes Smelter Sample Blends for the Niton XLt

-- Blend not included in accuracy evaluation.

Further review of extreme concentrations in the data sets indicated that matrices from the Leviathan Mine site may have also affected the accuracy of the XRF measurements for a number of elements. Specifically, one or more of the high outliers and extreme values depicted on Figure E-14 for arsenic, copper, nickel, and zinc were reported from samples collected at the Leviathan Mine site (blends 34, 35, 55, and 58). Chapter 2 indicates that the matrices from this site were clay soils that also included precipitates and solids from acid mine leachate and wastewater retention ponds. Many of the blends contained extreme concentrations of iron (in the range of 100,000 to 250,000 ppm). Although the complexity of this matrix produced a number of extreme values in the data sets, Table 7-8 indicates minimal impacts to median RPDs when compared with other sampling sites.

Other sampling sites and blends appeared to produce high RPDs for some elements on an isolated basis, or produced minor increases in mean or median RPDs. For example, Torch Lake blends produced high outliers for iron and selenium, and slight increases in RPDs for both iron and nickel. These effects appeared to be minor, however, and no other generalized trends in XRF accuracy versus the sample collection site or soil/sediment type could be discerned, given that the ranges of RPDs observed for some target elements were very broad. The spread in the accuracy results is illustrated on the box and whisker plot in Figure E-14. The plot shows that the distributions of RPDs were sufficiently broad to preclude the identification of any extreme values for antimony, mercury, or vanadium.

#### 7.6 Primary Objective 6 — Sample Throughput

Niton provided a single instrument operator during the field demonstration to perform all activities associated with sample preparation, instrumental analysis, and data reduction. The Niton XLt instrument operator was able to analyze all 326 demonstration samples in 3 days at the demonstration site. Once the XLt instrument had been set up and operations had been streamlined, the Niton instrument operator was able to analyze a maximum of 146 samples during an extended work day. Without an extended work day, it was estimated that the Niton instrument operator would have averaged about 91 samples per day.

This estimated sample throughput for a normal working day was much higher than that observed for the other instruments that participated in the demonstration (average of 66 samples per day). This was a particularly significant achievement given that the other developers utilized two-person or even three-person field teams. A detailed discussion of the time required to complete the various steps of sample analysis using the XLt is included as part of the labor cost analysis in Section 8.3.

## 7.7 Primary Objective 7 — Technology Costs

The evaluations pertaining to this primary objective are fully described in Chapter 8, Economic Analysis.

#### 7.8 Secondary Objective 1 — Training Requirements

The instrument operator must be suitably trained to safely set up and operate the instrument to successfully use XRF and obtain the level of data quality required for specific projects. The amount of training required depends on the complexity of the instrument and the associated software.

Niton recommends that the operator have a high school diploma and basic operational training. Field or laboratory technicians are generally qualified to operate this instrument. Additional understanding of soil chemical and physical properties would be valuable for preparing site-specific calibrations and for conducting specialty analyses. The operator of the instrument during the demonstration held an M.S. degree in earth sciences, with less than 1 year of experience in operation of the XLt.

Niton offers free training on the use of field-portable XRF analyzers for lead and other elements. Most classes are 1 day unless otherwise indicated. Classes are offered often (two to six classes per month) at varying locations throughout the U.S. The course materials include instrument theory, operation, and application. In addition, the course material includes radiation safety training, which some states require for licensing on these instruments. The course covers the following topics:

- Radiation safety
- X-ray fluorescence theory
- Hands-on training for lead-in-paint testing
- Hands-on-analysis of coatings for lead and other elements
- On-site analysis of dust wipes, soil (EPA Method 6200), and paint chips
- On-site analysis of worker exposure cassettes for airborne lead(National Institute of Occupational Safety and Health [NIOSH] Method #7702)
- On-site measurement of total suspended particulate (TSP) and fine particulate matter (PM₁₀) filters for air monitoring

Participants are encouraged to bring samples to class to analyze as part of the hands-on exercise for the training. Niton also offers site-specific training by request and will customize the training to the field conditions, matrices, analytes, reporting limits, and data quality levels required for individual project objectives.

Niton has not established written standard operating procedures (SOP) for the preparation or analysis of soil or sediment samples using the XLt. However, the instrument is accompanied by a clear and detailed operating manual that presents the general steps in analyzing soil and other environmental media. Instrument software is also helpful in directing users with intuitive operating menus. Niton and its distributors offer on-site training and telephone support to instrument users on an informal, as needed basis.

In addition to the general instrument operational instruction and training, the operator and data manager must become familiar with Niton's data acquisition software loaded onto the instrument. Niton provides a copy of the NDT PC software for each instrument. Although a PC is not required to acquire data, a laptop PC can be useful because the smaller instrument display can be projected onto the larger PC screen for easier viewing. In addition, data can be simultaneously recorded and stored in the PC, thereby maximizing data collection efficiency while minimizing the potential for lost data or transcription errors.

# 7.9 Secondary Objective 2 — Health and Safety

Included in the health and safety evaluation were the potential risks from: (1) potential radiation hazards from the instrument itself, and (2) exposure to any reagents used in preparing and analyzing the samples. However, the evaluation did not include potential risks from exposure to site-specific hazardous materials, such as sample contaminants, or to physical safety hazards. These factors were excluded because of the wide and unpredictable range of sites and conditions that could be encountered in the field during an actual project application of the instrument.

The XLt contains a miniature silver anode x-ray tube. However, each instrument is equipped with trigger locks and safety measures designed to minimize possible exposure to the x-ray radiation. Niton reports that risks from exposure to radiation are minimal; the radiation measured around the instrument during operation has been recorded at 0.1 milliRems per hour.

The second potential source of risk to XRF instrument operators is exposure to reagent chemicals. However, for the XLt, there are no risks from reagents because no chemical reagents are required for sample preparation.

#### 7.10 Secondary Objective 3 — Portability

Portability depends on the size, weight, number of components, and power requirements of the instrument, and the reagent required. The size of the instrument, including physical dimensions and weight, is presented in Table 6-1. The number of components, power requirements, support structures, and reagent requirements were also recorded. Two distinctions were made during the demonstration regarding portability:

- (1) The instrument was considered fully portable if the dimensions were such that the instrument could be easily brought directly to the sample location.
- (2) The instrument was considered transportable if the dimensions and power requirements were such that the instrument could be moved to a location near the sampling location, but required a larger and more stable environment (for example, a site trailer with AC power and stable conditions).

Based on its dimensions and power requirements, the XLt is defined as fully portable. It is a hand-held unit that can be carried directly to the sampling location for analysis of samples. The XLt is suitable for all types of field analysis, ranging from "point-and-shoot" readings on undisturbed soil surfaces to processed soil samples in plastic bags or sample cups. With an additional instrument stand, the XLt can also be used in a hands-free, bench-top mode. This instrument stand was used during the demonstration.

## 7.11 Secondary Objective 4 — Durability

Durability was evaluated by gathering information on the instrument's warranty and the expected lifespan of the radioactive source or x-ray tube. The ability to upgrade software or hardware also was evaluated. Weather resistance was evaluated by examining the instrument for exposed electrical connections and openings that may allow water to penetrate (for portable instruments only).

The outer construction of the XLt consists of a hardtooled plastic that is durable, weatherproof, and impact-resistant. The instrument is intrinsically tight and 100 percent waterproof. It can be submerged or dropped in water with no damage to the inner workings of the instrument. The external PC can be attached via USB port and cable. If the PC data acquisition system is used, it is recognized that the PC may not be weatherproof and should be used only in a protected environment. In addition, the instrument can be outfitted with wireless communications to further aid in data transfer. However, these modes of operation were not assessed during the demonstration.

Niton provides a 24-month limited warranty for the XLt instrument. The warranty does not cover batteries or accessories. Since x-ray tube sources are new to the world of portable instrumentation, no clear data have been obtained on the useful life that can be assumed. The average lifespan of an x-ray tube in a traditional bench-top device is 3 to 5 years. Therefore, it is generally assumed among developers of portable XRF instrumentation that the useful life of an x-ray tube in these systems will be about 3-5 years.

#### 7.12 Secondary Objective 5 — Availability

Niton was founded in 1987 and has two offices in the U.S. and one office in Germany. Niton reports sales of more than 1,000 new instruments each year both in the U.S. and abroad. The XLt is also available for purchase or rental from a nationwide network of distributors, and many can provide on-site training. The instrument can be repaired, maintained, and calibrated by the distributors or at the factory in Massachusetts. Niton also operates a telephone helpline in both the U.S. and Europe.

## Chapter 8 Economic Analysis

This chapter provides cost information for the Niton XLt 700 Series XRF analyzer. Cost elements that were addressed included instrument purchase or rental, supplies, labor, and ancillary items. Sources of cost information included input from the technology developer and suppliers as well as observations during the field demonstration. Comparisons are provided to average costs for other XRF technologies and for conventional fixedlaboratory analysis to provide some perspective on the relative cost of using the XLt.

#### 8.1 Equipment Costs

Capital equipment costs include either purchase or rental of the XLt and any ancillary equipment that is needed for specific analyses. (See Chapter 6 for a description of available accessories.) Price information for the analyzer and accessories was obtained from Niton as well as from licensed Niton distributors.

The XLt (with miniature x-ray tube) sells for a base cost of \$32,500. This cost includes the standard accessories described in Chapter 6 of this report. The instrument is shipped in a Pelican case, which holds the instrument, environmental test stand, covers, and communication cables. With the addition of the PERFECT software, the cost to purchase the equipment increases to \$35,000. The Bluetooth[©] wireless communication package adds \$1,000 to the equipment cost. A laptop computer may also be used to run the instrument and manage the analytical data at an additional cost.

The rental cost of the XLt instrument varies based on model type and the degree of instrument and software customization (such as site-specific calibrations or quantitation algorithms), as required for specific applications. This economic analysis assumed that the instrument was applied "off the shelf" and that no additional customization costs were included. The XLt analyzer that was used in the demonstration can be rented for \$1,500 per week or \$5,500 per month. Thus, purchase of the instrument could be justified as more cost-effective than rental for field activities that involve more than about 5 months of total field analysis time. The purchase price and shipping cost for the XLt compare favorably with the average costs for all XRF instruments that participated in the demonstration, as shown in Table 8-1.

Cost Element	Niton XLt	XRF Demonstration Average ¹
Shipping	\$240	\$410
Capital Cost (Purchase)	\$32,500	\$54,300
Weekly Rental	\$1,500	\$2,813
Autosampler (for Overnight Analysis)	N/A	N/A

Notes:

¹ Average for all eight instruments in the demonstration

N/A Not available or not applicable for this comparison

The x-ray tube in the XLt is protected by a 2 year warranty and is expected to last for 3 to 5 years, assuming that the instrument is operated 2,000 hours per year. The replacement cost, including a new power supply, would be about \$4,500.

#### 8.2 Supply Costs

The supplies that were included in the cost estimate include sample containers, Mylar[®] film, scoops, snap rings, and disposable gloves. The rate of consumption of these supplies was based on observations during the field demonstration. Unit prices for these supplies were based on price quotes from independent vendors.

During the field demonstration, the XLt was operated for 3 days to complete the analysis of the demonstration sample set (326 samples). The supplies required to process samples were similar for all XRF instruments that participated in the demonstration and were estimated to cost about \$245 for 326 samples or \$0.75 per sample.

#### 8.3 Labor Costs

Labor costs were estimated based on the total time required by the field team to complete the analysis of all 326 samples and the number of people in the field team, while making allowances for field team members that had responsibilities other than sample processing during the demonstration. For example, some developers sent sales representatives to the demonstration to communicate with visitors and provide outreach services; this type of staff time was not included in the labor cost analysis.

While overall labor costs were based on the total time required to process samples, the time required to complete each definable activity was also measured during the field demonstration. These activities included:

- Initial set-up and calibration
- Sample preparation
- Sample analysis
- Daily shutdown and startup
- End of project packing

The "total processing time per sample" was calculated as the sum of all these activities assuming that the activities were conducted sequentially; therefore, it represents how much time it would take a single trained analyst to complete these activities. However, the "total processing time per sample" does not include activities that were less definable in terms of the amount of time taken (such as data management and procurement of supplies) and is therefore not a true total.

The time to complete each activity using the XLt is compared with the average of all XRF instruments in Table 8-2 and with the range of all XRF instruments in Figure 8-1. Specifically, the XLt compared favorably against the other XRF instruments, exhibiting lower-than-average times for all activities except for daily shutdown and startup.

# Table 8-2. Time Required to CompleteAnalytical Activities1

Activity	Niton XLt	Average
Initial Set up and		
Calibration	30	54
Sample Preparation	2.0	3.1
Sample Analysis	2.5	6.7
Daily Shut		
Down/Start Up	10	10
End of Project		
Packing	10	43
Total Processing		
Time per Sample	4.7	10.0

Notes:

¹ All estimates are in minutes

² Average for all eight XRF instruments in the demonstration

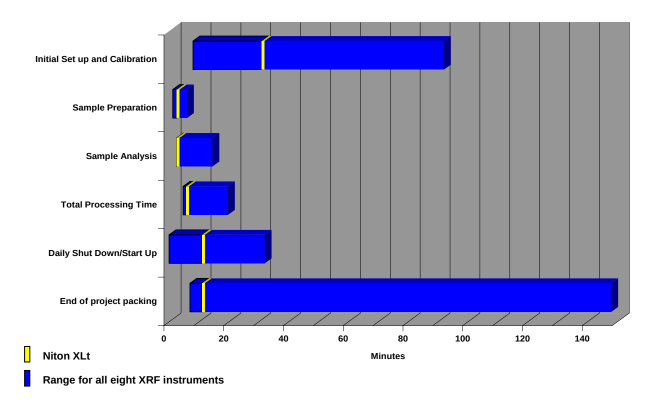


Figure 8-1. Comparison of activity times for the XLt versus other XRF instruments.

The Niton field team expended about 29 labor hours to complete all sample processing activities during the field demonstration using the XLt. This was much lower than the overall average of 69 hours for all instruments that participated in the demonstration. The primary reasons that labor hours were lower for the XLt include:

- Instrument run times (2 minutes) were significantly less than other instruments.
- The instrument operation was simple enough that a single technician performed all sample preparation and analysis activities during the demonstration.
- The software-based automation of the XLt allowed the operator to reduce the data for completed sample batches on a laptop PC while the instrument was processing a new batch.

Overall, the XLt exhibited the fastest sample processing time and the lowest labor hours of all the instruments participating in the demonstration.

## 8.4 Comparison of XRF Analysis and Reference Laboratory Costs

Two scenarios were evaluated to compare the cost for XRF analysis using the XLt with the cost of fixedlaboratory analysis using the reference methods. Both scenarios assumed that 326 samples were to be analyzed, as in the field demonstration. The first scenario assumed that only one element was to be measured in a metal-specific project or application (for example, lead in soil, paint, or other solids) for comparison to laboratory per-metal unit costs. The second scenario assumed that 13 elements were to be analyzed, as in the field demonstration, for comparison to laboratory costs for a full suite of metals. Typical unit costs for fixed-laboratory analysis using the reference methods were estimated using average costs from Tetra Tech's basic ordering agreement with six national laboratories. These unit costs assume a standard turnaround time of 21 days and standard hard copy and electronic data deliverables that summarize results and raw analytical data. No costs were included for field labor that would be specifically associated with off-site fixed laboratory analysis, such as sample packaging and shipment.

The cost for XRF analysis using the XLt was based on equipment rental for 1 week, along with labor and supplies estimates established during the field demonstration. Labor costs were added for drying, grinding, and homogenizing the samples (estimated at 10 minutes per sample) since these additional steps in sample preparation are required for XRF analysis but not for analysis in a fixed laboratory. A typical cost for managing investigation-derived waste (IDW), including general trash, personal protective equipment, wipes, and soil, was also added to the cost of XRF analysis because IDW costs are included in the unit cost for fixed-laboratory analysis. The IDW management cost was based on the average IDW disposal cost per instrument during the demonstration because IDW generation did not vary significantly between instruments. Since the cost for

XRF analysis of one element or multiple elements does not vary significantly (all target elements are determined simultaneously when a sample is analyzed), the XRF analysis cost was not adjusted for one element versus 13 elements.

Table 8-3 summarizes the costs for the XLt versus the cost for analysis in a fixed laboratory. This comparison shows that the XLt compares favorably to a fixed laboratory in terms of overall cost, particularly when a large number of elements are to be determined. Use of the XLt will likely produce additional cost savings because analytical results will be available within a few hours after samples are collected, thereby expediting project decisions and reducing or eliminating the need for additional mobilizations.

The total cost for the XLt in the example scenario (326 samples) was estimated at \$5,706. This estimate compares with the average of \$8,932 for all XRF instruments that participated in the demonstration. However, it should be noted that bench-top instruments, which typically cost more than hand-held instruments like the XLt, were included in the calculation of the average cost for all XRF instruments. The XLt cost was slightly lower than the three other hand-held instruments participating in the demonstration.

			Unit	
Niton XLt	Quantity	Item	Rate	Total
Shipping	1	Roundtrip	\$240	\$240
Weekly Rental	1	Week	\$1,500	\$1,500
Supplies	326	Sample	\$0.75	\$245
Labor	83	Hours	\$43.75	\$3,631
IDW	N/A	N/A	N/A	\$90
Total Niton XLt Analysis Cost				\$5,706
Fixed Laboratory (1 element)				
(EPA Method 6010, ICP-AES)	326	Sample	\$21	\$6,846
<b>Total Fixed Laboratory Costs (1 element)</b>				\$6,846
Fixed Laboratory (13 elements)				
Mercury (EPA Method 7471, CVAA)	326	Sample	\$36	\$11,736
All other Elements (EPA Method 6010, ICP-				
AES)	326	Sample	\$160	\$52,160
Total Fixed Laboratory Costs (13 elements)				\$63,896

 Table 8-3. Comparison of XRF Technology and Reference Method Costs

## Chapter 9 Summary of Technology Performance

The preceding chapters of this report document that the evaluation design succeeded in providing detailed performance data for the Niton XLt 700 Series XRF analyzer. The evaluation design incorporated 13 target elements, 70 distinct sample blends, and a total of 326 samples. The blends included both soil and sediment samples from nine sampling locations. A rigorous program of sample preparation and characterization, reference laboratory analysis, QA/QC oversight, and data reduction supported the evaluation of XRF instrument performance.

One important aspect of the demonstration was the sample blending and processing procedures (including drying, sieving, grinding, and homogenization) that significantly reduced uncertainties associated with the demonstration sample set. These procedures minimized heterogeneity impacts on method precision and on the comparability between XRF data and reference laboratory data. Thus, before XRF is used for largescale data collection, project teams are encouraged to assess the effects of sampling uncertainty on data quality and to adopt appropriate sample preparation protocols, particularly if the project will involve comparisons to other methods (such as off-site laboratories). An initial pilot-scale method evaluation, carried out in cooperation with an instrument vendor, can yield site-specific standard operating procedures for sample preparation and analysis that ensure the XRF method will meet data quality needs, such as accuracy and sensitivity requirements. A pilot study can also help the project team develop an initial understanding of the degree of correlation between field and laboratory data. This type of study is particularly appropriate for sampling programs that will involve complex soil or sediment matrices with high concentrations of multiple elements because the demonstration found that XRF performance was more variable under these conditions. Initial pilot studies can also be used to

develop site-specific calibrations, in accordance with EPA Method 6200, that adjust instrument algorithms to compensate for matrix effects.

The findings of the evaluation of the XLt analyzer for each primary and secondary objective are summarized in Tables 9-1 and 9-2. A performance comparison between the XLt and the combined performance of all eight vendors participating in the XRF technology evaluation program is provided in Figure 9-1. The comparisons in Figure 9-1 indicate that relative to the program as a whole, the XLt showed:

- Equivalent or better MDLs for 8 of the 13 target metals (exceptions included antimony, cadmium, copper, nickel, and selenium).
- Equivalent or better accuracy (RPDs) for 10 of the 13 target metals (exceptions included antimony, mercury, and nickel). Moreover, when RPDs for antimony are calculated versus sample spike levels rather than reference laboratory data (which may be biased low), accuracy for antimony improves to better than the program as whole.
- Equivalent or better precision (RSDs) for 9 of the 13 target metals (exceptions included antimony, copper, nickel, and silver).

As a hand-held instrument, the XLt is fully portable and can be operated in the hand-held mode at a sampling site. The reasons for the better-thanaverage sensitivity, accuracy, and precision are not known with any certainty but may relate to Niton's well-established method procedures, calibration protocols, and quantification algorithms for environmental applications.

<ul> <li>P1: Method Detection Limits</li> <li>Mean MDLs for the 12 target elements ranged as follows (iron was included in the MDL evaluation): <ul> <li>MDLs of 1 to 20 ppm: arsenic, lead, mercury, selenium, an vanadium.</li> <li>MDLs of 20 to 50 ppm: copper, silver, and zinc.</li> <li>MDLs of 50 to 100 ppm: chromium and nickel.</li> <li>MDLs for antimony (216 ppm) and cadmium (105 ppm) were based limited data (two MDL samples) and may be biased high because of high concentrations of other elements in these samples.</li> </ul> </li> </ul>	d
<ul> <li>Detection Limits</li> <li>included in the MDL evaluation): <ul> <li>MDLs of 1 to 20 ppm: arsenic, lead, mercury, selenium, an vanadium.</li> <li>MDLs of 20 to 50 ppm: copper, silver, and zinc.</li> <li>MDLs of 50 to 100 ppm: chromium and nickel.</li> <li>MDLs of greater than 100 ppm: antimony and cadmium.</li> </ul> </li> <li>MDLs for antimony (216 ppm) and cadmium (105 ppm) were based limited data (two MDL samples) and may be biased high because of high concentrations of other elements in these samples.</li> <li>No significant differences were noted between MDLs for soil and set</li> </ul>	d
<ul> <li>vanadium.</li> <li>MDLs of 20 to 50 ppm: copper, silver, and zinc.</li> <li>MDLs of 50 to 100 ppm: chromium and nickel.</li> <li>MDLs of greater than 100 ppm: antimony and cadmium.</li> <li>MDLs for antimony (216 ppm) and cadmium (105 ppm) were based limited data (two MDL samples) and may be biased high because of high concentrations of other elements in these samples.</li> <li>No significant differences were noted between MDLs for soil and set</li> </ul>	
<ul> <li>MDLs of 20 to 50 ppm: copper, silver, and zinc.</li> <li>MDLs of 50 to 100 ppm: chromium and nickel.</li> <li>MDLs of greater than 100 ppm: antimony and cadmium.</li> <li>MDLs for antimony (216 ppm) and cadmium (105 ppm) were based limited data (two MDL samples) and may be biased high because of high concentrations of other elements in these samples.</li> <li>No significant differences were noted between MDLs for soil and set</li> </ul>	
<ul> <li>MDLs of 50 to 100 ppm: chromium and nickel.</li> <li>MDLs of greater than 100 ppm: antimony and cadmium.</li> <li>MDLs for antimony (216 ppm) and cadmium (105 ppm) were based limited data (two MDL samples) and may be biased high because of high concentrations of other elements in these samples.</li> <li>No significant differences were noted between MDLs for soil and set</li> </ul>	
<ul> <li>MDLs of greater than 100 ppm: antimony and cadmium.</li> <li>MDLs for antimony (216 ppm) and cadmium (105 ppm) were based limited data (two MDL samples) and may be biased high because of high concentrations of other elements in these samples.</li> <li>No significant differences were noted between MDLs for soil and set</li> </ul>	
<ul> <li>MDLs for antimony (216 ppm) and cadmium (105 ppm) were based limited data (two MDL samples) and may be biased high because of high concentrations of other elements in these samples.</li> <li>No significant differences were noted between MDLs for soil and set</li> </ul>	
<ul> <li>limited data (two MDL samples) and may be biased high because of high concentrations of other elements in these samples.</li> <li>No significant differences were noted between MDLs for soil and set</li> </ul>	1
<ul><li>high concentrations of other elements in these samples.</li><li>No significant differences were noted between MDLs for soil and set</li></ul>	
• No significant differences were noted between MDLs for soil and set	the
• The calculated MDLs were comparable to the XLt's statistical LOD	s and
were generally below reference MDL data from EPA Method 6200.	
P2: Accuracy and • Median RPDs between the XLt and reference laboratory data reveal	ed the
Comparability following, with lower RPDs indicating greater accuracy:	
• RPDs of 1 to 10 percent: copper and selenium.	
<ul> <li>RPDs of 10 to 25 percent: arsenic, cadmium, chromium, ire</li> </ul>	on, lead,
silver, and zinc.	
• RPDs of 25 to 50 percent: nickel and vanadium.	
• RPDs greater than 50 percent: antimony and mercury.	
• Data review indicated that the reference laboratory results for some	-
demonstration samples may be biased low for antimony due to the v	-
of the spiking compounds used. Data review indicated that the refer	
laboratory results for some spiked demonstration samples may be bi	
low for antimony due to the volatility of the spiking compounds use	
RPDs for antimony were quite high when the XLt data was compare	
reference laboratory data (median RPD of 125 percent), but improve	
considerably when compared to certified spike values (median RPD percent). Thus, the XLt appeared to be more accurate with respect t	
true concentration of antimony than the reference laboratory.	0 the
	rations
RPDs increased (that is, accuracy declined) with increasing concent of silver and vanadium. RPDs for arsenic and cadmium were also s	
high in some high concentration matrixes.	Keweu
<ul> <li>Correlation plots relative to reference laboratory data indicated:</li> </ul>	
<ul> <li>High correlations for 9 of the 13 target elements.</li> </ul>	
<ul> <li>Positive biases for arsenic, lead, nickel, and silver.</li> </ul>	
<ul> <li>Negative biases for mercury and vanadium.</li> </ul>	
P3: Precision     • Median RSDs were good for all metals as follows:	
• RSDs of 0 to 5 percent: cadmium, iron, lead, selenium, and	zinc
<ul> <li>RSDs of 6 to 5 percent: catinitali, non, read, scientiali, and</li> <li>RSDs of 5 to 10 percent: antimony, arsenic, chromium, cor</li> </ul>	
mercury, nickel, silver, and vanadium.	,
• RSDs greater than 10 percent: none.	
<ul> <li>Median RSDs for the XLt were slightly lower than the reference lab</li> </ul>	oratory.

 Table 9-1. Summary of Niton XLt Performance – Primary Objectives

## Table 9-1. Summary of Niton XLt Performance – Primary Objectives

Objective	Performance Summary
P4: Effects of Sample Interferences	<ul> <li>High relative lead concentrations (more than 10 times) reduced accuracy for arsenic results. Median RPDs for arsenic increased from 13 percent to 50 percent, and a larger positive bias was observed in the arsenic results, as lead concentrations increased.</li> <li>High relative copper concentrations (more than 10 times) similarly reduced accuracy for nickel (median RPDs for nickel increased from 34 percent to 71 percent, and a larger positive bias was observed).</li> </ul>
P5: Effects of Soil Type	<ul> <li>Low relative accuracy was observed for multiple elements in blends of roaster slag from the Wickes Smelter site, which contained high overall element concentrations.</li> <li>High RPDs were also observed for multiple elements in blends from the Leviathan Mine site, which included precipitates with high iron concentrations.</li> </ul>
P6: Sample Throughput	<ul> <li>With an average sample preparation time of 2.0 minutes and an instrument analysis time of 2.5 minutes per sample, the total processing time was 4.7 minutes per sample.</li> <li>A maximum sample throughput of 146 samples per day was achieved during an extended work day. A more typical sample throughput was estimated to be 91 samples per day for an 8-hour work day.</li> <li>The total processing time was the lowest and the daily sample throughput was the highest of all eight instruments in the demonstration.</li> </ul>
P7: Costs	<ul> <li>The base instrument cost was \$32,500 (purchase) or \$1,500/week (rental), plus \$240 shipping. This cost includes peripherals such as an instrument stand, protective covers, communication cables, and 110 volt AC adapter.</li> <li>The XLt instrument operator expended approximately 29 labor hours to complete the processing of the demonstration sample set (326 samples). This was significantly lower than the average for all participating XRF instruments of 69 labor hours.</li> <li>Using the 1-week rental cost and adding labor and miscellaneous costs (\$485 for shipping and supplies), a total project cost of \$5,706 was estimated for a project the size of the demonstration. In comparison, the project cost averaged \$8,932 for all participating XRF instruments and the cost for fixed-laboratory analysis of all samples for 13 elements was \$63,896.</li> </ul>

Objective	Performance Summary
S1: Training Requirements	<ul> <li>Field or laboratory technicians with a high school diploma and basic operational training are generally qualified to operate the XLt.</li> <li>Niton offers free training on the use of field-portable XRF analyzers for lead and other elements. Most are 1-day classes offered at varying locations throughout the U.S. (two to six classes per month).</li> <li>Niton also offers site-specific training by request and will customize the training to the field conditions, matrices, analytes, reporting limits, and data quality objectives for a project.</li> </ul>
S2: Health and Safety	<ul> <li>The XLt is equipped with safety measures to minimize possible exposure to emissions from the x-ray tube. Niton reports that the resulting risks from radiation exposure are minimal (less than 0.1 milliRem per hour).</li> <li>No chemicals are used during sample preparation or analysis that would pose potential hazards.</li> </ul>
S3: Portability	<ul> <li>Based on dimensions, weight, and power requirements, the XLt is a fully portable instrument. It can be used as a hand-held unit to analyze undisturbed soil or bagged samples.</li> <li>With an additional instrument stand, the XLt can be used in a hands-free, bench-top mode.</li> </ul>
S4: Durability	<ul> <li>Niton instruments have a 24-month limited warranty for parts and labor.</li> <li>The vendor estimates that the useful life of the x-ray tube source is 3-5 years.</li> <li>The Niton XLt is impact-resistant and weatherproof. It is designed to operate under wet and dirty conditions and may be used in adverse weather conditions as it is dust and splash resistant.</li> </ul>
S5: Availability	<ul> <li>Niton produces and sells more than 1,000 instruments a year through offices in the U.S. and Germany.</li> <li>Rental instruments and supporting software are also available from Niton or from numerous distributors throughout the U.S.</li> </ul>

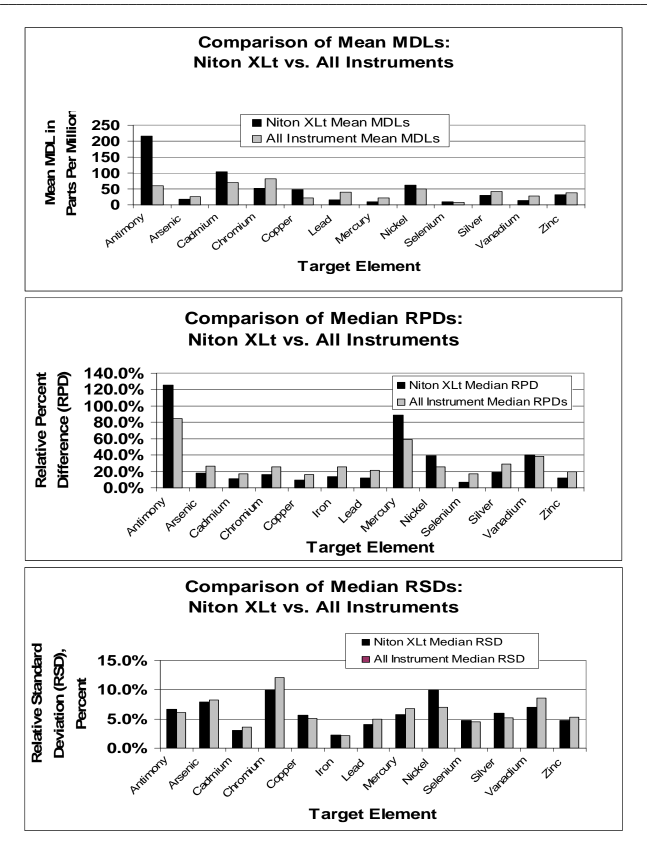


Figure 9-1. Method detection limits (sensitivity), accuracy, and precision of the XLt in comparison to the average of all eight XRF instruments.

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## Chapter 10 References

- Gilbert, R.O. 1987. Statistical Methods for Environmental Pollution Monitoring. Van Nostrand Reinhold, New York.
- Tetra Tech EM Inc. 2005. *Demonstration and Quality Assurance Plan*. Prepared for U.S. Environmental Protection Agency, Superfund Innovative Technology Evaluation Program. March.
- U.S. Environmental Protection Agency (EPA). 1996a. TN Spectrace TN 9000 and TN Pb Field Portable X-ray Fluorescence Analyzers. EPA/600/R-97/145. March.
- EPA. 1996b. Field Portable X-ray Fluorescence Analyzer HNU Systems SEFA-P. EPA/600/R-97/144. March.
- EPA. 1996c. Test Methods for Evaluating Solid Waste, Physical/Chemical Methods (SW-846). December.
- EPA. 1998a. Environmental Technology Verification Report; Field Portable X-ray Fluorescence Analyzer, Metorex X-Met 920-MP. EPA/600/R-97/151. March.
- EPA. 1998b. Environmental Technology Verification Report; Field Portable X-ray Fluorescence Analyzer, Niton XL Spectrum Analyzer. EPA/600/R-97/150. March.

- EPA. 1998c. Scitect MAP Spectrum Analyzer Field Portable X-Ray Fluorescence Analyzers. EPA/600/R-97/147. March.
- EPA. 1998d. *Metorex X-MET* 920-P and 940 Field Portable X-ray Fluorescence Analyzers. EPA/600/R-97/146. March.
- EPA. 1998e. EPA Method 6200, from "Test Methods for Evaluating Solid Waste, Physical/Chemical Methods (SW-846), Update IVA. December.
- EPA. 2000. Guidance for Data Quality Assessment: Practical Methods for Data Analysis. EPA QA/G-9 QA00 Update. EPA/600/R-96/084. July.
- EPA. 2004a. Innovative Technology Verification Report: Field Measurement Technology for Mercury in Soil and Sediment - Metorex's X-MET[®] 2000 X-Ray Fluorescence Technology. EPA/600/R-03/149. May.
- EPA. 2004b. Innovative Technology Verification Report: Field Measurement Technology for Mercury in Soil and Sediment - Niton's XLi/XLt 700 Series X-Ray Fluorescence Analyzers. EPA/600/R-03/148. May.
- EPA. 2004c. USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review. Final. OSWER 9240.1-45. EPA 540-R-04-004. October.

## APPENDIX A

## **VERIFICATION STATEMENT**

# **UNITED STATES ENVIRONMENTAL PROTECTION AGENCY**

Office of Research and Development Washington, DC 20460



## SITE Monitoring and Measurement Technology Program Verification Statement

TECHNOLOGY TYPE:	X-ray Fluorescence (XRF) Analyzer
APPLICATION:	MEASUREMENT OF TRACE ELEMENTS IN SOIL AND SEDIMENT
TECHNOLOGY NAME:	XLt 700 Series XRF Analyzer
COMPANY:	Niton Analyzer
COMPANY:	Niton Analyzers, A Division of Thermoelectron
ADDRESS:	900 Middlesex Turnpike, Building #8
Telephone:	Billerica, MA 01821 (800) 875-1578
Fax:	978-670-7430
Email:	dmercuro@niton.com
Internet:	www.niton.com

## VERIFICATION PROGRAM DESCRIPTION

The U.S. Environmental Protection Agency (EPA) created the Superfund Innovative Technology Evaluation (SITE) Monitoring and Measurement Technology (MMT) Program to facilitate deployment of innovative technologies through performance verification and information dissemination. The goal of this program is to further environmental protection by substantially accelerating the acceptance and use of improved and cost-effective technologies. The program assists and informs those involved in designing, distributing, permitting, and purchasing environmental technologies. This document summarizes the results of a demonstration of the Niton XLt 700 Series portable x-ray fluorescence (XRF) analyzer for the analysis of 13 target elements in soil and sediment, including antimony, arsenic, cadmium, chromium, copper, iron, lead, mercury, nickel, selenium, silver, vanadium, and zinc.

## **PROGRAM OPERATION**

Under the SITE MMT Program, with the full participation of the technology developers, EPA evaluates and documents the performance of innovative technologies by developing demonstration plans, conducting field tests, collecting and analyzing demonstration data, and preparing reports. The technologies are evaluated under rigorous quality assurance protocols to produce well-documented data of known quality. EPA's National Exposure Research Laboratory, which demonstrates field sampling, monitoring, and measurement technologies, selected Tetra Tech EM Inc. as the verification organization to assist in field testing technologies for measuring trace elements in soil and sediment using XRF technology.

## **DEMONSTRATION DESCRIPTION**

The field demonstration of eight XRF instruments to measure trace elements in soil and sediment was conducted from January 24 through 28, 2005, at the Kennedy Athletic, Recreational and Social (KARS) Park, which is part of the Kennedy Space Center on Merritt Island, Florida. A total of 326 samples were analyzed by each XRF instrument, including the XLt, during the field demonstration. These samples were derived from 70 different blends and spiked blends of soil and sediment collected from nine sites across the U.S. The sample blends were thoroughly dried, sieved, crushed, mixed, and characterized before they were used for the demonstration. Some

blends were also spiked to further adjust and refine the concentration ranges of the target elements. Between three and seven replicate samples of each blend were included in the demonstration sample set and analyzed by the technology developers during the field demonstration.

Shealy Environmental Services, Inc. (Shealy), of Cayce, South Carolina, was selected as the reference laboratory to generate comparative data in evaluation of XRF instrument performance. Shealy analyzed all demonstration samples (both environmental and spiked) concurrently with the developers during the field demonstration. The samples were analyzed by inductively coupled plasma–atomic emission spectroscopy (ICP-AES) using EPA SW-846 Method 3050B/6010B and by cold vapor atomic absorption spectroscopy (CVAA) using EPA SW-846 Method 7471A (mercury only).

This verification statement provides a summary of the evaluation results for the Niton XLt 700 Series XRF analyzer. More detailed discussion can be found in the *Innovative Technology Verification Report – XRF Technologies for Measuring Trace Elements in Soil and Sediment: Niton XLt XRF Analyzer* (EPA/540/R-06/004).

## **TECHNOLOGY DESCRIPTION**

XRF spectroscopy is an analytical technique that exposes a sample (soil, alloy metal, filters, other solids, and thin samples) to an x-ray source. The x-rays from the source have the appropriate excitation energy that causes elements in the sample to emit characteristic x-rays. A qualitative elemental analysis is possible from the characteristic energy, or wavelength, of the fluorescent x-rays emitted. A quantitative elemental analysis is possible from the number (intensity) of x-rays at a given wavelength.

The XLt is a field portable instrument that features a miniaturized x-ray tube source and a Peltier-cooled Si-PiN xray detector. The analyzer is factory-calibrated and programmed to analyze and report 25 elements. An optional software package called PERFECT (Programmable Excitation by Regulation of Filters, Energy, Current, and Time) is also available for analysis and reporting of light elements (including vanadium and chromium) and optimizing of the detection limits for specific applications. Other features include an integrated touch-screen display; integrated bar code reader and virtual keypad; remote operation and custom report generation capability from a Windowsbased PC; shielded bench-top test stand; and Bluetooth wireless communication to a laptop or personal data assistant (PDA).

In soil and sediment applications, the XLt is designed to be used as either a hand-held instrument for analysis of undisturbed or bagged soil, or as a bench-top instrument in a test stand for analysis of processed samples. The analyzer is factory-calibrated, and it also accepts user-generated empirical calibrations, if required, for specific data needs and applications.

#### **VERIFICATION OF PERFORMANCE**

**Method detection limit:** MDLs were calculated using seven replicate analyses from each of 12 low-concentration sample blends, according to the procedure described in Title 40 Code of Federal Regulations (CFR) Part 136, Appendix B, Revision 1.11. A mean MDL was further calculated for each element. The ranges into which the mean MDLs fell for the XLt are listed below (lower MDLs indicate higher sensitivity).

Relative Sensitivity	Mean MDL	Target Elements
High	1 – 20 ppm	Arsenic, Lead, Mercury, Selenium, and Vanadium.
Moderate	20 – 50 ppm	Copper, Silver, and Zinc.
Low	50 – 100 ppm	Chromium and Nickel.
Very Low	> 100 ppm	Antimony and Cadmium.

Notes: ppm = Parts per million. Iron was not included in the MDL evaluation.

Accuracy: The determination of accuracy was based on the agreement of the XLt results with the reference laboratory data. Accuracy was assessed by calculating the absolute relative percent difference (RPD) between the

mean XRF concentration and the mean reference laboratory concentration for each blend. Accuracy of the XLt was classified from high to very low for the different target elements as indicated in the table below based on the overall median RPDs calculated for the demonstration.

<b>Relative Accuracy</b>	Median RPD	Target Elements
High	0% - 10%	Copper and Selenium.
Moderate	10% - 25%	Arsenic, Cadmium, Chromium, Iron, Lead, Silver, and Zinc.
Low	25% - 50%	Nickel and Vanadium.
Very Low	> 50%	Antimony* and Mercury.

* Calculation of RPDs versus sample spike concentrations rather than reference laboratory results (due to potential low bias in the reference laboratory results for antimony) improves accuracy from Very Low to Moderate.

Accuracy was also assessed through correlation plots between the mean XLt and mean reference laboratory concentrations for the different sample blends. Correlation coefficients ( $r^2$ ) for linear regression analysis of the plots are summarized below, along with the bias apparent from the plots in the XRF data versus the reference laboratory data.

	Antimony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury	Nickel	Selenium	Silver	Vanadium	Zinc
Correlation	0.88	0.95	0.98	0.95	0.87	0.94	0.95	0.98	0.98	0.99	0.71	0.51	0.76
Bias	High	High					High	Low	High			Low	High

Note: -- = No significant bias.

**Precision:** Replicates were analyzed for all sample blends. Precision was determined by calculating the standard deviation of the replicates, dividing by the average concentration of the replicates, and multiplying by 100% to give the relative standard deviation (RSD) for each blend. Precision of the XLt was classified from high to very low for the different target elements as indicated in the table below based on the overall median RSDs. These results indicated a higher level of precision in the XLt data than in the reference laboratory data for 11 of the 13 target elements.

<b>Relative Precision</b>	Median RSD	Target Elements
High	0% - 5%	Cadmium, Iron, Lead, Selenium, and Zinc.
Moderate	5% - 10%	Antimony, Arsenic, Chromium, Copper, Mercury, Nickel, Silver, and Vanadium.
Low	10% - 20%	None.
Very Low	> 20%	None.

**Effects of Interferences:** The RPDs from the accuracy evaluation were further grouped and compared for a few elements of concern (arsenic, nickel, copper, and zinc) based on the relative concentrations of potentially interfering elements. This evaluation found that high relative lead concentrations (> 10X) reduced the accuracy for arsenic results in the XLt data set, increasing the median RPDs for arsenic from 13% (in the "moderate" range) to 50% (in the "low" range) as the lead concentration increases. Similarly, increasing copper concentrations reduce accuracy for nickel, increasing median RPDs for nickel from 34% ("low") to 71% ("very low"). The interferences produced increasing positive biases in the arsenic and nickel results.

**Effects of Soil Characteristics:** The RPDs from the accuracy evaluation were also further evaluated relative to sampling site and soil type. This evaluation found low relative accuracy from the XLt for multiple elements in soils impacted by minerals processing, which contained high overall element concentrations or high iron concentrations. In particular, high RPD outliers were observed in the XLt data set for blends of roaster slag from the Wickes Smelter site, and for blends from the Leviathan Mine that were impacted by iron-containing precipitates.

**Sample Throughput:** The total processing time per sample was estimated at 4.7 minutes, which included 2.0 minutes of sample preparation and 2.5 minutes of instrument analysis time. The instrument analysis time and total processing time were the lowest among the eight instruments participating in the demonstration. A sample throughput of 91 samples per 8-hour work day was estimated. As noted above, however, the sample blends had undergone rigorous pre-processing before the demonstration. Sample throughput would have decreased if these sample preparation steps (grinding, drying, sieving) had been performed during the demonstration; these steps can add from 10 minutes to 2 hours to the sample preparation time.

**Costs:** A cost assessment identified a base purchase cost of \$32,500 and typical rental cost of \$1,500/week, plus \$240 shipping, for the XLt. A total cost of \$5,706 (with a labor cost of \$3,631 at \$43.75/hr) was estimated for a project similar to the demonstration (326 samples of soil and sediment). In comparison, the project cost averaged \$8,932 for all eight XRF instruments participating in the demonstration, and \$63,896 for fixed-laboratory analysis of all samples for the 13 target elements.

**Skills and training required:** Field or laboratory technicians with a high school diploma are generally qualified to operate the XLt. Niton offers free 1-day training course on the use of field portable XRF analyzers for lead and other elements. Niton and its distributors offer informal on-site training for specific customer and applications. In addition, toll-free telephone support is also available.

**Health and safety aspects:** Health risks associated with exposure to X-ray tube emissions from the XLt are minimized through shielding and other safety measures; the vendor reports that the resulting risks from radiation exposure are very minimal (0.1 mrem/hr). No other risks or potential hazards associated with instrument operation could be identified.

**Portability:** Based on dimensions, weight, and power requirements, the XLi is a fully portable instrument. It can be used as a hand-held unit to analyze undisturbed soil or bagged samples. With an available instrument stand, the XLi can be used in a hands-free, bench-top mode.

**Durability:** The instrument housing is impact-resistant and completely sealed for protection from moisture and dust. It is designed to operate under wet and dirty conditions and can be submerged in water. Niton instruments have a 24-month limited warranty for parts and labor. The useful life of the X-ray tube source is estimated by the vendor to be 3-5 years.

**Availability:** Instruments, accessories, and supporting software are available for purchase or rental from Niton's offices in the U.S. and Germany, or from numerous distributors throughout the U.S.

## **RELATIVE PERFORMANCE**

The overall performance of the XLt relative to the average of all eight XRF instruments that participated in the demonstration is shown below:

	Antimony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury	Nickel	Selenium	Silver	Vanadium	Zinc
Sensitivity	0	•	ο	•	ο	NC	•	•	0	ο	•	•	•
Accuracy	0	•	•	٠	•	٠	•	0	0	•	•	Same	•
Precision	0	Same	•	٠	0	Same	•	•	0	Same	0	٠	Same
Key:	•	Better	0	Worse	NC	No MD	L Calcula	ated.					

NOTICE: Verifications are based on an evaluation of technology performance under specific, predetermined criteria and the appropriate quality assurance procedures. EPA makes no expressed or implied warranties as to the performance of the technology and does not certify that a technology will always operate as verified. The end user is solely responsible for complying with any and all applicable federal, state, and local requirements.

## **APPENDIX B**

## **DEVELOPER DISCUSSION**

## **DEVELOPER DISCUSSION**

Thermo Electron NITON analyzers would like to thank the EPA and SITE program for the opportunity to demonstrate our instrument's effectiveness for trace elemental analysis in soil and sediments. Overall, it proved to be a useful exercise in determining our instrument's capabilities for field analysis. We would like to thank everyone involved as we found the staff incredibly helpful in ensuring that the performance of our analyzers was appropriately documented. Overall, Thermo Electron feels that the report adequately represents the utility of the NITON analyzers in testing soils and sediments for common contaminants.

Our published limits of detection (LOD) for the 13 elements included in this study all correspond well with the results in the report with the exception of chromium and cadmium. The method detection limits (MDL) for chromium and cadmium were both reported higher than anticipated. However, given that most of these samples had been intentionally spiked with high levels of various elements, it was not surprising that our performance was inferior to what we typically expect in "real" samples. In this study, samples were required to serve more than one purpose. Instead of merely attempting to determine a detection limit, one sample might be put to the task of LOD and multi-element interference analysis. For elements such as antimony and cadmium, this proved to be a problem for our analyzer.

Our analyzers report actual 3 sigma detection limits (<XXX) for a given element instead of a value and error. These LOD's are sample specific and are calculated by the analyzer's algorithm. The algorithm is based on the statistical analysis of the background and the relative element concentrations in the sample analyzed. The way in which the SITE group calculates the MDL for any given sample does not allow them to use the <LOD we provide. Therefore, it proved challenging to determine MDLs for some elements when very few of the samples had concentrations high enough to be reported as real numbers.

As the data indicates, a few elements showed a bias versus the laboratory data. These biases may exist because our analyzers are factory calibrated using NIST traceable standards. They contain levels typical of what is usually found in typical remediation situations. The supplied samples in this evaluation were "real" samples that had been altered and/or spiked to meet certain concentration goals. Whatever the case, system biases can easily be remedied by inserting correction factors directly into the analyzer, a process that is completed in a matter of seconds. Using our CorrectCal software, "on-the-fly" adjustment can be made as needed. Additionally, empirical calibrations can be created and loaded directly onto the analyzer. However, many previously characterized samples are required to set up an empirical calibration for 13 elements. For this evaluation we presented the analyzer in a way that accurately represents the standard off-the-shelf factory calibration and modes of use.

The best indicator of an analyzer's performance is its ability to provide precise results. As the data generated during the SITE program show, the median Relative Standard Deviation (RSD) for the XLt 700 was lower than the RSD calculated for the reference laboratory data for all of the 13 target elements. Given good precision, and the ability to make site specific adjustments, required accuracy can be easily achieved.

To request additional information or an on-site demonstration of this or any of our analyzers, please contact Thermo Electron at 978-670-7460 or visit our website at <u>www.niton.com</u>.

## **APPENDIX C**

DATA VALIDATION SUMMARY REPORT

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## **APPENDIX**

DATA VALIDATION REPORTS

## ABBREVIATIONS AND ACRONYMS

CCV	Continuing calibration verification
CVAA	Cold vapor atomic absorption
DVSR	Data validation summary report
EPA	U.S. Environmental Protection Agency
FAR	Federal acquisition regulations
ICP-AES	Inductively coupled plasma-atomic emission spectroscopy
ICS	Interference check sample
ICV	Initial calibration verification
LCS	Laboratory control sample
LCSD	Laboratory control sample duplicate
MDL	Method detection limit
mg/kg	Milligram per kilogram
MS	Matrix spike
MSD	Matrix spike duplicate
PARCC	Precision, accuracy, representativeness, completeness, and comparability
PQL	Practical quantitation limit
QA/QC	Quality assurance and quality control
QAPP	Quality assurance project plan
QC	Quality control
RSD	Relative standard deviation
RPD	Relative percent difference
SDG	Sample delivery group
Shealy	Shealy Environmental Services, Inc.
SITE	Superfund Innovative Technology Evaluation
Tetra Tech	Tetra Tech EM Inc.
XRF	X-ray fluorescence

## **1.0 INTRODUCTION**

This data validation summary report (DVSR) summarizes the reference laboratory quality control (QC) data gathered during the x-ray fluorescence (XRF) technologies demonstration conducted under the U.S. Environmental Protection Agency (EPA) Superfund Innovative Technology Evaluation (SITE) program. The reference laboratory was procured following the federal acquisition regulations (FAR) and an extensive selection process. Shealy Environmental Services, Inc. (Shealy), of Cayce, South Carolina, was selected as the reference laboratory for this project. Thirteen target analytes were measured in reference samples and include antimony, arsenic, cadmium, chromium, copper, iron, lead, mercury, nickel, selenium, silver, vanadium, and zinc. The laboratory reported results for 22 metals at the request of EPA; however, for the purposes of meeting project objectives, only the data validation for the 13 target analytes is summarized in this document. The objective of the validation is to determine the validity of the reference data, as well as its usability in meeting the primary objective of comparing reference data to XRF data generated during the demonstration. Shealy provided the data to Tetra Tech EM Inc. (Tetra Tech) in electronic and hardcopy formats; a total of 13 sample delivery groups (SDG) contain all the data for this project.

The DVSR consists of seven sections, including this introduction. Section 2.0 presents the data validation methodology. Section 3.0 presents the results of the reference laboratory data validation. Section 4.0 summarizes the precision, accuracy, representativeness, completeness, and comparability (PARCC) evaluation. Section 5.0 presents conclusions about the overall evaluation of the reference data. Section 6.0 lists the references used to prepare this DVSR. Tables are presented following Section 6.0.

## 2.0 VALIDATION METHODOLOGY

Data validation is the systematic process for reviewing and qualifying data against a set of criteria to ensure that the reference data are adequate for the intended use. The data validation process assesses acceptability of the data by evaluating the critical indicator parameters of PARCC. The laboratory analytical data were validated according to the procedures outlined in the following documents:

- "USEPA Contract Laboratory Program National Functional Guidelines for Inorganic Data Review" (EPA 2004), hereinafter referred to as the "EPA guidance."
- "Demonstration and Quality Assurance Project Plan, XRF Technologies for Measuring Trace Elements in Soil and Sediment" (Tetra Tech 2005), hereinafter referred to as "the QAPP."

Data validation occurred in the following two stages: (1) a cursory review of analytical reports and quality assurance and quality control (QA/QC) information for 100 percent of the reference data and (2) full validation of analytical reports, QA/QC information, and associated raw data for 10 percent of the reference data as required by the QAPP (Tetra Tech 2005).

QA/QC criteria were reviewed in accordance with EPA guidance (EPA 2004) and the QAPP (Tetra Tech 2005). The cursory review for total metals consisted of evaluating the following requirements, as applicable:

• Holding times

- Initial and continuing calibrations
- Laboratory blank results
- Laboratory control sample (LCS) and laboratory control sample duplicates (LCSD) results
- Matrix spike (MS) and matrix spike duplicate (MSD) results
- Serial dilutions results

In addition to QA/QC criteria described above, the following criteria were reviewed during full validation:

- ICP interference check samples (ICS)
- Target analyte identification and quantitation
- Quantitation limit verification

Section 3.0 presents the results of the both the cursory review and full validation.

During data validation, worksheets were produced for each SDG that identify any QA/QC issues resulting in data qualification. Data validation findings were written in 13 individual data validation reports (one for each SDG). Data qualifiers were assigned to the results in the electronic database in accordance with EPA guidelines (EPA 2004). In addition to data validation qualifiers, comment codes were added to the database to indicate the primary reason for the validation qualifier. Table 1 defines data validation qualifiers and comment codes that are applied to the data set. Details about specific QC issues can be found in the individual SDG data validation reports and accompanying validation worksheets provided in the Appendix.

The overall objective of data validation is to ensure that the quality of the reference data set is adequate for the intended use, as defined by the QAPP (Tetra Tech 2005) for the PARCC parameters. Table 2 provides the QC criteria as defined by the QAPP. PARCC parameters were assessed by completing the following tasks:

- Reviewing precision and accuracy of laboratory QC data
- Reviewing the overall analytical process, including holding time, calibration, analytical or matrix performance, and analyte identification and quantitation
- Assigning qualifiers to affected data when QA/QC criteria were not achieved
- Reviewing and summarizing implications of the frequency and severity of qualifiers in the validated data

Prior to the XRF demonstration, soil and sediment samples were collected from nine locations across the U.S. and then blended, dried, sieved, and homogenized in the characterization laboratory to produce a set of 326 reference samples. Each of these samples were subsequently analyzed by both the reference

laboratory and all participating technology vendors. As such, 326 prepared soil/sediment samples were delivered to Shealy for the measurement of total metals. The analytical program included the following analyses and methods:

- Total metal for 22 analytes by inductively coupled plasma atomic emission spectroscopy (ICP-AES) according to EPA Methods 3050B/6010B (EPA 1996)
- Total mercury by cold vapor atomic absorption spectroscopy (CVAA) according to EPA Method 7471A (EPA 1996)

## 3.0 DATA VALIDATION RESULTS

The parameters listed in Section 2.0 were evaluated during cursory review and full validation of analytical reports for all methods, as applicable. Each of the validation components discussed in this section is summarized as follows:

- Acceptable All criteria were met and no data were qualified on that basis
- Acceptable with qualification Most criteria were met, but at least one data point was qualified as estimated because of issues related to the review component

Since no data were rejected, all data were determined to be either acceptable or acceptable with qualification. Sections 3.1 through 3.9 discuss each review component and the results of each. Tables that summarize the data validation findings follow Section 6.0 of this DVSR. Only qualified data are included in the tables. No reference laboratory data were rejected during the validation process. As such, all results are acceptable with the qualification noted in the sections that follow.

## 3.1 Holding Time

Acceptable. The technical holding times were defined as the maximum time allowable between sample collection and, as applicable, sample extraction, preparation, or analysis. The holding times used for validation purposes were recommended in the specific analytical methods (EPA 1996) and were specified in the QAPP (Tetra Tech 2005).

Because the soil and sediment samples were prepared prior to submission to the reference laboratory, and because the preparation included drying to remove moisture, no chemical or physical (for example ice) preservation was required. The holding time for sample digestion was 180 days for the ICP-AES analyses and 28 days for mercury. All sample digestions and analyses were conducted within the specified holding times. No data were qualified based on holding time exceedances. This fact contributes to the high technical quality of the reference data.

## 3.2 Calibration

Acceptable. Laboratory instrument calibration requirements were established to ensure that analytical instruments could produce acceptable qualitative and quantitative data for all target analytes. Initial calibration demonstrates that the instrument is capable of acceptable performance at the beginning of an analytical run, while producing a linear curve. Continuing calibration demonstrates that the instrument is capable of repeating the performance established during the initial calibration (EPA 1996).

For total metal analyses (ICP-AES and CVAA), initial calibration review included evaluating criteria for the curve's correlation coefficient and initial calibration verification (ICV) percent recoveries. The ICV percent recoveries verify that the analytical system is operating within the established calibration criteria at the beginning of an analytical run. The continuing calibration review included evaluation of the criteria for continuing calibration verification (CCV) percent recoveries. The CCV percent recoveries verify that the analytical system is operating within the established calibration of the criteria for continuing calibration verification (CCV) percent recoveries. The CCV percent recoveries verify that the analytical system is operating within the established calibration throughout the analytical run.

All ICV and CCV percent recoveries associated with the reference data were within acceptable limits of 90 to 110 percent. As such, no data were qualified or rejected because of calibration exceedances. This fact contributes to the high technical quality of the data.

## 3.3 Laboratory Blanks

Acceptable with qualification. No field blanks were required by the QAPP, since samples were prepared after collection and before submission to the reference laboratory. However, laboratory blanks were prepared and analyzed to evaluate the existence and magnitude of contamination resulting from laboratory activities. Blanks prepared and analyzed in the laboratory consisted of calibration and preparation blanks. If a problem with any blank existed, all associated data were carefully evaluated to assess whether the sample data were affected. At a minimum, calibration blanks were analyzed for every 10 analyses conducted on each instrument. Preparation blanks were prepared at a frequency of one per preparation batch per matrix or every 20 samples, whichever is greater (EPA 1996).

When laboratory blank contamination was identified, sample results were compared to the practical quantitation limit (PQL) and the maximum blank value as required by the validation guidelines (EPA 2004). Most of the blank detections were positive results (i.e. greater than the method detection limit [MDL]), but less than the PQL. In these instances, if associated sample results were also less than the PQL, they were qualified as undetected (U); with the comment code "b." In these same instances, if the associated sample results were greater than the PQL, the reviewer used professional judgment to determine if the sample results were adversely affected. If so, then the results were qualified as estimated with the potential for being biased high (J+). If not, then no qualification was required.

In a few cases, the maximum blank value exceeded the PQL. In these cases, all associated sample results less than the PQL were qualified as undetected (U) with the comment code "b." In cases where the associated sample results were greater than the PQL, but less than the blank concentration, the results were also qualified as undetected (U); with the comment code "b." If the associated sample results were greater than both the PQL and the blank value, the reviewer used professional judgment to determine if sample results were adversely affected. If so, then the results were qualified as estimated with the potential for being biased high (J+); with the comment code "b." Sample results significantly above the blank were not qualified.

In addition to laboratory blank contamination, negative drift greater than the magnitude of the PQL was observed in some laboratory blanks. Associated sample data were qualified as undetected (U) if the results were less than the PQL. Professional judgement was used to determine if the negative drift adversely affected associated sample results greater than the PQL. If so, then sample results were qualified as estimated with the potential for being biased low (J-) due to the negative drift of the instrument baseline; with the comment code "b."

Of all target analyte data, 2.6 percent of the data was qualified as undetected because of laboratory blank contamination (U, b), and less than 1 percent of the data was qualified as estimated (either J+, b or J-, b). The low occurrence of results affected by blank contamination indicates that the general quality of the

analytical data was not significantly compromised by blank contamination. Table 3 provides all results that were qualified based on laboratory blanks.

## 3.4 Laboratory Control Samples

Acceptable. LCSs and LCSDs were prepared and analyzed with each batch of 20 or fewer samples of the same matrix. All percent recoveries were within the QC limits of 80 to 120 percent; all relative percent differences (RPD) between the LCD and LCSD values were less than the criterion of 20 percent. No data were qualified or rejected on the basis of LCS/LCSD results. This fact contributes to the high technical quality of the data.

## 3.5 Matrix Spike Samples

Acceptable with qualification. MS and MSD samples were prepared and analyzed with each batch of 20 or fewer samples of the same matrix. All percent recoveries were within the QC limits of 75 to 125 percent, and all RPDs between the MS and MSD values were less than the criterion of 25 percent, except as discussed in the following paragraphs.

Sample results affected by MS and MSD percent recoveries issues were qualified as estimated and either biased high (J+) if the recoveries were greater than 125 percent; or qualified as estimated and biased low (J-) if the recoveries were less than 75 percent. In at least one case, the MS was higher than 125 percent and the MSD was lower than 75 percent; the associated results were qualified as estimated (J) with no distinction for potential bias. All data qualified on the basis of MS and MSD recovery were also assigned the comment code "e." Of all target analyte data, less than 1 percent was qualified as estimated and biased low (J-, e). Antimony and silver were the most frequently qualified sample results. Based on experience, antimony and silver soil recoveries are frequently low using the selected methods. Table 4 provides the results that were qualified based on MS/MSD results.

The precision between MS and MSD results were generally acceptable. If the RPD between MS and MSD results were greater than 25 percent, the data were already qualified based on exceedance of the acceptance window for recovery. Therefore, no additional qualification was required for MS/MSD precision.

No data were rejected on the basis of MS/MSD results. The relatively low occurrence of data qualification due to MS/MSD recoveries and RPDs contribute to the high technical quality of the data.

## 3.6 Serial Dilution Results

Acceptable with qualification. Serial dilutions were conducted and analyzed by Shealy at a frequency of 1 per batch of 20 samples. The serial dilution analysis can evaluate whether matrix interference exists and whether the accuracy of the analytical data is affected. For all target analyte data, less than 1 percent of the data was qualified as estimated and biased high (J+, j), while about 2 percent of the data were qualified as estimated and biased low (J-, j). Serial dilution results are used to determine whether characteristics of the digest matrix, such as viscosity or the presence of analytes at high concentrations, may interfere with the detected analytes. Qualifiers were applied to cases where interference was suspected. However, the low incidence of apparent matrix interference contributes to the high technical quality of the data. Table 5 provides the results that were qualified based on MS/MSD results.

## **3.7 ICP Interference Check Samples**

Acceptable. ICP results for each ICS were evaluated. The ICS verifies the validity of the laboratory's inter-element and background correction factors. High levels of certain elements (including aluminum, calcium, iron, and magnesium) can affect sample results if the inter-element and background correction factors have not been optimized. Incorrect correction factors may result in false positives, false negatives, or biased results. All ICS recoveries were within QC limits of 80 to 120 percent, and no significant biases were observed due to potential spectral interference. No data were qualified or rejected because of ICS criteria violations. This fact contributes to the high technical quality of the data.

## 3.8 Target Analyte Identification and Quantitation

Acceptable Identification is determined by measuring the characteristic wavelength of energy emitted by the analyte (ICP) or absorbed by the analyte (CVAA). External calibration standards are used to quantify the analyte concentration in the sample digest. Sample digest concentrations are converted to soil units (milligrams per kilogram) and corrected for percent moisture. For 10 percent of the samples, results were recalculated to verify the accuracy of reporting. All results were correctly calculated by the laboratory, except for one mercury result, whose miscalculation was the result of an error in entering the dilution factor. Shealy immediately resolved this error and corrected reports were provided. Since the result was corrected, no qualification was required. No other reporting errors were observed.

For inorganic analyses, analytical instruments can make reliable qualitative identification of analytes at concentrations below the PQL. Detected results below the PQL are considered quantitatively uncertain. Sample results below the PQL were reported by the laboratory with a "J" qualifier. No additional qualification was required.

## 3.9 Quantitation Limit Verification

Acceptable. Reference laboratory quantitation limits were specified in the QAPP (Tetra Tech 2005). Circumstances that affected quantitation were limited and included dilution and percent moisture factors. Since the samples were prepared prior to submission to the reference laboratory, moisture content was very low and had little impact on quantitation limits. The laboratory did correct all quantitation limits for moisture content. Due to the presence of percent-level analytes in some samples, dilutions were required. However, the required PQLs for the reference laboratory were high enough that even with dilution and moisture content factors applied, the reporting limits did not exceed those of the XRF instruments. This allows for effective comparison of results between the reference laboratory and XRF instruments.

## 4.0 PRECISION, ACCURACY, REPRESENTATIVENESS, COMPLETENESS, AND COMPARABILITY EVALUATION SUMMARY

All analytical data were reviewed for PARCC parameters to validate reference data. The following sections discuss the overall data quality, including the PARCC parameters, as determined by the data validation.

## 4.1 Precision

Precision is a measure of the reproducibility of an experimental value without considering a true or referenced value. The primary indicators of precision were the MS/MSD RPD and LCS/LCSD RPD between the duplicate results. Precision criteria of less than 20 percent RPD for LCS/LCSD and 25 percent for MS/MSD were generally met for all duplicate pairs. No data were qualified based on duplicate precision of MS/MSD or LCS/LCSD pairs that were not already qualified for other reasons. Such low occurrence of laboratory precision problems supports the validity, usability, and defensibility of the data.

## 4.2 Accuracy

Accuracy assesses the proximity of an experimental value to a true or referenced value. The primary accuracy indicators were the recoveries of MS and LCS spikes. Accuracy is expressed as percent recovery. Overall, about 8 percent of the data was qualified as estimated and no data were rejected because of accuracy problems. The low frequency of accuracy problems supports the validity, usability, and defensibility of the data.

## 4.3 Representativeness

Representativeness refers to how well sample data accurately reflect true environmental conditions. The QAPP was carefully designed to ensure that actual environmental samples be collected by choosing representative sites across the US from which sample material was collected. The blending and homogenization was executed according to the approved QAPP (Tetra Tech 2005).

#### 4.4 Completeness

Completeness is defined as the percentage of measurements that are considered to be valid. The validity of sample results is evaluated through the data validation process. Sample results that are rejected and any missing analyses are considered incomplete. Data that are qualified as estimated (J) or undetected estimated (UJ) are considered valid and usable. Data qualified as rejected (R) are considered unusable for all purposes. Since no data were rejected in this data set, a completeness of 100 percent was achieved. A total of 4,238 target analyte results were evaluated. The completeness goal stated in the QAPP (Tetra Tech 2005) was 90 percent.

## 4.5 Comparability

Comparability is a qualitative parameter that expresses the confidence with which one data set may be compared to another. Widely-accepted SW-846 methods were used for this project. It is recognized that direct comparison of the reference laboratory data (using ICP-AES and CVAA techniques) to the XRF measurements may result in discrepancies due to differences in the preparation and measurement techniques; however, the reference laboratory data is expected to provide an acceptable basis for comparison to XRF measurement results in accordance with the project objectives.

Comparability of the data was also achieved by producing full data packages, by using a homogenous matrix, standard quantitation limits, standardized data validation procedures, and by evaluating the PARCC parameters uniformly. In addition, the use of specified and well-documented analyses, approved laboratories, and the standardized process of data review and validation have resulted in a high degree of comparability for the data.

#### 5.0 CONCLUSIONS FOR DATA QUALITY AND DATA USABILITY

Although some qualifiers were added to the data, a final review of the data set with respect to the data quality parameters discussed in Section 4.0 indicates that the data are of overall good quality. No analytical data were rejected. The data quality is generally consistent with project objectives for producing data of suitable quality for comparison to XRF data. All supporting documentation and data are available upon request, including cursory review and full validation reports as well as the electronic database that contains sample results.

#### 6.0 **REFERENCES**

- Tetra Tech EM, Inc. (Tetra Tech). 2005. "Demonstration and Quality Assurance Project Plan, XRF Technologies for Measuring Trace Elements in Soil and Sediment." March.
- U.S. Environmental Protection Agency (EPA). 1996. "Test Methods for Evaluating Solid Waste", Third Edition (SW-846). With promulgated revisions. December.
- EPA. 2004. "USEPA Contract Laboratory Program National Functional Guidelines For Inorganic Data Review". October.

## TABLES

## TABLE 1: DATA VALIDATION QUALIFIERS AND COMMENT CODES

Qualifier	Definition
No Qualifier	Indicates that the data are acceptable both qualitatively and quantitatively.
U	Indicates compound was analyzed for but not detected above the concentration listed. The value listed is the sample quantitation limit.
J	Indicates an estimated concentration value. The result is considered qualitatively acceptable, but quantitatively unreliable.
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.
UJ	Indicates an estimated quantitation limit. The compound was analyzed for, but was considered non-detected.
R	The data are unusable (compound may or may not be present). Resampling and reanalysis is necessary for verification.
Comment Code	Definition
a	Surrogate recovery exceeded (not applicable to this data set)
b	Laboratory method blank and common blank contamination
с	Calibration criteria exceeded
d	Duplicate precision criteria exceeded
e	Matrix spike or laboratory control sample recovery exceeded
f	Field blank contamination (not applicable to this data set)
g	Quantification below reporting limit
h	Holding time exceeded
i	Internal standard criteria exceeded (not applicable to this data set)
j	Other qualification (will be specified in report)

#### TABLE 2: QC CRITERIA

Parameter	Method	QC Check	Frequency	Criterion	Corrective Action
			Reference	Method	
Target Metals (12 ICP metals and Hg)	3050B/6010B and 7471A	Method and instrument blanks	One per analytical batch of 20 or less	Less than the reporting limit	<ol> <li>Check calculations</li> <li>Assess and eliminate source of contamination</li> <li>Reanalyze blank</li> <li>Inform Tetra Tech project manager</li> <li>Flag affected results</li> </ol>
		MS/MSD	One per analytical batch of 20 or less	75 to 125 percent recovery RPD $\leq 25$	<ol> <li>Check calculations</li> <li>Check LCS/LCSD and digest duplicate results to determine whether they meet criterion</li> <li>Inform Tetra Tech project manager</li> <li>Flag affected results</li> </ol>
		LCS/LCSD	One per analytical batch of 20 or less	80 to 120 percent recovery RPD ≤ 20	<ol> <li>Check calculations</li> <li>Check instrument operating conditions and adjust as necessary</li> <li>Check MS/MSD and digest duplicate results to determine whether they meet criterion</li> <li>Inform Tetra Tech project manager</li> <li>Redigest and reanalyze the entire batch of samples</li> <li>Flag affected results</li> </ol>
		Performance audit samples	One per analytical batch of 20 or less	Within acceptance limits	<ol> <li>Evaluated by Tetra Tech QA chemist</li> <li>Inform laboratory and recommend changes</li> <li>Flag affected results</li> </ol>
Percent moisture		Laboratory duplicates	One per analytical batch of 20 or less	RPD ≤ 20	<ol> <li>Check calculations</li> <li>Reanalyze sample batch</li> <li>Inform Tetra Tech project manager</li> <li>Flag affected results</li> </ol>

Sample ID	Analyte	Result	Unit	Validation Qualifier	Comment Code
AS-SO-04-XX	Selenium	6.2	mg/kg	U	b
AS-SO-06-XX	Antimony	2.4	mg/kg	UJ	b, e
AS-SO-10-XX	Selenium	1.1	mg/kg	U	b
AS-SO-11-XX	Selenium	1.1	mg/kg	U	b
AS-SO-13-XX	Antimony	2.4	mg/kg	UJ	b, e
BN-SO-18-XX	Silver	0.94	mg/kg	U	b
BN-SO-28-XX	Silver	0.77	mg/kg	U	b
BN-SO-31-XX	Silver	0.97	mg/kg	U	b
BN-SO-35-XX	Silver	0.85	mg/kg	U	b
KP-SE-01-XX	Mercury	0.053	mg/kg	U	b
KP-SE-11-XX	Mercury	0.079	mg/kg	U	b
KP-SE-12-XX	Mercury	0.06	mg/kg	U	b
KP-SE-14-XX	Mercury	0.065	mg/kg	U	b
KP-SE-17-XX	Mercury	0.082	mg/kg	U	b
KP-SE-19-XX	Mercury	0.044	mg/kg	U	b
KP-SE-25-XX	Mercury	0.096	mg/kg	U	b
KP-SE-25-XX	Selenium	0.26	mg/kg	U	b
KP-SE-28-XX	Mercury	0.056	mg/kg	U	b
KP-SE-30-XX	Mercury	0.1	mg/kg	U	b
KP-SE-30-XX	Selenium	0.24	mg/kg	U	b
KP-SO-02-XX	Mercury	0.043	mg/kg	U	b
KP-SO-02-XX	Selenium	0.42	mg/kg	U	b
KP-SO-03-XX	Cadmium	0.074	mg/kg	U	b
KP-SO-03-XX	Mercury	0.044	mg/kg	U	b
KP-SO-04-XX	Cadmium	0.046	mg/kg	U	b
KP-SO-04-XX	Mercury	0.018	mg/kg	U	b
KP-SO-04-XX	Selenium	0.28	mg/kg	U	b
KP-SO-05-XX	Cadmium	0.13	mg/kg	U	b
KP-SO-05-XX	Mercury	0.044	mg/kg	U	b
KP-SO-05-XX	Selenium	0.24	mg/kg	U	b
KP-SO-06-XX	Arsenic	0.73	mg/kg	J-	b
KP-SO-06-XX	Mercury	0.059	mg/kg	U	b
KP-SO-07-XX	Arsenic	2	mg/kg	J-	b
KP-SO-07-XX	Mercury	0.027	mg/kg	U	b
KP-SO-07-XX	Selenium	0.21	mg/kg	U	b
KP-SO-09-XX	Cadmium	0.094	mg/kg	U	b
KP-SO-09-XX	Mercury	0.046	mg/kg	U	b

#### TABLE 3: DATA QUALIFICATION: LABORATORY METHOD BLANK CONTAMINATION

Sample ID	Analyte	Result	Unit	Validation Qualifier	Comment Code
KP-SO-10-XX	Arsenic	0.7	mg/kg	J-	b
KP-SO-10-XX	Mercury	0.028	mg/kg	U	b
KP-SO-10-XX	Selenium	0.22	mg/kg	U	b
KP-SO-13-XX	Arsenic	1.4	mg/kg	J-	b
KP-SO-13-XX	Cadmium	0.045	mg/kg	U	b
KP-SO-13-XX	Mercury	0.037	mg/kg	U	b
KP-SO-15-XX	Arsenic	0.76	mg/kg	J-	b
KP-SO-15-XX	Mercury	0.029	mg/kg	U	b
KP-SO-16-XX	Cadmium	0.063	mg/kg	U	b
KP-SO-16-XX	Mercury	0.016	mg/kg	U	b
KP-SO-18-XX	Arsenic	0.56	mg/kg	J-	b
KP-SO-18-XX	Mercury	0.016	mg/kg	U	b
KP-SO-20-XX	Arsenic	1.5	mg/kg	J-	b
KP-SO-20-XX	Mercury	0.03	mg/kg	U	b
KP-SO-21-XX	Cadmium	0.098	mg/kg	U	b
KP-SO-21-XX	Mercury	0.042	mg/kg	U	b
KP-SO-22-XX	Arsenic	0.7	mg/kg	J-	b
KP-SO-22-XX	Mercury	0.027	mg/kg	U	b
KP-SO-23-XX	Cadmium	0.048	mg/kg	U	b
KP-SO-23-XX	Mercury	0.017	mg/kg	U	b
KP-SO-24-XX	Arsenic	1.4	mg/kg	J-	b
KP-SO-24-XX	Mercury	0.017	mg/kg	U	b
KP-SO-26-XX	Cadmium	0.061	mg/kg	U	b
KP-SO-26-XX	Mercury	0.013	mg/kg	U	b
KP-SO-26-XX	Selenium	0.22	mg/kg	U	b
KP-SO-27-XX	Arsenic	1.3	mg/kg	J-	b
KP-SO-27-XX	Cadmium	0.05	mg/kg	U	b
KP-SO-27-XX	Mercury	0.021	mg/kg	U	b
KP-SO-29-XX	Arsenic	1.5	mg/kg	J-	b
KP-SO-29-XX	Mercury	0.013	mg/kg	U	b
KP-SO-31-XX	Mercury	0.017	mg/kg	U	b
KP-SO-32-XX	Arsenic	1.6	mg/kg	J-	b
KP-SO-32-XX	Cadmium	0.045	mg/kg	U	b
KP-SO-32-XX	Mercury	0.014	mg/kg	U	b
LV-SE-02-XX	Mercury	0.02	mg/kg	U	b
LV-SE-10-XX	Mercury	0.023	mg/kg	U	b
LV-SE-11-XX	Selenium	1.3	mg/kg	U	b

### TABLE 3: DATA QUALIFICATION: LABORATORY METHOD BLANK CONTAMINATION (Continued)

Sample ID	Analyte	Result	Unit	Validation Qualifier	Comment Code
LV-SE-14-XX	Mercury	0.056	mg/kg	U	b
LV-SE-21-XX	Mercury	0.048	mg/kg	U	b
LV-SE-24-XX	Mercury	0.053	mg/kg	U	b
LV-SE-29-XX	Selenium	1.2	mg/kg	U	b
LV-SE-32-XX	Mercury	0.052	mg/kg	U	b
RF-SE-07-XX	Mercury	0.091	mg/kg	U	b
RF-SE-08-XX	Silver	0.39	mg/kg	U	b
RF-SE-10-XX	Silver	0.34	mg/kg	U	b
RF-SE-12-XX	Mercury	0.099	mg/kg	U	b
RF-SE-23-XX	Copper	0.2	mg/kg	U	b
RF-SE-23-XX	Zinc	0.6	mg/kg	U	b
RF-SE-33-XX	Silver	0.33	mg/kg	U	b
RF-SE-36-XX	Mercury	0.081	mg/kg	U	b
RF-SE-36-XX	Selenium	1	mg/kg	U	b
RF-SE-45-XX	Cadmium	0.52	mg/kg	U	b
RF-SE-53-XX	Cadmium	0.57	mg/kg	U	b
SB-SO-03-XX	Antimony	1.2	mg/kg	UJ	b, e
SB-SO-12-XX	Silver	2.1	mg/kg	UJ	b
SB-SO-13-XX	Silver	2.2	mg/kg	UJ	b
SB-SO-15-XX	Silver	1.6	mg/kg	UJ	b
SB-SO-17-XX	Silver	2.3	mg/kg	UJ	b, e
SB-SO-18-XX	Antimony	1.2	mg/kg	UJ	b, e
SB-SO-30-XX	Selenium	1.3	mg/kg	J+	b
SB-SO-32-XX	Silver	0.1	mg/kg	UJ	b, e
SB-SO-37-XX	Silver	2	mg/kg	UJ	b
SB-SO-46-XX	Silver	2.2	mg/kg	UJ	b, e
SB-SO-48-XX	Silver	0.1	mg/kg	UJ	b, e
SB-SO-53-XX	Antimony	1.2	mg/kg	UJ	b, e
TL-SE-01-XX	Mercury	0.074	mg/kg	U	b
TL-SE-03-XX	Mercury	0.32	mg/kg	J-	b
TL-SE-03-XX	Silver	0.94	mg/kg	U	b
TL-SE-04-XX	Mercury	0.26	mg/kg	J-	b
TL-SE-10-XX	Mercury	0.19	mg/kg	J-	b
TL-SE-11-XX	Mercury	0.021	mg/kg	U	b
TL-SE-12-XX	Mercury	0.22	mg/kg	J-	b
TL-SE-14-XX	Mercury	0.08	mg/kg	U	b
TL-SE-15-XX	Mercury	0.28	mg/kg	J-	b

### TABLE 3: DATA QUALIFICATION: LABORATORY METHOD BLANK CONTAMINATION (Continued)

Sample ID	Analyte	Result	Unit	Validation Qualifier	Comment Code
TL-SE-15-XX	Silver	1	mg/kg	U	b
TL-SE-18-XX	Mercury	0.025	mg/kg	U	b
TL-SE-19-XX	Mercury	0.32	mg/kg	J-	b
TL-SE-19-XX	Silver	1.1	mg/kg	U	b
TL-SE-20-XX	Mercury	0.26	mg/kg	J-	b
TL-SE-22-XX	Mercury	0.082	mg/kg	U	b
TL-SE-23-XX	Mercury	0.41	mg/kg	J-	b
TL-SE-23-XX	Silver	1.3	mg/kg	U	b
TL-SE-24-XX	Mercury	0.26	mg/kg	J-	b
TL-SE-24-XX	Silver	1.3	mg/kg	U	b
TL-SE-25-XX	Mercury	0.44	mg/kg	J-	b
TL-SE-25-XX	Silver	0.94	mg/kg	U	b
TL-SE-26-XX	Mercury	0.24	mg/kg	J-	b
TL-SE-27-XX	Mercury	0.02	mg/kg	U	b
TL-SE-29-XX	Mercury	0.076	mg/kg	U	b
TL-SE-31-XX	Mercury	0.57	mg/kg	J-	b
TL-SE-31-XX	Silver	1.2	mg/kg	U	b
WS-SO-06-XX	Mercury	0.07	mg/kg	U	b
WS-SO-08-XX	Mercury	0.063	mg/kg	U	b
WS-SO-10-XX	Mercury	0.058	mg/kg	U	b
WS-SO-12-XX	Mercury	0.068	mg/kg	UJ	b, e
WS-SO-17-XX	Mercury	0.069	mg/kg	UJ	b, e
WS-SO-20-XX	Mercury	0.06	mg/kg	U	b
WS-SO-23-XX	Mercury	0.05	mg/kg	U	b
WS-SO-30-XX	Mercury	0.069	mg/kg	UJ	b, e
WS-SO-31-XX	Selenium	1.2	mg/kg	U	b
WS-SO-35-XX	Mercury	0.071	mg/kg	UJ	b, e

### TABLE 3: DATA QUALIFICATION: LABORATORY METHOD BLANK CONTAMINATION (Continued)

Notes:

mg/kg = Milligrams per kilogram

b = Data were qualified based on blank contamination

e = Data were additionally qualified based on matrix spike/matrix spike duplicate exceedances

J+ = Result is estimated and potentially biased high

J- = Result is estimated and potentially biased low

UJ = Result is undetected at estimated quantitation limits

Sample ID	Analyte	Result	Unit	Validation Qualifier	Validation Code
AS-SO-01-XX	Antimony	3.8	mg/kg	J-	e
AS-SO-02-XX	Antimony	<2.6	mg/kg	UJ	e
AS-SO-03-XX	Mercury	3.7	mg/kg	J-	е
AS-SO-03-XX	Silver	480	mg/kg	J-	е
AS-SO-04-XX	Antimony	<6.4	mg/kg	UJ	е
AS-SO-05-XX	Mercury	2.5	mg/kg	J-	е
AS-SO-05-XX	Silver	330	mg/kg	J-	е
AS-SO-06-XX	Antimony	2.4	mg/kg	UJ	b, e
AS-SO-07-XX	Antimony	3.6	mg/kg	J-	е
AS-SO-08-XX	Mercury	2.5	mg/kg	J-	e
AS-SO-08-XX	Silver	280	mg/kg	J-	e
AS-SO-09-XX	Antimony	<2.6	mg/kg	UJ	e
AS-SO-10-XX	Antimony	1.9	mg/kg	J-	e
AS-SO-11-XX	Antimony	3.7	mg/kg	J-	e
AS-SO-12-XX	Antimony	<2.6	mg/kg	UJ	e
AS-SO-13-XX	Antimony	2.4	mg/kg	UJ	b, e
BN-SO-01-XX	Antimony	<1.3	mg/kg	UJ	e
BN-SO-01-XX	Silver	<1.3	mg/kg	UJ	e
BN-SO-05-XX	Antimony	160	mg/kg	J-	e
BN-SO-07-XX	Antimony	110	mg/kg	J-	e
BN-SO-07-XX	Silver	990	mg/kg	J+	e
BN-SO-09-XX	Antimony	750	mg/kg	J-	e
BN-SO-09-XX	Silver	100	mg/kg	J-	e
BN-SO-10-XX	Antimony	<1.3	mg/kg	UJ	e
BN-SO-10-XX	Silver	<1.3	mg/kg	UJ	e
BN-SO-11-XX	Antimony	4	mg/kg	J-	e
BN-SO-11-XX	Silver	140	mg/kg	J-	e
BN-SO-12-XX	Antimony	750	mg/kg	J-	e
BN-SO-12-XX	Silver	210	mg/kg	J-	e
BN-SO-14-XX	Antimony	3.5	mg/kg	J-	e
BN-SO-14-XX	Silver	140	mg/kg	J-	e
BN-SO-15-XX	Antimony	<1.3	mg/kg	UJ	е
BN-SO-15-XX	Silver	<1.3	mg/kg	UJ	e
BN-SO-16-XX	Antimony	120	mg/kg	J-	e
BN-SO-16-XX	Arsenic	1100	mg/kg	J+	e
BN-SO-19-XX	Antimony	150	mg/kg	J-	е
BN-SO-21-XX	Antimony	150	mg/kg	J-	e

Sample ID	Analyte	Result	Unit	Validation Qualifier	Validation Code
BN-SO-21-XX	Arsenic	1300	mg/kg	J+	e
BN-SO-23-XX	Antimony	<1.2	mg/kg	UJ	e
BN-SO-23-XX	Silver	130	mg/kg	J-	e
BN-SO-24-XX	Antimony	810	mg/kg	J-	e
BN-SO-24-XX	Silver	140	mg/kg	J-	e
BN-SO-25-XX	Antimony	82	mg/kg	J-	e, j
BN-SO-25-XX	Arsenic	700	mg/kg	J	e, j
BN-SO-26-XX	Antimony	150	mg/kg	J-	e
BN-SO-29-XX	Antimony	150	mg/kg	J-	е
BN-SO-32-XX	Antimony	160	mg/kg	J-	е
BN-SO-33-XX	Antimony	100	mg/kg	J-	e
CN-SO-01-XX	Antimony	13	mg/kg	J-	e
CN-SO-02-XX	Mercury	270	mg/kg	J-	е
CN-SO-03-XX	Mercury	34	mg/kg	J-	е
CN-SO-04-XX	Antimony	13	mg/kg	J-	е
CN-SO-05-XX	Mercury	280	mg/kg	J-	e
CN-SO-06-XX	Mercury	40	mg/kg	J-	e
CN-SO-07-XX	Mercury	36	mg/kg	J-	e
CN-SO-08-XX	Antimony	15	mg/kg	J-	e
CN-SO-09-XX	Mercury	260	mg/kg	J-	e
CN-SO-10-XX	Antimony	13	mg/kg	J-	e
CN-SO-11-XX	Antimony	17	mg/kg	J-	e
KP-SE-01-XX	Lead	310	mg/kg	J-	e
KP-SE-01-XX	Silver	< 0.26	mg/kg	UJ	e
KP-SE-08-XX	Lead	300	mg/kg	J-	e
KP-SE-08-XX	Silver	< 0.27	mg/kg	UJ	e
KP-SE-11-XX	Lead	310	mg/kg	J-	e
KP-SE-11-XX	Silver	< 0.27	mg/kg	UJ	e
KP-SE-12-XX	Lead	320	mg/kg	J-	e
KP-SE-12-XX	Silver	< 0.26	mg/kg	UJ	e
KP-SE-14-XX	Lead	680	mg/kg	J-	e, j
KP-SE-14-XX	Silver	< 0.26	mg/kg	UJ	е
KP-SE-17-XX	Lead	300	mg/kg	J-	e
KP-SE-17-XX	Silver	< 0.27	mg/kg	UJ	e
KP-SE-25-XX	Lead	310	mg/kg	J-	e
KP-SE-25-XX	Silver	< 0.27	mg/kg	UJ	e
KP-SE-30-XX	Lead	300	mg/kg	J-	e

Sample ID	Analyte	Result	Unit	Validation Qualifier	Validation Code
KP-SE-30-XX	Silver	<0.27	mg/kg	UJ	e
KP-SO-04-XX	Antimony	94	mg/kg	J+	e
KP-SO-06-XX	Antimony	8.1	mg/kg	J+	e
KP-SO-07-XX	Antimony	17	mg/kg	J+	e
KP-SO-10-XX	Antimony	6.1	mg/kg	J+	e
KP-SO-13-XX	Antimony	16	mg/kg	J+	e
KP-SO-15-XX	Antimony	6.3	mg/kg	J+	e
KP-SO-16-XX	Antimony	93	mg/kg	J+	e
KP-SO-18-XX	Antimony	6.7	mg/kg	J+	e
KP-SO-20-XX	Antimony	19	mg/kg	J+	e
KP-SO-22-XX	Antimony	8.3	mg/kg	J+	e
KP-SO-23-XX	Antimony	86	mg/kg	J+	e
KP-SO-24-XX	Antimony	17	mg/kg	J+	e
KP-SO-26-XX	Antimony	90	mg/kg	J+	e
KP-SO-27-XX	Antimony	15	mg/kg	J+	e
KP-SO-29-XX	Antimony	18	mg/kg	J+	е
KP-SO-32-XX	Antimony	16	mg/kg	J+	e
LV-SE-01-XX	Antimony	<1.5	mg/kg	UJ	e
LV-SE-02-XX	Antimony	<1.3	mg/kg	UJ	e
LV-SE-02-XX	Lead	20	mg/kg	J-	e
LV-SE-02-XX	Silver	<1.3	mg/kg	UJ	e
LV-SE-05-XX	Mercury	2.6	mg/kg	J-	e
LV-SE-06-XX	Mercury	610	mg/kg	J-	e
LV-SE-07-XX	Antimony	<6.7	mg/kg	UJ	e
LV-SE-08-XX	Antimony	<1.3	mg/kg	UJ	e
LV-SE-09-XX	Lead	14	mg/kg	J-	e
LV-SE-10-XX	Antimony	<1.3	mg/kg	UJ	e
LV-SE-10-XX	Lead	25	mg/kg	J-	e
LV-SE-10-XX	Silver	<1.3	mg/kg	UJ	e
LV-SE-11-XX	Antimony	<1.4	mg/kg	UJ	e
LV-SE-12-XX	Lead	19	mg/kg	J-	e
LV-SE-13-XX	Mercury	640	mg/kg	J-	е
LV-SE-14-XX	Antimony	<1.5	mg/kg	UJ	е
LV-SE-15-XX	Antimony	290	mg/kg	J+	е
LV-SE-15-XX	Silver	300	mg/kg	J-	е
LV-SE-16-XX	Antimony	<1.3	mg/kg	UJ	e
LV-SE-17-XX	Antimony	280	mg/kg	J+	e

		D K	<b>T</b> T •/	Validation	Validation
Sample ID	Analyte	Result	Unit	Qualifier	Code
LV-SE-17-XX	Lead	17	mg/kg	J-	e
LV-SE-17-XX	Silver	200	mg/kg	J-	e
LV-SE-18-XX	Antimony	<6.7	mg/kg	UJ	e
LV-SE-19-XX	Lead	17	mg/kg	J-	e
LV-SE-20-XX	Antimony	140	mg/kg	J+	e
LV-SE-20-XX	Silver	75	mg/kg	J-	e
LV-SE-21-XX	Antimony	<1.5	mg/kg	UJ	e
LV-SE-22-XX	Antimony	<1.3	mg/kg	UJ	e
LV-SE-22-XX	Lead	22	mg/kg	J-	e
LV-SE-22-XX	Silver	<1.3	mg/kg	UJ	e
LV-SE-23-XX	Antimony	<6.6	mg/kg	UJ	e
LV-SE-24-XX	Antimony	<1.5	mg/kg	UJ	e
LV-SE-25-XX	Antimony	<1.3	mg/kg	UJ	e
LV-SE-25-XX	Lead	23	mg/kg	J-	e
LV-SE-25-XX	Silver	<1.3	mg/kg	UJ	e
LV-SE-26-XX	Lead	25	mg/kg	J-	e
LV-SE-27-XX	Lead	16	mg/kg	J-	e
LV-SE-28-XX	Antimony	<1.3	mg/kg	UJ	e
LV-SE-29-XX	Antimony	<1.4	mg/kg	UJ	e
LV-SE-30-XX	Antimony	<1.3	mg/kg	UJ	e
LV-SE-31-XX	Antimony	<1.3	mg/kg	UJ	e
LV-SE-31-XX	Lead	49	mg/kg	J-	e
LV-SE-31-XX	Silver	<1.3	mg/kg	UJ	e
LV-SE-32-XX	Antimony	<1.4	mg/kg	UJ	e
LV-SE-33-XX	Lead	21	mg/kg	J-	e
LV-SE-35-XX	Antimony	<1.3	mg/kg	UJ	e
LV-SE-35-XX	Lead	22	mg/kg	J-	e
LV-SE-35-XX	Silver	<1.3	mg/kg	UJ	e
LV-SE-36-XX	Lead	21	mg/kg	J-	е
LV-SE-38-XX	Lead	15	mg/kg	J-	e
LV-SE-39-XX	Lead	22	mg/kg	J-	е
LV-SE-41-XX	Mercury	610	mg/kg	J-	e
LV-SE-42-XX	Lead	22	mg/kg	J-	е
LV-SE-43-XX	Antimony	160	mg/kg	J+	e
LV-SE-43-XX	Silver	60	mg/kg	J-	e
LV-SE-45-XX	Antimony	<6.7	mg/kg	UJ	e
LV-SE-47-XX	Antimony	<1.3	mg/kg	UJ	e

Sample ID	Analyte	Result	Unit	Validation Qualifier	Validation Code
LV-SE-48-XX	Antimony	<6.6	mg/kg	UJ	e
LV-SE-50-XX	Lead	24	mg/kg	J-	e
LV-SE-51-XX	Antimony	210	mg/kg	J+	e
LV-SE-51-XX	Silver	250	mg/kg	J-	e
LV-SO-03-XX	Mercury	48	mg/kg	J-	e
LV-SO-03-XX	Silver	210	mg/kg	J-	e
LV-SO-04-XX	Mercury	130	mg/kg	J-	e
LV-SO-04-XX	Silver	<1.2	mg/kg	UJ	e
LV-SO-34-XX	Mercury	130	mg/kg	J-	e
LV-SO-34-XX	Silver	<1.2	mg/kg	UJ	e
LV-SO-37-XX	Mercury	130	mg/kg	J-	e
LV-SO-40-XX	Mercury	46	mg/kg	J-	e
LV-SO-40-XX	Silver	210	mg/kg	J-	e
LV-SO-49-XX	Mercury	52	mg/kg	J-	e
LV-SO-49-XX	Silver	220	mg/kg	J-	е
RF-SE-02-XX	Antimony	<1.3	mg/kg	UJ	е
RF-SE-03-XX	Antimony	<1.2	mg/kg	UJ	е
RF-SE-04-XX	Antimony	3.2	mg/kg	J+	е
RF-SE-04-XX	Silver	12	mg/kg	J-	е
RF-SE-05-XX	Antimony	4.1	mg/kg	J+	е
RF-SE-05-XX	Silver	7.4	mg/kg	J-	е
RF-SE-06-XX	Antimony	<1.3	mg/kg	UJ	е
RF-SE-13-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-14-XX	Antimony	4.4	mg/kg	J+	e
RF-SE-14-XX	Silver	13	mg/kg	J-	e
RF-SE-15-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-19-XX	Antimony	3.7	mg/kg	J+	e
RF-SE-19-XX	Silver	14	mg/kg	J-	e
RF-SE-22-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-24-XX	Antimony	<1.3	mg/kg	UJ	е
RF-SE-25-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-26-XX	Antimony	2.2	mg/kg	J+	e
RF-SE-26-XX	Silver	7.2	mg/kg	J-	e
RF-SE-27-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-28-XX	Antimony	<1.2	mg/kg	UJ	e
RF-SE-30-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-31-XX	Antimony	<1.3	mg/kg	UJ	e

Sample ID	Analyte	Result	Unit	Validation Qualifier	Validation Code
RF-SE-32-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-34-XX	Antimony	2.9	mg/kg	J+	e
RF-SE-34-XX	Silver	10	mg/kg	J-	e
RF-SE-38-XX	Antimony	<1.2	mg/kg	UJ	e
RF-SE-39-XX	Antimony	2.9	mg/kg	J+	e
RF-SE-39-XX	Silver	8.2	mg/kg	J-	e
RF-SE-42-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-43-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-44-XX	Antimony	2.7	mg/kg	J+	e
RF-SE-44-XX	Silver	7.2	mg/kg	J-	e
RF-SE-45-XX	Antimony	<1.3	mg/kg	UJ	е
RF-SE-49-XX	Antimony	<1.2	mg/kg	UJ	e
RF-SE-52-XX	Antimony	3.4	mg/kg	J+	e
RF-SE-52-XX	Silver	11	mg/kg	J-	е
RF-SE-53-XX	Antimony	<1.3	mg/kg	UJ	е
RF-SE-55-XX	Antimony	<1.2	mg/kg	UJ	e
RF-SE-56-XX	Antimony	3.5	mg/kg	J+	e
RF-SE-56-XX	Silver	8.3	mg/kg	J-	e
RF-SE-57-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-58-XX	Antimony	<1.3	mg/kg	UJ	e
RF-SE-59-XX	Antimony	<1.3	mg/kg	UJ	e
SB-SO-01-XX	Antimony	180	mg/kg	J	e
SB-SO-02-XX	Antimony	44	mg/kg	J-	e, j
SB-SO-02-XX	Silver	<1.2	mg/kg	UJ	e
SB-SO-03-XX	Antimony	1.2	mg/kg	UJ	b, e
SB-SO-04-XX	Silver	<1.3	mg/kg	UJ	e
SB-SO-05-XX	Antimony	1.6	mg/kg	J-	e
SB-SO-06-XX	Antimony	1.7	mg/kg	J-	e
SB-SO-07-XX	Antimony	45	mg/kg	J	e
SB-SO-08-XX	Antimony	5.4	mg/kg	J-	е
SB-SO-09-XX	Antimony	<1.3	mg/kg	UJ	е
SB-SO-09-XX	Silver	160	mg/kg	J-	е
SB-SO-10-XX	Antimony	62	mg/kg	J	е
SB-SO-11-XX	Antimony	5.7	mg/kg	J-	е
SB-SO-12-XX	Antimony	620	mg/kg	J	е
SB-SO-13-XX	Antimony	430	mg/kg	J	е
SB-SO-14-XX	Antimony	4.1	mg/kg	J-	e

		D K	<b>T</b> T <b>'</b> 4	Validation	Validation
Sample ID	Analyte	Result	Unit	Qualifier	Code
SB-SO-15-XX	Antimony	600	mg/kg	J-	j, e
SB-SO-16-XX	Antimony	170	mg/kg	J	e
SB-SO-17-XX	Antimony	800	mg/kg	J+	e
SB-SO-17-XX	Silver	2.3	mg/kg	UJ	b, e
SB-SO-18-XX	Antimony	1.2	mg/kg	UJ	b, e
SB-SO-19-XX	Antimony	310	mg/kg	J	e
SB-SO-20-XX	Antimony	<1.3	mg/kg	UJ	e
SB-SO-20-XX	Silver	140	mg/kg	J-	e
SB-SO-21-XX	Antimony	4.9	mg/kg	J	e
SB-SO-22-XX	Antimony	10	mg/kg	J	e, j
SB-SO-23-XX	Antimony	48	mg/kg	J-	e
SB-SO-23-XX	Silver	<0.26	mg/kg	UJ	e
SB-SO-24-XX	Antimony	180	mg/kg	J	e
SB-SO-25-XX	Antimony	6.8	mg/kg	J+	e
SB-SO-26-XX	Antimony	61	mg/kg	J	e
SB-SO-27-XX	Antimony	6.7	mg/kg	J+	e
SB-SO-28-XX	Antimony	42	mg/kg	J-	e
SB-SO-28-XX	Silver	< 0.26	mg/kg	UJ	e
SB-SO-29-XX	Silver	<1.2	mg/kg	UJ	e
SB-SO-30-XX	Antimony	3.2	mg/kg	J-	e
SB-SO-31-XX	Antimony	<1.3	mg/kg	UJ	e
SB-SO-31-XX	Silver	160	mg/kg	J-	e, j
SB-SO-32-XX	Antimony	46	mg/kg	J-	e
SB-SO-32-XX	Silver	0.1	mg/kg	UJ	b, e
SB-SO-33-XX	Antimony	350	mg/kg	J	e
SB-SO-33-XX	Silver	2	mg/kg	J	e
SB-SO-34-XX	Silver	<1.3	mg/kg	UJ	e
SB-SO-35-XX	Antimony	6	mg/kg	J+	e
SB-SO-36-XX	Silver	<1.2	mg/kg	UJ	e
SB-SO-37-XX	Antimony	340	mg/kg	J	е
SB-SO-38-XX	Antimony	<1.3	mg/kg	UJ	е
SB-SO-39-XX	Antimony	4.7	mg/kg	J-	е
SB-SO-40-XX	Antimony	2.2	mg/kg	J-	e
SB-SO-41-XX	Antimony	<1.3	mg/kg	UJ	e
SB-SO-42-XX	Antimony	4.6	mg/kg	J-	e
SB-SO-43-XX	Antimony	40	mg/kg	J-	e
SB-SO-43-XX	Silver	< 0.26	mg/kg	UJ	e

Sample ID	Analyte	Result	Unit	Validation Qualifier	Validation Code
SB-SO-44-XX	Antimony	6.8	mg/kg	J+	e
SB-SO-45-XX	Antimony	180	mg/kg	J	e
SB-SO-45-XX	Silver	2.1	mg/kg	J-	e
SB-SO-46-XX	Antimony	740	mg/kg	J+	e
SB-SO-46-XX	Silver	2.2	mg/kg	UJ	b, e
SB-SO-47-XX	Antimony	<1.3	mg/kg	UJ	e
SB-SO-48-XX	Antimony	39	mg/kg	J-	e
SB-SO-48-XX	Silver	0.1	mg/kg	UJ	b, e
SB-SO-49-XX	Silver	<1.2	mg/kg	UJ	e
SB-SO-50-XX	Antimony	57	mg/kg	J	e
SB-SO-51-XX	Antimony	<1.3	mg/kg	UJ	e
SB-SO-52-XX	Antimony	150	mg/kg	J	e
SB-SO-53-XX	Antimony	1.2	mg/kg	UJ	b, e
SB-SO-54-XX	Lead	5.2	mg/kg	J-	e
SB-SO-54-XX	Silver	<0.5	mg/kg	UJ	e
SB-SO-55-XX	Antimony	340	mg/kg	J	e
SB-SO-55-XX	Silver	2.2	mg/kg	J	e
SB-SO-56-XX	Silver	<1.2	mg/kg	UJ	e
TL-SE-01-XX	Antimony	<1.2	mg/kg	UJ	e
TL-SE-01-XX	Lead	48	mg/kg	J-	e
TL-SE-01-XX	Silver	5.7	mg/kg	J-	e
TL-SE-05-XX	Antimony	100	mg/kg	J+	e
TL-SE-05-XX	Silver	180	mg/kg	J-	e
TL-SE-09-XX	Antimony	100	mg/kg	J+	e
TL-SE-09-XX	Silver	170	mg/kg	J-	e
TL-SE-11-XX	Antimony	<1.2	mg/kg	UJ	e
TL-SE-11-XX	Lead	54	mg/kg	J-	e
TL-SE-11-XX	Silver	5.5	mg/kg	J-	e
TL-SE-13-XX	Antimony	95	mg/kg	J+	j, e
TL-SE-13-XX	Silver	160	mg/kg	J	j, e
TL-SE-14-XX	Antimony	<1.2	mg/kg	UJ	e
TL-SE-14-XX	Lead	50	mg/kg	J-	e
TL-SE-14-XX	Silver	5.7	mg/kg	J-	e
TL-SE-18-XX	Antimony	<1.2	mg/kg	UJ	e
TL-SE-18-XX	Lead	46	mg/kg	J-	e
TL-SE-18-XX	Silver	6.3	mg/kg	J-	e
TL-SE-22-XX	Antimony	<1.2	mg/kg	UJ	e

Sample ID	Analyte	Result	Unit	Validation Qualifier	Validation Code
TL-SE-22-XX	Lead	54	mg/kg	J-	
TL-SE-22-XX	Silver	6.5	mg/kg	J-	e e
TL-SE-27-XX	Antimony	<1.2		J- UJ	-
			mg/kg	J-	e
TL-SE-27-XX	Lead	51	mg/kg	J- J-	e
TL-SE-27-XX	Silver	7.8	mg/kg		e
TL-SE-29-XX	Antimony	<1.2	mg/kg	UJ	e
TL-SE-29-XX	Lead	51	mg/kg	J-	е
TL-SE-29-XX	Silver	5.9	mg/kg	J-	е
WS-SO-01-XX	Antimony	41	mg/kg	J-	e
WS-SO-01-XX	Mercury	5.8	mg/kg	J	e, j
WS-SO-01-XX	Silver	69	mg/kg	J-	e
WS-SO-02-XX	Antimony	130	mg/kg	J-	e
WS-SO-02-XX	Silver	150	mg/kg	J-	e
WS-SO-03-XX	Antimony	8.9	mg/kg	J-	e
WS-SO-03-XX	Mercury	0.86	mg/kg	J-	e
WS-SO-04-XX	Antimony	45	mg/kg	J-	e
WS-SO-04-XX	Silver	76	mg/kg	J-	e
WS-SO-05-XX	Antimony	8.6	mg/kg	J-	e
WS-SO-05-XX	Silver	0.76	mg/kg	J-	e
WS-SO-07-XX	Silver	400	mg/kg	J-	e
WS-SO-09-XX	Antimony	7.1	mg/kg	J-	e
WS-SO-09-XX	Mercury	0.89	mg/kg	J-	e
WS-SO-10-XX	Silver	<1.3	mg/kg	UJ	e
WS-SO-11-XX	Silver	340	mg/kg	J-	e
WS-SO-12-XX	Antimony	<1.3	mg/kg	UJ	e
WS-SO-12-XX	Mercury	0.068	mg/kg	UJ	b, e
WS-SO-13-XX	Antimony	200	mg/kg	J-	e
WS-SO-13-XX	Silver	170	mg/kg	J-	е
WS-SO-14-XX	Antimony	8.4	mg/kg	J-	е
WS-SO-14-XX	Mercury	0.74	mg/kg	J-	е
WS-SO-15-XX	Antimony	48	mg/kg	J-	е
WS-SO-15-XX	Silver	90	mg/kg	J-	e
WS-SO-16-XX	Antimony	110	mg/kg	J-	e
WS-SO-16-XX	Silver	150	mg/kg	J-	e
WS-SO-17-XX	Antimony	<1.3	mg/kg	UJ	e
WS-SO-17-XX	Mercury	0.069	mg/kg	UJ	b, e
WS-SO-18-XX	Antimony	130	mg/kg	J-	e

Sample ID	Analyte	Result	Unit	Validation Qualifier	Validation Code
WS-SO-18-XX	Silver	140	mg/kg	J-	e
WS-SO-19-XX	Antimony	150	mg/kg	J-	e
WS-SO-19-XX	Silver	160	mg/kg	J-	e
WS-SO-20-XX	Silver	<1.3	mg/kg	UJ	e
WS-SO-21-XX	Antimony	120	mg/kg	J-	e
WS-SO-21-XX	Silver	150	mg/kg	J-	e
WS-SO-22-XX	Antimony	41	mg/kg	J-	e
WS-SO-22-XX	Silver	72	mg/kg	J-	e
WS-SO-23-XX	Silver	<1.3	mg/kg	UJ	e
WS-SO-24-XX	Antimony	97	mg/kg	J-	e
WS-SO-24-XX	Silver	140	mg/kg	J-	e
WS-SO-25-XX	Silver	450	mg/kg	J-	e
WS-SO-26-XX	Antimony	7.6	mg/kg	J-	e
WS-SO-26-XX	Mercury	0.83	mg/kg	J-	e
WS-SO-27-XX	Antimony	<1.3	mg/kg	UJ	e
WS-SO-27-XX	Mercury	0.11	mg/kg	J-	e
WS-SO-28-XX	Antimony	120	mg/kg	J-	e
WS-SO-28-XX	Silver	130	mg/kg	J-	e
WS-SO-29-XX	Antimony	120	mg/kg	J-	e
WS-SO-29-XX	Silver	140	mg/kg	J-	e
WS-SO-30-XX	Antimony	1.2	mg/kg	J-	e
WS-SO-30-XX	Mercury	0.069	mg/kg	UJ	b, e
WS-SO-31-XX	Antimony	7.2	mg/kg	J-	e
WS-SO-31-XX	Mercury	0.85	mg/kg	J-	e
WS-SO-32-XX	Antimony	190	mg/kg	J-	e
WS-SO-32-XX	Silver	190	mg/kg	J-	e
WS-SO-33-XX	Antimony	6.9	mg/kg	J-	e
WS-SO-33-XX	Mercury	0.87	mg/kg	J-	e
WS-SO-34-XX	Antimony	45	mg/kg	J-	e
WS-SO-34-XX	Silver	78	mg/kg	J-	e
WS-SO-35-XX	Antimony	<1.3	mg/kg	UJ	e
WS-SO-35-XX	Mercury	0.071	mg/kg	UJ	b, e
WS-SO-36-XX	Antimony	120	mg/kg	J-	e
WS-SO-36-XX	Silver	120	mg/kg	J-	e
WS-SO-37-XX	Antimony	120	mg/kg	J-	e
WS-SO-37-XX	Silver	140	mg/kg	J-	e

Notes:

- < = Less than
- mg/kg = Milligram per kilogram
- b = Data were qualified based on blank contamination
- e = Data were additionally qualified based on matrix spike/matrix spike duplicate exceedances
- j = Data were additionally qualified based on serial dilution exceedances
- J = Result is estimated and biased could not be determined
- J+ = Result is estimated and potentially biased high
- J- = Result is estimated and potentially biased low
- UJ = Result is undetected at estimated quantitation limit

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Sample ID	Analyte	Result	Unit	Qualifier	Code
AS-SO-09-XX	Arsenic	25	mg/kg	J-	<u>J</u>
AS-SO-09-XX	Cadmium	100	mg/kg	J-	J
AS-SO-09-XX	Chromium	390	mg/kg	J-	j
AS-SO-09-XX	Copper	250	mg/kg	J-	j
AS-SO-09-XX	Iron	94000	mg/kg	J-	j
AS-SO-09-XX	Lead	3200	mg/kg	J-	j
AS-SO-09-XX	Nickel	170	mg/kg	J-	j
AS-SO-09-XX	Silver	9.6	mg/kg	J-	j
AS-SO-09-XX	Vanadium	65	mg/kg	J-	j
AS-SO-09-XX	Zinc	6800	mg/kg	J-	j
BN-SO-11-XX	Mercury	24	mg/kg	J-	j
BN-SO-25-XX	Antimony	82	mg/kg	J-	e, j
BN-SO-25-XX	Arsenic	700	mg/kg	J	e, j
BN-SO-25-XX	Cadmium	370	mg/kg	J-	j
BN-SO-25-XX	Chromium	64	mg/kg	J-	j
BN-SO-25-XX	Copper	930	mg/kg	J-	j
BN-SO-25-XX	Iron	16000	mg/kg	J-	j
BN-SO-25-XX	Lead	5400	mg/kg	J-	j
BN-SO-25-XX	Nickel	88	mg/kg	J-	j
BN-SO-25-XX	Selenium	19	mg/kg	J-	j
BN-SO-25-XX	Silver	48	mg/kg	J-	j
BN-SO-25-XX	Vanadium	28	mg/kg	J-	j
BN-SO-25-XX	Zinc	2900	mg/kg	J-	j
KP-SE-14-XX	Antimony	11	mg/kg	J-	j
KP-SE-14-XX	Chromium	46	mg/kg	J-	j
KP-SE-14-XX	Copper	2.7	mg/kg	J+	j
KP-SE-14-XX	Iron	520	mg/kg	J-	i
KP-SE-14-XX	Lead	680	mg/kg	J-	e, j
KP-SE-14-XX	Nickel	23	mg/kg	J-	i
LV-SE-29-XX	Lead	7.2	mg/kg	J+	i
LV-SE-29-XX	Mercury	1.5	mg/kg	J-	i
LV-SE-35-XX	Arsenic	31	mg/kg	J-	i
LV-SE-35-XX	Chromium	74	mg/kg	J-	i
LV-SE-35-XX	Iron	24000	mg/kg	J-	i
LV-SE-35-XX	Nickel	170	mg/kg	J-	i
LV-SE-35-XX	Vanadium	55	mg/kg	J-	i
LV-SE-35-XX	Zinc	67	mg/kg	J-	i
LV-SO-34-XX	Antimony	870	mg/kg	J-	i

#### TABLE 5: DATA QUALIFICATION: SERIAL DILUTION EXCEEDANCES

### TABLE 5: DATA QUALIFICATION: SERIAL DILUTION EXCEEDANCES (Continued)

Sample ID	Analyte	Result	Unit	Validation Qualifier	Comment Code
LV-SO-34-XX	Arsenic	110	mg/kg	J-	i
LV-SO-34-XX	Cadmium	2300	mg/kg	J-	i
LV-SO-34-XX	Chromium	2200	mg/kg	J-	j
LV-SO-34-XX	Iron	20000	mg/kg	J-	i
LV-SO-34-XX	Lead	3700	mg/kg	J-	i
LV-SO-34-XX	Nickel	1900	mg/kg	J-	i
LV-SO-34-XX	Selenium	220	mg/kg	J-	i
LV-SO-34-XX	Vanadium	230	mg/kg	J-	i
LV-SO-34-XX	Zinc	48	mg/kg	J-	i
RF-SE-16-XX	Antimony	85	mg/kg	J-	i
RF-SE-16-XX	Arsenic	72	mg/kg	J-	i
RF-SE-16-XX	Cadmium	310	mg/kg	J-	i
RF-SE-16-XX	Chromium	820	mg/kg	J-	i
RF-SE-16-XX	Copper	73	mg/kg	J-	i
RF-SE-16-XX	Iron	16000	mg/kg	J-	i
RF-SE-16-XX	Lead	24	mg/kg	J-	i
RF-SE-16-XX	Nickel	1700	mg/kg	J-	i
RF-SE-16-XX	Silver	130	mg/kg	J-	i
RF-SE-16-XX	Vanadium	32	mg/kg	J-	i
RF-SE-16-XX	Zinc	760	mg/kg	J-	j
RF-SE-24-XX	Arsenic	130	mg/kg	J+	j
RF-SE-24-XX	Cadmium	6.5	mg/kg	J+	j
RF-SE-24-XX	Chromium	74	mg/kg	J+	j
RF-SE-24-XX	Copper	860	mg/kg	J+	j
RF-SE-24-XX	Iron	24000	mg/kg	J+	j
RF-SE-24-XX	Lead	410	mg/kg	J+	j
RF-SE-24-XX	Nickel	170	mg/kg	J+	j
RF-SE-24-XX	Silver	3.8	mg/kg	J+	j
RF-SE-24-XX	Vanadium	46	mg/kg	J+	j
RF-SE-24-XX	Zinc	1400	mg/kg	J-	j
SB-SO-02-XX	Antimony	44	mg/kg	J-	e, j
SB-SO-02-XX	Arsenic	23	mg/kg	J-	j
SB-SO-02-XX	Lead	22	mg/kg	J-	j
SB-SO-02-XX	Mercury	130	mg/kg	J+	j
SB-SO-15-XX	Antimony	600	mg/kg	J-	j, e
SB-SO-15-XX	Arsenic	170	mg/kg	J-	j
SB-SO-15-XX	Chromium	91	mg/kg	J-	j
SB-SO-15-XX	Copper	30	mg/kg	J-	j
SB-SO-15-XX	Iron	51000	mg/kg	J-	j

### TABLE 5: DATA QUALIFICATION: SERIAL DILUTION EXCEEDANCES (Continued)

Sample ID	Analyte	Result	Unit	Validation Qualifier	Comment Code
SB-SO-15-XX	Lead	40	mg/kg	J-	j
SB-SO-15-XX	Nickel	100	mg/kg	J-	j
SB-SO-15-XX	Vanadium	52	mg/kg	J-	j
SB-SO-15-XX	Zinc	36	mg/kg	J-	j
SB-SO-22-XX	Antimony	10	mg/kg	J	e, j
SB-SO-22-XX	Zinc	64	mg/kg	J-	j
SB-SO-31-XX	Arsenic	8	mg/kg	J-	j
SB-SO-31-XX	Nickel	3200	mg/kg	J-	j
SB-SO-31-XX	Selenium	28	mg/kg	J-	j
SB-SO-31-XX	Silver	160	mg/kg	J-	e, j
SB-SO-31-XX	Zinc	3900	mg/kg	J-	j
TL-SE-13-XX	Antimony	95	mg/kg	J+	j, e
TL-SE-13-XX	Chromium	36	mg/kg	J+	j
TL-SE-13-XX	Copper	4400	mg/kg	J+	j
TL-SE-13-XX	Iron	22000	mg/kg	J+	j
TL-SE-13-XX	Lead	1100	mg/kg	J+	j
TL-SE-13-XX	Silver	160	mg/kg	J	j, e
TL-SE-13-XX	Vanadium	59	mg/kg	J+	j
WS-SO-01-XX	Mercury	5.8	mg/kg	J	e, j
WS-SO-33-XX	Arsenic	450	mg/kg	J-	j
WS-SO-33-XX	Cadmium	11	mg/kg	J-	j
WS-SO-33-XX	Chromium	120	mg/kg	J-	j
WS-SO-33-XX	Copper	150	mg/kg	J-	j
WS-SO-33-XX	Iron	28000	mg/kg	J-	j
WS-SO-33-XX	Lead	3700	mg/kg	J-	j
WS-SO-33-XX	Nickel	65	mg/kg	J-	j
WS-SO-33-XX	Silver	13	mg/kg	J-	j
WS-SO-33-XX	Vanadium	53	mg/kg	J-	j
WS-SO-33-XX	Zinc	830	mg/kg	J-	j

Notes:

mg/kg = Milligram per kilogram

e = Data were additionally qualified based on matrix spike/matrix spike duplicate exceedances

j = Data were qualified based on serial dilution exceedances

J = Result is estimated and biased could not be determined

J+ = Result is estimated and potentially biased high

J- = Result is estimated and potentially biased low

#### APPENDIX D

#### DEVELOPER AND REFERENCE LABORATORY DATA

Blend No.	Sample ID	Source of Data	St	)	As		Cd		Cr	Cu	Fe	Pb	Hg	7
1	KP-SO-06-XX	Reference Laboratory	8.1	$\mathbf{J}+$	1	J-	0.1	U	290	26	1,400	620	0.059	U
1	KP-SO-10-XX	Reference Laboratory	6.1	$\mathbf{J}+$	1	J-	0.1	U	300	26	1,600	560	0.028	U
1	KP-SO-15-XX	Reference Laboratory	6.3	$\mathbf{J}+$	1	J-	0.1	U	340	26	1,600	510	0.029	U
1	KP-SO-18-XX	Reference Laboratory	6.7	$\mathbf{J}+$	1	J-	0.1	U	250	24	1,200	500	0.016	U
1	KP-SO-22-XX	Reference Laboratory	8.3	$\mathbf{J}+$	1	J-	0.1	U	260	29	1,300	650	0.027	U
1	KP-SO-06-XC	Xcalibur XRF Services	8		68		25		306	17	1,292	478		
1	KP-SO-10-XC	Xcalibur XRF Services	12		111				323	14	1,287	527	2	
1	KP-SO-15-XC	Xcalibur XRF Services	6		84				298	16	1,407	541		
1	KP-SO-18-XC	Xcalibur XRF Services	8		76				277	22	1,206	518		
1	KP-SO-22-XC	Xcalibur XRF Services	6		57		23		326	16	1,389	551		
2	KP-SO-07-XX	Reference Laboratory	17	$\mathbf{J}+$	2	J-	0.1	U	170	48	990	1,200	0.027	U
2	KP-SO-13-XX	Reference Laboratory	16	$\mathbf{J}+$	1	J-	0.045	U	180	52	980	1,200	0.037	U
2	KP-SO-20-XX	Reference Laboratory	19	$\mathbf{J}+$	2	J-	0.1	U	160	46	910	1,300	0.03	U
2	KP-SO-24-XX	Reference Laboratory	17	$\mathbf{J}+$	1	J-	0.1	U	160	49	900	1,100	0.017	U
2	KP-SO-27-XX	Reference Laboratory	15	$\mathbf{J}+$	1	J-	0.05	U	170	45	970	1,200	0.021	U
2	KP-SO-29-XX	Reference Laboratory	18	$\mathbf{J}+$	2	J-	0.1	U	150	42	870	1,200	0.013	U
2	KP-SO-32-XX	Reference Laboratory	16	$\mathbf{J}+$	2	J-	0.045	U	180	50	970	1,200	0.014	U
2	KP-SO-07-XC	Xcalibur XRF Services	16		83		26		198	39	906	1,192		
2	KP-SO-13-XC	Xcalibur XRF Services	14		98				219	39	1,001	1,248	3	
2	KP-SO-20-XC	Xcalibur XRF Services	20		73				193	35	933	1,156		
2	KP-SO-24-XC	Xcalibur XRF Services	21		85				205	33	930	1,150		
2	KP-SO-27-XC	Xcalibur XRF Services	23		43		26		182	43	922	1,252		
2	KP-SO-29-XC	Xcalibur XRF Services	14		44		30		217	37	964	1,279		
2	KP-SO-32-XC	Xcalibur XRF Services	21		0				216	40	914	1,168		
3	KP-SO-04-XX	Reference Laboratory	94	$\mathbf{J}+$	3		0.046	U	180	200	1,300	5,800	0.018	U
3	KP-SO-16-XX	Reference Laboratory	93	$\mathbf{J}+$	3		0.063	U	200	230	1,400	6,100	0.016	U
3	KP-SO-23-XX	Reference Laboratory	86	$\mathbf{J}+$	3		0.048	U	180	190	1,300	5,300	0.017	U
3	KP-SO-26-XX	Reference Laboratory	90	$\mathbf{J}+$	4		0.061	U	210	230	1,500	6,500	0.013	U
3	KP-SO-31-XX	Reference Laboratory	88		28		0.1	U	140	200	1,100	5,700	0.017	U
3	KP-SO-04-XC	Xcalibur XRF Services	71		176		36		163	172	1,057	5,603	5	
3	KP-SO-16-XC	Xcalibur XRF Services	80		70		62		168	149	981	5,618	3	
3	KP-SO-23-XC	Xcalibur XRF Services			77		82		185	183	1,079	5,944	3	
3	KP-SO-26-XC	Xcalibur XRF Services	82		63				171	176	1,088	6,142	5	
3	KP-SO-31-XC	Xcalibur XRF Services	74		0		24		146	156	903	5,458		

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
1	KP-SO-06-XX	Reference Laboratory	140	0.25 U	0.25 U	2 J	11
1	KP-SO-10-XX	Reference Laboratory	150	0.22 U	0.25 U	2 J	12
1	KP-SO-15-XX	Reference Laboratory	170	0.25 U	0.25 U	2 J	15
1	KP-SO-18-XX	Reference Laboratory	120	0.25 U	0.25 U	2 J	11
1	KP-SO-22-XX	Reference Laboratory	130	0.25 U	0.25 U	2 J	11
1	KP-SO-06-XC	Xcalibur XRF Services	138	38		0	41
1	KP-SO-10-XC	Xcalibur XRF Services	144		13	0	0
1	KP-SO-15-XC	Xcalibur XRF Services	160			0	33
1	KP-SO-18-XC	Xcalibur XRF Services	138			0	21
1	KP-SO-22-XC	Xcalibur XRF Services	141	14	14	0	37
2	KP-SO-07-XX	Reference Laboratory	87	0.21 U	0.25 U	1 J	26
2	KP-SO-13-XX	Reference Laboratory	90	0.25 U	0.25 U	1 J	24
2	KP-SO-20-XX	Reference Laboratory	79	0.25 U	0.25 U	1 J	25
2	KP-SO-24-XX	Reference Laboratory	78	0.25 U	0.25 U	1 J	22
2	KP-SO-27-XX	Reference Laboratory	87	0.25 U	0.25 U	1 J	24
2	KP-SO-29-XX	Reference Laboratory	73	0.25 U	0.25 U	1 J	22
2	KP-SO-32-XX	Reference Laboratory	88	0.51	0.25 U	1 J	24
2	KP-SO-07-XC	Xcalibur XRF Services	93		13	0	67
2	KP-SO-13-XC	Xcalibur XRF Services	92		4	0	75
2	KP-SO-20-XC	Xcalibur XRF Services	86	11	5	0	56
2	KP-SO-24-XC	Xcalibur XRF Services	87			0	60
2	KP-SO-27-XC	Xcalibur XRF Services	107	35	3	0	78
2	KP-SO-29-XC	Xcalibur XRF Services	95	23	12	0	55
2	KP-SO-32-XC	Xcalibur XRF Services	83	50	6	0	60
3	KP-SO-04-XX	Reference Laboratory	93	0.28 U	0.16 J	1 J	45
3	KP-SO-16-XX	Reference Laboratory	100	0.25 U	0.16 J	1 J	47
3	KP-SO-23-XX	Reference Laboratory	91	0.25 U	0.13 J	1 J	41
3	KP-SO-26-XX	Reference Laboratory	110	0.22 U	0.17 J	1 J	52
3	KP-SO-31-XX	Reference Laboratory	68	0.25 U	0.4	2 J	38
3	KP-SO-04-XC	Xcalibur XRF Services	77	25		34	82
3	KP-SO-16-XC	Xcalibur XRF Services	70	6	11	0	80
3	KP-SO-23-XC	Xcalibur XRF Services	101	54	20	0	79
3	KP-SO-26-XC	Xcalibur XRF Services	87	59	4	0	94
3	KP-SO-31-XC	Xcalibur XRF Services	74	51	12	0	68

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg	
4	KP-SO-02-XX	Reference Laboratory	410	10	0.1	6	780	1,700	18,000	0.043	U
4	KP-SO-03-XX	Reference Laboratory	360	9	0.074 U	5	670	1,600	19,000	0.044	U
4	KP-SO-05-XX	Reference Laboratory	410	12	0.13 U	6	780	2,000	24,000	0.044	U
4	KP-SO-09-XX	Reference Laboratory	420	11	0.094 U	5	780	1,800	22,000	0.046	U
4	KP-SO-21-XX	Reference Laboratory	370	10	0.098 U	5	700	1,700	19,000	0.042	U
4	KP-SO-02-XC	Xcalibur XRF Services	233	0	32		614	1,047	18,603	15	
4	KP-SO-03-XC	Xcalibur XRF Services	242	94			641	1,084	19,642	12	
4	KP-SO-05-XC	Xcalibur XRF Services	237	0			609	1,130	19,166	12	
4	KP-SO-09-XC	Xcalibur XRF Services	240	49			641	1,070	19,206	11	
4	KP-SO-21-XC	Xcalibur XRF Services	252	86			666	1,167	20,448	13	
5	WS-SO-06-XX	Reference Laboratory	1.3 U	48	1.9	120	50	28,000	110	0.07	U
5	WS-SO-08-XX	Reference Laboratory	1.3	45	2	120	47	26,000	71	0.063	U
5	WS-SO-12-XX	Reference Laboratory	1.3 UJ	43	1.8	110	45	25,000	65	0.068	UJ
5	WS-SO-17-XX	Reference Laboratory	1.3 UJ	47	1.9	120	49	28,000	70	0.069 1	UJ
5	WS-SO-27-XX	Reference Laboratory	1.3 UJ	49	2	120	51	28,000	72	0.11 J	J-
5	WS-SO-30-XX	Reference Laboratory	1.2 J-	51	2	130	53	29,000	81	0.069	UJ
5	WS-SO-35-XX	Reference Laboratory	1.3 UJ	49	2	130	51	28,000	74	0.071	UJ
5	WS-SO-06-XC	Xcalibur XRF Services	1	105	31	61	68	21,856	202		
5	WS-SO-08-XC	Xcalibur XRF Services	1	108		79	73	21,786	180		
5	WS-SO-12-XC	Xcalibur XRF Services	1	113		59	69	21,623	164	3	
5	WS-SO-17-XC	Xcalibur XRF Services	1	102	25	49	70	21,219	181	5	
5	WS-SO-27-XC	Xcalibur XRF Services	3	81		64	61	18,393	172		
5	WS-SO-30-XC	Xcalibur XRF Services	1	60	41	39	56	19,246	149		
5	WS-SO-35-XC	Xcalibur XRF Services	1	94	30		61	19,086	152		
6	WS-SO-03-XX	Reference Laboratory	8.9 J-	500	12	140	170	32,000	4,300	0.86 J	J-
6	WS-SO-05-XX	Reference Laboratory	8.6 J-	440	12	140	160	31,000	4,000	0.76 J	J-
6	WS-SO-09-XX	Reference Laboratory	7.1 J-	480	12	130	160	30,000	4,000	0.89 J	J-
6	WS-SO-14-XX	Reference Laboratory	8.4 J-	430	11	120	150	28,000	3,700	0.74 J	J-
6	WS-SO-26-XX	Reference Laboratory	7.6 J-	520	12	140	160	30,000	4,000	0.83 J	J-
6	WS-SO-31-XX	Reference Laboratory	7.2 J-	520	12	140	170	32,000	4,200	0.85 J	J-
6	WS-SO-33-XX	Reference Laboratory	6.9 J-	450 J-	11 J-	120 J-	150 J-	28,000 J-	3,700 J-	0.87 J	J-
6	WS-SO-03-XC	Xcalibur XRF Services	36	473	34	52	199	23,201	9,334		
6	WS-SO-05-XC	Xcalibur XRF Services	41	537	10	56	187	23,725	9,860		
6	WS-SO-09-XC	Xcalibur XRF Services	38	490	12	72	195	23,021	9,696		
6	WS-SO-14-XC	Xcalibur XRF Services	39	427		64	182	22,896	9,433	7	
6	WS-SO-26-XC	Xcalibur XRF Services	23	297	36	51	150	19,241	6,110	27	
6	WS-SO-31-XC	Xcalibur XRF Services	26	252			158	19,974	7,009		
6	WS-SO-33-XC	Xcalibur XRF Services	25	195	28	45	165	20,042	6,809	9	

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	i	Se		Ag	r	V		Zn	
4	KP-SO-02-XX	Reference Laboratory	4		0.42	U	0.82		0	J	100	
4	KP-SO-03-XX	Reference Laboratory	3		0.25	U	0.73		0	J	92	
4	KP-SO-05-XX	Reference Laboratory	4		0.24	U	0.82		0	J	110	
4	KP-SO-09-XX	Reference Laboratory	3		0.25	U	0.84		0	J	110	
4	KP-SO-21-XX	Reference Laboratory	4		0.25	U	0.76		0	J	100	
4	KP-SO-02-XC	Xcalibur XRF Services	2		275		42		0		176	
4	KP-SO-03-XC	Xcalibur XRF Services	2		230		31		0		123	
4	KP-SO-05-XC	Xcalibur XRF Services	3		231		13		0		154	
4	KP-SO-09-XC	Xcalibur XRF Services	2		240		25		0		178	
4	KP-SO-21-XC	Xcalibur XRF Services	3		211		27		0		196	
5	WS-SO-06-XX	Reference Laboratory	61		1.3	U	0.93	J	56		230	
5	WS-SO-08-XX	Reference Laboratory	58		1.3	U	0.86	J	52		220	
5	WS-SO-12-XX	Reference Laboratory	55		1.3	U	0.94	J	49		210	
5	WS-SO-17-XX	Reference Laboratory	59		1.3	U	0.89	J	56		230	
5	WS-SO-27-XX	Reference Laboratory	61		1.3	U	0.9	J	57		230	
5	WS-SO-30-XX	Reference Laboratory	65		1.3	U	1	J	58		240	
5	WS-SO-35-XX	Reference Laboratory	62		1.3	U	1	J	57		240	
5	WS-SO-06-XC	Xcalibur XRF Services	39		34		10		23		380	
5	WS-SO-08-XC	Xcalibur XRF Services	57		15		11		24		387	
5	WS-SO-12-XC	Xcalibur XRF Services	39		5		10		48		385	
5	WS-SO-17-XC	Xcalibur XRF Services	38				10		15		384	
5	WS-SO-27-XC	Xcalibur XRF Services	55				6		36		345	
5	WS-SO-30-XC	Xcalibur XRF Services	35						28		345	
5	WS-SO-35-XC	Xcalibur XRF Services	70				6		55		348	
6	WS-SO-03-XX	Reference Laboratory	75		1.6		15		58		930	
6	WS-SO-05-XX	Reference Laboratory	71		1.3	U	15		57		900	
6	WS-SO-09-XX	Reference Laboratory	70		1.3	U	14		56		870	
6	WS-SO-14-XX	Reference Laboratory	64		1.3	U	13		50		820	
6	WS-SO-26-XX	Reference Laboratory	70		1.3	U	14		56		900	
6	WS-SO-31-XX	Reference Laboratory	72		1.2	U	15		60		950	
6	WS-SO-33-XX	Reference Laboratory	65	J-	1.3	U	13	J-	53	J-	830	J-
6	WS-SO-03-XC	Xcalibur XRF Services	71				27		23		1,191	
6	WS-SO-05-XC	Xcalibur XRF Services	58		43		38		36		1,132	
6	WS-SO-09-XC	Xcalibur XRF Services	55		19		30		13		1,190	
6	WS-SO-14-XC	Xcalibur XRF Services	67				24		46		1,134	
6	WS-SO-26-XC	Xcalibur XRF Services	60		16		18		31		934	
6	WS-SO-31-XC	Xcalibur XRF Services	35				29		64		996	
6	WS-SO-33-XC	Xcalibur XRF Services	67				15		0		1,020	

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	St	)	As	Cd	Cr	Cu	Fe	Pb	Hg
7	WS-SO-01-XX	Reference Laboratory	41	J-	1900	47	100	590	32,000	18,000	5.8 J
7	WS-SO-04-XX	Reference Laboratory	45	J-	2000	50	94	640	34,000	20,000	6.5
7	WS-SO-15-XX	Reference Laboratory	48	J-	2300	56	82	720	37,000	24,000	5.8
7	WS-SO-22-XX	Reference Laboratory	41	J-	1900	47	84	620	33,000	17,000	4.8
7	WS-SO-34-XX	Reference Laboratory	45	J-	2000	50	91	660	36,000	22,000	5.4
7	WS-SO-01-XC	Xcalibur XRF Services	144		2,917	67	85	725	29,297	42,266	
7	WS-SO-04-XC	Xcalibur XRF Services	152		2,719	102	81	658	25,925	39,796	
7	WS-SO-15-XC	Xcalibur XRF Services	140		2,780	44	64	734	29,457	42,430	
7	WS-SO-22-XC	Xcalibur XRF Services	137		1,689	76	63	695	27,447	37,445	90
7	WS-SO-34-XC	Xcalibur XRF Services	136		2,066	47	60	597	23,244	30,445	65
8	WS-SO-02-XX	Reference Laboratory	130	J-	4200	98	49	1,300	44,000	35,000	17
8	WS-SO-16-XX	Reference Laboratory	110	J-	3900	91	59	1,300	42,000	24,000	15
8	WS-SO-18-XX	Reference Laboratory	130	J-	4100	95	63	1,300	44,000	37,000	17
8	WS-SO-21-XX	Reference Laboratory	120	J-	3900	90	43	1,200	40,000	43,000	14
8	WS-SO-24-XX	Reference Laboratory	97	J-	3600	81	54	1,100	38,000	27,000	16
8	WS-SO-29-XX	Reference Laboratory	120	J-	3800	90	51	1,200	40,000	42,000	15
8	WS-SO-37-XX	Reference Laboratory	120	J-	4100	95	63	1,300	42,000	26,000	14
8	WS-SO-02-XC	Xcalibur XRF Services	178		5,187	145	101	1,266	33,774	64,837	
8	WS-SO-16-XC	Xcalibur XRF Services	170		4,682	122	98	1,259	34,523	61,339	30
8	WS-SO-18-XC	Xcalibur XRF Services	194		3,655	120		1,211	31,756	59,680	46
8	WS-SO-21-XC	Xcalibur XRF Services	177		4,194	114		1,144	32,103	56,773	43
8	WS-SO-24-XC	Xcalibur XRF Services	202		4,123	82		932	24,425	40,706	116
8	WS-SO-29-XC	Xcalibur XRF Services	213		3,683	98	51	1,052	27,220	48,582	100
8	WS-SO-37-XC	Xcalibur XRF Services	202		4,497	62		1,071	28,622	48,690	28
9	WS-SO-13-XX	Reference Laboratory	200	J-	5800	150	53	1800	47,000	45,000	11
9	WS-SO-19-XX	Reference Laboratory	150	J-	5000	130	66	1500	39,000	24,000	12
9	WS-SO-28-XX	Reference Laboratory	120	J-	4200	100	54	1200	33,000	30,000	11
9	WS-SO-32-XX	Reference Laboratory	190	J-	5500	140	54	1700	44,000	30,000	11
9	WS-SO-36-XX	Reference Laboratory	120	J-	3800	92	51	1100	30,000	45,000	13
9	WS-SO-13-XC	Xcalibur XRF Services	208		7,772	139	67	1,511	35,738	47,796	
9	WS-SO-19-XC	Xcalibur XRF Services	178		4,948	184		1,559	39,189	48,951	84
9	WS-SO-28-XC	Xcalibur XRF Services	258		5,176	130		1,207	27,159	39,360	90
9	WS-SO-32-XC	Xcalibur XRF Services	237		5,696	131		1,331	29,668	41,616	52
9	WS-SO-36-XC	Xcalibur XRF Services	260		6,570	107		1,260	29,661	39,863	48

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
7	WS-SO-01-XX	Reference Laboratory	66	1.3 U	69 J-	42	3,000
7	WS-SO-04-XX	Reference Laboratory	62	1.3 U	76 J-	44	3,100
7	WS-SO-15-XX	Reference Laboratory	58	1.3 U	90 J-	52	3,400
7	WS-SO-22-XX	Reference Laboratory	57	1.3 U	72 J-	44	3,000
7	WS-SO-34-XX	Reference Laboratory	60	1.3 U	78 J-	47	3,200
7	WS-SO-01-XC	Xcalibur XRF Services	42		105	0	4,227
7	WS-SO-04-XC	Xcalibur XRF Services	38		99	0	4,061
7	WS-SO-15-XC	Xcalibur XRF Services	79		104	0	4,265
7	WS-SO-22-XC	Xcalibur XRF Services	67		86	0	4,070
7	WS-SO-34-XC	Xcalibur XRF Services	65		59	0	3,534
8	WS-SO-02-XX	Reference Laboratory	57	1.3 U	150 J-	36	6,000
8	WS-SO-16-XX	Reference Laboratory	60	1.1 J	150 J-	35	5,700
8	WS-SO-18-XX	Reference Laboratory	62	1.9	140 J-	36	5,900
8	WS-SO-21-XX	Reference Laboratory	51	1.6	150 J-	33	5,500
8	WS-SO-24-XX	Reference Laboratory	54	2.1	140 J-	30	5,200
8	WS-SO-29-XX	Reference Laboratory	55	1.7	140 J-	33	5,500
8	WS-SO-37-XX	Reference Laboratory	63	3	140 J-	34	5,800
8	WS-SO-02-XC	Xcalibur XRF Services	95		195	0	7,030
8	WS-SO-16-XC	Xcalibur XRF Services	85		182	17	6,880
8	WS-SO-18-XC	Xcalibur XRF Services	42		199	41	6,777
8	WS-SO-21-XC	Xcalibur XRF Services	42		179	0	6,552
8	WS-SO-24-XC	Xcalibur XRF Services	90	126	105	12	5,451
8	WS-SO-29-XC	Xcalibur XRF Services	80		144	0	5,972
8	WS-SO-37-XC	Xcalibur XRF Services	39	44	117	0	5,885
9	WS-SO-13-XX	Reference Laboratory	75	3.7	170 J-	24	9,000
9	WS-SO-19-XX	Reference Laboratory	74	3.7	160 J-	20	7,700
9	WS-SO-28-XX	Reference Laboratory	59	2.3	130 J-	16	6,100
9	WS-SO-32-XX	Reference Laboratory	73	3.7	190 J-	23	8,500
9	WS-SO-36-XX	Reference Laboratory	55	1.7	120 J-	15	5,700
9	WS-SO-13-XC	Xcalibur XRF Services	112		206	0	9,997
9	WS-SO-19-XC	Xcalibur XRF Services	43		223	0	10,289
9	WS-SO-28-XC	Xcalibur XRF Services	90		137	0	8,400
9	WS-SO-32-XC	Xcalibur XRF Services	97		163	73	8,948
9	WS-SO-36-XC	Xcalibur XRF Services	100	14	144	77	8,632

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg
10	BN-SO-01-XX	Reference Laboratory	1.3 UJ	38	0.94	120	32	24,000	63	0.13
10	BN-SO-10-XX	Reference Laboratory	1.3 UJ	50	1.2	110	35	24,000	140	0.14
10	BN-SO-15-XX	Reference Laboratory	1.3 UJ	34	0.82	110	29	22,000	56	0.15
10	BN-SO-18-XX	Reference Laboratory	1.3 U	37	0.89	110	29	22,000	59	0.13
10	BN-SO-28-XX	Reference Laboratory	1.5	35	0.87	100	28	22,000	58	0.16
10	BN-SO-31-XX	Reference Laboratory	1.3	41	1	140	33	26,000	65	0.14
10	BN-SO-35-XX	Reference Laboratory	1.4	37	0.98	120	30	23,000	60	0.15
10	BN-SO-01-XC	Xcalibur XRF Services	10	47	11	86	26	17,877	115	
10	BN-SO-10-XC	Xcalibur XRF Services	4	64		64	37	18,087	126	
10	BN-SO-15-XC	Xcalibur XRF Services	8	79	25	82	31	18,208	93	
10	BN-SO-18-XC	Xcalibur XRF Services	8	74	33	101	30	18,517	84	
10	BN-SO-28-XC	Xcalibur XRF Services	0	89		92	39	18,150	88	2
10	BN-SO-31-XC	Xcalibur XRF Services	7	84	25	85	46	18,130	163	
10	BN-SO-35-XC	Xcalibur XRF Services	9	89		78	45	18,569	134	5
11	BN-SO-02-XX	Reference Laboratory	11	140	50	90	170	28,000	840	0.37
11	BN-SO-04-XX	Reference Laboratory	9.1	120	42	79	140	24,000	700	0.36
11	BN-SO-17-XX	Reference Laboratory	9.3	110	39	79	140	23,000	680	0.39
11	BN-SO-22-XX	Reference Laboratory	7.3	98	34	65	110	20,000	590	0.37
11	BN-SO-27-XX	Reference Laboratory	9.6	110	39	78	130	24,000	660	0.38
11	BN-SO-02-XC	Xcalibur XRF Services	12	164	26	43	149	17,560	1,155	
11	BN-SO-04-XC	Xcalibur XRF Services	13	160	39	56	146	18,022	1,154	
11	BN-SO-17-XC	Xcalibur XRF Services	13	130	46	62	162	18,512	1,176	
11	BN-SO-22-XC	Xcalibur XRF Services	4	203	73	50	158	18,691	1,137	
11	BN-SO-27-XC	Xcalibur XRF Services	23	155	64	36	156	18,679	1,192	
12	BN-SO-03-XX	Reference Laboratory	65	620	290	120	840	25,000	4,700	1.6
12	BN-SO-06-XX	Reference Laboratory	60	600	280	94	810	24,000	4,500	2
12	BN-SO-08-XX	Reference Laboratory	57	570	270	100	750	22,000	4,300	2
12	BN-SO-13-XX	Reference Laboratory	65	320	150	98	410	17,000	2,400	1.6
12	BN-SO-20-XX	Reference Laboratory	57	540	260	88	730	22,000	4,100	1.6
12	BN-SO-30-XX	Reference Laboratory	64	630	300	100	860	26,000	4,800	1.6
12	BN-SO-34-XX	Reference Laboratory	68	630	290	110	830	25,000	4,700	2
12	BN-SO-03-XC	Xcalibur XRF Services	61	684	298	75	850	21,467	6,485	
12	BN-SO-06-XC	Xcalibur XRF Services	66	751	284	104	869	21,431	6,565	
12	BN-SO-08-XC	Xcalibur XRF Services	63	731	319	91	851	21,601	6,593	11
12	BN-SO-13-XC	Xcalibur XRF Services	70	715	326	60	848	21,842	6,781	9
12	BN-SO-20-XC	Xcalibur XRF Services	62	752	321	76	837	21,577	6,406	
12	BN-SO-30-XC	Xcalibur XRF Services	63	597	307	61	837	21,784	6,448	
12	BN-SO-34-XC	Xcalibur XRF Services	62	652	312	44	873	22,434	6,660	

#### Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
10	BN-SO-01-XX	Reference Laboratory	63	1.3 U	1.3 UJ	55	92
10	BN-SO-10-XX	Reference Laboratory	54	1.2 J	1.3 UJ	55	110
10	BN-SO-15-XX	Reference Laboratory	58	1.3 U	1.3 UJ	49	89
10	BN-SO-18-XX	Reference Laboratory	59	1.3	0.94 U	46	88
10	BN-SO-28-XX	Reference Laboratory	54	1.3 U	0.77 U	48	81
10	BN-SO-31-XX	Reference Laboratory	71	1.3 U	0.97 U	54	94
10	BN-SO-35-XX	Reference Laboratory	63	1.2 J	0.85 U	50	87
10	BN-SO-01-XC	Xcalibur XRF Services	54		11	0	155
10	BN-SO-10-XC	Xcalibur XRF Services	69		11	15	172
10	BN-SO-15-XC	Xcalibur XRF Services	60			34	157
10	BN-SO-18-XC	Xcalibur XRF Services	85	2	18	0	169
10	BN-SO-28-XC	Xcalibur XRF Services	66		8	0	159
10	BN-SO-31-XC	Xcalibur XRF Services	71			46	206
10	BN-SO-35-XC	Xcalibur XRF Services	61		6	44	193
11	BN-SO-02-XX	Reference Laboratory	54	4.3	7.6	60	470
11	BN-SO-04-XX	Reference Laboratory	48	2.9	6.5	50	400
11	BN-SO-17-XX	Reference Laboratory	47	2.7	6.3	49	390
11	BN-SO-22-XX	Reference Laboratory	40	2.8	5.4	43	330
11	BN-SO-27-XX	Reference Laboratory	46	3.7	6.1	52	380
11	BN-SO-02-XC	Xcalibur XRF Services	32	6		29	665
11	BN-SO-04-XC	Xcalibur XRF Services	33			0	643
11	BN-SO-17-XC	Xcalibur XRF Services	57	72	14	42	646
11	BN-SO-22-XC	Xcalibur XRF Services	59			14	669
11	BN-SO-27-XC	Xcalibur XRF Services	34		14	24	661
12	BN-SO-03-XX	Reference Laboratory	100	17	42	48	2,300
12	BN-SO-06-XX	Reference Laboratory	92	15	41	48	2,300
12	BN-SO-08-XX	Reference Laboratory	94	14	38	39	2,200
12	BN-SO-13-XX	Reference Laboratory	71	9.2	21	37	1,200
12	BN-SO-20-XX	Reference Laboratory	84	14	37	44	2,100
12	BN-SO-30-XX	Reference Laboratory	99	17	44	50	2,400
12	BN-SO-34-XX	Reference Laboratory	100	17	42	49	2,300
12	BN-SO-03-XC	Xcalibur XRF Services	83	67	33	61	3,299
12	BN-SO-06-XC	Xcalibur XRF Services	72		44	44	3,454
12	BN-SO-08-XC	Xcalibur XRF Services	79	6	41	0	3,331
12	BN-SO-13-XC	Xcalibur XRF Services	69	114	65	0	3,358
12	BN-SO-20-XC	Xcalibur XRF Services	86	65	31	0	3,356
12	BN-SO-30-XC	Xcalibur XRF Services	76	101	37	0	3,325
12	BN-SO-34-XC	Xcalibur XRF Services	82		58	0	3,406

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg
13	BN-SO-07-XX	Reference Laboratory	110 J-	990 J+	520	82	1,400	23,000	6,900	3.4
13	BN-SO-16-XX	Reference Laboratory	120 J-	1,100 J+	570	86	1,500	25,000	8,100	3.4
13	BN-SO-21-XX	Reference Laboratory	150 J-	1,300 J+	660	110	1,700	30,000	8,900	3.6
13	BN-SO-25-XX	Reference Laboratory	82 J-	700 J	370 J-	64 J-	930 J-	16,000 J-	5,400 J-	3.8
13	BN-SO-33-XX	Reference Laboratory	100 J-	1,100	640	100	1,600	27,000	8,000	4
13	BN-SO-07-XC	Xcalibur XRF Services	100	1,133	601		1,456	23,645	8,902	
13	BN-SO-16-XC	Xcalibur XRF Services	99	1,134	561	74	1,457	23,360	8,699	
13	BN-SO-21-XC	Xcalibur XRF Services	105	1,120	586	57	1,466	24,645	8,942	
13	BN-SO-25-XC	Xcalibur XRF Services	103	1,016	592	105	1,369	22,952	8,762	
13	BN-SO-33-XC	Xcalibur XRF Services	100	1,243	599	39	1,527	25,053	8,705	
14	BN-SO-05-XX	Reference Laboratory	160 J-	1,600	850	86	2,200	26,000	12,000	5
14	BN-SO-19-XX	Reference Laboratory	150 J-	1,600	860	79	2,200	26,000	12,000	5
14	BN-SO-26-XX	Reference Laboratory	150 J-	1,700	900	82	2,400	27,000	12,000	5.4
14	BN-SO-29-XX	Reference Laboratory	150 J-	1,600	880	86	2,300	26,000	12,000	5.4
14	BN-SO-32-XX	Reference Laboratory	160 J-	1,600	860	84	2,300	26,000	12,000	5.4
14	BN-SO-05-XC	Xcalibur XRF Services	136	1,935	808	87	2,193	26,751	10,794	
14	BN-SO-19-XC	Xcalibur XRF Services	131	1,673	811	63	2,199	27,009	10,790	
14	BN-SO-26-XC	Xcalibur XRF Services	134	1,719	821	45	2,194	26,274	10,750	
14	BN-SO-29-XC	Xcalibur XRF Services	130	1,751	834		2,254	27,470	10,894	
14	BN-SO-32-XC	Xcalibur XRF Services	132	1,843	852	45	2,187	26,292	10,562	
15	CN-SO-01-XX	Reference Laboratory	13 J-	13	21	190	700	38,000	1,200	0.13
15	CN-SO-04-XX	Reference Laboratory	13 J-	11	21	200	680	37,000	1,200	0.14
15	CN-SO-08-XX	Reference Laboratory	15 J-	15	25	210	740	43,000	1,300	0.16
15	CN-SO-10-XX	Reference Laboratory	13 J-	13	22	200	760	39,000	1,200	0.12
15	CN-SO-11-XX	Reference Laboratory	17 J-	16	30	240	860	47,000	1,600	0.15
15	CN-SO-01-XC	Xcalibur XRF Services	16	99	20	158	731	34,316	1,738	
15	CN-SO-04-XC	Xcalibur XRF Services	19	111		179	741	34,495	1,901	
15	CN-SO-08-XC	Xcalibur XRF Services	19	40	19	179	737	32,236	1,790	
15	CN-SO-10-XC	Xcalibur XRF Services	14	83	20	170	752	33,088	1,755	
15	CN-SO-11-XC	Xcalibur XRF Services	21	93		155	729	33,039	1,806	
16	AS-SO-02-XX	Reference Laboratory	2.6 UJ	18	50	180	140	48,000	1,600	0.76
16	AS-SO-06-XX	Reference Laboratory	2.4 UJ	19	52	190	130	52,000	1,600	0.74
16	AS-SO-10-XX	Reference Laboratory	1.9 J-	18	48	180	110	45,000	1,400	0.78
16	AS-SO-11-XX	Reference Laboratory	3.7 J-	22	63	230	150	52,000	2,100	0.72
16	AS-SO-13-XX	Reference Laboratory	2.4 UJ	20	57	200	150	52,000	1,700	0.79
16	AS-SO-02-XC	Xcalibur XRF Services	14	85	64	146	311	33,843	2,497	10
16	AS-SO-06-XC	Xcalibur XRF Services	12	85	33	225	319	36,284	2,390	15
16	AS-SO-10-XC	Xcalibur XRF Services	12	34	63	172	330	36,503	2,495	17
16	AS-SO-11-XC	Xcalibur XRF Services	14	42	62	154	349	38,108	2,669	12
16	AS-SO-13-XC	Xcalibur XRF Services	13	41	47	175	350	36,984	2,579	8

#### Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni		Se	;	Ag	5	V	7	Zn	
13	BN-SO-07-XX	Reference Laboratory	120		26		70		41		4,000	
13	BN-SO-16-XX	Reference Laboratory	130		29		77		44		4,400	
13	BN-SO-21-XX	Reference Laboratory	160		35		88		52		5,100	
13	BN-SO-25-XX	Reference Laboratory	88	J-	19	J-	48	J-	28	J-	2,900	J-
13	BN-SO-33-XX	Reference Laboratory	150		34		81		48		5,100	
13	BN-SO-07-XC	Xcalibur XRF Services	100		47		75		31		5,643	
13	BN-SO-16-XC	Xcalibur XRF Services	71		15		73		41		5,547	
13	BN-SO-21-XC	Xcalibur XRF Services	103		20		76		0		5,498	
13	BN-SO-25-XC	Xcalibur XRF Services	99				66		0		5,524	
13	BN-SO-33-XC	Xcalibur XRF Services	117				84		0		5,746	
14	BN-SO-05-XX	Reference Laboratory	160		48		110		39		6,700	
14	BN-SO-19-XX	Reference Laboratory	160		48		120		39		6,700	
14	BN-SO-26-XX	Reference Laboratory	160		49		120		40		7,000	
14	BN-SO-29-XX	Reference Laboratory	160		48		120		41		6,800	
14	BN-SO-32-XX	Reference Laboratory	160		48		120		39		6,700	
14	BN-SO-05-XC	Xcalibur XRF Services	113		100		123		0		7,367	
14	BN-SO-19-XC	Xcalibur XRF Services	134		72		106		0		7,318	
14	BN-SO-26-XC	Xcalibur XRF Services	107				117		0		7,323	
14	BN-SO-29-XC	Xcalibur XRF Services	126				128		0		7,482	
14	BN-SO-32-XC	Xcalibur XRF Services	125				115		30		7,359	
15	CN-SO-01-XX	Reference Laboratory	240		2.2		12		21		3,100	
15	CN-SO-04-XX	Reference Laboratory	240		1.5		12		22		2,900	
15	CN-SO-08-XX	Reference Laboratory	280		1.3	U	15		26		3,200	
15	CN-SO-10-XX	Reference Laboratory	240		1.9		14		22		3,000	
15	CN-SO-11-XX	Reference Laboratory	320		1.3	U	16		27		3,500	
15	CN-SO-01-XC	Xcalibur XRF Services	198				18		48		3,910	
15	CN-SO-04-XC	Xcalibur XRF Services	183				23		0		3,742	
15	CN-SO-08-XC	Xcalibur XRF Services	184		55		3		51		3,732	
15	CN-SO-10-XC	Xcalibur XRF Services	185				15		56		3,451	
15	CN-SO-11-XC	Xcalibur XRF Services	217				3		0		3,451	
16	AS-SO-02-XX	Reference Laboratory	91		2.6	U	4.5		42		3,300	
16	AS-SO-06-XX	Reference Laboratory	93		2.6	U	4.8		44		3,500	
16	AS-SO-10-XX	Reference Laboratory	84		1.1	U	4.4		42		3,000	
16	AS-SO-11-XX	Reference Laboratory	120		1.1	U	5.6		54		3,800	
16	AS-SO-13-XX	Reference Laboratory	100		3		5.2		50		3,800	
16	AS-SO-02-XC	Xcalibur XRF Services	92				5		60		3,543	
16	AS-SO-06-XC	Xcalibur XRF Services	86						0		3,424	
16	AS-SO-10-XC	Xcalibur XRF Services	95				0		0		3,321	
16	AS-SO-11-XC	Xcalibur XRF Services	86				10		0		3,440	
16	AS-SO-13-XC	Xcalibur XRF Services	86				8		0		3,539	

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	S	b	As	Cd		Cr	Cu	Fe	Pb	Hg
17	AS-SO-01-XX	Reference Laboratory	3.8	J-	26	100		420	250	100,000	3,200	1.4
17	AS-SO-04-XX	Reference Laboratory	6.4	UJ	22	110		480	260	110,000	3,300	1.3
17	AS-SO-07-XX	Reference Laboratory	3.6	J-	21	97		380	240	88,000	2,900	1.4
17	AS-SO-09-XX	Reference Laboratory	2.6	UJ	25 J-	100	J-	390 J-	250 J-	94,000 J-	3,200 J-	1.4
17	AS-SO-12-XX	Reference Laboratory	2.6	UJ	29	120		440	270	93,000	3,300	1.4
17	AS-SO-01-XC	Xcalibur XRF Services	4		74	67		371	591	74,049	6,123	53
17	AS-SO-04-XC	Xcalibur XRF Services			46	75		364	667	79,399	6,244	56
17	AS-SO-07-XC	Xcalibur XRF Services			45	105		415	623	78,896	6,402	51
17	AS-SO-09-XC	Xcalibur XRF Services	6		46	78		523	629	83,038	6,597	47
17	AS-SO-12-XC	Xcalibur XRF Services	6		85	84		313	686	82,623	6,547	44
18	SB-SO-03-XX	Reference Laboratory	1.2	UJ	9	0.51	U	150	48	38,000	18	62
18	SB-SO-06-XX	Reference Laboratory	1.7	J-	8	0.51	U	140	44	35,000	16	55
18	SB-SO-14-XX	Reference Laboratory	4.1	J-	9	0.51	U	150	46	37,000	17	55
18	SB-SO-38-XX	Reference Laboratory	1.3	UJ	10	0.51	U	150	57	37,000	18	56
18	SB-SO-41-XX	Reference Laboratory	1.3	UJ	9	0.51	U	160	58	40,000	19	54
18	SB-SO-47-XX	Reference Laboratory	1.3	UJ	8	0.51	U	140	44	34,000	16	58
18	SB-SO-51-XX	Reference Laboratory	1.3	UJ	9	0.51	U	160	50	40,000	18	54
18	SB-SO-03-XC	Xcalibur XRF Services	6		50	10		104	28	27,814	85	11
18	SB-SO-06-XC	Xcalibur XRF Services	0		63			115	20	27,713	88	15
18	SB-SO-14-XC	Xcalibur XRF Services	3		82			130	37	27,854	52	16
18	SB-SO-38-XC	Xcalibur XRF Services	4		72			148	27	27,016	75	19
18	SB-SO-41-XC	Xcalibur XRF Services	9		63			160	28	26,866	87	22
18	SB-SO-47-XC	Xcalibur XRF Services	15		77			183	29	27,284	0	9
18	SB-SO-51-XC	Xcalibur XRF Services	4		60	48		119	33	27,183	106	29
19	SB-SO-05-XX	Reference Laboratory	1.6	J-	9	0.51	U	140	46	35,000	16	540
19	SB-SO-18-XX	Reference Laboratory	1.2	UJ	10	0.51	U	150	46	38,000	17	280
19	SB-SO-30-XX	Reference Laboratory	3.2	J-	7	0.51	U	94	27	22,000	10	290
19	SB-SO-40-XX	Reference Laboratory	2.2	J-	9	0.51	U	120	40	33,000	15	280
19	SB-SO-53-XX	Reference Laboratory	1.2	UJ	10	0.51	U	140	44	37,000	17	270
19	SB-SO-05-XC	Xcalibur XRF Services	9		70	29		132	5	25,964	0	219
19	SB-SO-18-XC	Xcalibur XRF Services	0		82	34		68	23	23,181	68	155
19	SB-SO-30-XC	Xcalibur XRF Services	0		65	31		62	39	25,684	64	187
19	SB-SO-40-XC	Xcalibur XRF Services	0		72			153	37	25,849	0	205
19	SB-SO-53-XC	Xcalibur XRF Services	13		46			106	31	26,290	41	195

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se		Ag		V	Zn
17	AS-SO-01-XX	Reference Laboratory	180	2.6	U	9.3		66	6,900
17	AS-SO-04-XX	Reference Laboratory	200	6.2	U	12		72	7,400
17	AS-SO-07-XX	Reference Laboratory	160	2.7		8.9		63	6,300
17	AS-SO-09-XX	Reference Laboratory	170 J-	2.6	U	9.6	J-	65 J-	6,800 J-
17	AS-SO-12-XX	Reference Laboratory	190	2.6	U	3.2		73	7,500
17	AS-SO-01-XC	Xcalibur XRF Services	130			3		88	2,484
17	AS-SO-04-XC	Xcalibur XRF Services	153			4		0	2,195
17	AS-SO-07-XC	Xcalibur XRF Services	159	24		9		0	2,196
17	AS-SO-09-XC	Xcalibur XRF Services	119			9		0	1,858
17	AS-SO-12-XC	Xcalibur XRF Services	133			7		37	2,034
18	SB-SO-03-XX	Reference Laboratory	210	1.3	U	1.3	U	67	90
18	SB-SO-06-XX	Reference Laboratory	200	1.3	U	1.3	U	63	82
18	SB-SO-14-XX	Reference Laboratory	210	1.3	U	1.3	U	66	95
18	SB-SO-38-XX	Reference Laboratory	210	1.3	U	1.3	U	68	91
18	SB-SO-41-XX	Reference Laboratory	230	1.3	U	1.3	U	71	96
18	SB-SO-47-XX	Reference Laboratory	200	1.3	U	1.3	U	62	82
18	SB-SO-51-XX	Reference Laboratory	230	1.3	U	1.3	U	74	93
18	SB-SO-03-XC	Xcalibur XRF Services	179	54		4		90	109
18	SB-SO-06-XC	Xcalibur XRF Services	197					36	98
18	SB-SO-14-XC	Xcalibur XRF Services	173			6		77	108
18	SB-SO-38-XC	Xcalibur XRF Services	196					35	114
18	SB-SO-41-XC	Xcalibur XRF Services	171					0	98
18	SB-SO-47-XC	Xcalibur XRF Services	168					0	114
18	SB-SO-51-XC	Xcalibur XRF Services	167			6		55	102
19	SB-SO-05-XX	Reference Laboratory	200	1.3	U	1.3	U	61	80
19	SB-SO-18-XX	Reference Laboratory	210	1.3	U	1.3	U	70	84
19	SB-SO-30-XX	Reference Laboratory	120	1.3	J+	1.3	U	43	50
19	SB-SO-40-XX	Reference Laboratory	180	1.3	U	1.3	U	58	74
19	SB-SO-53-XX	Reference Laboratory	200	1.3	U	1.3	U	64	81
19	SB-SO-05-XC	Xcalibur XRF Services	165					62	92
19	SB-SO-18-XC	Xcalibur XRF Services	139					0	89
19	SB-SO-30-XC	Xcalibur XRF Services	144	13		8		147	90
19	SB-SO-40-XC	Xcalibur XRF Services	154			16		42	106
19	SB-SO-53-XC	Xcalibur XRF Services	158			8		74	88

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sł	)	As	Cd		Cr	Cu	Fe	Pb	Hg
20	SB-SO-08-XX	Reference Laboratory	5.4	J-	13	0.51	U	120	39	32,000	17	730
20	SB-SO-11-XX	Reference Laboratory	5.7	J-	13	0.51	U	140	46	36,000	20	810
20	SB-SO-21-XX	Reference Laboratory	4.9	J	13	0.51	U	130	43	34,000	18	740
20	SB-SO-39-XX	Reference Laboratory	4.7	J-	13	0.51	U	140	46	34,000	19	790
20	SB-SO-42-XX	Reference Laboratory	4.6	J-	13	0.51	U	140	45	35,000	18	740
20	SB-SO-08-XC	Xcalibur XRF Services	10		63			109	25	24,137	0	656
20	SB-SO-11-XC	Xcalibur XRF Services	11		71			97	26	24,094	0	650
20	SB-SO-21-XC	Xcalibur XRF Services	0		72			51	18	22,026	0	485
20	SB-SO-39-XC	Xcalibur XRF Services	8		90			133	27	23,521	0	632
20	SB-SO-42-XC	Xcalibur XRF Services	13		120	26		132	5	24,055	0	617
21	SB-SO-22-XX	Reference Laboratory	10	J	18	0.51	U	120	37	29,000	22	3300
21	SB-SO-25-XX	Reference Laboratory	6.8	$\mathbf{J}+$	18	0.51	U	120	37	29,000	22	3000
21	SB-SO-27-XX	Reference Laboratory	6.7	$\mathbf{J}+$	18	0.51	U	120	37	29,000	22	3100
21	SB-SO-35-XX	Reference Laboratory	6	$\mathbf{J}+$	17	0.51	U	110	35	28,000	21	3100
21	SB-SO-44-XX	Reference Laboratory	6.8	$\mathbf{J}+$	18	0.51	U	120	37	29,000	22	3000
21	SB-SO-22-XC	Xcalibur XRF Services	10		124			51	16	18,480	0	1,057
21	SB-SO-25-XC	Xcalibur XRF Services	14		109	0		20	3	19,155	0	1,107
21	SB-SO-27-XC	Xcalibur XRF Services	15		112				20	19,141	0	1,128
21	SB-SO-35-XC	Xcalibur XRF Services	14		119			53	22	19,400	0	1,233
21	SB-SO-44-XC	Xcalibur XRF Services	20		116			61	25	19,543	0	1,283
22	SB-SO-23-XX	Reference Laboratory	48	J-	37	0.1	U	21	7	4,500	36	8500
22	SB-SO-28-XX	Reference Laboratory	42	J-	36	0.1	U	21	7	4,400	36	8800
22	SB-SO-32-XX	Reference Laboratory	46	J-	40	0.1	U	23	7.6	4,900	40	8900
22	SB-SO-43-XX	Reference Laboratory	40	J-	35	0.1	U	20	6.7	4,200	34	7600
22	SB-SO-48-XX	Reference Laboratory	39	J-	36	0.1	U	21	6.9	4,500	36	8200
22	SB-SO-23-XC	Xcalibur XRF Services	35		317	0		36	1	2,818	0	1,347
22	SB-SO-28-XC	Xcalibur XRF Services	39		347	12		25	0	2,961	0	1,494
22	SB-SO-32-XC	Xcalibur XRF Services	38		349	0			0	2,881	0	1,505
22	SB-SO-43-XC	Xcalibur XRF Services	34		425	40			1	2,850	0	1,494
22	SB-SO-48-XC	Xcalibur XRF Services	39		367	0			0	2,881	0	1,496
23	SB-SO-02-XX	Reference Laboratory	44	J-	23 J-	0.5	U	130	43	35,000	22 J-	130 J+
23	SB-SO-07-XX	Reference Laboratory	45	J	22	0.5	U	120	38	35,000	23	270
23	SB-SO-10-XX	Reference Laboratory	62	J	26	0.5	U	140	44	41,000	27	220
23	SB-SO-26-XX	Reference Laboratory	61	J	30	0.5	U	160	50	46,000	31	260
23	SB-SO-50-XX	Reference Laboratory	57	J	27	0.5	U	140	46	42,000	28	200
23	SB-SO-02-XC	Xcalibur XRF Services	49		99	35		115	30	28,994	49	74
23	SB-SO-07-XC	Xcalibur XRF Services	58		72			118	26	29,271	67	87
23	SB-SO-10-XC	Xcalibur XRF Services	53		52			122	38	29,497	74	69
23	SB-SO-26-XC	Xcalibur XRF Services	53		46	38		101	40	27,573	83	62
23	SB-SO-50-XC	Xcalibur XRF Services	50		74	24		134	39	28,740	0	74

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
20	SB-SO-08-XX	Reference Laboratory	180	1.3 U	1.3 U	57	70
20	SB-SO-11-XX	Reference Laboratory	200	1.3 U	1.3 U	66	84
20	SB-SO-21-XX	Reference Laboratory	190	1.3 U	1.3 U	58	75
20	SB-SO-39-XX	Reference Laboratory	200	1.3 U	1.3 U	62	77
20	SB-SO-42-XX	Reference Laboratory	200	1.3 U	1.3 U	65	78
20	SB-SO-08-XC	Xcalibur XRF Services	154			29	74
20	SB-SO-11-XC	Xcalibur XRF Services	134		1	96	72
20	SB-SO-21-XC	Xcalibur XRF Services	120			28	60
20	SB-SO-39-XC	Xcalibur XRF Services	135		18	53	77
20	SB-SO-42-XC	Xcalibur XRF Services	135			63	98
21	SB-SO-22-XX	Reference Laboratory	160	1.3 U	1.3 U	52	64 J-
21	SB-SO-25-XX	Reference Laboratory	160	1.3 U	1.3 U	54	63
21	SB-SO-27-XX	Reference Laboratory	170	1.3 U	1.3 U	54	65
21	SB-SO-35-XX	Reference Laboratory	160	1.3 U	1.3 U	50	62
21	SB-SO-44-XX	Reference Laboratory	170	1.3 U	1.3 U	53	64
21	SB-SO-22-XC	Xcalibur XRF Services	114			22	58
21	SB-SO-25-XC	Xcalibur XRF Services	119			121	67
21	SB-SO-27-XC	Xcalibur XRF Services	117	52		69	57
21	SB-SO-35-XC	Xcalibur XRF Services	121	25	12	96	60
21	SB-SO-44-XC	Xcalibur XRF Services	113		29	68	67
22	SB-SO-23-XX	Reference Laboratory	26	0.22 J	0.26 UJ	13	8
22	SB-SO-28-XX	Reference Laboratory	26	0.26 U	0.26 UJ	13	8
22	SB-SO-32-XX	Reference Laboratory	28	0.36	0.1 UJ	14	9
22	SB-SO-43-XX	Reference Laboratory	24	0.26 U	0.26 UJ	13	8
22	SB-SO-48-XX	Reference Laboratory	25	0.26 U	0.1 UJ	13	8
22	SB-SO-23-XC	Xcalibur XRF Services	6		7	147	18
22	SB-SO-28-XC	Xcalibur XRF Services	6		5	93	0
22	SB-SO-32-XC	Xcalibur XRF Services	6		15	156	0
22	SB-SO-43-XC	Xcalibur XRF Services	24		5	162	13
22	SB-SO-48-XC	Xcalibur XRF Services	6			135	0
23	SB-SO-02-XX	Reference Laboratory	180	1.2 U	1.2 UJ	59	88
23	SB-SO-07-XX	Reference Laboratory	170	1.4	1.6	53	86
23	SB-SO-10-XX	Reference Laboratory	200	2.8	1.8	59	100
23	SB-SO-26-XX	Reference Laboratory	220	3.4	1.8	68	110
23	SB-SO-50-XX	Reference Laboratory	200	2.9	1.8	61	100
23	SB-SO-02-XC	Xcalibur XRF Services	162		12	0	113
23	SB-SO-07-XC	Xcalibur XRF Services	160		18	0	114
23	SB-SO-10-XC	Xcalibur XRF Services	166	12	18	0	117
23	SB-SO-26-XC	Xcalibur XRF Services	173			43	103
23	SB-SO-50-XC	Xcalibur XRF Services	190		15	38	114

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	St	)	As		Cd		Cr	Cu		Fe	Pb	)	Hg	5
24	SB-SO-01-XX	Reference Laboratory	180	J	65		0.5	U	140	46		47,000	30		400	
24	SB-SO-16-XX	Reference Laboratory	170	J	64		0.5	U	140	45		47,000	30		480	
24	SB-SO-24-XX	Reference Laboratory	180	J	66		0.5	U	150	49		49,000	32		420	
24	SB-SO-45-XX	Reference Laboratory	180	J	63		0.5	U	140	45		47,000	30		450	
24	SB-SO-52-XX	Reference Laboratory	150	J	62		0.5	U	140	47		46,000	29		430	
24	SB-SO-01-XC	Xcalibur XRF Services	123		83				132	44		32,072	67		158	
24	SB-SO-16-XC	Xcalibur XRF Services	129		108		33		115	30		30,888	0		141	
24	SB-SO-24-XC	Xcalibur XRF Services	105		103				76	25		28,359	0		111	
24	SB-SO-45-XC	Xcalibur XRF Services	122		88		25		161	37		30,355	83		168	
24	SB-SO-52-XC	Xcalibur XRF Services	131		94				110	38		29,892	76		153	
25	SB-SO-13-XX	Reference Laboratory	430	J	160		1	U	140	46		61,000	36		850	
25	SB-SO-19-XX	Reference Laboratory	310	J	100		0.5	U	100	32		42,000	25		740	
25	SB-SO-33-XX	Reference Laboratory	350	J	110		0.5	U	100	33		45,000	28		870	
25	SB-SO-37-XX	Reference Laboratory	340	J	130		1	U	120	39		51,000	31		790	
25	SB-SO-55-XX	Reference Laboratory	340	J	120		0.5	U	120	37		49,000	29		900	
25	SB-SO-13-XC	Xcalibur XRF Services	280		194		28		92	32		35,473	58		339	
25	SB-SO-19-XC	Xcalibur XRF Services	210		174		24			22		31,909	0		220	
25	SB-SO-33-XC	Xcalibur XRF Services	259		195		30		92	19		34,068	0		293	
25	SB-SO-37-XC	Xcalibur XRF Services	258		172				170	35		36,020	76		310	
25	SB-SO-55-XC	Xcalibur XRF Services	278		195				99	35		36,018	105		338	
26	SB-SO-12-XX	Reference Laboratory	620	J	190		1	U	100	33		55,000	43		1,400	
26	SB-SO-15-XX	Reference Laboratory	600	J-	170 J	J-	1	U	91 J-	30	J-	51,000 J-	40	J-	1,100	
26	SB-SO-17-XX	Reference Laboratory	800	$\mathbf{J}+$	210		1	U	110	37		61,000	48		1,200	
26	SB-SO-46-XX	Reference Laboratory	740	$\mathbf{J}+$	190		1	U	120	35		57,000	47		670	
26	SB-SO-54-XX	Reference Laboratory	280		31		0.2	U	25	5.8		8,600	5	J-	560	
26	SB-SO-12-XC	Xcalibur XRF Services	428		266		27		132	20		39,828	91		570	
26	SB-SO-15-XC	Xcalibur XRF Services	419		298				133	28		39,562	0		524	
26	SB-SO-17-XC	Xcalibur XRF Services	257		215				32	22		33,967	0		337	
26	SB-SO-46-XC	Xcalibur XRF Services	417		259		29		133	24		39,119	80		504	
26	SB-SO-54-XC	Xcalibur XRF Services	425		260		28		204	28		39,886	84		495	
27	KP-SE-08-XX	Reference Laboratory	6.2		3		0.11	U	88	3.8		840	300	J-	0.089	U
27	KP-SE-11-XX	Reference Laboratory	5.6		3		0.11	U	96	4.1		940	310	J-	0.079	U
27	KP-SE-17-XX	Reference Laboratory	4.9		3		0.11	U	98	4.1		940	300	J-	0.082	U
27	KP-SE-25-XX	Reference Laboratory	6		3		0.11	U	99	4.3		960	310	J-	0.096	U
27	KP-SE-30-XX	Reference Laboratory	5.7		3		0.11	U	83	3.6		830	300	J-	0.1	U
27	KP-SE-08-XC	Xcalibur XRF Services	13		126				96	0		966	442			
27	KP-SE-11-XC	Xcalibur XRF Services	16		82				94	1		940	450			
27	KP-SE-17-XC	Xcalibur XRF Services	17		76		32		92	1		1,036	458		3	
27	KP-SE-25-XC	Xcalibur XRF Services	12		0				113	1		1,039	512		3	
27	KP-SE-30-XC	Xcalibur XRF Services	14		40		28		93	1		1,048	491			

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
24	SB-SO-01-XX	Reference Laboratory	190	1.8	2.3	65	95
24	SB-SO-16-XX	Reference Laboratory	190	1.9	2.2	65	97
24	SB-SO-24-XX	Reference Laboratory	200	2.5	2.3	67	95
24	SB-SO-45-XX	Reference Laboratory	190	2.8	2.1 J-	63	93
24	SB-SO-52-XX	Reference Laboratory	190	1.8	2.2	64	90
24	SB-SO-01-XC	Xcalibur XRF Services	170		16	57	96
24	SB-SO-16-XC	Xcalibur XRF Services	139		6	24	84
24	SB-SO-24-XC	Xcalibur XRF Services	130		22	0	67
24	SB-SO-45-XC	Xcalibur XRF Services	149		4	59	91
24	SB-SO-52-XC	Xcalibur XRF Services	151		20	28	96
25	SB-SO-13-XX	Reference Laboratory	180	4.4	2.2 UJ	74	70
25	SB-SO-19-XX	Reference Laboratory	120	2.5	1.8	51	51
25	SB-SO-33-XX	Reference Laboratory	130	3	2 J	52	56
25	SB-SO-37-XX	Reference Laboratory	150	2.5 U	2 UJ	63	58
25	SB-SO-55-XX	Reference Laboratory	140	2.5	2.2 J	61	60
25	SB-SO-13-XC	Xcalibur XRF Services	125		24	64	56
25	SB-SO-19-XC	Xcalibur XRF Services	121		27	0	57
25	SB-SO-33-XC	Xcalibur XRF Services	100		13	148	57
25	SB-SO-37-XC	Xcalibur XRF Services	132		29	179	68
25	SB-SO-55-XC	Xcalibur XRF Services	128		37	48	57
26	SB-SO-12-XX	Reference Laboratory	110	2.5 U	2.1 UJ	59	42
26	SB-SO-15-XX	Reference Laboratory	100 J-	3.4	1.6 UJ	52 J-	36 J-
26	SB-SO-17-XX	Reference Laboratory	120	2.8	2.3 UJ	60	42
26	SB-SO-46-XX	Reference Laboratory	120	2.6	2.2 UJ	57	41
26	SB-SO-54-XX	Reference Laboratory	20	0.5 U	0.5 UJ	11	6
26	SB-SO-12-XC	Xcalibur XRF Services	122			0	41
26	SB-SO-15-XC	Xcalibur XRF Services	109		23	40	32
26	SB-SO-17-XC	Xcalibur XRF Services	56			0	35
26	SB-SO-46-XC	Xcalibur XRF Services	111		13	33	34
26	SB-SO-54-XC	Xcalibur XRF Services	110			9	37
27	KP-SE-08-XX	Reference Laboratory	42	0.27 U	0.27 UJ	4	5
27	KP-SE-11-XX	Reference Laboratory	46	0.43	0.27 UJ	4	6
27	KP-SE-17-XX	Reference Laboratory	47	0.27 U	0.27 UJ	4	5
27	KP-SE-25-XX	Reference Laboratory	47	0.26 U	0.27 UJ	4	5
27	KP-SE-30-XX	Reference Laboratory	39	0.24 U	0.27 UJ	4	5
27	KP-SE-08-XC	Xcalibur XRF Services	39			26	0
27	KP-SE-11-XC	Xcalibur XRF Services	36		12	0	35
27	KP-SE-17-XC	Xcalibur XRF Services	42	2	5	0	25
27	KP-SE-25-XC	Xcalibur XRF Services	46	41	2	0	25
27	KP-SE-30-XC	Xcalibur XRF Services	47	41		32	18

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg	
28	KP-SE-01-XX	Reference Laboratory	3.2	2	0.1 U	J 34	2.2	480	310 J-	0.053	U
28	KP-SE-12-XX	Reference Laboratory	3.1	2	0.1 U	J 42	2.5	510	320 J-	0.06	U
28	KP-SE-14-XX	Reference Laboratory	11 J-	2	0.1 U	J 46 J-	2.7 J+	520 J-	680 J-	0.065	U
28	KP-SE-19-XX	Reference Laboratory	3	2	0.1 U	J 44	2.3	510	330	0.044	U
28	KP-SE-28-XX	Reference Laboratory	3.3	2	0.1 U	J 45	2.3	520	320	0.056	U
28	KP-SE-01-XC	Xcalibur XRF Services		70	11	72	8	651	405	3	
28	KP-SE-12-XC	Xcalibur XRF Services	4	27	26	59	1	509	343	2	
28	KP-SE-14-XC	Xcalibur XRF Services	5	107		61	1	635	423		
28	KP-SE-19-XC	Xcalibur XRF Services	13	81		45	1	640	452	2	
28	KP-SE-28-XC	Xcalibur XRF Services	5	36		70	1	646	453		
29	TL-SE-04-XX	Reference Laboratory	1.2 U	10	0.5 U	J 62	1,900	42,000	32	0.26	J-
29	TL-SE-10-XX	Reference Laboratory	1.2 U	10	0.5 U	J 64	2,000	43,000	35	0.19	J-
29	TL-SE-12-XX	Reference Laboratory	1.2 U	10	0.5 U	J 66	2,100	44,000	34	0.22	J-
29	TL-SE-15-XX	Reference Laboratory	1.2 U	9	0.5 U	J 54	1,800	36,000	28	0.28	J-
29	TL-SE-20-XX	Reference Laboratory	1.2 U	10	0.5 U	J 64	2,000	42,000	32	0.26	J-
29	TL-SE-24-XX	Reference Laboratory	1.2 U	11	0.5 U	J 67	2,100	43,000	37	0.26	J-
29	TL-SE-26-XX	Reference Laboratory	1.2 U	10	0.5 U	J 62	2,000	40,000	34	0.24	J-
29	TL-SE-04-XC	Xcalibur XRF Services	0	58			2,586	36,085	98		
29	TL-SE-10-XC	Xcalibur XRF Services	5	0		7	2,606	36,600	144		
29	TL-SE-12-XC	Xcalibur XRF Services	7	53			2,626	36,634	133		
29	TL-SE-15-XC	Xcalibur XRF Services	5	30		36	2,461	35,872	122		
29	TL-SE-20-XC	Xcalibur XRF Services	10	45	12		2,718	37,221	119		
29	TL-SE-24-XC	Xcalibur XRF Services	9	78			2,911	37,288	135		
29	TL-SE-26-XC	Xcalibur XRF Services	16	42		26	2,631	36,523	195	6	
30	TL-SE-03-XX	Reference Laboratory	2.5 U	9	1 U	J 91	1,600	63,000	12	0.32	J-
30	TL-SE-19-XX	Reference Laboratory	2.5 U	10	1 U	J 96	1,700	66,000	13	0.32	J-
30	TL-SE-23-XX	Reference Laboratory	2.5 U	9	1 U	J 92	1,600	64,000	12	0.41	J-
30	TL-SE-25-XX	Reference Laboratory	2.5 U	10	1 U	J 91	1,600	62,000	11	0.44	J-
30	TL-SE-31-XX	Reference Laboratory	2.5 U	10	1 U	110	1,800	74,000	13	0.57	J-
30	TL-SE-03-XC	Xcalibur XRF Services	11	67	12		1,879	52,922	139		
30	TL-SE-19-XC	Xcalibur XRF Services	6	32			2,405	56,855	60		
30	TL-SE-23-XC	Xcalibur XRF Services	10	74			2,272	57,909	106		
30	TL-SE-25-XC	Xcalibur XRF Services	11	27			2,281	58,350	115		
30	TL-SE-31-XC	Xcalibur XRF Services	9	79	16		2,096	55,112	160		

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	A	g	V	Zn
28	KP-SE-01-XX	Reference Laboratory	16	0.26 U	0.26	UJ	2 J	6
28	KP-SE-12-XX	Reference Laboratory	20	0.26 L	0.26	UJ	2 J	8
28	KP-SE-14-XX	Reference Laboratory	23 J-	0.26 L	0.26	UJ	3 J	7
28	KP-SE-19-XX	Reference Laboratory	22	0.26 L	0.26	U	2 J	7
28	KP-SE-28-XX	Reference Laboratory	22	0.26 L	0.26	U	2 J	6
28	KP-SE-01-XC	Xcalibur XRF Services	27	11			0	33
28	KP-SE-12-XC	Xcalibur XRF Services	20	47			0	25
28	KP-SE-14-XC	Xcalibur XRF Services	24	27	3		0	30
28	KP-SE-19-XC	Xcalibur XRF Services	23				21	38
28	KP-SE-28-XC	Xcalibur XRF Services	18		3		0	35
29	TL-SE-04-XX	Reference Laboratory	71	1.2 U	1.3		95	160
29	TL-SE-10-XX	Reference Laboratory	72	1.2 U	1.2	U	95	160
29	TL-SE-12-XX	Reference Laboratory	75	1.2 U	1.2	U	100	170
29	TL-SE-15-XX	Reference Laboratory	63	1.2 U	1	U	84	140
29	TL-SE-20-XX	Reference Laboratory	74	1.2 U	1.2	U	100	160
29	TL-SE-24-XX	Reference Laboratory	77	1.2 U	1.3	U	100	170
29	TL-SE-26-XX	Reference Laboratory	70	1.2 U	1.2	U	96	160
29	TL-SE-04-XC	Xcalibur XRF Services	116	5	13		0	183
29	TL-SE-10-XC	Xcalibur XRF Services	115				0	180
29	TL-SE-12-XC	Xcalibur XRF Services	117		15		0	165
29	TL-SE-15-XC	Xcalibur XRF Services	123		25		0	179
29	TL-SE-20-XC	Xcalibur XRF Services	89				0	174
29	TL-SE-24-XC	Xcalibur XRF Services	107		17		0	168
29	TL-SE-26-XC	Xcalibur XRF Services	100				54	192
30	TL-SE-03-XX	Reference Laboratory	110	2.5 U	0.94	U	140	200
30	TL-SE-19-XX	Reference Laboratory	120	2.5 U	1.1	U	150	210
30	TL-SE-23-XX	Reference Laboratory	110	2.5 U	1.3	U	150	200
30	TL-SE-25-XX	Reference Laboratory	110	2.5 U	0.94	U	150	200
30	TL-SE-31-XX	Reference Laboratory	130	2.5 U	1.2	U	170	230
30	TL-SE-03-XC	Xcalibur XRF Services	154		6		0	113
30	TL-SE-19-XC	Xcalibur XRF Services	137		41		0	98
30	TL-SE-23-XC	Xcalibur XRF Services	121		23		0	95
30	TL-SE-25-XC	Xcalibur XRF Services	115	3	54		0	93
30	TL-SE-31-XC	Xcalibur XRF Services	115		15		66	97

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	S	b	As	5	Cd		Cr		Cu	Fe		Pb		Hg	
31	TL-SE-01-XX	Reference Laboratory	1.2	UJ	9		0.5	U	110		1,400	19,000		48	J-	0.074	U
31	TL-SE-11-XX	Reference Laboratory	1.2	UJ	15		0.5	U	140		1,600	28,000		54	J-	0.021	U
31	TL-SE-14-XX	Reference Laboratory	1.2	UJ	10		0.27	J	110		1,500	18,000		50	J-	0.08	U
31	TL-SE-18-XX	Reference Laboratory	1.2	UJ	10		0.5	U	150		1,300	24,000		46	J-	0.025	U
31	TL-SE-22-XX	Reference Laboratory	1.2	UJ	11		0.5	U	150		1,700	26,000		54	J-	0.082	U
31	TL-SE-27-XX	Reference Laboratory	1.2	UJ	10		0.28	J	130		1,500	19,000		51	J-	0.02	U
31	TL-SE-29-XX	Reference Laboratory	1.2	UJ	11		0.22	J	140		1,600	23,000		51	J-	0.076	U
31	TL-SE-01-XC	Xcalibur XRF Services	0		68		19		124		1,462	26,263		154		5	
31	TL-SE-11-XC	Xcalibur XRF Services	7		86				102		1,883	34,010		145		5	
31	TL-SE-14-XC	Xcalibur XRF Services	3		58				87		1,900	33,304		141		3	
31	TL-SE-18-XC	Xcalibur XRF Services	4		51				98		1,876	32,929		193			
31	TL-SE-22-XC	Xcalibur XRF Services	9		37				149		1,865	33,004		181		4	
31	TL-SE-27-XC	Xcalibur XRF Services	12		16		13		141		1,932	32,418		187		18	
31	TL-SE-29-XC	Xcalibur XRF Services	10		45		1		93		1,809	31,056		119		1	
32	LV-SE-02-XX	Reference Laboratory	1.3	UJ	28		0.51	U	72		33	23,000		20	J-	0.02	U
32	LV-SE-10-XX	Reference Laboratory	1.3	UJ	34		0.51	U	84		42	28,000		25	J-	0.023	U
32	LV-SE-22-XX	Reference Laboratory	1.3	UJ	30		0.51	U	69		33	23,000		22	J-	1.1	
32	LV-SE-25-XX	Reference Laboratory	1.3	UJ	31		0.51	U	74		36	25,000		23	J-	1	
32	LV-SE-31-XX	Reference Laboratory	1.3	UJ	32		0.51	U	78		36	25,000		49	J-	1	
32	LV-SE-35-XX	Reference Laboratory	1.3	UJ	31	J-	0.51	U	74	J-	35	24,000	J-	22	J-	1.4	
32	LV-SE-50-XX	Reference Laboratory	2.5	U	29		1	U	74		34	24,000		24	J-	1.2	
32	LV-SE-02-XC	Xcalibur XRF Services	5		74		33		39		6	19,902		80		13	
32	LV-SE-10-XC	Xcalibur XRF Services	9		82		35				30	20,112		0			
32	LV-SE-22-XC	Xcalibur XRF Services	10		81				46		25	19,644		59		7	
32	LV-SE-25-XC	Xcalibur XRF Services	5		85		11		44		23	20,308		125		6	
32	LV-SE-31-XC	Xcalibur XRF Services	0		87		11				21	19,882		57			
32	LV-SE-35-XC	Xcalibur XRF Services	0		73				49		20	20,470		96			
32	LV-SE-50-XC	Xcalibur XRF Services	0		79		25				5	19,573		69		19	
33	LV-SE-12-XX	Reference Laboratory	2.6	U	190		1	U	55		34	72,000		19	J-	5.6	
33	LV-SE-26-XX	Reference Laboratory	2.6	U	220		1	U	64		39	83,000		25	J-	6	
33	LV-SE-33-XX	Reference Laboratory	2.6	U	170		1	U	52		31	66,000		21	J-	6.8	
33	LV-SE-39-XX	Reference Laboratory	2.6	U	190		1	U	58		35	74,000		22	J-	8	
33	LV-SE-42-XX	Reference Laboratory	2.7	U	170		1.1	U	50		30	65,000		22	J-	4.3	
33	LV-SE-12-XC	Xcalibur XRF Services			230				98		32	51,313		155			
33	LV-SE-26-XC	Xcalibur XRF Services	8		243				133		32	53,305		120			
33	LV-SE-33-XC	Xcalibur XRF Services	7		227						35	53,895		183			
33	LV-SE-39-XC	Xcalibur XRF Services	10		291						43	54,680		0			
33	LV-SE-42-XC	Xcalibur XRF Services	11		261				81		26	54,288		104			

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
31	TL-SE-01-XX	Reference Laboratory	180	1.2 U	5.7 J-	75	130
31	TL-SE-11-XX	Reference Laboratory	210	1.2 U	5.5 J-	85	140
31	TL-SE-14-XX	Reference Laboratory	180	1.2 U	5.7 J-	73	140
31	TL-SE-18-XX	Reference Laboratory	190	1.2 U	6.3 J-	70	120
31	TL-SE-22-XX	Reference Laboratory	210	1.2 U	6.5 J-	80	150
31	TL-SE-27-XX	Reference Laboratory	200	1.2 U	7.8 J-	67	140
31	TL-SE-29-XX	Reference Laboratory	200	1.2 U	5.9 J-	80	140
31	TL-SE-01-XC	Xcalibur XRF Services	156			0	97
31	TL-SE-11-XC	Xcalibur XRF Services	220		33	0	104
31	TL-SE-14-XC	Xcalibur XRF Services	198		21	64	107
31	TL-SE-18-XC	Xcalibur XRF Services	199		24	54	93
31	TL-SE-22-XC	Xcalibur XRF Services	206		26	0	107
31	TL-SE-27-XC	Xcalibur XRF Services	232	64	23	65	109
31	TL-SE-29-XC	Xcalibur XRF Services	209	37	26	44	104
32	LV-SE-02-XX	Reference Laboratory	160	3.8	1.3 UJ	53	65
32	LV-SE-10-XX	Reference Laboratory	200	4.7	1.3 UJ	66	77
32	LV-SE-22-XX	Reference Laboratory	170	5.2	1.3 UJ	51	66
32	LV-SE-25-XX	Reference Laboratory	170	5.1	1.3 UJ	56	70
32	LV-SE-31-XX	Reference Laboratory	180	5.1	1.3 UJ	58	70
32	LV-SE-35-XX	Reference Laboratory	170 J-	5	1.3 UJ	55 J-	67 J-
32	LV-SE-50-XX	Reference Laboratory	170	3.3	2.5 U	57	65
32	LV-SE-02-XC	Xcalibur XRF Services	115		11	55	115
32	LV-SE-10-XC	Xcalibur XRF Services	119		6	73	97
32	LV-SE-22-XC	Xcalibur XRF Services	106		7	23	127
32	LV-SE-25-XC	Xcalibur XRF Services	141		10	0	124
32	LV-SE-31-XC	Xcalibur XRF Services	111		8	0	113
32	LV-SE-35-XC	Xcalibur XRF Services	116		2	45	91
32	LV-SE-50-XC	Xcalibur XRF Services	113		21	82	100
33	LV-SE-12-XX	Reference Laboratory	71	3	2.6 U	72	66
33	LV-SE-26-XX	Reference Laboratory	83	6.1	2.6 U	86	75
33	LV-SE-33-XX	Reference Laboratory	66	2.8	2.6 U	67	59
33	LV-SE-39-XX	Reference Laboratory	74	5.1	2.6 U	74	66
33	LV-SE-42-XX	Reference Laboratory	67	3.4	2.7 U	64	57
33	LV-SE-12-XC	Xcalibur XRF Services	71			25	48
33	LV-SE-26-XC	Xcalibur XRF Services	72		3	27	37
33	LV-SE-33-XC	Xcalibur XRF Services	72		4	0	40
33	LV-SE-39-XC	Xcalibur XRF Services	102	22	5	0	31
33	LV-SE-42-XC	Xcalibur XRF Services	99		2	0	41

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	)	As	Cd		Cr	Cu	Fe	Pb	Hg	
34	LV-SE-09-XX	Reference Laboratory	6.7	U	450	2.7	U	48	34	150,000	14 J-	6	
34	LV-SE-19-XX	Reference Laboratory	6.7	U	500	2.7	U	55	37	160,000	17 J-	7.2	
34	LV-SE-27-XX	Reference Laboratory	6.7	U	530	2.7	U	56	39	180,000	16 J-	11	
34	LV-SE-36-XX	Reference Laboratory	6.7	U	550	2.7	U	60	40	180,000	21 J-	8.5	
34	LV-SE-38-XX	Reference Laboratory	6.7	U	480	2.7	U	52	36	160,000	15 J-	7.9	
34	LV-SE-09-XC	Xcalibur XRF Services	0		513			270	40	105,797	0		
34	LV-SE-19-XC	Xcalibur XRF Services	11		517	24		384	2	104,569	166		
34	LV-SE-27-XC	Xcalibur XRF Services	10		488	18		86	3	103,211	257		
34	LV-SE-36-XC	Xcalibur XRF Services	15		545				36	107,069	0		
34	LV-SE-38-XC	Xcalibur XRF Services	0		503	0		92	4	105,744	0		
35	LV-SE-07-XX	Reference Laboratory	6.7	UJ	780	2.7	U	57	48	200,000	11	5.5	
35	LV-SE-18-XX	Reference Laboratory	6.7	UJ	800	2.7	U	61	49	210,000	11	5.4	
35	LV-SE-23-XX	Reference Laboratory	6.6	UJ	660	2.6	U	53	40	170,000	8	5	
35	LV-SE-45-XX	Reference Laboratory	6.7	UJ	650	2.7	U	50	40	170,000	8	5.6	
35	LV-SE-48-XX	Reference Laboratory	6.6	UJ	680	2.6	U	52	42	180,000	9	7.3	
35	LV-SE-07-XC	Xcalibur XRF Services	12		692	12		314	47	132,559	270		
35	LV-SE-18-XC	Xcalibur XRF Services	11		670			702	2	133,997	356		
35	LV-SE-23-XC	Xcalibur XRF Services	22		705	9		346	28	129,948	192		
35	LV-SE-45-XC	Xcalibur XRF Services	35		642				54	120,254	0		
35	LV-SE-48-XC	Xcalibur XRF Services	28		585				1	124,294	445	109	
36	LV-SE-01-XX	Reference Laboratory	1.5	UJ	6	0.76		4	18	1,100	17	0.098	U
36	LV-SE-14-XX	Reference Laboratory	1.5	UJ	5	0.74		4	16	980	14	0.056	U
36	LV-SE-21-XX	Reference Laboratory	1.5	UJ	7	0.84		4	19	970	18	0.048	U
36	LV-SE-24-XX	Reference Laboratory	1.5	UJ	5	0.68		4	15	840	14	0.053	U
36	LV-SE-32-XX	Reference Laboratory	1.4	UJ	6	0.87		4	16	860	14	0.052	U
36	LV-SE-01-XC	Xcalibur XRF Services	118		91	28			1	416	43	3	
36	LV-SE-14-XC	Xcalibur XRF Services	80		70				14	446	22		
36	LV-SE-21-XC	Xcalibur XRF Services	98		72				0	455	22		
36	LV-SE-24-XC	Xcalibur XRF Services	98		66				0	437	27	2	
36	LV-SE-32-XC	Xcalibur XRF Services	107		73				1	433	32		
37	LV-SE-08-XX	Reference Laboratory	1.3	UJ	30	0.52	U	54	23	23,000	55	5.2	
37	LV-SE-16-XX	Reference Laboratory	1.3	UJ	29	0.52	U	53	22	22,000	53	5.4	
37	LV-SE-28-XX	Reference Laboratory	1.3	UJ	31	0.52	U	59	25	25,000	59	5.4	
37	LV-SE-30-XX	Reference Laboratory	1.3	UJ	30	0.52	U	58	25	24,000	58	6.3	
37	LV-SE-47-XX	Reference Laboratory	1.3	UJ	31	0.52	U	56	23	23,000	57	4.9	
37	LV-SE-08-XC	Xcalibur XRF Services	0		69	36			6	20,743	137	8	
37	LV-SE-16-XC	Xcalibur XRF Services	1		83	52			5	20,064	177	12	
37	LV-SE-28-XC	Xcalibur XRF Services	1		71			40	5	20,981	165		
37	LV-SE-30-XC	Xcalibur XRF Services	6		57	34		58	5	20,597	193	7	
37	LV-SE-47-XC	Xcalibur XRF Services	0		93				23	19,257	145		

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
34	LV-SE-09-XX	Reference Laboratory	55	6.7 U	6.7 U	100	51 J
34	LV-SE-19-XX	Reference Laboratory	65	5.9 J	6.7 U	110	55 J
34	LV-SE-27-XX	Reference Laboratory	64	6.7 U	6.7 U	120	58 J
34	LV-SE-36-XX	Reference Laboratory	70	11	6.7 U	120	60 J
34	LV-SE-38-XX	Reference Laboratory	75	6.7 U	6.7 U	100	54 J
34	LV-SE-09-XC	Xcalibur XRF Services	74		13	0	
34	LV-SE-19-XC	Xcalibur XRF Services	74			0	
34	LV-SE-27-XC	Xcalibur XRF Services	75	111	14	0	
34	LV-SE-36-XC	Xcalibur XRF Services	73		10	0	
34	LV-SE-38-XC	Xcalibur XRF Services	74	10		0	
35	LV-SE-07-XX	Reference Laboratory	58	10	6.7 U	130	24 J
35	LV-SE-18-XX	Reference Laboratory	60	12	6.7 U	140	52 J
35	LV-SE-23-XX	Reference Laboratory	50 J	9.6	6.6 U	120	18 J
35	LV-SE-45-XX	Reference Laboratory	50 J	8.2	6.7 U	120	19 J
35	LV-SE-48-XX	Reference Laboratory	50 J	7.6	6.6 U	120	30 J
35	LV-SE-07-XC	Xcalibur XRF Services	55			0	
35	LV-SE-18-XC	Xcalibur XRF Services	54			0	
35	LV-SE-23-XC	Xcalibur XRF Services	57	5		0	
35	LV-SE-45-XC	Xcalibur XRF Services	65	9		15	0
35	LV-SE-48-XC	Xcalibur XRF Services	111			0	
36	LV-SE-01-XX	Reference Laboratory	49	1.5 U	1.5 U	2 J	14 J
36	LV-SE-14-XX	Reference Laboratory	46	1.5 U	1.5 U	1 J	12 J
36	LV-SE-21-XX	Reference Laboratory	49	1.5 U	1.5 U	2 J	14 J
36	LV-SE-24-XX	Reference Laboratory	44	1.5 U	1.5 U	1 J	12 J
36	LV-SE-32-XX	Reference Laboratory	47	1.4 U	1.4 U	1 J	19
36	LV-SE-01-XC	Xcalibur XRF Services	21		9	0	22
36	LV-SE-14-XC	Xcalibur XRF Services	25			0	53
36	LV-SE-21-XC	Xcalibur XRF Services	17			0	0
36	LV-SE-24-XC	Xcalibur XRF Services	31		4	0	0
36	LV-SE-32-XC	Xcalibur XRF Services	15		19	0	29
37	LV-SE-08-XX	Reference Laboratory	110	4.8	1.3 U	44	61
37	LV-SE-16-XX	Reference Laboratory	110	5	1.3 U	42	59
37	LV-SE-28-XX	Reference Laboratory	120	5.8	1.3 U	48	65
37	LV-SE-30-XX	Reference Laboratory	120	5.6	1.3 U	48	66
37	LV-SE-47-XX	Reference Laboratory	120	4.2	1.3 U	45	65
37	LV-SE-08-XC	Xcalibur XRF Services	122	6	4	46	116
37	LV-SE-16-XC	Xcalibur XRF Services	84		0	32	103
37	LV-SE-28-XC	Xcalibur XRF Services	102		9	16	100
37	LV-SE-30-XC	Xcalibur XRF Services	108		15	36	92
37	LV-SE-47-XC	Xcalibur XRF Services	96		2	32	107

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	1	As	Cd		Cr		Cu	Fe	Pb	Hg	Ni
38	LV-SE-11-XX	Reference Laboratory	1.4	UJ	150	6.6		120		270	42,000	7	2.8	870
38	LV-SE-29-XX	Reference Laboratory	1.4	UJ	150	6.3		120		260	42,000	7 J+	1.5 J-	860
38	LV-SE-44-XX	Reference Laboratory	1.4	U	140	6.1		120		250	40,000	8	1.5	830
38	LV-SE-46-XX	Reference Laboratory	0.88	U	110	5		92		200	32,000	6	1.4	660
38	LV-SE-52-XX	Reference Laboratory	1.4	U	160	6.8		130		280	44,000	8	21	910
38	LV-SE-11-XC	Xcalibur XRF Services	17		218			57		212	30,708	0		608
38	LV-SE-29-XC	Xcalibur XRF Services	13		228					202	30,725	0		636
38	LV-SE-44-XC	Xcalibur XRF Services			195					176	26,884			508
38	LV-SE-46-XC	Xcalibur XRF Services	9		199			41		200	27,483	0		567
38	LV-SE-52-XC	Xcalibur XRF Services	11		167					186	29,383	96		551
39	RF-SE-07-XX	Reference Laboratory	1.3	U	12	0.5	U	92		81	17,000	24	0.091 U	180
39	RF-SE-12-XX	Reference Laboratory	1.2	U	14	0.5	U	100		110	20,000	25	0.099 U	210
39	RF-SE-23-XX	Reference Laboratory	0.25	U	0 U	0.1	U	0	U	0.2 U	4 J	0 U	2.4	2 U
39	RF-SE-36-XX	Reference Laboratory	1.2	U	12	0.5	U	91		82	17,000	22	0.081 U	180
39	RF-SE-42-XX	Reference Laboratory	1.3	UJ	14	0.56		110		95	19,000	28	0.084 U	210
39	RF-SE-45-XX	Reference Laboratory	1.3	UJ	15	0.52	U	110		100	21,000	33	0.084 U	220
39	RF-SE-53-XX	Reference Laboratory	1.3	UJ	14	0.57	U	110		95	19,000	28	0.084 U	210
39	RF-SE-07-XC	Xcalibur XRF Services	0		63	10		48		56	15,548	108		132
39	RF-SE-12-XC	Xcalibur XRF Services	0		72			70		52	16,025	96	7	135
39	RF-SE-23-XC	Xcalibur XRF Services	0		54			80		72	16,718	97	7	155
39	RF-SE-36-XC	Xcalibur XRF Services	0		63			88		67	15,862	56	4	124
39	RF-SE-42-XC	Xcalibur XRF Services	4		34			51		75	17,031	60	8	165
39	RF-SE-45-XC	Xcalibur XRF Services	9		64	11		118		59	16,455	119	4	141
39	RF-SE-53-XC	Xcalibur XRF Services	0		73	30		70		55	16,043		86	144
40	RF-SE-03-XX	Reference Laboratory	1.2	UJ	27	1.3		93		200	17,000	88	0.48	150
40	RF-SE-28-XX	Reference Laboratory	1.2	UJ	31	1.5		100		220	18,000	99	0.57	160
40	RF-SE-38-XX	Reference Laboratory	1.2	UJ	27	1.2		90		190	16,000	83	0.41	140
40	RF-SE-49-XX	Reference Laboratory	1.2	UJ	31	1.5		100		220	18,000	97	0.43	170
40	RF-SE-55-XX	Reference Laboratory	1.2	UJ	24	1.1		91		180	15,000	75	0.42	140
40	RF-SE-03-XC	Xcalibur XRF Services	1		80	51				210	15,187	188	9	117
40	RF-SE-28-XC	Xcalibur XRF Services	5		53			106		211	15,903	204	7	116
40	RF-SE-38-XC	Xcalibur XRF Services	10		65	12		80		228	16,972	197		123
40	RF-SE-49-XC	Xcalibur XRF Services	1		64			73		200	15,568	178		117
40	RF-SE-55-XC	Xcalibur XRF Services	1		99			41		209	16,258	219	8	118

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se		Ag		V		Zn	
38	LV-SE-11-XX	Reference Laboratory	870	1.3	U	1.4	U	35		200	
38	LV-SE-29-XX	Reference Laboratory	860	1.2	U	1.4	U	35		200	
38	LV-SE-44-XX	Reference Laboratory	830	1.4	U	1.4	U	34		190	
38	LV-SE-46-XX	Reference Laboratory	660	0.88	U	0.88	U	27		150	
38	LV-SE-52-XX	Reference Laboratory	910	1.4	U	1.4	U	38		210	
38	LV-SE-11-XC	Xcalibur XRF Services	608			7		0		191	
38	LV-SE-29-XC	Xcalibur XRF Services	636			0		0		202	
38	LV-SE-44-XC	Xcalibur XRF Services	508					0		202	
38	LV-SE-46-XC	Xcalibur XRF Services	567					0		189	
38	LV-SE-52-XC	Xcalibur XRF Services	551			0		61		187	
39	RF-SE-07-XX	Reference Laboratory	180	1.3	U	1.3	U	34		130	
39	RF-SE-12-XX	Reference Laboratory	210	1.2	U	1.2	U	38		140	
39	RF-SE-23-XX	Reference Laboratory	2 U	0.25	U	0.37		3	U	1	U
39	RF-SE-36-XX	Reference Laboratory	180	1	U	1.2	U	34		120	
39	RF-SE-42-XX	Reference Laboratory	210	1.3	U	1.3	U	40		140	
39	RF-SE-45-XX	Reference Laboratory	220	1.3	U	1.3	U	43		150	
39	RF-SE-53-XX	Reference Laboratory	210	1.3	U	1.3	U	40		140	
39	RF-SE-07-XC	Xcalibur XRF Services	132			25		0		171	
39	RF-SE-12-XC	Xcalibur XRF Services	135			4		35		190	
39	RF-SE-23-XC	Xcalibur XRF Services	155	6				49		219	
39	RF-SE-36-XC	Xcalibur XRF Services	124			4		0		227	
39	RF-SE-42-XC	Xcalibur XRF Services	165	14				33		248	
39	RF-SE-45-XC	Xcalibur XRF Services	141			14		0		217	
39	RF-SE-53-XC	Xcalibur XRF Services	144			3		26		185	
40	RF-SE-03-XX	Reference Laboratory	150	1.2	U	1.2	U	40		300	
40	RF-SE-28-XX	Reference Laboratory	160	1.2	U	1.2	U	44		320	
40	RF-SE-38-XX	Reference Laboratory	140	1.2	U	1.2	U	39		300	
40	RF-SE-49-XX	Reference Laboratory	170	1.2	U	1.2	U	43		330	
40	RF-SE-55-XX	Reference Laboratory	140	1.2	U	1.2	U	35		280	
40	RF-SE-03-XC	Xcalibur XRF Services	117			6		42		481	
40	RF-SE-28-XC	Xcalibur XRF Services	116					61		486	
40	RF-SE-38-XC	Xcalibur XRF Services	123			3		0		552	
40	RF-SE-49-XC	Xcalibur XRF Services	117			8		9		530	
40	RF-SE-55-XC	Xcalibur XRF Services	118					0		534	

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	S	b	A	s	С	d	C	r	Cu	l	Fe		P	b	Hg	5
41	RF-SE-06-XX	Reference Laboratory	1.3	UJ	70		3.6		90		490		20,000		230		1.1	
41	RF-SE-13-XX	Reference Laboratory	1.3	UJ	76		3.7		92		530		21,000		230		1.2	
41	RF-SE-27-XX	Reference Laboratory	1.3	UJ	64		3.1		78		440		18,000		200		1.2	
41	RF-SE-31-XX	Reference Laboratory	1.3	UJ	39		1.8		63		250		12,000		120		1.1	
41	RF-SE-58-XX	Reference Laboratory	1.3	UJ	71		3.6		89		500		21,000		230		1.2	
41	RF-SE-06-XC	Xcalibur XRF Services	2		96						468		16,459		356			
41	RF-SE-13-XC	Xcalibur XRF Services	7		92		22		78		485		16,918		452			
41	RF-SE-27-XC	Xcalibur XRF Services			92		2		97		485		17,224		381		5	
41	RF-SE-31-XC	Xcalibur XRF Services	2		123				59		523		17,807		392			
41	RF-SE-58-XC	Xcalibur XRF Services	6		89						520		17,583		501			
42	RF-SE-02-XX	Reference Laboratory	1.3	UJ	110		5.4		93		740		24,000		330		1.6	
42	RF-SE-22-XX	Reference Laboratory	1.3	UJ	99		4.7		84		670		22,000		300		1.7	
42	RF-SE-25-XX	Reference Laboratory	1.3	UJ	88		4		78		580		19,000		270		1.5	
42	RF-SE-30-XX	Reference Laboratory	1.3	UJ	89		4.3		78		610		21,000		290		1.5	
42	RF-SE-57-XX	Reference Laboratory	1.3	UJ	89		4.5		79		610		21,000		300		1.5	
42	RF-SE-02-XC	Xcalibur XRF Services	2		143		4		58		599		17,633		478			
42	RF-SE-22-XC	Xcalibur XRF Services	10		179				79		649		17,877		467		7	
42	RF-SE-25-XC	Xcalibur XRF Services	2		107		22		80		649		18,000		497			
42	RF-SE-30-XC	Xcalibur XRF Services	7		125		25		101		637		18,543		542			
42	RF-SE-57-XC	Xcalibur XRF Services	2		123						637		17,762		505			
43	RF-SE-15-XX	Reference Laboratory	1.3	UJ	120		6.2		72		820		23,000		390		2.6	
43	RF-SE-24-XX	Reference Laboratory	1.3	UJ	130	$\mathbf{J}+$	6.5	$\mathbf{J}+$	74	$\mathbf{J}+$	860	$\mathbf{J}+$	24,000	$\mathbf{J}+$	410	$\mathbf{J}+$	2.3	
43	RF-SE-32-XX	Reference Laboratory	1.3	UJ	120		5.1		64		770		20,000		330		2.8	
43	RF-SE-43-XX	Reference Laboratory	1.3	UJ	130		5.7		68		840		22,000		350		2.7	
43	RF-SE-59-XX	Reference Laboratory	1.3	UJ	140		5.9		73		890		23,000		380		0.085	U
43	RF-SE-15-XC	Xcalibur XRF Services	6		180				49		887		19,197		661		8	
43	RF-SE-24-XC	Xcalibur XRF Services	3		207				75		889		19,447		654			
43	RF-SE-32-XC	Xcalibur XRF Services	3		148		21				901		19,415		703			
43	RF-SE-43-XC	Xcalibur XRF Services	10		168		38				938		19,857		678			
43	RF-SE-59-XC	Xcalibur XRF Services	12		152				36		928		19,383		686			

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
41	RF-SE-06-XX	Reference Laboratory	150	1.3 U	1.3 U	44	740
41	RF-SE-13-XX	Reference Laboratory	160	1.3 U	1.3	45	790
41	RF-SE-27-XX	Reference Laboratory	130	1.3 U	1.3 U	39	670
41	RF-SE-31-XX	Reference Laboratory	86	1.3 U	1.3 U	28	420
41	RF-SE-58-XX	Reference Laboratory	150	1.3 U	1.3 U	46	770
41	RF-SE-06-XC	Xcalibur XRF Services	108		24	50	1,092
41	RF-SE-13-XC	Xcalibur XRF Services	122		22	0	1,173
41	RF-SE-27-XC	Xcalibur XRF Services	100	24		14	1,090
41	RF-SE-31-XC	Xcalibur XRF Services	115		5	33	1,156
41	RF-SE-58-XC	Xcalibur XRF Services	107		25	23	1,231
42	RF-SE-02-XX	Reference Laboratory	180	1.3 U	2.7	50	1,100
42	RF-SE-22-XX	Reference Laboratory	160	1.3 U	2.3	44	990
42	RF-SE-25-XX	Reference Laboratory	140	1.5	1.7	40	890
42	RF-SE-30-XX	Reference Laboratory	150	1.3 U	1.9	44	960
42	RF-SE-57-XX	Reference Laboratory	150	2	2.2	44	1,000
42	RF-SE-02-XC	Xcalibur XRF Services	105		9	0	1,295
42	RF-SE-22-XC	Xcalibur XRF Services	150	4		0	1,445
42	RF-SE-25-XC	Xcalibur XRF Services	140	23		0	1,371
42	RF-SE-30-XC	Xcalibur XRF Services	131		10	0	1,452
42	RF-SE-57-XC	Xcalibur XRF Services	112		3	0	1,465
43	RF-SE-15-XX	Reference Laboratory	160	1.4	3.6	45	1,300
43	RF-SE-24-XX	Reference Laboratory	170 J+	1.3 U	3.8 J+	46 J+	1,400 J-
43	RF-SE-32-XX	Reference Laboratory	140	1.3 U	4.2	36	1,100
43	RF-SE-43-XX	Reference Laboratory	150	1.3 U	4	40	1,200
43	RF-SE-59-XX	Reference Laboratory	160	1.3 U	4.5	42	1,300
43	RF-SE-15-XC	Xcalibur XRF Services	125		4	0	1,768
43	RF-SE-24-XC	Xcalibur XRF Services	140	14		22	1,861
43	RF-SE-32-XC	Xcalibur XRF Services	127		16	0	1,857
43	RF-SE-43-XC	Xcalibur XRF Services	127		19	40	1,939
43	RF-SE-59-XC	Xcalibur XRF Services	136		22	48	1,868

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg
44	RF-SE-05-XX	Reference Laboratory	4.1 J+	160	9.1	69	1,000	26,000	450	2.6
44	RF-SE-26-XX	Reference Laboratory	2.2 J+	140	8.4	64	990	23,000	440	2.5
44	RF-SE-39-XX	Reference Laboratory	2.9 J+	160	9.3	73	1,100	26,000	490	2.2
44	RF-SE-44-XX	Reference Laboratory	2.7 J+	140	8.2	64	970	24,000	420	2.3
44	RF-SE-56-XX	Reference Laboratory	3.5 J+	180	9.6	75	1200	27,000	490	2.2
44	RF-SE-05-XC	Xcalibur XRF Services	3	163	15	40	1,115	20,304	729	7
44	RF-SE-26-XC	Xcalibur XRF Services	9	225	30	43	1,133	20,882	683	
44	RF-SE-39-XC	Xcalibur XRF Services	12	209	24	86	1,157	21,647	769	
44	RF-SE-44-XC	Xcalibur XRF Services	7	231		46	1,171	21,394	657	5
44	RF-SE-56-XC	Xcalibur XRF Services	12	212		45	1,138	20,900	694	
45	RF-SE-04-XX	Reference Laboratory	3.2 J+	230	12	42	1,500	27,000	730	4.2
45	RF-SE-14-XX	Reference Laboratory	4.4 J+	260	12	47	1,700	30,000	800	4.7
45	RF-SE-19-XX	Reference Laboratory	3.7 J+	250	13	48	1,700	30,000	800	3.9
45	RF-SE-34-XX	Reference Laboratory	2.9 J+	210	10	39	1,400	24,000	660	4.5
45	RF-SE-52-XX	Reference Laboratory	3.4 J+	220	11	42	1,500	26,000	720	4.1
45	RF-SE-04-XC	Xcalibur XRF Services	4	303			1,637	22,522	1,023	
45	RF-SE-14-XC	Xcalibur XRF Services	5	274	27	51	1,635	23,684	1,118	
45	RF-SE-19-XC	Xcalibur XRF Services	15	284		47	1,673	22,966	1,145	
45	RF-SE-34-XC	Xcalibur XRF Services	17	343	12		1,727	23,564	1,069	
45	RF-SE-52-XC	Xcalibur XRF Services	16	311	25	49	1,720	23,725		1,100
46	BN-SO-11-XX	Reference Laboratory	4 J-	2,900	720	820	120	23,000	56	24 J-
46	BN-SO-14-XX	Reference Laboratory	3.5 J-	2,800	690	800	120	22,000	51	26
46	BN-SO-23-XX	Reference Laboratory	1.2 UJ	2,800	700	800	120	23,000	52	31
46	BN-SO-11-XC	Xcalibur XRF Services	5	4,644	951	519	431	24,935	0	25
46	BN-SO-14-XC	Xcalibur XRF Services	8	4,470	850	507	406	24,994	61	43
46	BN-SO-23-XC	Xcalibur XRF Services	0	4,327	818	533	423	25,464	77	37
47	BN-SO-09-XX	Reference Laboratory	750 J-	97	2,700	2,900	100	22,000	4,700	0.39
47	BN-SO-12-XX	Reference Laboratory	750 J-	89	2,600	2,800	96	21,000	4,500	0.34
47	BN-SO-24-XX	Reference Laboratory	810 J-	97	2,900	3,000	100	23,000	4,900	0.37
47	BN-SO-09-XC	Xcalibur XRF Services	374	60	3,128	1,613	76	15,548	7,409	
47	BN-SO-12-XC	Xcalibur XRF Services	366	52	3,078	1,605	67	15,511	7,237	
47	BN-SO-24-XC	Xcalibur XRF Services	348	70	2,870	1,563	75	15,803	7,395	

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
44	RF-SE-05-XX	Reference Laboratory	150	3.1	7.4 J-	48	1,800
44	RF-SE-26-XX	Reference Laboratory	140	2.8	7.2 J-	42	1,700
44	RF-SE-39-XX	Reference Laboratory	150	2.6	8.2 J-	49	1,900
44	RF-SE-44-XX	Reference Laboratory	140	2.4	7.2 J-	44	1,600
44	RF-SE-56-XX	Reference Laboratory	160	1.8	8.3 J-	51	1,900
44	RF-SE-05-XC	Xcalibur XRF Services	105		11	50	2,313
44	RF-SE-26-XC	Xcalibur XRF Services	107	1		0	2,307
44	RF-SE-39-XC	Xcalibur XRF Services	94		17	12	2,279
44	RF-SE-44-XC	Xcalibur XRF Services	107		6	0	2,351
44	RF-SE-56-XC	Xcalibur XRF Services	112		7	40	2,222
45	RF-SE-04-XX	Reference Laboratory	130	2.8	12 J-	46	2,400
45	RF-SE-14-XX	Reference Laboratory	140	3	13 J-	51	2,600
45	RF-SE-19-XX	Reference Laboratory	140	4.1	14 J-	52	2,700
45	RF-SE-34-XX	Reference Laboratory	120	1.9	10 J-	42	2,200
45	RF-SE-52-XX	Reference Laboratory	130	2	11 J-	47	2,300
45	RF-SE-04-XC	Xcalibur XRF Services	117		42	48	2,884
45	RF-SE-14-XC	Xcalibur XRF Services	105		25	34	3,004
45	RF-SE-19-XC	Xcalibur XRF Services	126		33	0	2,922
45	RF-SE-34-XC	Xcalibur XRF Services	125		12	0	2,971
45	RF-SE-52-XC	Xcalibur XRF Services	137		9	0	2,916
46	BN-SO-11-XX	Reference Laboratory	2,900	140	140 J-	150	3,900
46	BN-SO-14-XX	Reference Laboratory	2,800	130	140 J-	150	3,800
46	BN-SO-23-XX	Reference Laboratory	2,800	130	130 J-	150	3,800
46	BN-SO-11-XC	Xcalibur XRF Services	2,533	260	166	48	5,392
46	BN-SO-14-XC	Xcalibur XRF Services	2,429	208	142	75	5,294
46	BN-SO-23-XC	Xcalibur XRF Services	2,460	206	135	76	5,306
47	BN-SO-09-XX	Reference Laboratory	1,500	290	100 J-	340	81
47	BN-SO-12-XX	Reference Laboratory	1,400	290	210 J-	310	74
47	BN-SO-24-XX	Reference Laboratory	1,600	300	140 J-	350	81
47	BN-SO-09-XC	Xcalibur XRF Services	1,158	717	520	99	104
47	BN-SO-12-XC	Xcalibur XRF Services	1,194	624	512	40	110
47	BN-SO-24-XC	Xcalibur XRF Services	1,169	611	447	75	116

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg
48	SB-SO-09-XX	Reference Laboratory	1.3 UJ	9	0.51 U	130	120	35,000	19	30
48	SB-SO-20-XX	Reference Laboratory	1.3 UJ	11	0.51 U	170	150	44,000	24	10
48	SB-SO-31-XX	Reference Laboratory	1.3 UJ	8 J-	0.51 U	140	130	38,000	21	32
48	SB-SO-09-XC	Xcalibur XRF Services	3	67	10	160	414	38,474	32	34
48	SB-SO-20-XC	Xcalibur XRF Services	11	62	13	64	379	32,246	0	
48	SB-SO-31-XC	Xcalibur XRF Services	0	68		104	393	35,657	0	
49	SB-SO-29-XX	Reference Laboratory	1.2 U	9	0.5 U	140	130	41,000	19	7.9 J
49	SB-SO-36-XX	Reference Laboratory	1.2 U	8	0.5 U	120	100	33,000	15	36
49	SB-SO-56-XX	Reference Laboratory	1.2 U	10	0.5 U	150	140	42,000	20	9
49	SB-SO-29-XC	Xcalibur XRF Services	0	74	9	75	408	36,179	41	15
49	SB-SO-36-XC	Xcalibur XRF Services	0	52	28	150	424	37,462	55	19
49	SB-SO-56-XC	Xcalibur XRF Services	0	95	17	144	419	38,650	40	26
50	SB-SO-04-XX	Reference Laboratory	940	13	2,800	2,800	100	38,000	21	40
50	SB-SO-34-XX	Reference Laboratory	980	12	2,500	2,500	91	34,000	18	36
50	SB-SO-49-XX	Reference Laboratory	700	12	2,500	2,400	89	33,000	18	36
50	SB-SO-04-XC	Xcalibur XRF Services	522	121	3,100	1,800	80	24,853	68	34
50	SB-SO-34-XC	Xcalibur XRF Services		121	2,907	1,691	72	23,843	20	484
50	SB-SO-49-XC	Xcalibur XRF Services	528	85	3,174	1,770	71	24,585	51	44
51	WS-SO-07-XX	Reference Laboratory	3.8	53	1.9	640	4,400	25,000	1,700	0.26
51	WS-SO-11-XX	Reference Laboratory	1.2 U	46	1.4	570	3,900	19,000	1,500	0.27
51	WS-SO-25-XX	Reference Laboratory	1.2 U	59	3.1	730	4,900	24,000	1,900	0.25
51	WS-SO-07-XC	Xcalibur XRF Services	16	95		499	9,740	20,630	5,119	
51	WS-SO-11-XC	Xcalibur XRF Services	17	138		476	9,332	19,876	4,887	
51	WS-SO-25-XC	Xcalibur XRF Services	13	44		379	6,663	16,469	3,193	9
52	WS-SO-10-XX	Reference Laboratory	1.3 U	83	1.8	67	76	19,000	1,900	0.058 U
52	WS-SO-20-XX	Reference Laboratory	1.3 U	100	1.9	81	90	23,000	2,300	0.06 U
52	WS-SO-23-XX	Reference Laboratory	1.3 U	110	2.1	82	96	23,000	2,500	0.05 U
52	WS-SO-10-XC	Xcalibur XRF Services	22	80	11	93	286	26,434	5,426	
52	WS-SO-20-XC	Xcalibur XRF Services	23	56	40	52	288	25,411	4,817	20
52	WS-SO-23-XC	Xcalibur XRF Services	21	44	50	80	271	24,797	4,721	18
53	AS-SO-03-XX	Reference Laboratory	1.2 U	14	1,300	33	6,200	15,000	160	3.7 J-
53	AS-SO-05-XX	Reference Laboratory	1.2 U	9	900	23	4,500	11,000	110	2.5 J-
53	AS-SO-08-XX	Reference Laboratory	1.2 U	10	930	24	4,600	11,000	120	2.5 J-
53	AS-SO-03-XC	Xcalibur XRF Services	4	101	1,167	59	8,843	11,957	208	
53	AS-SO-05-XC	Xcalibur XRF Services	1	67	1,184		9,977	12,322	211	
53	AS-SO-08-XC	Xcalibur XRF Services	1	55	1,217		9,488	12,255	218	5

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	A	g	V	Zn
48	SB-SO-09-XX	Reference Laboratory	2900	26	160	J-	120	3,600
48	SB-SO-20-XX	Reference Laboratory	3700	30	140	J-	160	4,500
48	SB-SO-31-XX	Reference Laboratory	3200 J-	28 J-	160	J-	140	3,900 J-
48	SB-SO-09-XC	Xcalibur XRF Services	2,715	28	387		41	4,097
48	SB-SO-20-XC	Xcalibur XRF Services	2,375	31	300		0	3,934
48	SB-SO-31-XC	Xcalibur XRF Services	2,499	21	335		144	4,005
49	SB-SO-29-XX	Reference Laboratory	200	160	1.2	UJ	400	3,900
49	SB-SO-36-XX	Reference Laboratory	160	130	1.2	UJ	320	3,200
49	SB-SO-56-XX	Reference Laboratory	210	160	1.2	UJ	410	4,100
49	SB-SO-29-XC	Xcalibur XRF Services	146	381	3		199	4,175
49	SB-SO-36-XC	Xcalibur XRF Services	166	369	4		225	4,103
49	SB-SO-56-XC	Xcalibur XRF Services	169	383	5		97	4,111
50	SB-SO-04-XX	Reference Laboratory	3,300	390	1.3	UJ	58	86
50	SB-SO-34-XX	Reference Laboratory	3,000	360	1.3	UJ	52	77
50	SB-SO-49-XX	Reference Laboratory	2,800	330	1.2	UJ	52	72
50	SB-SO-04-XC	Xcalibur XRF Services	2,718	912			0	70
50	SB-SO-34-XC	Xcalibur XRF Services	2,565	855			76	62
50	SB-SO-49-XC	Xcalibur XRF Services	2,648	960	65		96	63
51	WS-SO-07-XX	Reference Laboratory	260	1.2 U	400	J-	48	180
51	WS-SO-11-XX	Reference Laboratory	240	1.2 U	340	J-	43	160
51	WS-SO-25-XX	Reference Laboratory	300	1.2 U	450	J-	54	200
51	WS-SO-07-XC	Xcalibur XRF Services	437	9	542		39	306
51	WS-SO-11-XC	Xcalibur XRF Services	420		537		9	327
51	WS-SO-25-XC	Xcalibur XRF Services	310		378		0	279
52	WS-SO-10-XX	Reference Laboratory	290	280	1.3	UJ	260	1,900
52	WS-SO-20-XX	Reference Laboratory	350	340	1.3	UJ	320	2,300
52	WS-SO-23-XX	Reference Laboratory	380	360	1.3	UJ	330	2,500
52	WS-SO-10-XC	Xcalibur XRF Services	328	1,175	19		151	3,742
52	WS-SO-20-XC	Xcalibur XRF Services	312	998	8		106	3,763
52	WS-SO-23-XC	Xcalibur XRF Services	319	1,017	16		117	3,736
53	AS-SO-03-XX	Reference Laboratory	520	200	480	J-	29	350
53	AS-SO-05-XX	Reference Laboratory	370	140	330	J-	23	250
53	AS-SO-08-XX	Reference Laboratory	380	140	280	J-	23	260
53	AS-SO-03-XC	Xcalibur XRF Services	607	334	366		13	532
53	AS-SO-05-XC	Xcalibur XRF Services	675	307	544		0	599
53	AS-SO-08-XC	Xcalibur XRF Services	605	339	377		0	556

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg
54	LV-SO-03-XX	Reference Laboratory	1.6	42	590	600	130	24,000	94	48 J-
54	LV-SO-40-XX	Reference Laboratory	2.7	42	580	590	130	24,000	92	46 J-
54	LV-SO-49-XX	Reference Laboratory	7.4	43	600	610	130	25,000	98	52 J-
54	LV-SO-03-XC	Xcalibur XRF Services	6	61	670	346	451	34,877	173	
54	LV-SO-40-XC	Xcalibur XRF Services	4	75	730	393	449	35,004	153	13
54	LV-SO-49-XC	Xcalibur XRF Services	1	65	604	346	413	30,730	197	
55	LV-SO-04-XX	Reference Laboratory	860	120	2,400	2,300	98	22,000	4,000	130 J-
55	LV-SO-34-XX	Reference Laboratory	870 J-	110 J-	2,300 J-	2,200 J-	87	20,000 J-	3,700 J-	130 J-
55	LV-SO-37-XX	Reference Laboratory	590	84	1,700	1,600	66	16,000	2,800	130 J-
55	LV-SO-04-XC	Xcalibur XRF Services	347	90	3,054	1,351	61	20,656	8,958	37
55	LV-SO-34-XC	Xcalibur XRF Services	335	52	2,985	1,443	48	20,387	8,837	38
55	LV-SO-37-XC	Xcalibur XRF Services	344	39	3,095	1,423	72	20,591	8,918	21
56	CN-SO-03-XX	Reference Laboratory	22	87	63	17	72	15,000	130	34 J-
56	CN-SO-06-XX	Reference Laboratory	20	91	64	18	74	16,000	130	40 J-
56	CN-SO-07-XX	Reference Laboratory	20	90	63	19	72	17,000	130	36 J-
56	CN-SO-03-XC	Xcalibur XRF Services	11	169	77		93	16,552	272	10
56	CN-SO-06-XC	Xcalibur XRF Services	12	201	68		82	15,546	256	7
56	CN-SO-07-XC	Xcalibur XRF Services	8	163	94		93	17,267	294	15
57	CN-SO-02-XX	Reference Laboratory	230	19	820	290	140	22,000	490	270 J-
57	CN-SO-05-XX	Reference Laboratory	130	6	630	26	160	23,000	25	280 J-
57	CN-SO-09-XX	Reference Laboratory	120	6	580	21	140	19,000	23	260 J-
57	CN-SO-02-XC	Xcalibur XRF Services	74	79	677	38	336	22,137	0	98
57	CN-SO-05-XC	Xcalibur XRF Services	78	80	695	35	308	20,389	0	110
57	CN-SO-09-XC	Xcalibur XRF Services	78	51	667		296	20,290	43	98
58	LV-SE-06-XX	Reference Laboratory	30	23	160	540	30	18,000	1,600	610 J-
58	LV-SE-13-XX	Reference Laboratory	31	24	160	540	30	18,000	1,600	640 J-
58	LV-SE-41-XX	Reference Laboratory	30	21	150	480	26	16,000	1,500	610 J-
58	LV-SE-06-XC	Xcalibur XRF Services	40	87	269	341	3	17,343	3,778	245
58	LV-SE-13-XC	Xcalibur XRF Services	38	54	234	379	4	17,202	3,596	225
58	LV-SE-41-XC	Xcalibur XRF Services	45	74	235	365	16	18,059	3,754	249
59	LV-SE-05-XX	Reference Laboratory	92	20	440	840	39	16,000	14	2.6 J-
59	LV-SE-20-XX	Reference Laboratory	140 J+	31	680	1,400	60	22,000	21	2.8
59	LV-SE-43-XX	Reference Laboratory	160 J+	24	550	1,100	47	19,000	17	2.8
59	LV-SE-05-XC	Xcalibur XRF Services	90	93	657	652	225	25,232	101	
59	LV-SE-20-XC	Xcalibur XRF Services	93	74	745	714	231	24,839	63	
59	LV-SE-43-XC	Xcalibur XRF Services	103	59	743	709	233	24,954	107	8

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni		Se		A	g	V		Zn	
54	LV-SO-03-XX	Reference Laboratory	2,000		120		210	J-	120		3,700	
54	LV-SO-40-XX	Reference Laboratory	1,900		120		210	J-	120		3,700	
54	LV-SO-49-XX	Reference Laboratory	2,000		120		220	J-	120		3,800	
54	LV-SO-03-XC	Xcalibur XRF Services	1,795		320		204		41		4,677	
54	LV-SO-40-XC	Xcalibur XRF Services	1,815		291		223		99		4,664	
54	LV-SO-49-XC	Xcalibur XRF Services	1,650		270		191		81		4,653	
55	LV-SO-04-XX	Reference Laboratory	2,000		230		1.2	UJ	260		53	
55	LV-SO-34-XX	Reference Laboratory	1,900	J-	220	J-	1.2	UJ	230	J-	48	J-
55	LV-SO-37-XX	Reference Laboratory	1,400		170		1.2	U	180		37	
55	LV-SO-04-XC	Xcalibur XRF Services	1,834		502				107		57	
55	LV-SO-34-XC	Xcalibur XRF Services	1,791		549		19		105		38	
55	LV-SO-37-XC	Xcalibur XRF Services	1,875		586		22		133		63	
56	CN-SO-03-XX	Reference Laboratory	74		36		90		30		58	
56	CN-SO-06-XX	Reference Laboratory	76		38		94		32		59	
56	CN-SO-07-XX	Reference Laboratory	75		37		91		33		58	
56	CN-SO-03-XC	Xcalibur XRF Services	81		22		114		33		132	
56	CN-SO-06-XC	Xcalibur XRF Services	78		46		99		23		140	
56	CN-SO-07-XC	Xcalibur XRF Services	102		41		108		29		140	
57	CN-SO-02-XX	Reference Laboratory	530		190		68		160		1,900	
57	CN-SO-05-XX	Reference Laboratory	360		190		78		160		2,200	
57	CN-SO-09-XX	Reference Laboratory	330		170		74		140		2,100	
57	CN-SO-02-XC	Xcalibur XRF Services	327		406		82		75		3,552	
57	CN-SO-05-XC	Xcalibur XRF Services	308		385		73		67		3,580	
57	CN-SO-09-XC	Xcalibur XRF Services	289		388		92		78		3,451	
58	LV-SE-06-XX	Reference Laboratory	360		160		110		480		52	
58	LV-SE-13-XX	Reference Laboratory	360		160		110		470		51	
58	LV-SE-41-XX	Reference Laboratory	320		150		99		420		46	
58	LV-SE-06-XC	Xcalibur XRF Services	299		406		148		194		64	
58	LV-SE-13-XC	Xcalibur XRF Services	305		343		139		207		79	
58	LV-SE-41-XC	Xcalibur XRF Services	331		329		139		217		64	
59	LV-SE-05-XX	Reference Laboratory	400		340		49		340		1,800	
59	LV-SE-20-XX	Reference Laboratory	660		500		75	J-	530		2,800	
59	LV-SE-43-XX	Reference Laboratory	530		420		60	J-	430		2,300	
59	LV-SE-05-XC	Xcalibur XRF Services	411		1,189		65		151		3,269	
59	LV-SE-20-XC	Xcalibur XRF Services	427		1,255		74		211		3,354	
59	LV-SE-43-XC	Xcalibur XRF Services	420		1,210		77		173		3,300	

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg
60	LV-SE-15-XX	Reference Laboratory	290 J+	32	1,300	83	2,300	22,000	18	500
60	LV-SE-17-XX	Reference Laboratory	280 J+	31	1,300	79	2,200	21,000	17 J-	490
60	LV-SE-51-XX	Reference Laboratory	210 J+	26	1,100	72	2,000	19,000	15	470
60	LV-SE-15-XC	Xcalibur XRF Services	150	70	1,365	69	2,440	18,435	47	247
60	LV-SE-17-XC	Xcalibur XRF Services	152	72	1,362		2,459	18,517	80	236
60	LV-SE-51-XC	Xcalibur XRF Services	119	129	1,186	45	2,275	17,652	0	212
61	TL-SE-05-XX	Reference Laboratory	100 J+	34	0.34 J	40	4,900	24,000	1,200	980
61	TL-SE-09-XX	Reference Laboratory	100 J+	33	0.24 J	39	4,800	23,000	1,200	820
61	TL-SE-13-XX	Reference Laboratory	95 J+	31	0.45 J	36 J+	4,400 J+	22,000 J+	1,100 J+	990
61	TL-SE-05-XC	Xcalibur XRF Services	134	53			6,233	22,153	2,327	424
61	TL-SE-09-XC	Xcalibur XRF Services	128	65			6,272	21,976	2,295	434
61	TL-SE-13-XC	Xcalibur XRF Services	125	87			6,013	22,026	2,354	450
62	TL-SE-06-XX	Reference Laboratory	1.2 U	86	350	34	2000	22,000	1,700	2.2
62	TL-SE-17-XX	Reference Laboratory	1.2 U	85	340	33	2100	21,000	1,700	2.6
62	TL-SE-28-XX	Reference Laboratory	1.2 U	89	360	34	2100	22,000	1,700	2.8
62	TL-SE-06-XC	Xcalibur XRF Services	16	67	463		2,895	22,639	4,143	
62	TL-SE-17-XC	Xcalibur XRF Services	17	63	501		2,729	21,993	4,379	
62	TL-SE-28-XC	Xcalibur XRF Services	21	126	520	20	2,659	21,638	4,711	
63	TL-SE-07-XX	Reference Laboratory	30	11	48	66	2200	37,000	13	40
63	TL-SE-21-XX	Reference Laboratory	33	13	51	73	2300	44,000	15	120
63	TL-SE-30-XX	Reference Laboratory	31	11	47	64	2200	36,000	14	100
63	TL-SE-07-XC	Xcalibur XRF Services	48	74	58	16	3,700	53,232	73	41
63	TL-SE-21-XC	Xcalibur XRF Services	50	69	42	13	3,736	53,772	117	19
63	TL-SE-30-XC	Xcalibur XRF Services	52	54	44		3,573	52,769	129	23
64	TL-SE-02-XX	Reference Laboratory	77	15	160	64	3,100	32,000	12	400
64	TL-SE-08-XX	Reference Laboratory	66	10	180	74	3,200	45,000	11	350
64	TL-SE-16-XX	Reference Laboratory	73	15	170	69	3,100	38,000	13	420
64	TL-SE-02-XC	Xcalibur XRF Services	101	55	159	43	4,258	45,180	144	152
64	TL-SE-08-XC	Xcalibur XRF Services	123	71	214		5,452	54,821	73	163
64	TL-SE-16-XC	Xcalibur XRF Services	120	33	213	12	5,324	52,189	43	192

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
60	LV-SE-15-XX	Reference Laboratory	230	92	300 J-	180	62
60	LV-SE-17-XX	Reference Laboratory	220	89	200 J-	170	58
60	LV-SE-51-XX	Reference Laboratory	200	76	250 J-	160	54
60	LV-SE-15-XC	Xcalibur XRF Services	187	76	513	94	27
60	LV-SE-17-XC	Xcalibur XRF Services	184	95	577	127	53
60	LV-SE-51-XC	Xcalibur XRF Services	166	70	437	69	0
61	TL-SE-05-XX	Reference Laboratory	54	130	180 J-	66	100
61	TL-SE-09-XX	Reference Laboratory	53	130	170 J-	63	100
61	TL-SE-13-XX	Reference Laboratory	49	120	160 J	59 J+	96
61	TL-SE-05-XC	Xcalibur XRF Services	116	190	224	0	86
61	TL-SE-09-XC	Xcalibur XRF Services	120	216	254	0	104
61	TL-SE-13-XC	Xcalibur XRF Services	103	180	227	0	99
62	TL-SE-06-XX	Reference Laboratory	44	45	56	78	83
62	TL-SE-17-XX	Reference Laboratory	43	44	56	78	81
62	TL-SE-28-XX	Reference Laboratory	44	45	57	81	83
62	TL-SE-06-XC	Xcalibur XRF Services	90	83	86	0	105
62	TL-SE-17-XC	Xcalibur XRF Services	71	31	85	0	116
62	TL-SE-28-XC	Xcalibur XRF Services	65	50	89	13	117
63	TL-SE-07-XX	Reference Laboratory	94	120	63	110	160
63	TL-SE-21-XX	Reference Laboratory	100	140	67	120	170
63	TL-SE-30-XX	Reference Laboratory	93	120	62	100	160
63	TL-SE-07-XC	Xcalibur XRF Services	152	404	117	0	95
63	TL-SE-21-XC	Xcalibur XRF Services	118	421	99	0	94
63	TL-SE-30-XC	Xcalibur XRF Services	122	483	130	58	115
64	TL-SE-02-XX	Reference Laboratory	99	44	120	110	160
64	TL-SE-08-XX	Reference Laboratory	100	39	130	120	170
64	TL-SE-16-XX	Reference Laboratory	100	44	120	110	160
64	TL-SE-02-XC	Xcalibur XRF Services	181	77	171	65	91
64	TL-SE-08-XC	Xcalibur XRF Services	155	92	227	0	93
64	TL-SE-16-XC	Xcalibur XRF Services	176	66	227	0	91

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb	As	Cd	Cr	Cu	Fe	Pb	Hg
65	RF-SE-01-XX	Reference Laboratory	12	230	40	280	63	14,000	22	47
65	RF-SE-09-XX	Reference Laboratory	10	260	45	310	71	16,000	26	45
65	RF-SE-11-XX	Reference Laboratory	11	240	43	300	72	15,000	25	52
65	RF-SE-17-XX	Reference Laboratory	11	250	43	300	67	15,000	26	20
65	RF-SE-29-XX	Reference Laboratory	13	280	49	330	75	17,000	26	20
65	RF-SE-37-XX	Reference Laboratory	11	260	45	320	72	16,000	27	22
65	RF-SE-50-XX	Reference Laboratory	8.9	230	40	280	65	14,000	23	19
65	RF-SE-01-XC	Xcalibur XRF Services	8	406	49	158	205	15,474	33	7
65	RF-SE-09-XC	Xcalibur XRF Services	11	424	69	232	216	15,814	77	19
65	RF-SE-11-XC	Xcalibur XRF Services	10	433	52	213	209	15,982	27	7
65	RF-SE-17-XC	Xcalibur XRF Services	11	406	59	270	220	16,378	82	16
65	RF-SE-29-XC	Xcalibur XRF Services	0	418	69	192	212	16,296	70	17
65	RF-SE-37-XC	Xcalibur XRF Services	0	447	66	219	216	16,332	32	6
65	RF-SE-50-XC	Xcalibur XRF Services	4	437	62	219	211	16,381	37	14
66	RF-SE-08-XX	Reference Laboratory	14	460	67	510	1,800	18,000	580	29
66	RF-SE-10-XX	Reference Laboratory	12	400	58	440	1,500	16,000	510	27
66	RF-SE-33-XX	Reference Laboratory	13	440	64	490	1,700	18,000	570	28
66	RF-SE-08-XC	Xcalibur XRF Services	21	737	65	337	2,066	12,810	1,023	16
66	RF-SE-10-XC	Xcalibur XRF Services	21	784	75	355	2,205	13,872	987	8
66	RF-SE-33-XC	Xcalibur XRF Services	19	709	73	324	2,138	13,548	1,068	11
67	RF-SE-16-XX	Reference Laboratory	85 J-	72 J-	310 J-	820 J-	73 J-	16,000 J-	24 J-	260
67	RF-SE-41-XX	Reference Laboratory	100	82	360	950	85	18,000	25	230
67	RF-SE-48-XX	Reference Laboratory	100	87	380	1,000	90	19,000	27	250
67	RF-SE-16-XC	Xcalibur XRF Services	52	114	421	664	112	14,873	44	102
67	RF-SE-41-XC	Xcalibur XRF Services	52	109	463	617	97	14,638	74	107
67	RF-SE-48-XC	Xcalibur XRF Services	57	142	432	640	104	14,899	36	111
68	RF-SE-18-XX	Reference Laboratory	320	810	770	950	78	16,000	860	600
68	RF-SE-35-XX	Reference Laboratory	300	740	700	860	70	15,000	780	650
68	RF-SE-54-XX	Reference Laboratory	320	880	840	1,000	86	18,000	920	670
68	RF-SE-18-XC	Xcalibur XRF Services	160	1,208	899	615	30	11,932	1,267	221
68	RF-SE-35-XC	Xcalibur XRF Services	164	1,239	942	681	44	12,298	1,329	242
68	RF-SE-54-XC	Xcalibur XRF Services	165	1,238	918	621	37	12,444	1,346	243

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
65	RF-SE-01-XX	Reference Laboratory	200	21	37	29	1,700
65	RF-SE-09-XX	Reference Laboratory	220	23	42	32	1,900
65	RF-SE-11-XX	Reference Laboratory	210	20	40	29	1,800
65	RF-SE-17-XX	Reference Laboratory	210	22	40	30	1,800
65	RF-SE-29-XX	Reference Laboratory	240	26	44	35	2,100
65	RF-SE-37-XX	Reference Laboratory	220	23	44	32	1,900
65	RF-SE-50-XX	Reference Laboratory	200	20	38	29	1,700
65	RF-SE-01-XC	Xcalibur XRF Services	169		46	0	3,293
65	RF-SE-09-XC	Xcalibur XRF Services	182		42	48	3,500
65	RF-SE-11-XC	Xcalibur XRF Services	192		27	0	3,405
65	RF-SE-17-XC	Xcalibur XRF Services	189	31	49	0	3,473
65	RF-SE-29-XC	Xcalibur XRF Services	186		62	0	3,421
65	RF-SE-37-XC	Xcalibur XRF Services	178		43	0	3,423
65	RF-SE-50-XC	Xcalibur XRF Services	183	59	52	0	3,463
66	RF-SE-08-XX	Reference Laboratory	250	42	0.39 U	120	120
66	RF-SE-10-XX	Reference Laboratory	220	39	0.34 U	100	110
66	RF-SE-33-XX	Reference Laboratory	240	41	0.33 U	120	130
66	RF-SE-08-XC	Xcalibur XRF Services	233			59	81
66	RF-SE-10-XC	Xcalibur XRF Services	252	28	13	61	107
66	RF-SE-33-XC	Xcalibur XRF Services	247	6	5	37	105
67	RF-SE-16-XX	Reference Laboratory	1,700 J-	1.2 U	130 J-	32 J-	760 J-
67	RF-SE-41-XX	Reference Laboratory	1,900	1.2 U	140	39	830
67	RF-SE-48-XX	Reference Laboratory	2,000	2.2	150	40	880
67	RF-SE-16-XC	Xcalibur XRF Services	1,663		156	21	1,373
67	RF-SE-41-XC	Xcalibur XRF Services	1,618		166	0	1,401
67	RF-SE-48-XC	Xcalibur XRF Services	1,608		169	54	1,377
68	RF-SE-18-XX	Reference Laboratory	390	140	140	390	120
68	RF-SE-35-XX	Reference Laboratory	350	140	150	340	110
68	RF-SE-54-XX	Reference Laboratory	420	160	180	410	120
68	RF-SE-18-XC	Xcalibur XRF Services	276	265	258	142	164
68	RF-SE-35-XC	Xcalibur XRF Services	340	257	268	142	176
68	RF-SE-54-XC	Xcalibur XRF Services	307	268	260	132	153

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Sb		As	Cd	Cr	Cu	Fe	Pb	Hg
69	RF-SE-20-XX	Reference Laboratory	550		1300	540	94	93	20,000	28	0.48
69	RF-SE-46-XX	Reference Laboratory	270		590	240	44	40	8,900	13	0.45
69	RF-SE-51-XX	Reference Laboratory	480		1100	450	77	77	17,000	23	0.48
69	RF-SE-20-XC	Xcalibur XRF Services	219		1,760	576	43	144	16,036		13
69	RF-SE-46-XC	Xcalibur XRF Services	232		1,858	619	73	139	16,115	89	19
69	RF-SE-51-XC	Xcalibur XRF Services	232		1,774	595	76	136	16,229	76	12
70	RF-SE-21-XX	Reference Laboratory	1.3	U	62	1,700	76	1,000	16,000	2,100	320
70	RF-SE-40-XX	Reference Laboratory	1.3	U	70	1,900	85	1,100	18,000	2,400	280
70	RF-SE-47-XX	Reference Laboratory	1.3	U	72	1,900	90	1,200	19,000	2,400	320
70	RF-SE-21-XC	Xcalibur XRF Services	19		85	2,180	50	1,060	13,404	3,814	95
70	RF-SE-40-XC	Xcalibur XRF Services	20		122	2,314	91	1,101	13,281	3,973	93
70	RF-SE-47-XC	Xcalibur XRF Services	19		116	2,295	53	1,093	13,555	3,841	92

Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

#### Appendix D. Analytical Data Summary, Xcalibur ElvaX and Reference Laboratory (Continued)

Blend No.	Sample ID	Source of Data	Ni	Se	Ag	V	Zn
69	RF-SE-20-XX	Reference Laboratory	1,400	380	59	36	1,400
69	RF-SE-46-XX	Reference Laboratory	650	170	26	16	650
69	RF-SE-51-XX	Reference Laboratory	1,200	320	48	30	1,200
69	RF-SE-20-XC	Xcalibur XRF Services	1,079	828	80	0	2,094
69	RF-SE-46-XC	Xcalibur XRF Services	1,056	912	69	0	2,115
69	RF-SE-51-XC	Xcalibur XRF Services	1,075	818	84	0	2,054
70	RF-SE-21-XX	Reference Laboratory	220	440	120	130	100
70	RF-SE-40-XX	Reference Laboratory	250	480	100	150	120
70	RF-SE-47-XX	Reference Laboratory	250	510	120	150	120
70	RF-SE-21-XC	Xcalibur XRF Services	179	1,232	305	58	107
70	RF-SE-40-XC	Xcalibur XRF Services	193	1,299	325	52	61
70	RF-SE-47-XC	Xcalibur XRF Services	211	1,321	312	59	117

Notes:

All concentrations reported in milligrams per kilogram (mg/kg), or parts per million (ppm).

Sample results for which "0" or no value was reported were considered nondetections as reported by Xcalibur. Reference laboratory data qualifiers were as follows:

J Estimated concentration.

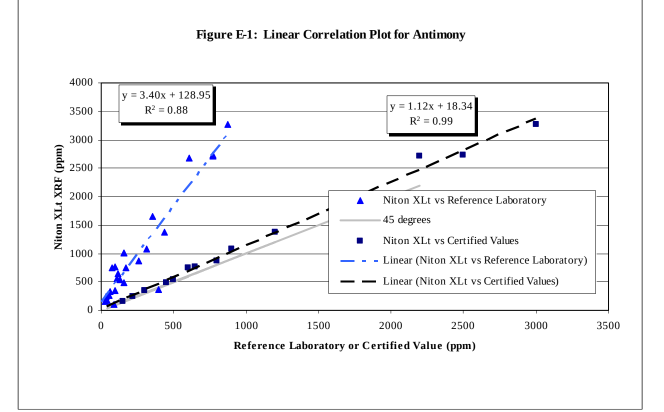
J+ Concentration is considered estimated and biased high.

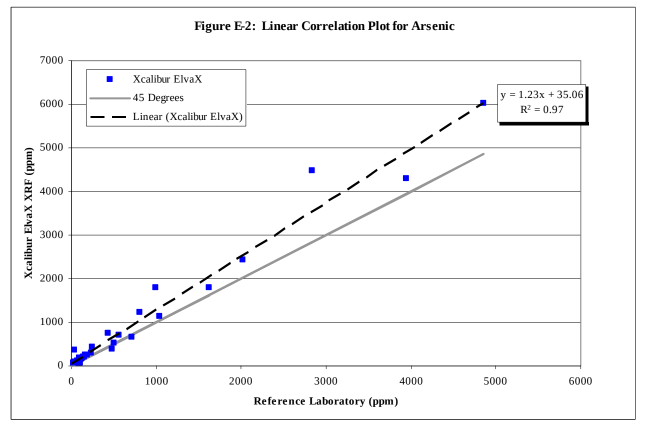
J- Concentration is considered estimated and biased low.

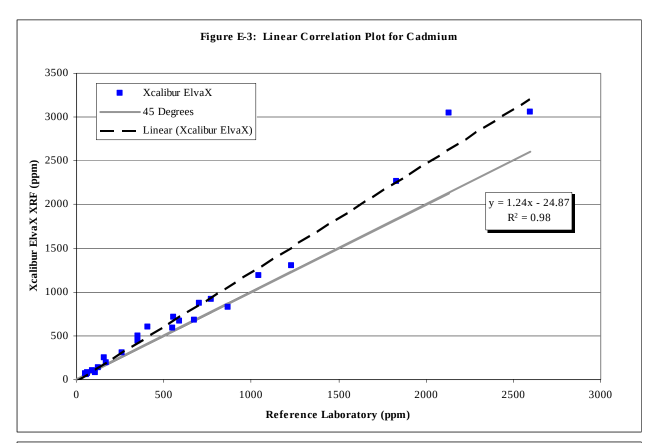
U Analyte is not detected; the associated concentration value is the sample reporting limit.

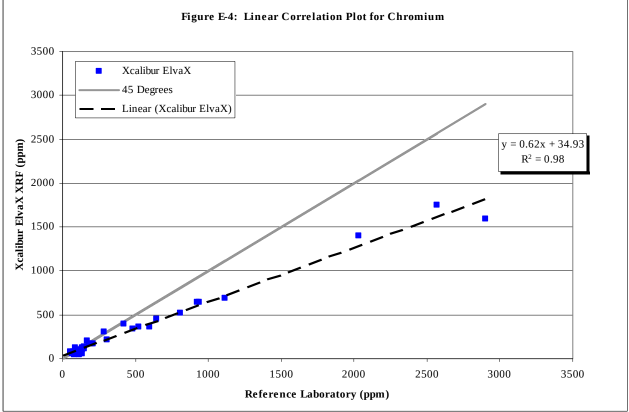
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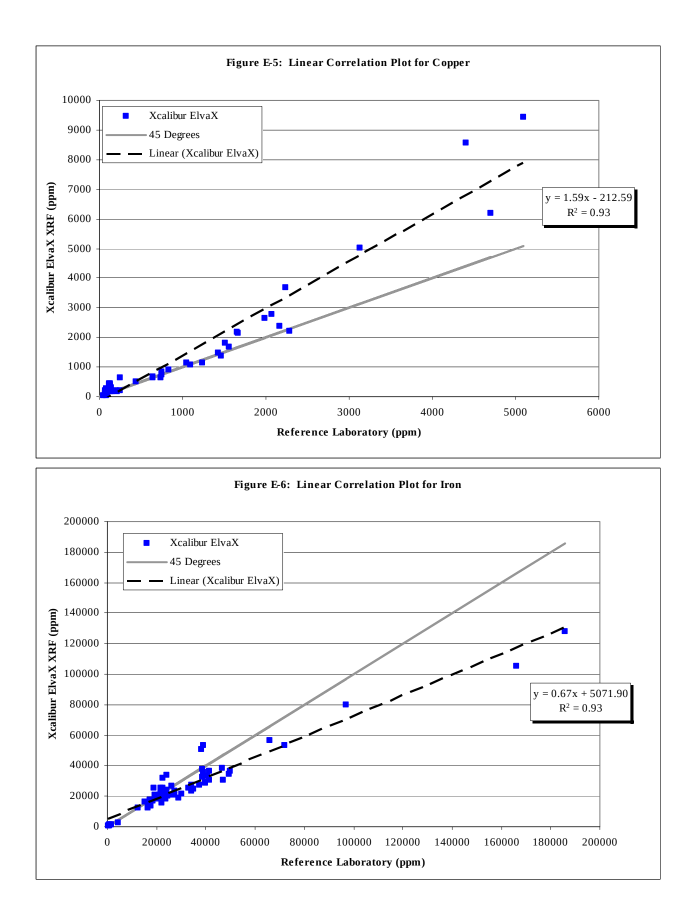
### STATISTICAL DATA SUMMARIES

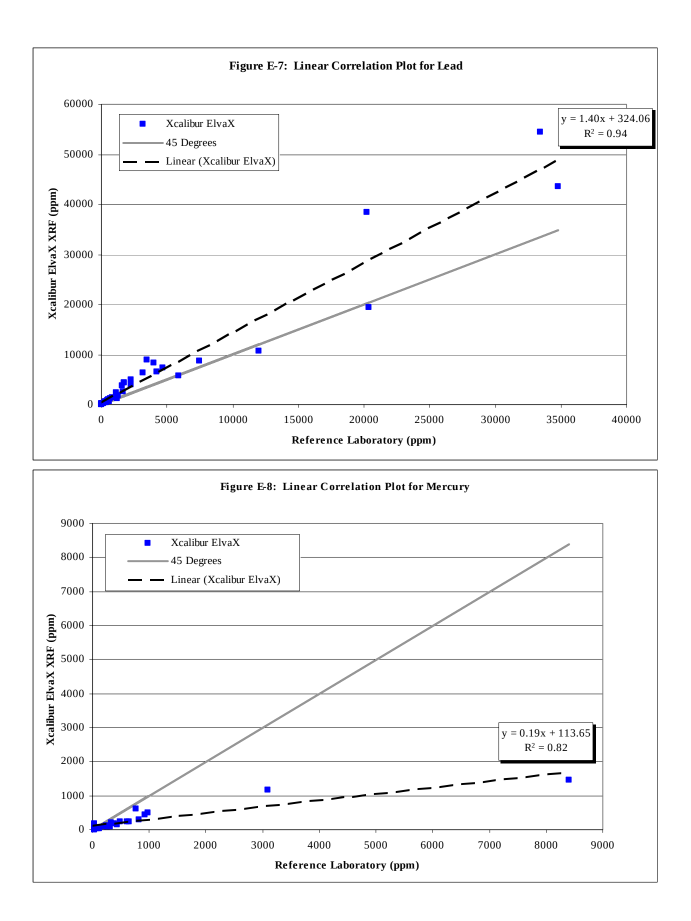


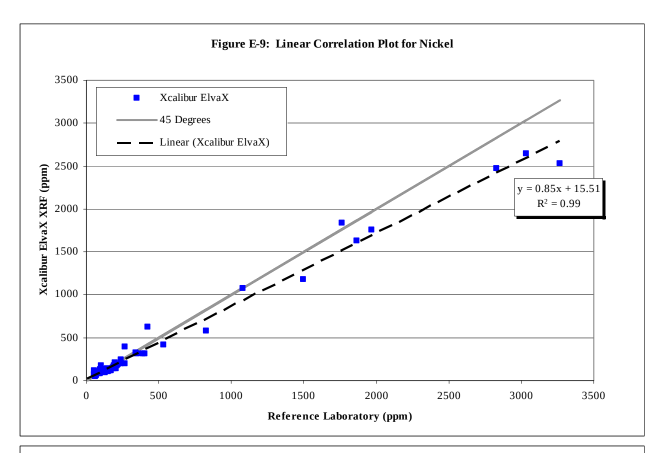


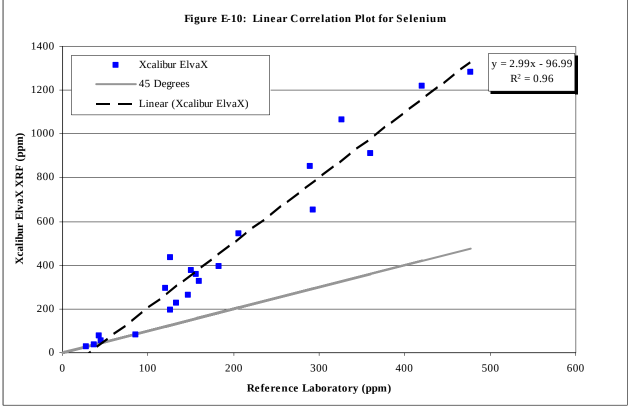


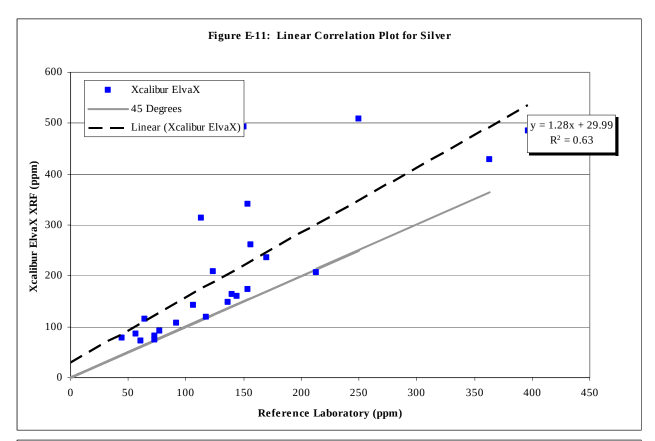


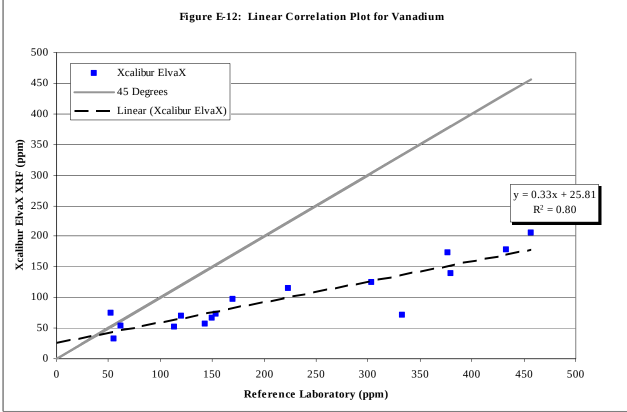


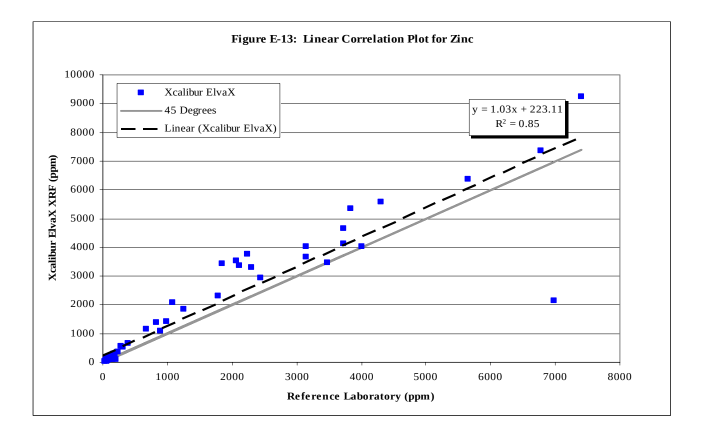


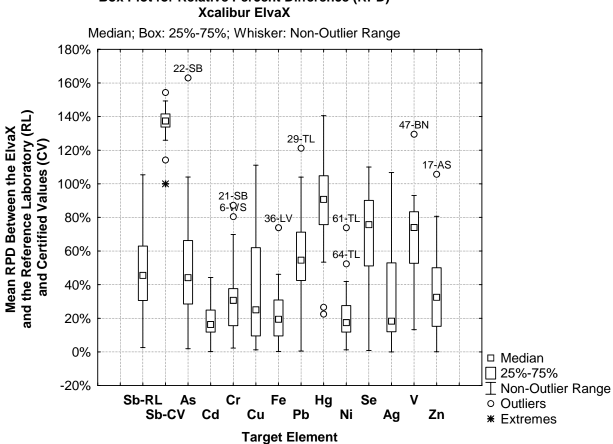












### Box Plot for Relative Percent Difference (RPD)

#### Notes:

The "box" in each box plot presents the range of RPD values that lie between the 25th and 75th percentiles (that is, the "quartiles") of the full RPD population for each element. In essence, the box displays the "interquartile range" of RPD values. The square data point within each box represents the median RPD for the population. The "whiskers" emanating from the top and bottom of each box represent the largest and smallest data points, respectively, that are within 1.5 times the interquartile range. Values outside the whiskers are identified as outliers and extremes.

Some of the more significant extremes and outliers are labeled with the associated Blend numbers and sample site abbreviations (see the footnotes of Table E-5 for definitions). Also refer to Appendix D for the results and sampling site associated with each Blend number.

#### Figure E-14. Box and Whiskers Plot for Mean RPD Values Showing Outliers and Extremes for Target Elements, Xcalibur ElvaX Data Set.

	Conc		Ant	imony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury
Matrix	Range	Statistic	Ref Lab	ERA Spike							
Soil	Level 1	Number	9	1	15	7	23	16	5	7	5
		Minimum	2.6%	154.4%	33.7%	0.2%	2.3%	9.9%	0.3%	8.3%	26.6%
		Maximum	105.4%	154.4%	163.0%	29.4%	87.2%	111.1%	46.2%	75.3%	133.5%
		Mean	43.7%	154.4%	62.6%	17.3%	28.4%	61.2%	24.7%	51.3%	99.0%
		Median	47.0%	154.4%	53.8%	16.4%	21.2%	62.7%	25.5%	50.5%	107.9%
	Level 2	Number	5	1	4	7	4	8	13	4	7
		Minimum	10.2%	6.2%	8.4%	0.4%	35.6%	1.3%	1.1%	0.5%	22.5%
		Maximum	70.2%	6.2%	22.2%	35.2%	49.6%	15.7%	33.9%	88.5%	99.6%
		Mean	32.2%	6.2%	15.6%	13.5%	41.2%	7.0%	13.8%	40.4%	75.2%
		Median	31.7%	6.2%	15.9%	12.4%	39.9%	5.7%	10.3%	36.2%	90.4%
	Level 3	Number	3	2	4	2	2	2	13	8	2
		Minimum	43.9%	0.7%	8.4%	10.1%	37.6%	59.7%	2.1%	2.2%	91.0%
		Maximum	77.4%	6.3%	45.0%	16.3%	58.1%	64.4%	40.3%	87.1%	140.5%
		Mean	64.4%	3.5%	23.4%	13.2%	47.9%	62.0%	24.6%	50.9%	115.7%
		Median	71.9%	3.5%	20.1%	13.2%	47.9%	62.0%	29.1%	55.5%	115.7%
	Level 4	Number							7	5	
		Minimum							18.9%	5.0%	
		Maximum							43.6%	62.3%	
		Mean							28.5%	29.6%	
		Median							30.9%	22.3%	
	All Soil	Number	17	4	23	16	29	26	38	24	14
		Minimum	2.6%	0.7%	8.4%	0.2%	2.3%	1.3%	0.3%	0.5%	22.5%
		Maximum	105.4%	154.4%	163.0%	35.2%	87.2%	111.1%	46.2%	88.5%	140.5%
		Mean	44.0%	41.9%	47.6%	15.1%	31.5%	44.6%	21.6%	44.8%	89.5%
		Median	43.9%	6.2%	45.0%	13.9%	29.4%	26.6%	24.1%	47.0%	96.2%

Table E-1. Evaluation of Accuracy - Relative Percent Difference versus Reference Laboratory Data Calculated for the Xcalibur ElvaX

	Conc		Nickel	Selenium	Silver	Vanadium	Zinc
Matrix	Range	Statistic		~			
Soil	Level 1	Number	24	2	3	3	20
		Minimum	1.5%	0.9%	2.6%	13.2%	0.1%
		Maximum	35.4%	3.7%	16.2%	50.4%	80.6%
		Mean	17.7%	2.3%	10.1%	33.0%	30.9%
		Median	16.7%	2.3%	11.6%	35.4%	21.6%
	Level 2	Number	5	5	3	3	6
		Minimum	6.1%	51.2%	0.0%	47.6%	0.8%
		Maximum	39.1%	86.3%	15.4%	77.4%	52.2%
		Mean	28.2%	72.5%	7.7%	65.2%	31.5%
		Median	30.9%	72.7%	7.6%	70.6%	35.2%
	Level 3	Number	6	4	7	4	9
		Minimum	3.7%	75.7%	3.6%	64.2%	0.3%
		Maximum	25.4%	106.0%	106.7%	129.5%	105.7%
		Mean	15.4%	89.6%	35.1%	87.7%	26.6%
		Median	13.6%	88.3%	16.5%	78.6%	22.2%
	Level 4	Number					
		Minimum					
		Maximum					
		Mean					
		Median					
	All Soil	Number	35	11	13	10	35
		Minimum	1.5%	0.9%	0.0%	13.2%	0.1%
		Maximum	39.1%	106.0%	106.7%	129.5%	105.7%
		Mean	18.8%	66.0%	23.0%	64.6%	29.9%
		Median	17.0%	75.7%	12.4%	67.4%	22.3%

 Table E-1. Evaluation of Accuracy - Relative Percent Difference versus Reference Laboratory Data Calculated for the Xcalibur ElvaX (Continued)

	Conc		Antimony		Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury
Matrix	Range	Statistic	Ref Lab	ERA Spike							
Sediment	Level 1	Number	3	3	17	3	6	8	3	12	3
		Minimum	30.6%	4.1%	1.9%	11.8%	4.8%	4.6%	10.9%	5.7%	80.5%
		Maximum	45.5%	100.0%	104.0%	44.3%	30.7%	101.8%	73.9%	121.2%	103.7%
		Mean	39.9%	38.5%	55.9%	23.4%	19.4%	43.0%	34.7%	67.0%	91.3%
		Median	43.8%	11.5%	42.5%	14.0%	20.6%	33.0%	19.2%	68.1%	89.8%
	Level 2	Number	4	4	4	4	3	4	19	3	4
		Minimum	27.0%	2.6%	2.2%	22.5%	34.0%	1.2%	1.2%	42.5%	70.9%
		Maximum	55.7%	18.8%	52.7%	37.1%	35.9%	8.3%	34.7%	66.4%	106.7%
		Mean	39.9%	9.9%	28.7%	29.7%	34.8%	4.8%	15.0%	56.2%	84.0%
		Median	38.4%	9.2%	29.9%	29.6%	34.6%	4.8%	14.0%	59.9%	79.2%
	Level 3	Number	3	3	2	3	3	10	4	3	3
		Minimum	59.9%	0.1%	41.1%	5.6%	36.2%	7.3%	16.2%	51.0%	72.3%
		Maximum	63.0%	4.2%	57.3%	21.0%	46.8%	48.7%	30.9%	88.7%	92.4%
		Mean	61.7%	2.3%	49.2%	14.8%	40.2%	26.7%	23.3%	73.7%	84.4%
		Median	62.3%	2.7%	49.2%	17.7%	37.8%	27.2%	23.0%	81.2%	88.5%
	Level 4	Number							6		
		Minimum							12.4%		
		Maximum							44.8%		
		Mean							28.5%		
		Median							30.6%		
	All Sediment	Number	10	10	23	10	12	22	32	18	10
		Minimum	27.0%	0.1%	1.9%	5.6%	4.8%	1.2%	1.2%	5.7%	70.9%
		Maximum	63.0%	100.0%	104.0%	44.3%	46.8%	101.8%	73.9%	121.2%	106.7%
		Mean	46.4%	16.2%	50.6%	23.3%	28.5%	28.7%	20.4%	66.3%	86.3%
		Median	45.6%	6.2%	42.5%	21.7%	32.4%	25.1%	17.6%	66.9%	84.5%

## Table E-1. Evaluation of Accuracy - Relative Percent Difference versus Reference Laboratory Data Calculated for the Xcalibur ElvaX (Continued)

	Conc		Nickel	Selenium	Silver	Vanadium	Zinc
Matrix	Range	Statistic					
Sediment	Level 1	Number	18	3	4	0	17
		Minimum	3.6%	6.5%	16.1%	NC	2.1%
		Maximum	73.9%	59.5%	57.1%	NC	70.9%
		Mean	26.3%	28.7%	42.6%	NC	34.6%
		Median	22.1%	20.0%	48.6%	NC	33.9%
	Level 2	Number	6	4	4	3	5
		Minimum	2.9%	42.6%	15.5%	55.1%	34.9%
		Maximum	22.8%	110.0%	51.1%	87.0%	63.4%
		Mean	15.3%	72.0%	31.9%	72.1%	47.8%
		Median	17.4%	67.7%	30.4%	74.2%	50.8%
	Level 3	Number	4	3	3	3	4
		Minimum	1.2%	91.7%	50.3%	75.8%	18.6%
		Maximum	36.0%	98.5%	93.9%	93.1%	60.1%
		Mean	18.5%	95.9%	70.8%	84.1%	35.0%
		Median	18.5%	97.4%	68.3%	83.3%	30.6%
	Level 4	Number					
		Minimum					
		Maximum					
		Mean					
		Median					
	All Sediment	Number	28	10	11	6	26
		Minimum	1.2%	6.5%	15.5%	55.1%	2.1%
		Maximum	73.9%	110.0%	93.9%	93.1%	70.9%
		Mean	22.8%	66.2%	46.4%	78.1%	37.2%
		Median	20.0%	69.1%	50.3%	79.5%	37.2%

# Table E-1. Evaluation of Accuracy - Relative Percent Difference versus Reference Laboratory Data Calculated for the Xcalibur ElvaX (Continued)

	Conc		Antimony		Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury
Matrix	Range	Statistic	Ref Lab	ERA Spike							
All Samples	Xcalibur ElvaX	Number	27	14	46	26	41	48	70	42	24
		Minimum	2.6%	0.1%	1.9%	0.2%	2.3%	1.2%	0.3%	0.5%	22.5%
		Maximum	105.4%	154.4%	163.0%	44.3%	87.2%	111.1%	73.9%	121.2%	140.5%
		Mean	44.9%	23.6%	49.1%	18.3%	30.6%	37.3%	21.1%	54.0%	88.2%
		Median	45.5%	6.2%	44.2%	16.3%	30.7%	25.1%	19.5%	54.6%	90.7%
All Samples	All Instruments	Number	206	110	320	209	338	363	558	392	192
		Minimum	0.1%	0.1%	0.2%	0.1%	0.1%	0.2%	0.0%	0.1%	0.0%
		Maximum	181.5%	162.0%	182.8%	168.1%	151.7%	111.1%	190.1%	135.2%	158.1%
		Mean	80.6%	62.7%	36.6%	29.6%	30.8%	24.6%	35.4%	30.9%	62.5%
		Median	84.3%	70.6%	26.2%	16.7%	26.0%	16.2%	26.0%	21.5%	58.6%

### Table E-1. Evaluation of Accuracy - Relative Percent Difference versus Reference Laboratory Data Calculated for the Xcalibur ElvaX (Continued)

# Table E-1. Evaluation of Accuracy - Relative Percent Difference versus Reference Laboratory Data Calculated for the Xcalibur ElvaX (Continued)

	Conc		Nickel	Selenium	Silver	Vanadium	Zinc
Matrix	Range	Statistic					
All Samples	Xcalibur ElvaX	Number	63	21	24	16	61
		Minimum	1.2%	0.9%	0.0%	13.2%	0.1%
		Maximum	73.9%	110.0%	106.7%	129.5%	105.7%
		Mean	20.6%	66.1%	33.7%	69.6%	33.0%
		Median	17.5%	75.7%	18.3%	74.0%	32.5%
All Samples	All Instruments	Number	403	195	177	218	471
		Minimum	0.3%	0.0%	0.0%	0.1%	0.0%
		Maximum	146.5%	127.1%	129.7%	129.5%	138.0%
		Mean	31.0%	32.0%	36.0%	42.2%	26.3%
		Median	25.4%	16.7%	28.7%	38.3%	19.4%

Notes: All RPDs presented in this table are absolute values.

	No samples reported by the reference laboratory in this concentration range.
Conc	Concentration.
ERA	Environmental Resource Associates, Inc.
NC	Not calculated because of a lack of XRF data.
Number	Number of demonstration samples evaluated.
Ref Lab	Reference laboratory (Shealy Environmental Services, Inc.).
RPD	Relative percent difference.
XRF	X-ray fluorescence.

	Conc									
Matrix	Range	Statistic	Antimony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury
Soil	Low	Number	9	15	7	23	16	5	7	5
		Minimum	3.0%	6.4%	4.7%	6.5%	2.0%	1.8%	1.9%	26.2%
		Maximum	24.2%	51.3%	35.2%	48.5%	20.5%	7.7%	25.2%	137.1%
		Mean	10.7%	21.4%	20.8%	23.4%	7.4%	4.8%	9.3%	54.3%
		Median	6.7%	16.1%	20.5%	20.9%	6.2%	4.4%	7.0%	37.7%
	Medium	Number	5	4	7	4	8	13	4	7
		Minimum	1.6%	5.9%	1.8%	2.5%	1.2%	0.7%	3.5%	7.1%
		Maximum	11.0%	34.7%	9.4%	14.1%	11.3%	11.7%	23.9%	18.1%
		Mean	5.3%	14.0%	4.1%	6.9%	5.2%	3.5%	9.0%	13.3%
		Median	3.0%	7.7%	2.2%	5.5%	3.8%	2.7%	4.2%	12.6%
	High	Number	3	4	2	2	2	13	8	2
		Minimum	1.8%	3.5%	4.5%	1.7%	6.0%	1.5%	0.7%	4.6%
		Maximum	19.0%	21.8%	4.5%	3.2%	19.5%	15.5%	19.3%	8.0%
		Mean	8.1%	14.3%	4.5%	2.4%	12.8%	5.5%	5.0%	6.3%
		Median	3.6%	16.0%	4.5%	2.4%	12.8%	3.8%	2.6%	6.3%
	Very High	Number						7	5	
		Minimum						2.8%	1.1%	
		Maximum						12.2%	15.8%	
		Mean						5.7%	8.7%	
		Median						4.5%	10.4%	
	All Soil	Number	17	23	16	29	26	38	24	14
		Minimum	1.6%	3.5%	1.8%	1.7%	1.2%	0.7%	0.7%	4.6%
		Maximum	24.2%	51.3%	35.2%	48.5%	20.5%	15.5%	25.2%	137.1%
		Mean	8.6%	18.9%	11.4%	19.7%	7.1%	4.8%	7.7%	26.9%
		Median	6.5%	14.8%	6.3%	17.7%	6.0%	4.1%	4.6%	15.7%

 Table E-2.
 Evaluation of Precision - Relative Standard Deviations Calculated for the Xcalibur ElvaX

	Conc						
Matrix	Range	Statistic	Nickel	Selenium	Silver	Vanadium	Zinc
Soil	Low	Number	24	2	3	3	20
		Minimum	2.9%	18.9%	8.6%	43.4%	1.8%
		Maximum	37.7%	35.1%	21.4%	52.3%	24.4%
		Mean	14.3%	27.0%	13.8%	48.4%	9.4%
		Median	10.3%	27.0%	11.5%	49.4%	8.1%
	Medium	Number	5	5	3	3	6
		Minimum	2.6%	2.0%	7.2%	7.7%	0.4%
		Maximum	17.7%	13.6%	10.9%	40.8%	7.3%
		Mean	8.1%	6.5%	8.5%	23.9%	3.2%
		Median	6.3%	5.2%	7.3%	23.3%	2.3%
	High	Number	6	4	7	4	9
		Minimum	1.6%	5.8%	7.9%	13.5%	0.3%
		Maximum	6.8%	9.1%	23.8%	42.0%	10.8%
		Mean	3.5%	7.9%	16.8%	28.3%	4.0%
		Median	2.6%	8.3%	19.2%	28.9%	1.8%
	Very High	Number					
		Minimum					
		Maximum					
		Mean					
		Median					
	All Soil	Number	35	11	13	10	35
		Minimum	1.6%	2.0%	7.2%	7.7%	0.3%
		Maximum	37.7%	35.1%	23.8%	52.3%	24.4%
		Mean	11.6%	10.7%	14.2%	33.0%	7.0%
		Median	7.9%	8.5%	11.5%	39.9%	6.4%

 Table E-2. Evaluation of Precision - Relative Standard Deviations Calculated for the Xcalibur ElvaX (Continued)

	Conc									
Matrix	Range	Statistic	Antimony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury
Sediment	Low	Number	3	17	3	6	8	3	12	3
		Minimum	3.8%	6.3%	7.4%	9.0%	2.3%	3.4%	2.9%	35.5%
		Maximum	9.0%	41.3%	16.0%	37.2%	18.1%	9.8%	47.0%	44.0%
		Mean	6.0%	19.2%	10.5%	27.0%	7.8%	6.0%	16.2%	40.8%
		Median	5.1%	18.5%	8.1%	30.0%	6.1%	4.9%	12.5%	42.8%
	Medium	Number	4	4	4	3	4	19	3	4
		Minimum	3.5%	3.7%	3.6%	4.7%	1.9%	0.4%	1.3%	1.9%
		Maximum	10.7%	7.2%	7.0%	16.0%	3.3%	8.2%	3.9%	12.1%
		Mean	6.7%	5.0%	5.4%	8.7%	2.4%	2.5%	2.8%	6.5%
		Median	6.3%	4.6%	5.5%	5.4%	2.3%	2.2%	3.2%	5.9%
	High	Number	3	2	3	3	10	4	3	3
		Minimum	1.6%	1.4%	2.3%	3.7%	2.3%	0.9%	2.2%	3.0%
		Maximum	13.2%	2.9%	7.9%	5.7%	13.1%	9.8%	6.5%	5.2%
		Mean	6.1%	2.2%	4.5%	4.8%	5.6%	3.9%	3.8%	4.5%
		Median	3.4%	2.2%	3.2%	5.0%	4.3%	2.4%	2.7%	5.2%
	Very High	Number						6		
		Minimum						1.4%		
		Maximum						6.2%		
		Mean						3.3%		
		Median						3.2%		
	All Sediment	Number	10	23	10	12	22	32	18	10
		Minimum	1.6%	1.4%	2.3%	3.7%	1.9%	0.4%	1.3%	1.9%
		Maximum	13.2%	41.3%	16.0%	37.2%	18.1%	9.8%	47.0%	44.0%
		Mean	6.3%	15.2%	6.6%	16.9%	5.8%	3.1%	11.9%	16.1%
		Median	5.2%	14.1%	6.5%	12.5%	4.3%	2.5%	6.4%	6.5%

 Table E-2. Evaluation of Precision - Relative Standard Deviations Calculated for the Xcalibur ElvaX (Continued)

	Conc						
Matrix	Range	Statistic	Nickel	Selenium	Silver	Vanadium	Zinc
Sediment	Low	Number	18	3	4	0	17
		Minimum	0.9%	16.0%	2.2%	NC	1.2%
		Maximum	35.4%	48.6%	13.6%	NC	31.5%
		Mean	11.1%	27.3%	8.6%	NC	10.2%
		Median	9.9%	17.3%	9.4%	NC	8.3%
	Medium	Number	6	4	4	3	5
		Minimum	4.0%	2.3%	3.4%	7.2%	1.1%
		Maximum	10.5%	11.4%	15.5%	30.0%	5.2%
		Mean	6.4%	8.2%	7.5%	21.0%	3.2%
		Median	5.9%	9.6%	5.6%	25.7%	3.3%
	High	Number	4	3	3	3	4
		Minimum	1.1%	2.8%	2.0%	4.4%	1.3%
		Maximum	8.7%	6.1%	13.8%	16.9%	2.1%
		Mean	3.4%	4.2%	6.3%	9.0%	1.7%
		Median	1.8%	3.6%	3.2%	5.6%	1.8%
	Very High	Number					
		Minimum					
		Maximum					
		Mean					
		Median					
	All Sediment	Number	28	10	11	6	26
		Minimum	0.9%	2.3%	2.0%	4.4%	1.1%
		Maximum	35.4%	48.6%	15.5%	30.0%	31.5%
		Mean	9.0%	12.7%	7.6%	15.0%	7.6%
		Median	8.2%	9.6%	7.0%	12.1%	5.8%

 Table E-2. Evaluation of Precision - Relative Standard Deviations Calculated for the Xcalibur ElvaX (Continued)

	Conc									
Matrix	Range	Statistic	Antimony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury
All Samples	Xcalibur ElvaX	Number	27	46	26	41	48	70	42	24
		Minimum	1.6%	1.4%	1.8%	1.7%	1.2%	0.4%	0.7%	1.9%
		Maximum	24.2%	51.3%	35.2%	48.5%	20.5%	15.5%	47.0%	137.1%
		Mean	7.8%	17.1%	9.6%	18.9%	6.5%	4.0%	9.5%	22.4%
		Median	5.4%	14.1%	6.5%	17.2%	4.9%	3.2%	6.0%	12.5%
All Samples	All Instruments	Number	206	320	209	338	363	558	392	192
		Minimum	0.5%	0.2%	0.4%	0.6%	0.1%	0.1%	0.2%	1.0%
		Maximum	97.7%	71.7%	92.8%	116.3%	58.3%	101.8%	115.6%	137.1%
		Mean	8.9%	11.2%	8.2%	15.9%	7.5%	5.2%	9.3%	14.3%
		Median	6.1%	8.2%	3.6%	12.1%	5.1%	2.2%	4.9%	6.8%

 Table E-2. Evaluation of Precision - Relative Standard Deviations Calculated for the Xcalibur ElvaX (Continued)

 Table E-2. Evaluation of Precision - Relative Standard Deviations Calculated for the Xcalibur ElvaX (Continued)

	Conc						
Matrix	Range	Statistic	Nickel	Selenium	Silver	Vanadium	Zinc
All Samples	Xcalibur ElvaX	Number	63	21	24	16	61
		Minimum	0.9%	2.0%	2.0%	4.4%	0.3%
		Maximum	37.7%	48.6%	23.8%	52.3%	31.5%
		Mean	10.4%	11.7%	11.2%	26.2%	7.2%
		Median	8.0%	8.9%	9.4%	24.5%	6.1%
All Samples	All Instruments	Number	403	195	177	218	471
		Minimum	0.3%	0.1%	0.6%	0.4%	0.1%
		Maximum	164.2%	98.8%	125.3%	86.1%	192.9%
		Mean	10.8%	7.2%	10.3%	12.5%	8.0%
		Median	7.0%	4.5%	5.2%	8.5%	5.3%

Notes:

	No samples reported by the reference laboratory in this concentration range.
Conc	Concentration.
NC	Not calculated because of a lack of XRF data.
Number	Number of demonstration samples evaluated.
RSD	Relative standard deviation.
XRF	X-ray fluorescence.

Matrix	Statistic	Antimony	Arsenic	Cadmium	Chromium	Copper	Iron	Lead	Mercury
All Soil	Number	17	23	15	34	26	38	33	16
	Minimum	3.6%	1.4%	0.9%	1.4%	0.0%	1.6%	0.0%	0.0%
	Maximum	38.0%	45.8%	21.4%	137.0%	21.0%	46.2%	150.0%	50.7%
	Mean	14.3%	11.7%	11.1%	14.3%	10.1%	10.2%	17.6%	13.8%
	Median	9.8%	12.4%	9.0%	10.6%	9.1%	8.7%	13.2%	6.6%
All Sediment	Number	7	24	10	26	21	31	22	10
	Minimum	2.9%	2.4%	2.9%	4.6%	1.8%	2.7%	0.0%	2.8%
	Maximum	33.6%	36.7%	37.5%	35.5%	38.8%	37.5%	41.1%	48.0%
	Mean	14.4%	10.7%	11.4%	9.8%	9.7%	9.9%	11.6%	14.3%
	Median	9.1%	9.2%	8.2%	7.5%	8.9%	8.1%	7.4%	6.9%
All Samples	Number	24	47	25	60	47	69	55	26
_	Minimum	2.9%	1.4%	0.9%	1.4%	0.0%	1.6%	0.0%	0.0%
	Maximum	38.0%	45.8%	37.5%	137.0%	38.8%	46.2%	150.0%	50.7%
	Mean	14.3%	11.2%	11.2%	12.4%	9.9%	10.1%	15.2%	14.0%
	Median	9.5%	9.5%	9.0%	8.4%	8.9%	8.5%	8.6%	6.6%

 Table E-3. Evaluation of Precision - Relative Standard Deviations Calculated for the Reference Laboratory

 Table E-3. Evaluation of Precision - Relative Standard Deviations Calculated for the Reference Laboratory (Continued)

Matrix	Statistic	Nickel	Selenium	Silver	Vanadium	Zinc
All Soil	Number	35	13	13	21	35
	Minimum	0.0%	0.0%	2.3%	0.0%	1.0%
	Maximum	44.9%	22.7%	37.1%	18.1%	46.5%
	Mean 11.4%		8.9%	12.4%	8.4%	10.4%
	Median	10.0%	7.1%	7.5%	6.6%	9.1%
All Sediment	Number	27	12	10	17	27
	Minimum	0.6%	1.3%	1.0%	2.2%	1.4%
	Maximum	35.8%	37.3%	21.3%	21.9%	35.8%
	Mean	9.4%	10.0%	9.4%	8.4%	8.9%
	Median	7.3%	7.6%	6.6%	8.1%	6.9%
All Samples	Number	62	25	23	38	62
	Minimum	0.0%	0.0%	1.0%	0.0%	1.0%
	Maximum	44.9%	37.3%	37.1%	21.9%	46.5%
	Mean	10.6%	9.4%	11.1%	8.4%	9.8%
	Median	8.2%	7.4%	7.1%	7.2%	7.4%

istic Lead	Effects on	Arsenic	Coppe	r Effects on	Nickel	Nickel	Effects on	Copper
<5	5 - 10	>10	<5	5 - 10	>10	<5	5 - 10	>10
29	7	10	44	5	14	39	1	8
mum -163.0% imum 8.1% n -55.0%	-33.7% 22.2% -11.7%	-104.0% 53.8% -22.9%	-24.1% 39.1% 14.0%	-3.6% 33.9% 12.1%	-73.9% 27.7% -17.3%	-108.7% 53.0% -22.2%	70.7% 70.7% 70.7%	-111.1% 32.4% -40.3%
ian -45.1%	-11.7%	-31.9%	14.0% 16.7%	12.1%	-17.5%	-22.2%	70.7%	-40.3% -44.9%
imum2.2%imum163.0%n55.5%ian45.1%	8.4% 33.7% 18.0% 21.5%	1.9% 104.0% 52.2% 50.6%	1.2% 39.1% 18.4% 17.2%	2.9% 33.9% 14.7% 15.7%	4.0% 73.9% 29.6% 27.6%	1.2% 108.7% 31.4% 21.0%	70.7% 70.7% 70.7% 70.7%	22.8% 111.1% 61.7% 49.4%
imum ND imum 1314 n 299 ian 152	1163 54373 19071 8802	2326 38476 8482 4699	ND 1085 193 73	852 2137 1372 1143	682 9436 3473 2298	ND 629 163 128	307 307 307 307 307	1070 2644 1888 1793
imum         67           imum         4480           n         452           imum         179	163 6033 2068	59 2434 310	48 2644 476	78 244 152	58 629 172	29 9436 1669	37 37 37 37	60 438 213 122
imum	4480	448060334522068	4480603324344522068310	44806033243426444522068310476	4480         6033         2434         2644         244           452         2068         310         476         152	44806033243426442446294522068310476152172	4480603324342644244629943645220683104761521721669	44806033243426442446299436374522068310476152172166937

Table E-4. Evaluation of the Effects of Interferent Elements on RPDs (Accuracy) of Other Target Elements¹

Parameter	Statistic	Zinc	Effects on	Copper	Сорре	er Effects o	n Zinc
Interferent/Element Ratio		<5	5 - 10	>10	<5	5 - 10	>10
Number of Samples		35	2	11	49	3	9
RPD of Target Element ²	Minimum Maximum	-64.4% 70.7%	-9.9% 6.1%	-111.1% -23.3%	-80.6% 105.7%	-47.1% 70.9%	-65.0% 56.2%
	Mean Median	-4.2% -7.3%	-1.9% -1.9%	-23.3% -87.9% -99.1%	-24.4% -25.3%	13.8% 17.6%	-0.4% 2.3%
RPD of Target Element	Minimum	1.2%	6.1%	23.3%	0.1%	17.6%	2.3%
(Absolute Value) ²	Maximum Mean	70.7% 23.0%	9.9% 8.0%	111.1% 87.9%	105.7% 32.0%	70.9% 45.2%	65.0% 34.7%
Interferent	Median Minimum	21.0%	8.0%	99.1% 1384	32.5% ND	47.1% 634	31.3% 1818
Concentration Range	Maximum Mean	7370 1267	9253 5169	5331 3447	2205 386	2187 1302	9436 4692
	Median	177	5169	3528	177	1085	3670
Target Element Concentration Range	Minimum Maximum	29 9436	177 1374	104 639	36 9253	95 165	92 562
	Mean Median	1760 909	775 775	336 332	1944 1085	120 99	183 103

 Table E-4. Evaluation of the Effects of Interferent Elements on RPDs (Accuracy) of Other Target Elements¹ (Continued)

Notes:

1. Concentrations are reported in units of milligrams per kilogram (mg/kg), or parts per million (ppm).

2. Table presents statistics for raw (unmodified) RPDs as well as absolute value RPDs.

< Less than.

> Greater than.

RPD Relative percent difference.

NC Not calculated because of a lack of XRF data.

ND Nondetect.

XRF X-ray fluorescence.

					Anti	imony		A	rsenic	C	admium
				Referer	nce Laboratory	Certif	ied Value	Reference	ce Laboratory	Referen	ce Laboratory
Matrix	Site	Matrix Description	Statistic	RPD	RPD ABS Val	RPD	RPD ABS Val	RPD	RPD ABS Val	RPD	RPD ABS Val
Soil	AS	Fine to medium sand	Number					1	1	3	3
		(steel processing)	Minimum					-82.6%	82.6%	-13.1%	0.2%
			Maximum					-82.6%	82.6%	25.0%	25.0%
			Mean					-82.6%	82.6%	4.0%	12.8%
			Median					-82.6%	82.6%	0.2%	13.1%
Soil	BN	Sandy loam, low	Number	4	4	1	1	7	7	5	5
		organic (ore residuals)	Minimum	-2.7%	2.7%	6.3%	6.3%	-63.7%	8.4%	-21.6%	5.3%
			Maximum	71.9%	71.9%	6.3%	6.3%	43.3%	63.7%	5.3%	21.6%
			Mean	23.6%	24.9%	6.3%	6.3%	-19.9%	32.3%	-9.8%	11.9%
			Median	12.6%	12.6%	6.3%	6.3%	-22.1%	33.7%	-10.1%	10.1%
Soil	CN	Sandy loam (burn pit	Number	2	2	2	2	1	1	2	2
		residue)	Minimum	67.6%	67.6%	6.2%	6.2%	-66.2%	66.2%	-23.0%	0.4%
			Maximum	70.2%	70.2%	154.4%	154.4%	-66.2%	66.2%	-0.4%	23.0%
			Mean	68.9%	68.9%	80.3%	80.3%	-66.2%	66.2%	-11.7%	11.7%
			Median	68.9%	68.9%	80.3%	80.3%	-66.2%	66.2%	-11.7%	11.7%
Soil &	KP	Soil: Fine to medium	Number	1	1						
Sediment		quartz sand.	Minimum	48.2%	48.2%						
		Sed.: Sandy loam, high organic.	Maximum	48.2%	48.2%						
			Mean	48.2%	48.2%						
	(Gun and skeet ranges.)	Median	48.2%	48.2%							

				Ch	romium	C	opper		Iron		Lead
				Referen	ce Laboratory	Reference	e Laboratory	Reference	e Laboratory	Reference	e Laboratory
	~	Matrix	~		RPD ABS		RPD ABS		RPD ABS		RPD ABS
Matrix	Site	Description	Statistic	RPD	Val	RPD	Val	RPD	Val	RPD	Val
Soil	AS	Fine to medium sand	Number	2	2	3	3	3	3	3	3
		(steel processing)	Minimum	6.1%	6.1%	-86.3%	59.7%	1.3%	1.3%	-67.0%	40.2%
			Maximum	11.6%	11.6%	-59.7%	86.3%	31.2%	31.2%	-40.2%	67.0%
			Mean	8.8%	8.8%	-76.5%	76.5%	17.4%	17.4%	-51.8%	51.8%
			Median	8.8%	8.8%	-83.7%	83.7%	19.7%	19.7%	-48.1%	48.1%
Soil	Soil BN	Sandy loam, low organic (ore residuals)	Number	5	5	6	6	7	7	6	6
			Minimum	31.6%	31.6%	-111.1%	2.0%	-10.3%	1.1%	-50.5%	10.9%
		Maximum	58.1%	58.1%	30.4%	111.1%	33.9%	33.9%	10.9%	50.5%	
			Mean	42.1%	42.1%	-17.3%	28.5%	11.3%	14.8%	-31.7%	35.3%
			Median	43.2%	43.2%	-6.6%	12.1%	5.7%	10.3%	-43.8%	43.8%
Soil	CN	Sandy loam (burn pit	Number	1	1	3	3	3	3	2	2
		residue)	Minimum	21.2%	21.2%	-72.5%	1.3%	-2.8%	1.9%	-71.2%	32.2%
			Maximum	21.2%	21.2%	1.3%	72.5%	19.8%	19.8%	-32.2%	71.2%
			Mean	21.2%	21.2%	-30.7%	31.6%	6.3%	8.2%	-51.7%	51.7%
			Median	21.2%	21.2%	-21.0%	21.0%	1.9%	2.8%	-51.7%	51.7%
Soil &	KP	Soil: Fine to medium	Number	4	4	2	2	6	6	6	6
Sediment		quartz sand.	Minimum	-20.1%	4.8%	15.7%	15.7%	-19.2%	0.3%	-43.0%	0.5%
		Sed.: Sandy loam, high	Maximum	8.9%	20.1%	22.8%	22.8%	46.2%	46.2%	8.3%	43.0%
		organic. (Gun and skeet ranges.)	Mean	-5.5%	10.0%	19.2%	19.2%	8.2%	18.3%	-5.6%	10.8%
		(Our and skeet ranges.)	Median	-5.4%	7.5%	19.2%	19.2%	3.9%	15.1%	0.8%	5.3%

Table E-5. Evaluation	of the Effects of Soil Type of	n RPDs (Accuracy) of Targe	et Elements (Continued)

				М	ercury	Ň	ickel	S	elenium	Silver	
				Referenc	e Laboratory	Reference	e Laboratory	Referen	ce Laboratory	Referen	ce Laboratory
		Matrix			RPD ABS		RPD ABS				
Matrix	Site	Description	Statistic	RPD	Val	RPD	Val	RPD	<b>RPD ABS Val</b>	RPD	<b>RPD ABS Val</b>
Soil	AS	Fine to medium sand	Number			3	3	1	1	1	1
		(steel processing)	Minimum			-39.1%	9.3%	-68.5%	68.5%	-16.5%	16.5%
			Maximum			25.8%	39.1%	-68.5%	68.5%	-16.5%	16.5%
			Mean			-1.3%	24.7%	-68.5%	68.5%	-16.5%	16.5%
			Median			9.3%	25.8%	-68.5%	68.5%	-16.5%	16.5%
Soil	BN	Sandy loam, low	Number	1	1	6	6	2	2	4	4
	organic (ore residuals)	Minimum	-26.6%	26.6%	-9.8%	9.8%	-75.7%	51.2%	-106.7%	0.0%	
			Maximum	-26.6%	26.6%	27.7%	27.7%	-51.2%	75.7%	0.0%	106.7%
			Mean	-26.6%	26.6%	16.5%	19.8%	-63.4%	63.4%	-29.2%	29.2%
			Median	-26.6%	26.6%	20.1%	20.1%	-63.4%	63.4%	-5.1%	5.1%
Soil	CN	Sandy loam (burn pit	Number	2	2	3	3	2	2	2	2
		residue)	Minimum	90.4%	90.4%	-14.8%	14.8%	-72.7%	0.9%	-15.4%	11.6%
			Maximum	107.9%	107.9%	30.9%	30.9%	0.9%	72.7%	-11.6%	15.4%
			Mean	99.2%	99.2%	14.6%	24.4%	-35.9%	36.8%	-13.5%	13.5%
			Median	99.2%	99.2%	27.6%	27.6%	-35.9%	36.8%	-13.5%	13.5%
Soil &	KP	Soil: Fine to medium	Number			3	3				
Sediment		quartz sand.	Minimum			-9.8%	1.5%				
		Sed.: Sandy loam, high	Maximum			12.0%	12.0%				
		organic. (Gun and skeet ranges.)	Mean			0.2%	7.8%				
		(Our and skeet ranges.)	Median			-1.5%	9.8%				

				Va	nadium		Zinc
				Reference	ce Laboratory	Reference	e Laboratory
		Matrix			RPD ABS		RPD ABS
Matrix	Site	Description	Statistic	RPD	Val	RPD	Val
Soil	AS	Fine to medium sand (steel processing)	Number			3	3
			Minimum			-65.0%	0.8%
			Maximum			105.7%	105.7%
			Mean			13.8%	57.1%
			Median			0.8%	65.0%
Soil	BN	Sandy loam, low	Number	2	2	7	7
		organic (ore residuals)	Minimum	77.4%	77.4%	-61.6%	8.3%
			Maximum	129.5%	129.5%	-8.3%	61.6%
			Mean	103.4%	103.4%	-36.8%	36.8%
			Median	103.4%	103.4%	-33.4%	33.4%
Soil	CN	Sandy loam (burn pit	Number	1	1	3	3
		residue)	Minimum	70.6%	70.6%	-80.6%	15.2%
			Maximum	70.6%	70.6%	-15.2%	80.6%
			Mean	70.6%	70.6%	-49.4%	49.4%
			Median	70.6%	70.6%	-52.2%	52.2%
Soil &	KP	Soil: Fine to medium	Number			2	2
Sediment		quartz sand.	Minimum			-57.3%	47.1%
		Sed.: Sandy loam, high	Maximum			-47.1%	57.3%
		organic. (Gun and skeet ranges.)	Mean			-52.2%	52.2%
		(Our and skeet ranges.)	Median			-52.2%	52.2%

Table E-5. Evaluation of the Effects of Soil Type on RPDs (Accuracy) of Target Elements (Continued)

					Ant	imony		A	rsenic	Cadmium	
				Referenc	e Laboratory	Certif	ied Value	Reference	e Laboratory	Referen	ce Laboratory
		Matrix			RPD ABS		RPD ABS		RPD ABS		
Matrix	Site	Description	Statistic	RPD	Val	RPD	Val	RPD	Val	RPD	<b>RPD ABS Val</b>
Sediment	LV	Clay/clay loam, salt	Number	4	4	4	4	12	12	4	4
		crust (iron and other	Minimum	35.9%	35.9%	-108.4%	9.9%	-31.8%	1.2%	-97.3%	59.2%
		precipitate)	Maximum	49.6%	49.6%	32.4%	108.4%	73.9%	73.9%	-59.2%	97.3%
			Mean	42.2%	42.2%	-15.1%	44.0%	16.5%	27.5%	-81.2%	81.2%
			Median	41.6%	41.6%	7.8%	28.9%	17.3%	28.4%	-84.2%	84.2%
Sediment	liment RF	Silty fine sand (tailings)	Number	4	4	4	4	12	12	5	5
			Minimum	-43.8%	43.8%	-2.6%	0.1%	-88.4%	25.7%	-37.1%	11.8%
			Maximum	63.0%	63.0%	98.9%	98.9%	-25.7%	88.4%	-11.8%	37.1%
			Mean	34.3%	56.2%	24.8%	26.1%	-44.8%	44.8%	-22.0%	22.0%
		Median	59.0%	59.0%	1.4%	2.7%	-41.8%	41.8%	-21.0%	21.0%	
Soil	SB	Coarse sand and gravel	Number	6	6			5	5	1	1
		(ore and waste rock)	Minimum	-68.5%	2.6%			-163.0%	39.2%	-16.3%	16.3%
			Maximum	43.9%	68.5%			-39.2%	163.0%	-16.3%	16.3%
		Mean	9.7%	32.5%			-76.5%	76.5%	-16.3%	16.3%	
			Median	23.0%	32.9%			-48.6%	48.6%	-16.3%	16.3%
Sediment	TL	Silt and clay (slag-	Number	3	3	3	3	2	2	2	2
		enriched)	Minimum	-45.7%	27.0%	10.2%	10.2%	-70.4%	1.9%	-34.3%	14.0%
			Maximum	-27.0%	45.7%	18.8%	18.8%	1.9%	70.4%	-14.0%	34.3%
			Mean	-39.4%	39.4%	13.5%	13.5%	-34.3%	36.2%	-24.1%	24.1%
			Median	-45.5%	45.5%	11.5%	11.5%	-34.3%	36.2%	-24.1%	24.1%
Soil	WS	Coarse sand and gravel	Number	3	3			7	7	3	3
		(roaster slag)	Minimum	-105.4%	37.6%			-66.4%	8.4%	-29.4%	12.2%
			Maximum	-37.6%	105.4%			47.4%	66.4%	-12.2%	29.4%
			Mean	-63.3%	63.3%			-14.3%	34.2%	-18.7%	18.7%
			Median	-47.0%	47.0%			-18.6%	22.2%	-14.7%	14.7%
	All	Samples	Number	27	27	14	14	46	46	26	26
			Minimum	-105.4%	2.6%	-2.6%	0.1%	-163.0%	1.9%	-44.3%	0.2%
			Maximum	77.4%	105.4%	154.4%	154.4%	53.8%	163.0%	25.0%	44.3%
			Mean	11.3%	44.9%	23.0%	23.5%	-41.4%	49.1%	-15.9%	18.3%
			Median	14.9%	45.5%	6.2%	6.2%	-41.0%	44.2%	-15.5%	16.3%

MatrixSiteSedimentLVSedimentRFSoilSB	Matrix Description Clay/clay loam, salt crust (iron and other precipitate) Silty fine sand (tailings)	Statistic Number Minimum Maximum Mean Median Number	Reference           RPD         3           70.9%         121.0%           93.5%         88.5%	Ce Laboratory           RPD ABS           Val           3           70.9%           121.0%           02.5%	<b>RPD</b> 11 -24.1%	e Laboratory RPD ABS Val 11 3.7%	<b>RPD</b> 5	e Laboratory RPD ABS Val 5	Reference RPD 4	ce Laboratory RPD ABS Val
MatrixSiteSedimentLVSedimentRFSoilSB	Description Clay/clay loam, salt crust (iron and other precipitate)	Number Minimum Maximum Mean Median	3 70.9% 121.0% 93.5%	Val 3 70.9% 121.0%	11 -24.1%	<b>Val</b> 11	5	Val		
Sediment LV Sediment RF Soil SB	Clay/clay loam, salt crust (iron and other precipitate)	Number Minimum Maximum Mean Median	3 70.9% 121.0% 93.5%	3 70.9% 121.0%	11 -24.1%	11	5			
Sediment RF	crust (iron and other precipitate)	Minimum Maximum Mean Median	70.9% 121.0% 93.5%	70.9% 121.0%	-24.1%			5	1	1 . 1
Sediment RF Soil SB	precipitate)	Maximum Mean Median	121.0% 93.5%	121.0%		3 7%			-	4
Sediment RF Soil SB		Mean Median	93.5%			5.770	-97.4%	6.5%	-68.3%	3.6%
Soil SB	Silty fine sand (tailings)	Median		02 50/	39.1%	39.1%	6.5%	97.4%	3.6%	68.3%
Soil SB	Silty fine sand (tailings)		88 5%	93.5%	8.9%	18.7%	-68.8%	71.3%	-27.4%	29.1%
Soil SB	Silty fine sand (tailings)	Number	00.570	88.5%	11.5%	14.3%	-84.1%	84.1%	-22.4%	22.4%
			8	8	13	13	13	13	10	10
		Minimum	11.7%	11.7%	-101.8%	1.2%	-5.3%	5.0%	-76.3%	42.5%
		Maximum	37.8%	37.8%	70.7%	101.8%	28.8%	28.8%	-42.5%	76.3%
		Mean	29.8%	29.8%	-11.0%	28.5%	13.9%	15.5%	-58.6%	58.6%
		Median	32.4%	32.4%	-8.3%	11.6%	16.2%	16.2%	-59.0%	59.0%
	Coarse sand and gravel	Number	10	10	4	4	12	12		
	(ore and waste rock)	Minimum	-34.8%	9.0%	-108.7%	22.8%	3.2%	3.2%		
		Maximum	87.2%	87.2%	53.0%	108.7%	44.0%	44.0%		
		Mean	21.8%	28.8%	-33.0%	70.9%	29.7%	29.7%		
		Median	19.5%	22.9%	-38.1%	76.1%	33.7%	33.7%		
	Silt and clay (slag-	Number	1	1	7	7	7	7	4	4
	enriched)	Minimum	15.6%	15.6%	-48.7%	18.2%	-34.7%	1.9%	-121.2%	66.4%
		Maximum	15.6%	15.6%	-18.2%	48.7%	15.7%	34.7%	-66.4%	121.2%
		Mean	15.6%	15.6%	-32.1%	32.1%	-9.0%	18.2%	-95.1%	95.1%
		Median	15.6%	15.6%	-28.6%	28.6%	-1.9%	15.7%	-96.4%	96.4%
	Coarse sand and gravel	Number	6	6	6	6	7	7	7	7
	(roaster slag)	Minimum	-41.9%	2.3%	-105.3%	5.4%	-16.4%	16.4%	-88.5%	22.3%
		Maximum	80.4%	80.4%	9.2%	105.3%	32.4%	32.4%	-22.3%	88.5%
		Mean	28.5%	42.4%	-28.3%	33.4%	19.3%	24.0%	-63.3%	63.3%
		Median	30.1%	38.7%	-7.6%	9.5%	23.8%	23.8%	-70.5%	70.5%
All	Samples	Number	41	41	48	48	70	70	42	42
		Minimum	-41.9%	2.3%	-111.1%	1.2%	-34.7%	0.3%	-121.2%	0.5%
		Maximum	87.2%	87.2%	70.7%	111.1%	73.9%	73.9%	10.9%	121.2%
		Mean	25.3%	30.6%	-23.3%	37.3%	14.4%	21.1%	-52.8%	54.0%
		Median	28.1%	30.7%	-11.4%	25.1%				

				Μ	ercury	N	ickel	Sel	enium	Silver	
				Referenc	e Laboratory	Reference	e Laboratory	Reference	e Laboratory	Referen	ce Laboratory
		Matrix			RPD ABS		RPD ABS		RPD ABS		
Matrix	Site	Description	Statistic	RPD	Val	RPD	Val	RPD	Val	RPD	<b>RPD ABS Val</b>
Sediment	LV	Clay/clay loam, salt	Number	3	3	11	11	5	5	4	4
		crust (iron and other	Minimum	70.9%	70.9%	-24.1%	3.7%	-97.4%	6.5%	-68.3%	3.6%
	precipitate)	Maximum	121.0%	121.0%	39.1%	39.1%	6.5%	97.4%	3.6%	68.3%	
			Mean	93.5%	93.5%	8.9%	18.7%	-68.8%	71.3%	-27.4%	29.1%
			Median	88.5%	88.5%	11.5%	14.3%	-84.1%	84.1%	-22.4%	22.4%
Sediment	RF	Silty fine sand (tailings)	Number	5	5	13	13	3	3	4	4
			Minimum	79.3%	79.3%	-2.9%	1.2%	-98.5%	56.8%	-93.9%	15.5%
			Maximum	106.7%	106.7%	34.5%	34.5%	-56.8%	98.5%	-15.5%	93.9%
			Mean	89.7%	89.7%	17.7%	18.1%	-82.3%	82.3%	-53.6%	53.6%
		Median	89.8%	89.8%	19.9%	19.9%	-91.7%	91.7%	-52.6%	52.6%	
Soil	SB	Coarse sand and gravel	Number	10	10	11	11	3	3	1	1
	(ore and waste rock)	Minimum	-133.5%	22.5%	-7.5%	7.5%	-86.5%	3.7%	-75.8%	75.8%	
		Maximum	140.5%	140.5%	35.4%	35.4%	3.7%	86.5%	-75.8%	75.8%	
			Mean	64.0%	90.7%	19.0%	20.4%	-56.4%	58.8%	-75.8%	75.8%
			Median	92.4%	96.2%	17.5%	17.5%	-86.3%	86.3%	-75.8%	75.8%
Sediment	TL	Silt and clay (slag-	Number	3	3	6	6	4	4	4	4
		enriched)	Minimum	72.3%	72.3%	-73.9%	3.6%	-110.0%	20.0%	-57.1%	32.1%
			Maximum	103.7%	103.7%	-3.6%	73.9%	-20.0%	110.0%	-32.1%	57.1%
			Mean	85.0%	85.0%	-35.5%	35.5%	-58.0%	58.0%	-45.7%	45.7%
			Median	79.1%	79.1%	-36.4%	36.4%	-51.1%	51.1%	-46.8%	46.8%
Soil	WS	Coarse sand and gravel	Number			7	7	1	1	4	4
		(roaster slag)	Minimum			-37.3%	4.0%	-106.0%	106.0%	-20.2%	10.5%
			Maximum			23.2%	37.3%	-106.0%	106.0%	-10.5%	20.2%
			Mean			-4.4%	18.6%	-106.0%	106.0%	-14.8%	14.8%
			Median			4.0%	16.3%	-106.0%	106.0%	-14.3%	14.3%
	All	Samples	Number	24	24	63	63	21	21	24	24
			Minimum	-133.5%	22.5%	-73.9%	1.2%	-110.0%	0.9%	-106.7%	0.0%
	1		Maximum	140.5%	140.5%	39.1%	73.9%	6.5%	110.0%	3.6%	106.7%
			Mean	74.8%	88.2%	6.9%	20.6%	-65.0%	66.1%	-33.4%	33.7%
			Median	90.1%	90.7%	13.6%	17.5%	-75.7%	75.7%	-18.3%	18.3%

				Va	nadium		Zinc
				Reference	ce Laboratory	Reference	e Laboratory
		Matrix			RPD ABS		RPD ABS
Matrix	Site	Description	Statistic	RPD	Val	RPD	Val
Sediment	LV	Clay/clay loam, salt	Number	5	5	8	8
		crust (iron and other	Minimum	47.6%	47.6%	-48.4%	2.1%
		precipitate)	Maximum	83.3%	83.3%	48.8%	48.8%
			Mean	65.2%	65.2%	-19.0%	31.2%
			Median	64.2%	64.2%	-27.4%	34.2%
Sediment RF S		Silty fine sand (tailings)	Number	3	3	13	13
			Minimum	74.2%	74.2%	-63.4%	17.6%
			Maximum	93.1%	93.1%	20.3%	63.4%
			Mean	84.8%	84.8%	-33.2%	39.0%
			Median	87.0%	87.0%	-38.4%	38.4%
Soil	SB	Coarse sand and gravel	Number	3	3	11	11
		(ore and waste rock)	Minimum	-35.4%	13.2%	-22.8%	0.1%
			Maximum	73.8%	73.8%	18.9%	22.8%
			Mean	17.2%	40.8%	-3.8%	9.3%
			Median	13.2%	35.4%	-0.3%	7.8%
Sediment	TL	Silt and clay (slag-	Number			7	7
		enriched)	Minimum			-31.3%	2.3%
			Maximum			70.9%	70.9%
			Mean			23.3%	35.2%
			Median			28.5%	31.3%
Soil	WS	Coarse sand and gravel	Number	2	2	7	7
		(roaster slag)	Minimum	50.4%	50.4%	-51.3%	11.8%
			Maximum	83.4%	83.4%	-11.8%	51.3%
			Mean	66.9%	66.9%	-32.5%	32.5%
			Median	66.9%	66.9%	-24.9%	24.9%
	All	Samples	Number	16	16	61	61
			Minimum	-35.4%	13.2%	-80.6%	0.1%
			Maximum	129.5%	129.5%	105.7%	105.7%
			Mean	65.2%	69.6%	-19.0%	33.0%
			Median	74.0%	74.0%	-22.8%	32.5%

 Table E-5. Evaluation of the Effects of Soil Type on RPDs (Accuracy) of Target Elements (Continued)

Site Abbreviations:

AS	Alton Steel Mill
BN	Burlington Northern Railroad/ASARCO East
CN	Naval Surface Warfare Center, Crane Division
KP	KARS Park – Kennedy Space Center
LV	Leviathan Mine/Aspen Creek
RF	Ramsey Flats – Silver Bow Creek
SB	Sulphur Bank Mercury Mine
TL	Torch Lake Superfund Site

WS Wickes Smelter Site

Other Notes:

	No samples reported by the reference laboratory in this concentration range.
Number	Number of demonstration samples evaluated.
RPD	Relative percent difference (raw value).

RPD ABS Val Relative percent difference (absolute value).