

SDG NARRATIVE

USEPA SDG # MYDA93 CASE # 51772 CONTRACT # 68HERH20D0011 SOW# SFAM01.1 LAB NAME: Alliance Technical Group, LLC LAB CODE: ACE LAB ORDER ID # P4299 MODIFIED ANALYSIS #3225.1, 3226.1

A. Number of Samples and Date of Receipt

20 Soil samples were delivered to the laboratory intact on 10/04/2024.

B. Parameters

Test requested for Metals CLP FULL = Aluminum, Antimony, Arsenic, Barium, Beryllium, Cadmium, Calcium, Chromium, Cobalt, Copper, Iron, Lead, Magnesium, Manganese, Nickel, Potassium, Selenium, Silver, Sodium, Thallium, Vanadium, Zinc.

Test requested for Metals CLP MS FULL = Antimony, Arsenic, Barium, Beryllium, Cadmium, Chromium, Cobalt, Copper, Lead, Nickel, Selenium, Silver, Thallium, Vanadium, Zinc.

C. Cooler Temp

Indicator Bottle: Presence/Absence

Cooler: 21.6°C

D. Detail Documentation (related to Sample Handling Shipping, Analytical Problem, Temp of Cooler etc):

Issue 1 : A "P" or "M" prefix was listed at the beginning of a CLP sample ID.

E. Corrective Action taken for above:

Resolution 1 : To maintain COC integrity, ASB requests no changes to the Sample IDs. The laboratory will note the issue in the SDG Narrative and proceed with the analysis of the samples.

F. Analytical Techniques:

All analyses were based on CLP Methodology by method SFAM01.1.



Inter Element correction factors (IECs) are determined annually and correction factor are applied during ICP-AES analysis.

G. Calculation:

Calculation for ICP-AES Soil Sample:

Conversion of Results from mg/L or ppm to mg/kg (Dry Weight Basis):

Concentration (mg/kg) = $C \times \frac{Vf}{W \times S} \times DF$

Where,

C = Instrument value in ppm (The average of all replicate exposures)
 Vf = Final digestion volume (mL)
 W = Initial aliquot amount (g) (Sample amount taken in prep)
 S = % Solids / 100 (Fraction of Percent Solids)
 DF = Dilution Factor

Example Calculation For Sample MYDA93 For Antimony:

If C = 0.0105351ppm Vf = 100 ml W = 1.14g S = 0.982(98.2/100)DF = 1

Concentration (mg/kg) = $0.0105351 \text{ x} \frac{100}{1.14 \text{ x} 0.982} \text{ x} 1$

= 1.88214 mg/kg

= 1.9 mg/kg (Reported Result with Signification)

Calculation for ICP-MS Soil Sample:

Conversion of Results from $\mu g / L$ or ppb to mg/kg :

Concentration (mg/kg) =
$$C \times Vf = Vf + 1000$$

W x S

Where,

C = Instrument value in ppb (The average of all replicate integrations)

Vf = Final digestion volume (mL)

W = Initial aliquot amount (g) (Fraction of Sample amount taken in prep)



S = % Solids / 100 (Fraction of Percent Solids) DF = Dilution Factor

Example Calculation For Sample MYDA93 For Antimony :

If C = 2.11 ppb Vf = 500 ml W = 1.14 g S = 0.982(98.2/100) DF = 1 Concentration (mg/kg) = $2.11 \text{ x} \underbrace{500}_{1.14 \text{ x} 0.982} \text{ x } 1 / 1000$ = 0.9424 mg/kg

= 0.94 mg/kg (Reported Result with Signification)

H. QA/QC

Calibrations met requirements. Interference check met requirements. Blank analyses did not indicate any presence of contamination. Laboratory Control sample was within control limits. AES Spike sample did meet requirements except for Antimony, Arsenic, Barium, Selenium. . Spike sample (MYDAB1SRE)did meet requirements except for Arsenic, Silver. Duplicate sample did meet requirements except for Barium. Serial Dilution did meet requirements.

Collision cell is being used to remove potential interferences. The analytes Na, Mg, Al, K, V, Cr, Mn, Fe, Co, Ni, Cu, Zn, As are being analyzed with collision cell and analytes Be, B, Ca, Ti, Se, Sr, Zr, Mo, Ag, Cd, Sn, Sb, Ba, Tl, Pb, U are being analyzed with Non-Collision Cell. Helium gas is used for the Collision Cell analysis.

Target Analyte	Associated Internal Standard
Antimony	159Tb
Arsenic	89Y
Barium	159Tb
Beryllium	6Li
Cadmium	159Tb
Chromium	45Sc

Internal Standard Association for ICP-MS analysis.



Cobalt	45Sc
Copper	45Sc
Lead	209Bi
Nickel	45Sc
Selenium	89Y
Silver	159Tb
Thallium	209Bi
Vanadium	45Sc
Zinc	45Sc

I certify that the data package is in compliance with the terms and conditions of the contract both technically and for completeness, for other than the conditions detailed above. Release of the data contained in this hard copy data package has been authorized by the Laboratory Director or his designee, as verified by the following signature.

Signature_____ Nam

Name: Nimisha Pandya

Date _____

Title: Document Control Officer

	MA: 3225.0	Title: ICP-MS with Modified Preparation Method and Analysis of Soils with Additional Laboratory QC
Method Source: SFAM01.1	Method: ICP-MS	
Matrix: Soil/Sediment		
Summary of Modification		
with additional modified LCS and Unless specifically modified by t	d Matrix Spikes and and his modification, all and	uples by EPA Draft Method 3050C (see below) alyze for the scheduled target analytes by ICP-MS. alyses, Quality Control (QC), and reporting nt EPA agreement remain unchanged and in full
I. Analyte Modifications		Not applicable 🔀
II. Calibration and QC Requirem	nents	Not applicable
 200.8) to report the resumption of the resumption of the resumption of the resumption of the repare and analyze and recovery limits do NOT and the repare a Matrix Spike set of the repare a Matrix Spike set of the repare and the repare set of the repare se	Ilts for these analyses. thod 3050C. additional Laboratory (apply to this LCS and n piked at three times th additional Matrix Spike	mined for routine soil analyses (i.e., Method The Laboratory is NOT required to perform an Control Sample (LCS) spiked at the CRQL. Percent o corrective actions are required. The levels specified in the SOW.
 Post-Digestion Spike req Post-Digestion Spike cor 	uirements apply to the	e 5x Matrix Spike only.
Post-Digestion Spike req	uirements apply to the rective actions apply to	e 5x Matrix Spike only.

IV. Special Reporting Requirements

The Laboratory shall:

- Ensure the SDG Narrative is updated as stated in the SOW, including any technical and administrative problems encountered and the resolution or corrective actions taken. These problems may include interference problems encountered during analysis, dilutions, re-analyses and/or re-preparations performed, and problems with the analysis of samples. Also include a discussion of any SOW Modified Analyses, including a copy of the approved modification form with the SDG Narrative.
- Initial analysis data are reported with a dilution factor of 1.0 and a final volume of 500 mL, per the SOW.
- Report the additional LCS as "LCSD" in the raw data and in the EDD with QCType "Laboratory_Control_Sample_Duplicate".
- Report the additional Matrix Spike with an "SRE" suffix in the raw data and EDD.
- Report any Post-Digestion Spike of the additional 5x Matrix Spike with an "ARE" suffix.

	09/04/2024	MA: 3226.0	Title: ICP-AES with Modified Preparation Method and Analysis of Soils with Additional Laboratory QC			
Metho	od Source: SFAM01.1	Method: ICP-AES				
Matrix	c: Soil/Sediment	1				
Summ	ary of Modification					
with a AES. U require	dditional modified LCS and Inless specifically modified	Matrix Spikes and ana by this modification, a	ples by EPA Draft Method 3050C (see below) alyze for the scheduled target analytes by ICP- III analyses, Quality Control (QC), and reporting at EPA agreement remain unchanged and in full			
I. Ana	alyte Modifications		Not applicable 🔀			
II. Cal	ibration and QC Requireme	ents	Not applicable			
•	Recovery limits do NOT a	pply to this LCS and no viked at two times the virements apply to the	•			
	T USE DIBESTION SPIKE CON		n Sh			
III. Pre	paration and Method Mod					
	paration and Method Mod	lifications	Not applicable			
	 boratory shall: Prepare and analyze the some sample thore Mix sample thore Add 10 mL 1:1 HM minutes. Add 5 mL concendigestion comple Concentrate sam Cool sample, add 1 mL aliquots of 3 Dilute to 100 mL 	lifications sample by EPA Draft Noughly and transfer 1.0 NO ₃ and 5 mL 1:1 HCl, trated HNO ₃ and reflute. ple to 5 mL or reflux v 2mL water and 3 mL 30% H ₂ O ₂ until efferve with water, centrifuge s can also be used for for both ICP-AES and	Not applicable Method 3050C as follows: D0 – 1.50 g to a digestion vessel. heat the sample at 95°C (±3°C) and reflux 10 -15 ax for 30 minutes at 95°C (±3°C), repeat until without boiling for 2 hours at 95°C (±3°C). 30% H ₂ O ₂ . Heat at 95°C (±3°C) and add additiona escence is minimal. e or filter as necessary prior to analysis. ICP-MS analysis. Separate Matrix Spikes and LCS			

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- Initial analysis data are reported with a dilution factor of 2.0 and a final volume of 100 mL, per the SOW.
- Report the additional LCS as "LCSD" in the raw data and in the EDD with QCType "Laboratory_Control_Sample_Duplicate".
- Ensure that up-to-date Interelement Correction Factors (IECs) are provided with the data package.

v	Element, Vavelength and Order	Use?	# IECs	IEC	k1	K2	Calc-in-fit
A	s 189.042 {479}	\boxtimes	1	Fe	-0.000064	0.000000	No
TI	190.856 {477}		5	Мо	-0.002450	0.000000	No
Ī				Co	0.002248	0.000000	No
			····	Ti	-0.000500	0.000000	No
				Mn	0.000370	0.000000	No
1				V	-0.012340	0.000000	No
Pt	220.353 {453}	M	6	Мо	-0.001480	0.000000	No
1				Al	-0.000075	0.000000	No
				Cu	0.001400	0.000000	No
1		••••••		Fe	0.000030	0.000000	No
1				Mn	0.000340	0.000000	No
				Ni	0.000630	0.000000	No
Se	196.090 {472}		3	Fe	-0.000308	0.000000	No
	1001000 (112)		1	Mn	0.000470	0.000000	No
			•	Co	-0.000630	0.000000	No
Sh	206.833 {463}	\boxtimes	4	Cr	0.010700	0.000000	No
	200.000 (100)			V	-0.001168	0.000000	No
				Mo	-0.002850	0.000000	No
				Ni	-0.002850		
Δ1	396.152 { 85}		4	å		0.000000	No
	493.409 { 68}		Nono	Мо	0.037230	0.000000	No
	234.861 {144}		None	Ma	0.000000	0.000000	
De	234.001 {144}	X	3	Mo	-0.000320	0.000000	No
				Fe	0.000010	0.000000	No
	214 420 (457)	57		Mn	-0.000047	0.000000	No
**********	214.438 {457}	<u> </u>	1	Fe	0.000040	0.000000	No
*****	373.690 { 90}		None				
****	267.716 {126}	<u>¤</u>	1	Mn	0.000160	0.000000	No
Co	228.616 {448}		2	Ti	0.001840	0.000000	No
				Мо	-0.001230	0.000000	No
Cu	324.754 {104}		4	Co	-0.000796	0.000000	No
ļ				Fe	-0.000100	0.000000	No
				Mn	0.000345	0.000000	No
				Ni	0.000895	0.000000	No
	259.837 {130}		None				
Mn	257.610 {131}		1	Ni	0.000897	0.000000	No
	279.079 {121}		None		[
Ni 2	31.604 {446}		None		I		1
Ag	328.068 {103}	\boxtimes	3	Fe	-0.000100	0.000000	No
				Mn	0.000146	0.000000	No
1				V	-0.000889	0.000000	No
Nat	318.326 { 41}		None			1	
V 29	92.402 {115}		2	Мо	-0.008480	0.000000	No
Î				Cr	-0.002220	0.000000	No
Zn 2	06.200 {464}		None				
	13.856 (158)		1	Ni	0.007280	0.000000	No
·	9.896 { 44 }		None			1	1
	7.495 {490}		2	Ni	0.001640	0.000000	No
1			_	Cu	-0.012530	0.000000	No
B 24	9.678 {135}		3	Co	0.002880	0.000000	No
17				V	-0.002000	0.000000	No
1			<u>i</u>	Fe	-0.002000	0.000000	No
Mo	202.030 {467}		None	16	-0.001300	0.000000	INU
	2.034 {485}		None	Mo	0.000000	0.000000	No
10 10			2	Mo	-0.008000	0.000000	No
1	1			Mn	0.002700	0.000000	No

	Element, Wavelength and Order	Use?	# IECs	IEC	k1	k2	Calc-in-fit?
	Si 251.611 {134}	\boxtimes	2	Мо	0.010520	0.000000	No
				Ti	0.005650	0.000000	No
	Sn 189.989 {478}		None		<u> </u>		
	Ti 336.121 {100}		1	Ni	-0.001000	0.000000	No
	Li 670.784 { 50}		None			1	110
	Y 224.306 {450}*		None				
I	Y 360.073 { 94}*		None				
Î	Y 371.030 { 91}*		None				
Ī	Y 224.306 {150}*		None				<u> </u>
	In 230.606 {446}*		None				
	Sr 407.771 { 83}		None				[[

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