

Rapid Methods

New Matrices

Improved Separations



eichrom 1955 University Lane Lisle, IL 60532, USA +1 (630)963-0320 www.eichrom.com

Eichrom Technologies Application Notes

AN-1700-TOC

Summary of Application Notes: The Application Notes presented in this book represent two page summaries of works published by respected scientist in the radiochemical community. These notes represent an extension to the basic methods available from the Eichrom Published Methods. The Application Notes contain some significant changes to the chemistry used in preparation and/or separation than the Eichrom methods. Application Notes will also have methodologies for different matrices or significant changes in sample size. These include new analytes or combination of analytes.

Application Notes

Number	Title
AN-1401	Rapid Determination of ²²⁶ Ra in Emergency Urine and Water
AN-1402	Rapid Determination of Sr in Emergency Milk Samples
AN-1403	Rapid Determination of Sr in 50g Soil Samples
AN-1404	Rapid Determination of Sr in 1-2 Liter Seawater Samples
AN-1405	Rapid Determination of Sr in Vegetation Samples
AN-1406	Rapid Determination of Actinides in Vegetation Samples
AN-1407	Rapid Determination of Sr in Animal Tissue Samples
AN-1408	Rapid Determination of Actinides in Animal Tissue Samples
AN-1409	Rapid Determination of Sr in Building Materials
AN-1410	Rapid Determination of Sr in Emergency Urine Samples
AN-1411	Rapid Determination of Sr in Emergency Water Samples
AN-1412	Rapid Determination of Actinides in Emergency Urine Samples
AN-1413	Rapid Determination of Actinides in Emergency Water Samples
AN-1414	Rapid Determination of ⁹⁰ Sr in Up to 40 Liter Seawater Samples
AN-1415	Rapid Determination of ²¹⁰ Po in Water Samples
AN-1416	Rapid Determination of Actinides and ²¹⁰ Po in Water
AN-1417	Rapid Determination of ^{226/228} Ra in Water Samples
AN-1418	Rapid Determination of ²²⁶ Ra in Water Samples
AN-1419	Rapid Determination of ²²⁶ Ra in Concrete and Brick
AN-1420	Rapid Determination of ²²⁶ Ra in Glass Fiber Air Filters
AN-1421	Rapid Determination of ²²⁶ Ra in 1g Soil Samples
AN-1422	Rapid Determination of ²²⁶ Ra in 5g Vegetation Samples
AN-1423	Rapid Determination of Pu, Np, and U in 1-8L Seawater Samples
AN-1424	Rapid Determination of Pu, Am and Cm in 80L Seawater Samples
AN-1425	Rapid Determination of Actinides in 10g Emergency Food Samples
AN-1426	Rapid Determination of Actinides in 100g Emergency Food Samples
AN-1427	Rapid Determination of Plutonium in Large Rice Samples
AN-1428	Rapid Determination of Actinides in Fecal Samples
AN-1429	Rapid Determination of Actinides in Asphalt samples
AN-1430	Rapid Determination of Actinides in Emergency Soil Samples
AN-1431	Rapid Determination of Determination of Actinides in 100g Soil Samples
AN-1432	Rapid Determination of Actinides in 1g Concrete and Brick Samples
AN-1433	Rapid Determination of Actinides in Emergency Air Filter Samples
AN-1434	Rapid Determination of Sr in Emergency Air Filter Samples
AN-1435	Rapid Determination of Np/Pu in 20-50g Soil Samples
AN-1436	Rapid Determination of Np/Pu in 20-75g Soil Samples (ICP-MS)
AN-1437	Rapid Determination of Actinides in Urine by ICP-MS + Alpha Spec
AN-1438	Rapid Determination of Np/Pu in Water Samples by ICP-MS

eichrom 1955 University Lane Lisle, IL 60532, USA +1 (630)963-0320 www.eichrom.com

Eichrom Technologies Application Notes

AN-1700-TOC

Number	Title
AN-1601	Method for ²²⁷ Ac in Geological Samples
AN-1602	Method for ²²⁷ Ac in Water Samples
AN-1603	Rapid Method for Actinides in Limestone and Marble
AN-1604	Rapid Method for ^{89/90} Sr in Limestone and Marble
AN-1605	Rapid Method for ^{89/90} Sr in Large Concrete Samples
AN-1606	Rapid Method for ⁹⁰ Sr in Large Concrete Samples
AN-1607	Rapid Method for Pu, Np, Am in Large Soil Samples
AN-1608	Rapid Method for U and Th in soil
AN-1609	Rapid Method for ³ H in water
AN-1610	Rapid Method for Ni-59/63 in Water
AN-1611	Rapid Method for Fe-55 in Water (TEVA)
AN-1612	Rapid Method for Fe-55 in Water (TRU)
AN-1613	Ga-68 Generator
AN-1614	Ac-225 Generator
AN-1615	Y-90 Generator
AN-1616a	Po-210/Bi-210 Generator
AN-1616b	Po-210 Generator
AN-1617	Th-227 and Ra-223 Generator
AN-1618	Th-228 and Th-231 Generators
AN-1619	Ra-224, Pb-212 Generators
AN-1620	Np-239 Generator
AN-1621	Th-234 Generator
AN-1622	Zr-89 Separation
AN-1623	Y-86 Separation
AN-1624	Options for Sr-89/90 Discrimination
AN-1701	CI– and I– using CL Resin
AN-1702	Converting Column Methods to Cartridges
AN-1703	Slurry Packing 2mL Eichrom Columns
AN-1801	Rapid Determination of 89/90Sr in Steel Samples
AN-1802	Rapid Determination of Pu in Steel Samples
AN-1803	Rapid Determination of 226Ra in Steel Samples
AN-1804	Rapid Determination of Pu/Np and Am/Cm in Granite Samples
AN-1805	Alpha Spectrometry Source Preparation: Rare Earth Fluoride Microprecipitation
AN-1806	Actinide/Rare Earth Separation (TEVA-SCN)
AN-1807	Alpha Spectrometry Source Preparation: Cerium Hydroxide Microprecipitation
AN-1808	Zirconium Separation on ZR Resin
AN-1809	Copper Separation on CU Resin
AN-1810	Cs Separation on AMP-PAN and KNiFC-PAN Resins
AN-1811	Ce Separation from Rare Earth Nitrate Solutions
AN-1812	Fe Separation from Rare Earth Chloride Solutions
AN-1813	Tc Separation on WBEC Resin

Rapid Determination of ²²⁶Ra in Emergency Urine and Water

AN-1401-10

Summary of Method ²²⁶Ra is isolated from 100mL urine samples or up to 1 liter water samples and measured by alpha spectrometry as described by Maxwell, et al.¹ Radium is precipitated from samples with calcium phosphate. The calcium phosphate precipitate is dissolved in hydrochloric acid, and cation exchange chromatography is used to purify radium and barium from matrix ions. Eichrom DGA Resin is used to remove other alpha emitting nuclides from radium. Samples are prepared for radium measurement by alpha spectrometry via barium sulfate microprecipitation onto Eichrom Resolve[®] Filters. Sample preparation, including alpha spectrometry source preparation, for batches of 12 samples can be completed by a single operator in as little as 3-4 hours, with >90% yield of Radium. Yields can be traced with ¹³³Ba by gamma spectrometry or ²²⁵Ra(²²⁹Th) by alpha spectrometry. If tracing with ²²⁵Ra, at least 8 hours of ingrowth time are required for the alpha emitting ²¹⁷At daughter of ²²⁵Ra prior to alpha spectrometry measurements.

Reagents

Cation Exchange Resin (Eichrom C8-B500-F-H) DGA Resin, Normal 2mL Cartridges (Eichrom DN-R50-S) Ammonium Hydroxide (Listed as 28% NH₃ or 56% NH₄OH) Nitric Acid (70%) Hydrochloric Acid (37%) Deionized Water Hydrogen Peroxide (30%) ¹³³Ba or ²²⁵Ra(²²⁹Th) Tracer*

1.25M Ca(NO₃)₂ 3.2M (NH₄)₂HPO₄ Barium Carrier (1mg/mL) Isopropyl Alcohol Ammonium Sulfate Denatured Ethanol

 *133Ba allows immediate counting.
 ²²⁵Ra(²²⁹Th) requires >8hrs ingrowth before alpha meas.
 Ba/Ra recoveries can differ by up to 10% in difficult matrices.

Equipment

Plastic Chromatography Column (Eichrom AC-50E-5M) Column Extension Funnel (Eichrom AC-20X-20M) Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filter in Disposable Funnel (Eichrom RF-DF-25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Hotplate 150mL Glass beakers Vacuum Pump Heat Lamp Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Gamma Spectrometry System (if ¹³³Ba tracer used)

Figure 1. Sample Preparation

100 mL urine or 1L water. Adjust to pH2 with HNO₃. Add tracer ¹³³Ba or ²²⁵Ra (²²⁹Th) +1 mL 1.25M Ca(NO₃)₃ +3mL 3.2 M (NH₄)₂HPO₄.**

**A calcium phosphate ppt. was chosen to minimize reagent background. A CaCO₃ ppt (AN1418) can help minimize ²²⁹Th in the final Ra fraction, when spiking directly with ²²⁹Th for ²²⁵Ra tracing.

Adjust to pH 10 with NH₄OH.

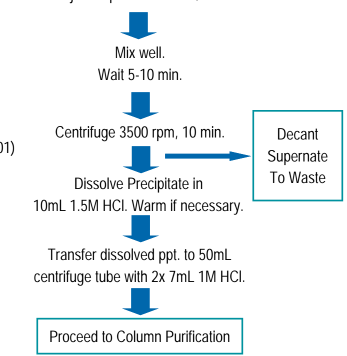
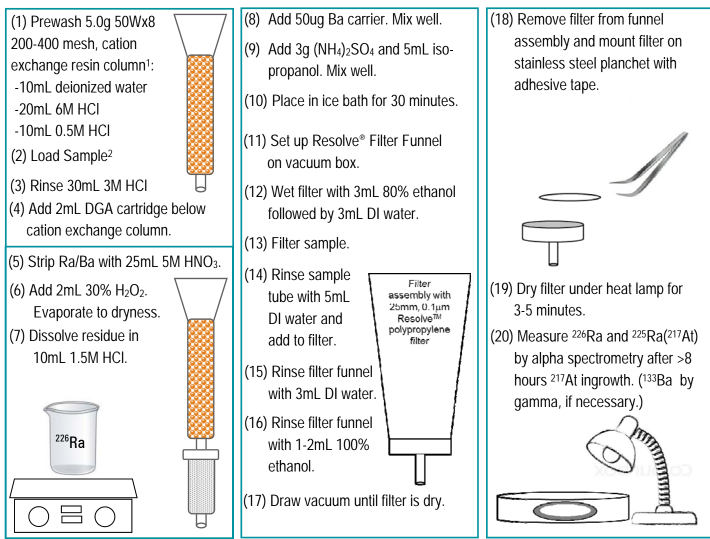


Figure 2. Column Purification and Alpha Source Preparation



¹If using ¹³³Ba tracer, 3.0g of cation exchange resin and proportionally smaller rinse volumes may be used. ²If tracing with ²²⁹Th, a 20mL 1M HCI-1M H₃PO₄ rinse following the sample load can improve purity of final ²²⁶Ra fraction.

Tracer					²²⁶ Ra Me	asu	red Value	
¹³³ Ba % Recovery			overy	²²⁶ Ra Reference	(mB	q/sa	mple)	
Replicates	Average		SD	Value (mBq/sample)	Average		SD	% Bias
6	93	<u>+</u>	3	73.7	76.5	<u>+</u>	4.7	3.9
6	98	<u>+</u>	3	18.4	17.9	<u>+</u>	0.8	-2.7
6	92	<u>+</u>	5	Blank*	0.15	<u>+</u>	0.12	

*Calculated MDA 15 mBq/L (4 hr count, 100 mL sample) *Calculated MDA 5 mBq/L (16 hr count, 100 mL sample)

References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchinson, Robin C. Utsey and Daniel R. McAlister, "Rapid Determination of ²²⁶Ra in Emergency Urine Samples," *J. Radioanal. Nucl. Chem.*, 300(3), 1159-1166 (2014).

Rapid Determination of Sr in Emergency Milk Samples

AN-1402-10

Summary of Method Strontium is separated and concentrated from 100mL milk samples using a calcium phosphate precipitation. The precipitate is dissolved with nitric acid and centrifuged to remove residual protein and fat. The supernate, containing Sr, is wet ashed with HNO₃-H₂O₂ and then heated in a muffle furnace at 550°C for 30-60 minutes to destroy any residual organic matter. The muffled residue is wet ashed again with HNO₃-H₂O₂ and dissolved in HNO₃-Al(NO₃)₃. Strontium is separated from matrix impurities and potentially interfering radionuclides in the sample using stacked 2mL and 1mL cartridges of Eichrom Sr Resin. Radiostrontium is measured on a low background gas flow proportional counter or liquid scintillation counter. Chemical yield of strontium is determined by gravimetric recovery of stable strontium or by ICP-AES measurement. Average chemical recovery of strontium is 75 \pm 17%. Measured values of ⁹⁰Sr agreed to within 3.2% and 0.5% of reference values for 20 minute count times and 60 minute count times, respectively. The lower limit of detection for 100mL samples with 20 minute count times is 0.5Bq/L and with 60 minute count times is 0.16Bq/L. A single operator can prepare

batches of 12-24 samples for ⁹⁰Sr measurement in less than 8 hours.

Reagents

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) Sr Resin, 1mL Cartridges (Eichrom SR1ML-R50-S) Ammonium Hydroxide (listed as 28% NH₃ or 56% NH₄OH) Nitric Acid (70%) Hydrogen Peroxide (30%) Deionized Water 1.25M Ca(NO₃)₂ 3.2M (NH₄)₂HPO₄ Strontium Carrier (10mg/mL) 2M Al(NO₃)₃ ⁹⁰Sr standard Oxalic acid

Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50mL and 250mL Centrifuge Tubes Centrifuge Cupped Stainless Steel Planchets (~5mL volume) Gas Flow Proportional Counter Muffle Furnace Hot Plate Analytical Balance 250mL Glass Beakers Vacuum Pump

Figure 1. Sample Preparation

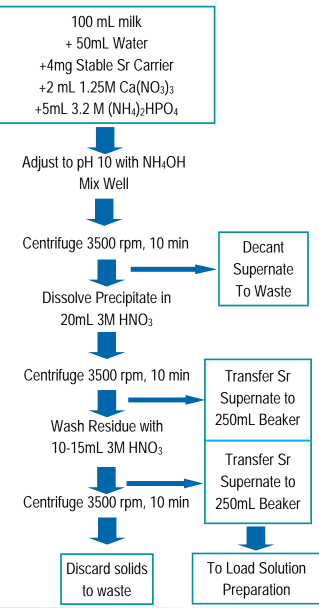
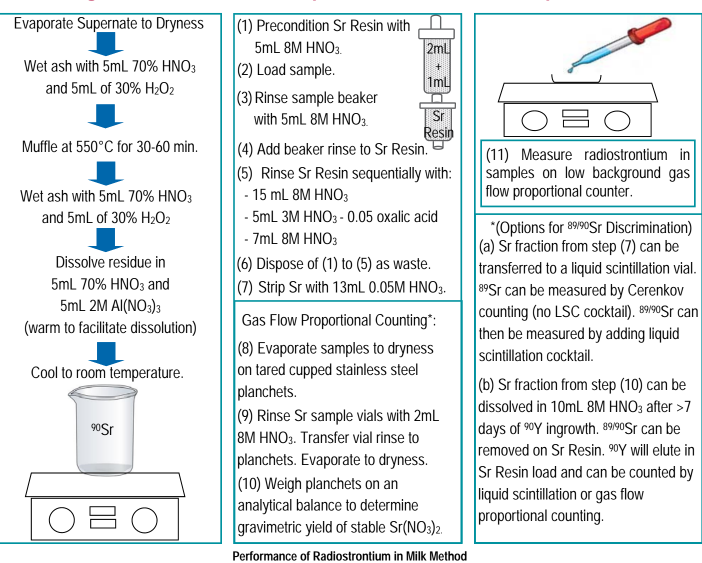


Figure 2. Load Solution Preparation and Strontium Separation



		1 01101111		00000		ou -
	20 Minute Coun	it Times] [60 Minute Cour
⁹⁰ Sr, reference	⁹⁰ Sr,measured	Uncertainty] [⁹⁰ Sr, reference	⁹⁰ Sr,measured
(Bq/L)	(Bq/L)	%, k = 2	% Bias		(Bq/L)	(Bq/L)
0	0.26	98.9	N/A	1 [0	0.11
0	0.26	81.9	N/A		0	0.27
2.86	2.66	24.1	-7.0	1 [2.86	3.09
2.86	3.96	24.7	38		2.86	3.11
2.86	3.31	20.2	15.7		2.86	2.67
2.86	2.67	18.7	-6.6		2.86	2.67
5.7	6.11	16.7	7.2	1 [5.7	5.85
5.7	5.71	13.1	0.2		5.7	5.75
5.7	5.16	13.9	-9.5		5.7	6.04
14.3	12.8	9.1	-11	1 [14.3	13.6
14.3	15.2	8.5	6.3		14.3	14.0
14.3	14.1	8.6	-1.4		14.3	14.2

int Times Uncertainty %, k = 2 % Bias 130 N/A 59 N/A 13.2 8.0 16.7 8.7 13.6 -6.6 11.3 -6.6 10.4 2.6 8.3 0.9 5.9 8.2 6.1 -4.9 6.1 -2.1 6.1 -0.7

References

1) Sherrod L. Maxwell, Brian K. Culligan, "Rapid method for the determination of radiostrontium in emergency milk samples," J. Radioanal. Nucl. Chem., 279(3), 757-760 (2009).

eichrom[®]

Rapid Determination of Sr in 50g Soil Samples

AN-1403-10

Summary of Method Strontium is separated and concentrated from 50 gram soil samples. Soils are leached with concentrated nitric and hydrochloric acid. The leachate is evaporated to dryness, and the residue is dissolved in 1M HCI. A ferric hydroxide-calcium phosphate precipitate concentrates strontium and removes matrix components leached from the soil. A calcium fluoride precipitate further concentrates and purifies the strontium fraction. Strontium is separated from matrix impurities and potentially interfering radionuclides in the sample using two stacked 2mL cartridges of Eichrom Sr Resin. Radiostrontium is measured on a low background gas flow proportional counter. Average chemical recovery of strontium, determined by gravimetric yield of stable strontium carrier, is $91 \pm 4\%$. Measured values of 90Sr agreed to within 2% of reference values for 90 minute count times. The minimum detectable activity for 90Sr in 50g samples with 90 minute count times is 0.41Bq/g. A single operator can prepare batches of 12

samples for the measurement of ⁹⁰Sr in less than 16 hours.

Reagents

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) Nitric Acid (70%) Hydrochloric Acid (37%) Hydrofluoric Acid (49%) or Sodium Fluoride Ammonium Hydroxide (listed as 28% NH₃ or 56% NH₄OH) Deionized Water 1.25M Ca(NO₃)₂ 3.2M (NH₄)₂HPO₄ Strontium Carrier (10mg/mL) 2M Al(NO₃)₃ Sr-90 standard Oxalic acid Boric acid

Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50mL and 250mL Centrifuge Tubes Centrifuge Cupped Stainless Steel Planchets (~5mL volume) Gas Flow Proportional Counter Muffle Furnace Hot Plate Analytical Balance 600mL Glass Beakers Vacuum Pump

Figure 1. Sample Preparation

Dry soil at 110°C. Blend and Size. Remove 50g aliquot into 600mL glass beaker .

Muffle at 550°C for 1-2 hours.

Add 6mg Sr Carrier*, 50mL 70% HNO₃, and 25mL 37% HCI.

*may need to adjust Sr carrier amount to account for native Sr content in soil.

Heat to dryness on hot plate, medium setting.

Add 50mL 70% HNO₃. Warm sample. Transfer solids and liquid to 250mL centrifuge tube.

Centrifuge 3500 rpm, 10 min. Transfer supernate to 600mL beaker.

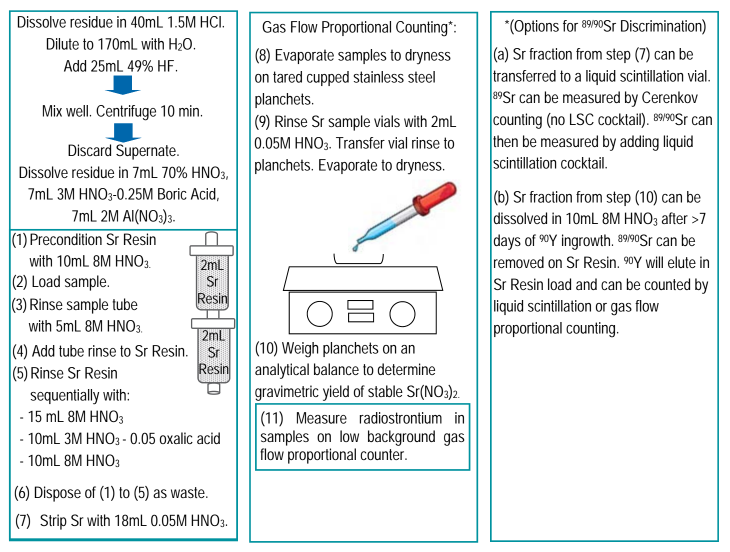
Add 25mL 70% HNO₃ to Solids. Mix and Centrifuge. Transfer supernate to same 600mL beaker. Repeat once. Discard solids to waste.

Evaporate supernate in 600mL beaker to dryness. Dissolve residue in 15-20mL 1M HCI. Transfer to 250mL centrifuge tube.

Dilute to 160mL. Add 1mL 1.25M Ca(NO₃)₃, 2mL 3.2M (NH₄)₂HPO₄, and 25mL 57% NH₄OH. Mix. Centrifuge. Decant supernate to waste.

Continue to load solution preparation.

Figure 2. Load Solution Preparation and Strontium Separation



Method Performance for 50g Soils Spiked with ⁹⁰Sr

Sample	⁹⁰ Sr Reference	⁹⁰ Sr Measured		Sr Carrier
replicates	Value (mBq/g)	Value (mBq/g)	% Bias	% Yield
7	5.92	5.95 <u>+</u> 0.22	5.0	94.0 <u>+</u> 2.6
7	11.8	11.5 <u>+</u> 0.7	-2.5	89.6 <u>+</u> 2.7
7	59.2	57.8 <u>+</u> 1.7	-2.4	89.3 <u>+</u> 4.7

MDA ⁹⁰Sr, 90 minute count, 50g Soil = 0.41 mBq/g

References

1) Sherrod L. Maxwell, Brian K. Culligan, Patrick J. Shaw "Rapid determination of radiostrontium in large soil samples," *J. Radioanal. Nucl. Chem., 295(2), 965-971* (2013).

AN-1404-10

Rapid Determination of Sr in **1-2 Liter Seawater Samples**

Strontium is separated and concentrated from 1-2L samples of seawater with a calcium Summary of Method phosphate precipitation, enhanced with 200mg of iron. Strontium is separated from matrix impurities and potentially interfering radionuclides in the sample using two stacked 2mL cartridges of Eichrom Sr Resin. Radiostrontium is measured on a low background gas flow proportional counter or liquid scintillation counter. Chemical yield of strontium is determined by gravimetric recovery of native stable strontium in the seawater or by ICP-AES measurement. Average chemical recovery of strontium is 89 + 5% for 1L samples and 82 + 4% for 2L samples. Measured values of ⁹⁰Sr agreed to within 1% and 4% of reference values, for 1L and 2L, respectively, with two hour count times. The minimum detectable activity for ⁹⁰Sr for 2L samples with a two hour count time is 9.1Bg/L. A single operator can prepare batches of 12-24 samples for measurement of radiostrontium in less than 8 hours.

Reagents

Equipment

Centrifuge

Hot Plate

Analytical Balance

Vacuum Pump

50mL Centrifuge Tubes

250-500mL Centrifuge Tubes

Gas Flow Proportional Counter

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) Nitric Acid (70%) Ammonium Hydroxide (listed as 28% NH₃ or 56% NH₄OH) **Deionized Water** Iron Carrier (50mg/mL Fe, as ferric nitrate) 3.2M (NH₄)₂HPO₄ $2M AI(NO_3)_3$ ⁹⁰Sr standard Oxalic acid

Figure 1. Sample Preparation

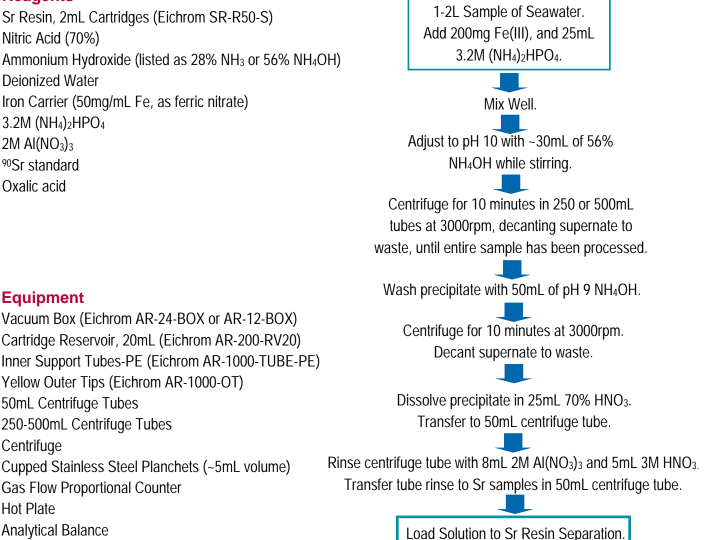
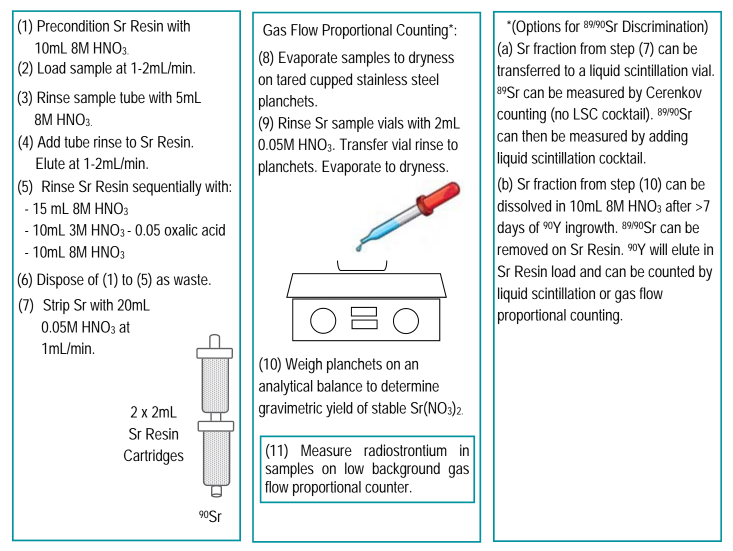


Figure 2. Strontium Resin Separation (Optional ⁹⁰Y Ingrowth)



Performance of ⁹⁰Sr Method for 1L and 2L Seawater Samples

Sample Replicates	Sample Volume, L	⁹⁰ Sr, Reference Value (mBq/L)	⁹⁰ Sr, Measured Value (mBq/L)	% Bias	Sr carrier % Recovery
11	1	148	150 <u>+</u> 11	1.2	89 <u>+</u> 5
4	2	148	154 <u>+</u> 5	4.2	82 <u>+</u> 4

2 hour count times

MDA = 9.1 mBq/L for 2L sample

References

1) Sherrod L. Maxwell, Brian K. Culligan, Robin C. Utsey, "Rapid determination of radiostrontium in seawater samples," *J. Radioanal. Nucl. Chem., 298(2), 867-875* (2013).

Rapid Determination of Sr in Vegetation Samples

AN-1405-10

Summary of Method Strontium is separated and concentrated from 5-10 gram vegetation samples. Samples are muffled in zirconium crucibles 2-4 hours to destroy organic content. The residue is wet ashed with HNO₃-H₂O₂ and then fused with 15g NaOH at 600°C for ten minutes. The fusion cakes are dissolved in water, transferred to 250mL centrifuge tubes and precipitated with calcium phosphate to facilitate matrix removal. Strontium is separated from matrix impurities and potentially interfering radionuclides in the sample using stacked 2mL and 1mL cartridges of Eichrom Sr Resin. Radiostrontium is measured on a low background gas flow proportional counter or liquid scintillation counter. Chemical yield of strontium is determined by gravimetric recovery of stable strontium or ICP-AES measurement. Average chemical recovery of strontium is $64 \pm 4\%$ for 5g samples and $70 \pm 8\%$ for 10g samples. Measured values of ⁹⁰Sr agreed to within 12% of reference values for 90 minute count times. The average time to complete the sample preparation is <8 hours.

Reagents

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S)Sr Resin, 1mL Cartridges (Eichrom SR1ML-R50-S)Nitric Acid (70%)Hydrogen Peroxide (30%)Deionized WaterIron Carrier (50mg/mL Fe, as ferric nitrate)Strontium Carrier (10mg/mL)1.25M Ca(NO_3)_23.2M (NH_4)_2HPO_42M Al(NO_3)_3% Sr standardOxalic acid

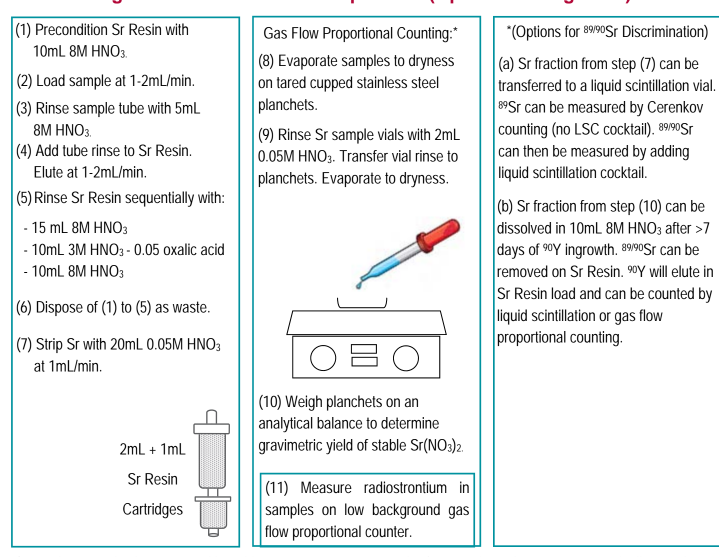
Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50mL and 250mL Centrifuge Tubes Centrifuge Cupped Stainless Steel Planchets (~5mL volume) Gas Flow Proportional Counter Muffle Furnace Hot Plate Analytical Balance 250mL Zirconium crucibles with zirconium lids Vacuum Pump

Figure 1. Sample Preparation

5-10g Vegetation sample in zirconium crucible Muffle at 600°C. 2 hours for 5g sample. 4 hours for 10g sample. Wet ash on hotplate with 5mL 70% HNO₃ and 5mL 30% H₂O₂. Fuse samples with 15g NaOH at 600°C for 10 minutes. Dissolve fusion cake with H₂O. Transfer to 250mL centrifuge tube. Add 125mg Fe and 4mg Sr. Dilute to 180mL. Add 4mL 1.25M Ca(NO₃)₂, 5mL 3.2M (NH₄)₂HPO₄. Mix. Cool in ice bath for 10min. Centrifuge at 3500rpm. Decant Supernate. Dissolve precipitate in 5mL warm 3M HNO₃, 7mL 70% HNO₃, and 7mL 2M AI(NO₃)₃.

Figure 2. Strontium Resin Separation (Optional ⁹⁰Y Ingrowth)



*Actinides may also be measured by adding a 2mL TEVA, TRU and DGA cartridges above Sr Resin and following the separation scheme in Eichrom application note AN-1406, "Rapid Determination of Actinides in Vegetation Samples."

Performance of ⁹⁰Sr Method 5-10g Vegetation Samples

Sample	Sample	⁹⁰ Sr, Reference	⁹⁰ Sr, Measured	04 D:	Sr carrier
Replicates	Mass, g	Value (Bq/g)	Value (Bq/g)	% Bias	% Recovery
6	5.0	0.255	0.285 <u>+</u> 0.03	12	64 <u>+</u> 4
2	10.0	0.156	0.156 <u>+</u> 0.001	0.0	69 <u>+</u> 2
2	10.0	0.110	0.109 <u>+</u> 0.003	-0.1	70 <u>+</u> 7

90 minute count times

References

1) Sherrod L. Maxwell, Brian K. Culligan, Gary W. Noyes, "Rapid separation of actinides and radiostrontium in vegetation samples," *J. Radioanal. Nucl. Chem., 286(1), 273-282* (2010).

Rapid Determination of Actinides in Vegetation Samples

AN-1406-10

Summary of Method U, Pu, Am and Cm are separated and concentrated from 5-10 gram vegetation samples. Samples are muffled in zirconium crucibles 2-4 hours to destroy organic content. The residue is wet ashed with HNO_3 - H_2O_2 and then fused with 15g NaOH at 600°C for ten minutes. The fusion cakes are dissolved in water, transferred to 250mL centrifuge tubes and precipitated twice to facilitate matrix removal. Actinides are separated on stacked 2mL

cartridges of Eichrom TEVA, TRU and DGA resins. Actinides are measured by alpha spectrometry following CeF₃ microprecipitation onto Eichrom Resolve[®] Filters. Chemical yields of tracers ranged from 90-101% for ²⁴²Pu, 84-93% for ²⁴³Am, and 81-87% for ²³²U. Measured values agreed to within 1-3% of reference values for Pu isotopes, 3-9% for Am and Cm isotopes, and 2-15% for U isotopes for 16 hour count times. A single operator can prepare batches of 12 samples for the measurement of actinides in less than 8 hours.

Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Iron Carrier (50mg/mL Fe, as ferric nitrate) Lanthanum and Cerium Carriers (10mg/mL) ²⁴²Pu (or ²³⁶Pu if meas. Np), ²⁴³Am and ²³²U tracers Oxalic acid/Ammonium oxalate Hydrofluoric Acid (49%) or Sodium Fluoride 3.2M (NH₄)₂HPO₄ 2M AI(NO₃)₃ 10% (w:w)TiCl₃ Boric acid Sodium Hydroxide Sodium Nitrite Denature Ethanol Sulfamic Acid Ascorbic Acid Hydrogen Peroxide (30%) Nitric Acid (70%) Hydrochloric Acid (37%) **Deionized Water** 1.25M Ca(NO₃)₂

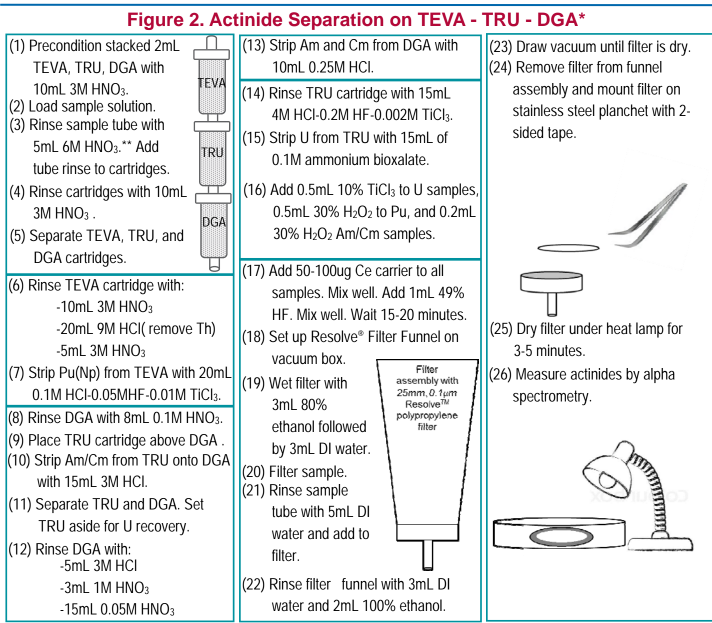
Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes 250mL Zirconium crucibles with zirconium lids Alpha Spectrometry System Centrifuge Muffle Furnace Hot Plate Heat Lamp Analytical Balance Vacuum Pump

Figure 1. Sample Preparation



Add 1mL 3.5M NaNO₂ and 1.5mL 70% HNO₃. Mix.



*Radiostrontium may also be measured by adding a 2mL + 1mL Sr Resin cartridge below DGA and following separation scheme in Eichrom application note AN-1405, "Rapid Determination of Sr in Vegetation Samples."

**Adding 50uL of 30% H₂O₂ to the tube rinse can help improve U recovery and decontamination in Pu/Np fractions.

	Performance of Actinides in Vegetation Method										
	5 gram Samples							10 gram	Samples		
		Reference	Measured		% Tracer			Reference	Measured		% Tracer
Nuclide	Replicates	(mBq/g)	(mBq/g)	% Bias	Recovery	Nuclide	Replicates	(mBq/g)	(mBq/g)	% Bias	Recovery
²³⁸ Pu	6	29.4	30.1 <u>+</u> 3.7	2.4	101 <u>+</u> 6	²³⁸ Pu	2	27.4	28.1 <u>+</u> 0.4	2.6	90 <u>+</u> 15
²³⁹ Pu	6	56.8	57.0 <u>+</u> 4.8	0.3	101 <u>+</u> 6	²³⁹ Pu	2	32.8	32.4 <u>+</u> 0.9	-1.2	90 <u>+</u> 15
²⁴¹ Am	6	48.0	48.5 <u>+</u> 4.6	1.0	93 <u>+</u> 7	²⁴¹ Am	2	31.2	30.8 <u>+</u> 0.0	-1.3	84 <u>+</u> 12
²⁴⁴ Cm	6	6.28	5.9 <u>+</u> 0.6	-6.1	93 <u>+</u> 7	²³⁴ U	2	41.6	41.3 <u>+</u> 1.3	-0.7	81 <u>+</u> 12
²³⁴ U	6	69.2	81 <u>+</u> 7	17	87 <u>+</u> 7	²³⁸ U	2	43.2	42.0 + 0.3	-2.8	81 <u>+</u> 12
²³⁸ U	6	71.8	83 <u>+</u> 10	16	87 <u>+</u> 7	-					

Performance of Actinides in Vegetation Method

References

1) Sherrod L. Maxwell, Brian K. Culligan, Gary W. Noyes, "Rapid separation of actinides and radiostrontium in vegetation samples," *J. Radioanal. Nucl. Chem., 286(1), 273-282* (2010).

Rapid Determination of Sr in Animal Tissue Samples

AN-1407-10

Summary of Method Strontium is separated and concentrated from up to 200g tissue samples. Samples are digested with aqua regia, wet ashed with HNO₃-H₂O₂ and muffled over night at 550°C to destroy organic content. Strontium is separated from matrix impurities and potentially interfering radionuclides in the sample using stacked 2mL and 1mL cartridges of Eichrom Sr Resin. Radiostrontium is measured on a low background gas flow proportional counter or liquid scintillation counter. Chemical yield of strontium is determined by gravimetric recovery of stable strontium or ICP-AES measurement. Average chemical recoveries of strontium are 74-89% for 200g samples of catfish, bass, red drum, mullet, sea trout. Average strontium recoveries for 100 gram samples of deer, hog, bream and shellfish are 83-96%. A single operator can complete the sample preparation, including 16 hours for muffling, for 12-24 samples in less than 24 hours.

Reagents

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) Sr Resin, 1mL Cartridges (Eichrom SR1ML-R50-S) Nitric Acid (70%) Hydrochloric Acid (37%) Hydrogen Peroxide (30%) Deionized Water Strontium Carrier (10mg/mL) Aluminum Nitrate, Nonahydrate Sr-90 standard Oxalic acid

Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Cupped Stainless Steel Planchets (~5mL volume) Gas Flow Proportional Counter Muffle Furnace Hot Plate Analytical Balance 600mL Glass Beakers Vacuum Pump

Figure 1. Sample Preparation

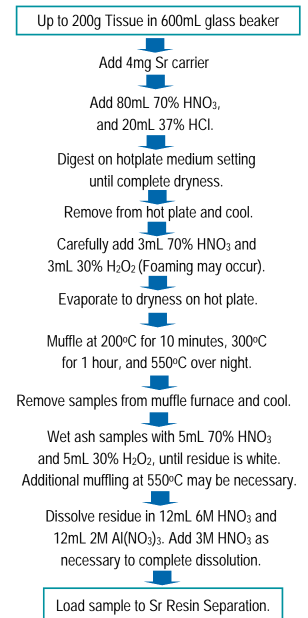
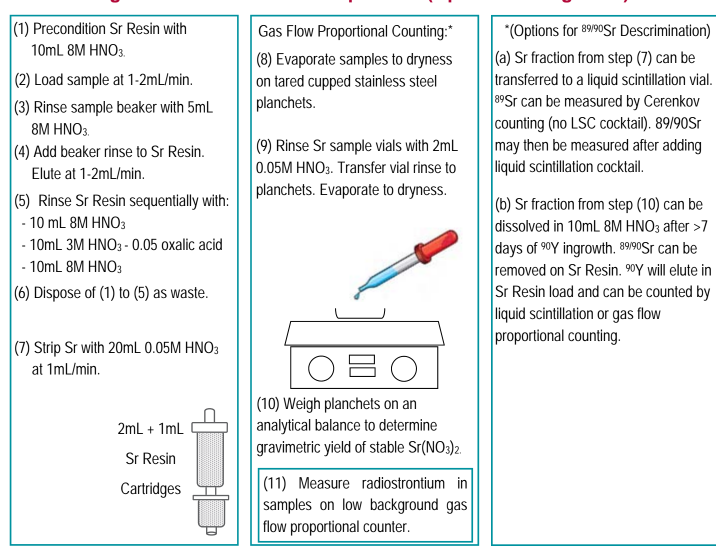


Figure 2. Strontium Resin Separation (Optional ⁹⁰Y Ingrowth)



Actinides may also be measured by adding 2mL TEVA, TRU and DGA cartridges above Sr Resin and following the separation scheme in Eichrom application note AN-1408, "Rapid Determination of Actinides in Animal Tissue Samples."

Sr Carrier Recovery for 100-200g Tissue Samples								
			% Recovery				% Recovery	
Sample	grams	replicate	Sr carrier	Sample	grams	replicate	Sr carrier	
Beef	100	6	96.3 <u>+</u> 0.5	Fish-Mullet	200	6	85.6 <u>+</u> 17	
Deer	100	59	83.4 <u>+</u> 3.5	Fish-Red Fish	200	6	77.7 <u>+</u> 21	
Fish-Bass	200	72	89.0 <u>+</u> 16	Fish-Sea Trout	200	6	74.4 <u>+</u> 25	
Fish-Bream	100	57	91.7 <u>+</u> 10	Hog	100	17	86.0 <u>+</u> 7.1	
Fish-Catfish	200	69	89.4 + 17	Shellfish	100	5	97.5 + 0.9	

References

1) Sherrod L. Maxwell, Donald M. Faison, "Rapid column extraction method for actinides and strontium in fish and other animal tissue samples," J. Radioanal. Nucl. Chem., 275(3), 605-612 (2007).

Rapid Determination of Actinides in Animal Tissue Samples

AN-1408-10

Summary of Method Uranium, Plutonium, and Americium-Curium are separated and concentrated from up to 200g tissue samples. Samples are digested with aqua regia, wet ashed with HNO₃-H₂O₂ and muffled over night at 550°C to destroy organic content. Actinides are separated from matrix impurities and potentially interfering radionuclides in the sample using stacked 2mL cartridges of Eichrom TEVA, TRU and DGA Resin. Actinides are measured via alpha spectrometry following cerium fluoride microprecipitation onto Eichrom Resolve[®] Filters. Average chemical recoveries of Pu for 100-200g samples are 93-101%. Typical americium recoveries are 93-105%. Typical uranium recoveries are 82-

96%. A single operator can complete the sample preparation for 12-24 samples, including 16 hours for muffling, in less than 24 hours.

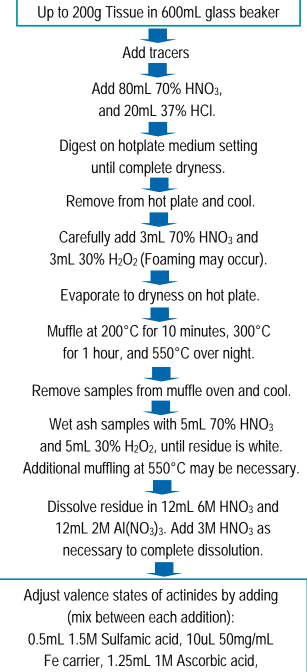
Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Hydrofluoric Acid (49%) or Sodium Fluoride Iron Carrier (50mg/mL Fe, as ferric nitrate) ²⁴²Pu (or ²³⁶Pu if meas. Np), ²⁴³Am and ²³²U tracers Oxalic acid/Ammonium oxalate Nitric Acid (70%) Hydrochloric Acid (37%) Hydrogen Peroxide (30%) **Deionized Water** Cerium Carrier (1mg/mL) $2M AI(NO_3)_3$ Sodium nitrite Sulfamic acid Ascorbic acid 10% (w:w) TiCl₃ **Denatured Ethanol**

Equipment

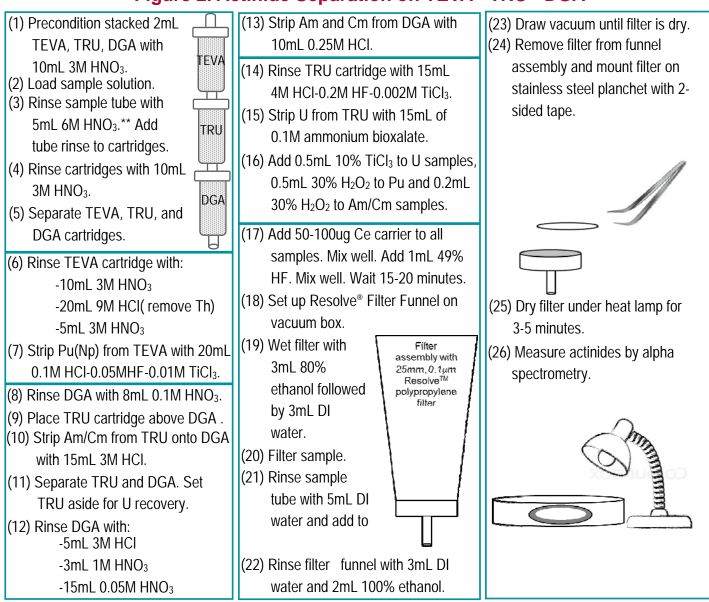
Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) Muffle Furnace Hot Plate Analytical Balance 600mL Glass Beakers Stainless Steel planchets with adhesive Vacuum Pump Alpha Spectrometry System Heat Lamp

Figure 1. Sample Preparation



1mL 3.5M NaNO₂, and 1.5mL 70% HNO₃.

Figure 2. Actinide Separation on TEVA - TRU - DGA*



*Radiostrontium may also be measured by adding a 2mL + 1mL Sr Resin cartridge below DGA and following separation scheme in Eichrom application note AN-1407, "Rapid Determination of Sr in Animal Tissue Samples." **Adding 50uL of 30% H₂O₂ to tube rinse can improve U recoveries and decontamination in Pu/Np samples.

Meth	Method Performance for 100-200g Tissue Samples							
			% Tracer Recovery					
Sample	mass, g	replicates	Pu-236	Am-243	U-232			
Beef	100	6	98.7 <u>+</u> 5.7	97.1 <u>+</u> 8.4	93.4 <u>+</u> 4.7			
Deer	100	59	99.3 <u>+</u> 12	93.4 <u>+</u> 10	90.4 <u>+</u> 8.0			
Fish-Bass	200	72	96.2 <u>+</u> 14	102 <u>+</u> 13	95.1 <u>+</u> 8.1			
Fish-Bream	100	57	96.6 <u>+</u> 12	98.4 <u>+</u> 7.7	91.1 <u>+</u> 6.3			
Fish-Catfish	200	69	98.3 <u>+</u> 12	103.7 <u>+</u> 7.6	89 <u>+</u> 12			
Hog	100	17	93 <u>+</u> 20	96.4 <u>+</u> 9.7	86 <u>+</u> 15			
Shelfish	100	5	101.3 <u>+</u> 2.2	97.4 <u>+</u> 7.1	81.7 <u>+</u> 3.2			

Reference Sherrod L. Maxwell, Donald M. Faison, "Rapid column extraction method for actinides and strontium in fish and other animal tissue samples," *J. Radioanal. Nucl. Chem., 275(3), 605-612* (2007).

Rapid Determination of Sr in Building Materials

AN-1409-10

Summary of Method Strontium is separated and concentrated from 1.5 gram samples of concrete or brick. Samples are finely ground and fused in a zirconium crucible for 15 minutes at 600°C with 15 grams of sodium hydroxide. The fusion cake is dissolved in water and strontium is concentrated and separated from the matrix using a calcium phosphate precipitate enhanced with iron. A secondary precipitation with calcium fluoride removes additional matrix and decreases the volume of precipitate. The calcium fluoride precipitate is dissolved with nitric acid-boric acid-aluminum nitrate to form the Sr Resin load solution. Strontium is separated from remaining matrix and potentially interfering radionuclides using stacked 2mL and 1mL Sr Resin cartridges. Batches of 12-24 samples can be prepared for analysis in less than 8 hours. Radiostrontium is measured by gas flow proportional counting or liquid scintillation counting. Chemical yield of strontium is determined by gravimetric yield of stable strontium or ICP-AES measurement.

Reagents

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) Sr Resin, 1mL Cartridges (Eichrom SR1ML-R50-S) Nitric Acid (70%) Hydrochloric Acid (37%) Hydrofluoric Acid (49%) or Sodium Fluoride Deionized Water 1.25M Ca(NO₃)₂ 3.2M (NH₄)₂HPO₄ 2M Al(NO₃)₃ Iron Carrier (50mg/mL Fe, as ferric nitrate) Strontium Carrier (10mg/mL) ⁹⁰Sr standard Oxalic acid Boric acid Sodium Hydroxide

Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50mL and 250mL Centrifuge Tubes Centrifuge Cupped Stainless Steel Planchets (~5mL volume) Gas Flow Proportional Counter Muffle Furnace Hot Plate Analytical Balance 250mL Zirconium crucibles with zirconium lids Vacuum Pump

Figure 1. Sample Preparation

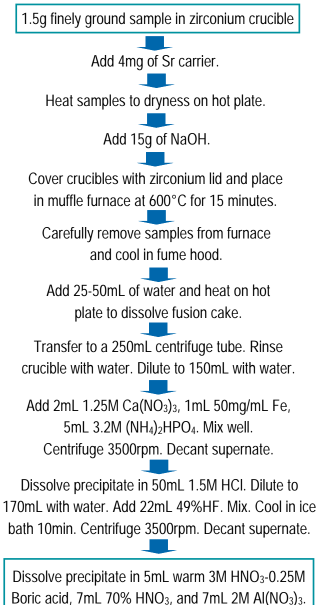
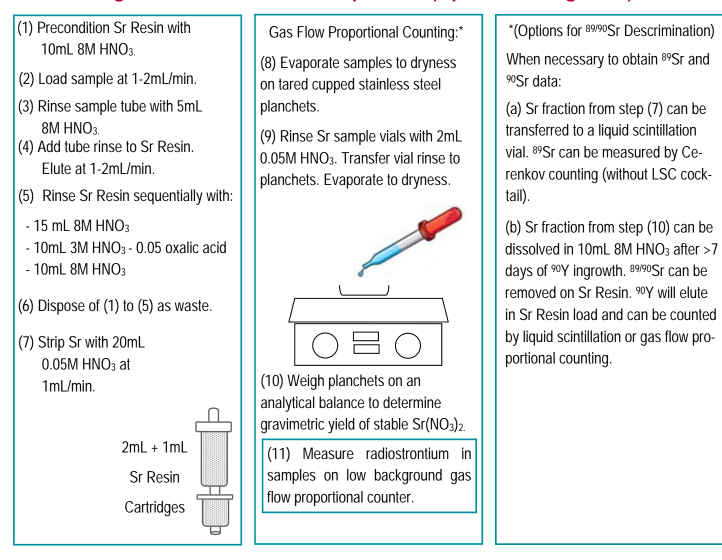


Figure 2. Strontium Resin Separation (Optional ⁹⁰Y Ingrowth)



References

1)"Rapid radiochemical method for total radiostrontium (Sr-90) in building materials for environmental remediation following radiological incidents," U.S. Environmental Protection Agency, National Analytical Radiation Environmental Laboratory, EPA 402-R14-001.

2)"Rapid method for sodium hydroxide fusion of concrete and brick matrices prior to americium, plutonium, strontium, radium, and uranium analyses for environmental remediation following radiological incidents,"U.S. Environmental Protection Agency, National Analytical Radiation Environmental Laboratory, EPA 402-R-14-004.

Rapid Determination of Sr in Emergency Urine Samples

AN-1410-10

Summary of Method Strontium is separated and concentrated from 100mL urine samples using calcium phosphate precipitation. An optional wet-ashing step with HNO₃-H₂O₂ destroys residual organic material. The precipitate or wet-ashed residue is dissolved in nitric acid and aluminum nitrate. Strontium is then separated from matrix impurities and potentially interfering radionuclides in the sample using a 2mL cartridge of Eichrom Sr Resin. Radiostrontium is measured on a low background gas flow proportional counter or liquid scintillation counter. Chemical yield of strontium is determined by gravimetric recovery of stable strontium or ICP-AES measurement. Typical chemical recovery of strontium is >80%. Measured values of ⁹⁰Sr agreed to within 1.7% of reference values for 10 minute count times, although longer count times can be used to improve detection limits and uncertainty. A single operator can complete the separation method for batches of 12-24 samples in as little as 3-4 hours.

Reagents

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) Nitric Acid (70%) Ammonium Hydroxide (listed as 28% NH₃ or 56% NH₄OH) Hydrogen Peroxide (30%) Deionized Water 1.25M Ca(NO₃)₂ 3.2M (NH₄)₂HPO₄ Sr Carrier (10mg/mL) 2M Al(NO₃)₃ ⁹⁰Sr standard Oxalic acid

Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50mL and 250mL Centrifuge Tubes Centrifuge Cupped Stainless Steel Planchets (~5mL volume) Gas Flow Proportional Counter Hot Plate Analytical Balance 250mL Glass Beakers Vacuum Pump

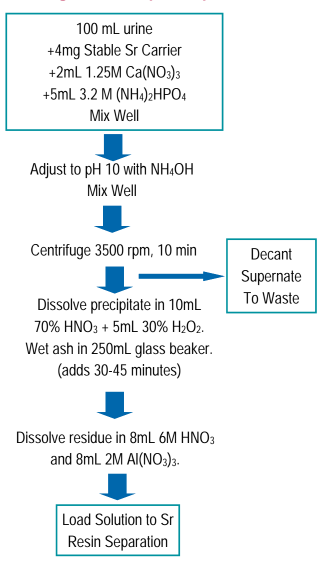
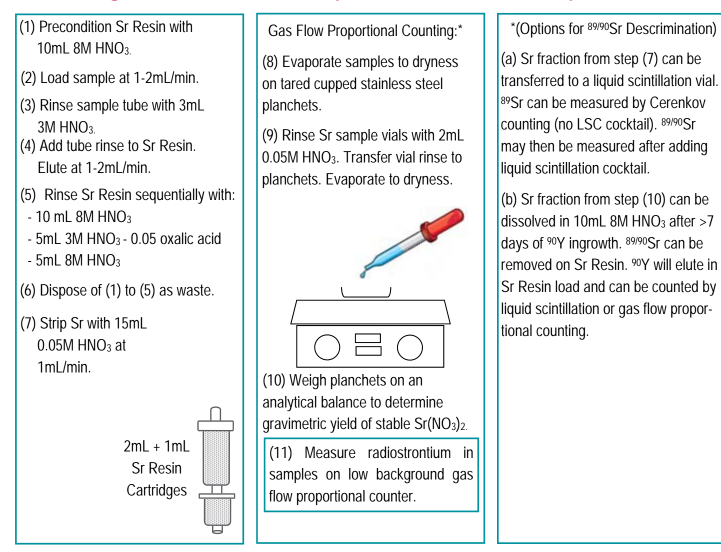


Figure 1. Sample Preparation

Figure 2. Load Solution Preparation and Strontium Separation



Actinides may also be measured by adding a 2mL TEVA, TRU and DGA cartridges above Sr Resin and following the separation scheme in Eichrom application note AN-1412, "Rapid Determination of Actinides in Emergency Urine Samples."

References

1) Sherrod L. Maxwell, Brian K. Culligan, "Rapid separation method for emergency water and urine samples," *J. Radioanal. Nucl. Chem., 279(3), 901-907* (2009).

Rapid Determination of Sr in Emergency Water Samples

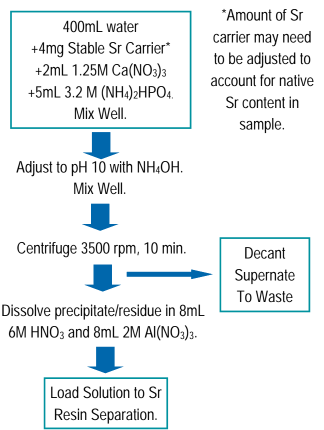
AN-1411-10

Summary of Method Strontium is separated and concentrated from up to 400mL water samples using calcium phosphate precipitation. The precipitate is dissolved in nitric acid and aluminum nitrate. Strontium is then separated from matrix impurities and potentially interfering radionuclides in the sample using a 2mL cartridge of Eichrom Sr Resin. Radiostrontium is measured on a low background gas flow proportional counter or liquid scintillation counter. Chemical yield of strontium is determined by gravimetric recovery of stable strontium or ICP-AES measurement. Typical chemical recovery of strontium is >80%. Measured values of ⁹⁰Sr agreed to within 14% of reference values for 10 minute count times, although longer count times can be used to improve detection limits and uncertainty. A single operator can complete the separation method for batches of 12-24 samples in as little as 3-4 hours.

Reagents

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) Nitric Acid (70%) Ammonium Hydroxide (listed as 28% NH₃ or 56% NH₄OH) Deionized Water 1.25M Ca(NO₃)₂ 3.2M (NH₄)₂HPO₄ Strontium Carrier (10mg/mL) 2M Al(NO₃)₃ ⁹⁰Sr standard Oxalic acid

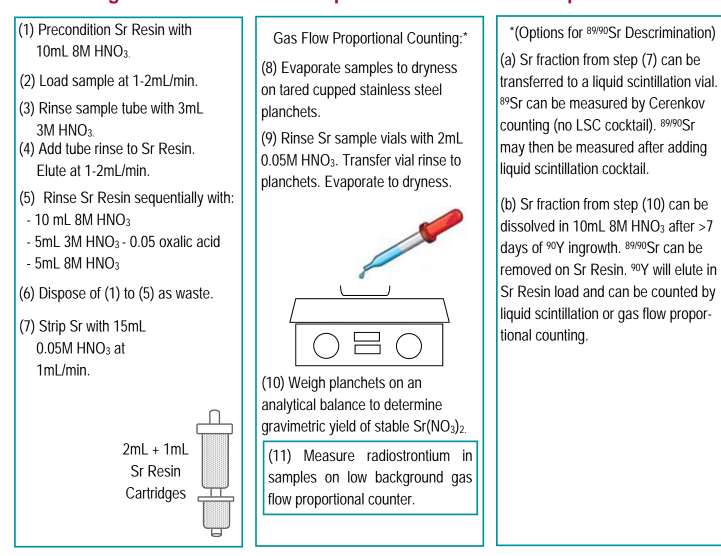
Figure 1. Sample Preparation



Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50mL and 250mL Centrifuge Tubes Centrifuge Cupped Stainless Steel Planchets (~5mL volume) Gas Flow Proportional Counter Analytical Balance Vacuum Pump

Figure 2. Load Solution Preparation and Strontium Separation



Actinides may also be measured by adding a 2mL TEVA, TRU and DGA cartridges above Sr Resin and following the separation scheme in Eichrom application note AN-1413, "Rapid Determination of Actinides in Emergency Water Samples."

References

1) Sherrod L. Maxwell, Brian K. Culligan, "Rapid separation method for emergency water and urine samples," *J. Radioanal. Nucl. Chem., 279(3), 901-907* (2009).

Cichrom^{*} Rapid Determination of Actinides in Emergency Urine Samples

Summary of Method Uranium, Plutonium, and Americium-Curium are separated and concentrated from 100mL urine samples using calcium phosphate precipitation. The precipitate is dissolved in HNO₃-H₂O₂ and wet ashed to destroy residual organic material. The wet-ashed residue is dissolved in nitric acid and aluminum nitrate. Actinides are separated from matrix impurities and potentially interfering radionuclides in the sample using 2mL cartridges of Eichrom TEVA and TRU Resins. Actinides are measured by alpha spectrometry following source preparation by cerium fluoride microprecipitation onto Eichrom Resolve[®] Filters. Chemical yields are determined by recovery of ²³²U, ²⁴³Am, and ²⁴²Pu (or ²³⁶Pu, if measuring ²³⁷Np) tracers. Typical chemical recoveries are >90%. A single operator can complete the separation method for batches of 12-24 samples in as little as 4-5 hours.

Reagents

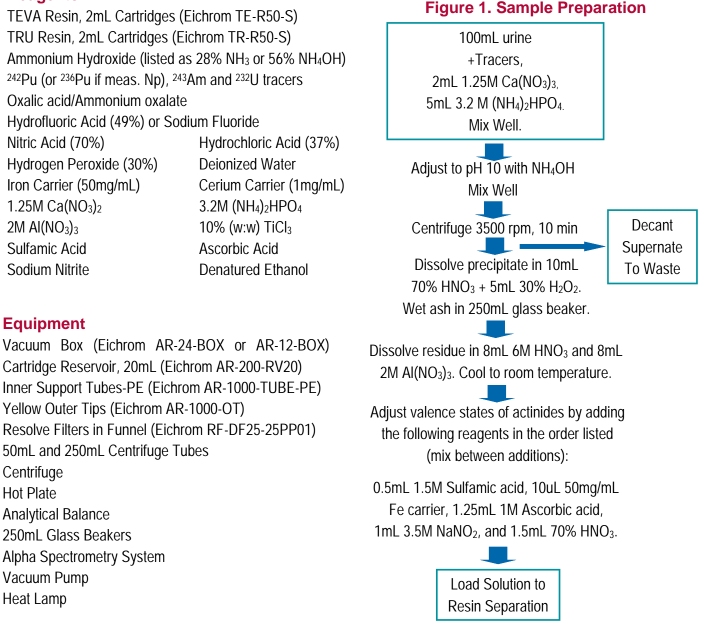
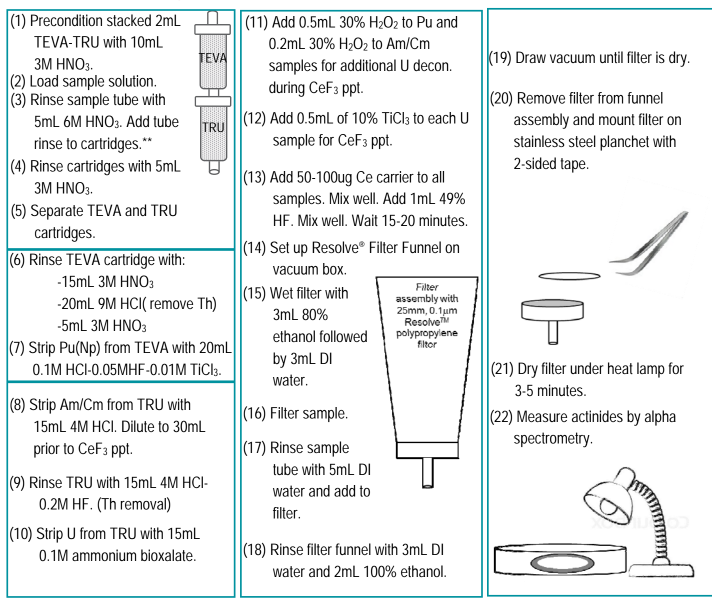


Figure 2. Actinide Separation on TEVA - TRU*



*Strontium may also be measured by adding a 2mL Sr Resin Cartridge below DGA and following the separation scheme in Eichrom application note AN-1410, "Rapid Determination of Sr in Emergency Urine Samples."

**Adding 50uL of 30% H_2O_2 to the 6M HNO₃ tube rinse can help improve U recoveries and decontamination in the Pu/Np fraction.

References

1) Sherrod L. Maxwell, Brian K. Culligan, "Rapid separation method for emergency water and urine samples," *J. Radioanal. Nucl. Chem., 279(3), 901-907* (2009).

Cichrom^{*} AN-1413-10 Rapid Determination of Actinides in Emergency Water Samples

Summary of Method Uranium, Plutonium and Americium-Curium are separated and concentrated from up to 400mL water samples using calcium phosphate precipitation. The precipitate is dissolved in nitric acid and aluminum nitrate. Actinides are separated from matrix impurities and potentially interfering radionuclides in the sample using 2mL cartridges of Eichrom TEVA and TRU Resins. Actinides are measured by alpha spectrometry following source preparation by cerium fluoride microprecipitation onto Eichrom Resolve[®] Filters. Chemical yields are determined by recovery of ²³²U, ²⁴³Am, and ²⁴²Pu (or ²³⁶Pu, if measuring ²³⁷Np) tracers. Typical chemical recoveries are >90%. A single operator can complete the separation method for batches of 12-24 samples in as little as 4-5 hours.

Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) Ammonium Hydroxide (listed as 28% NH₃ or 56% NH₄OH) Nitric Acid (70%) Hydrochloric Acid (37%) Hydrofluoric Acid (49%) or Sodium Fluoride **Deionized Water** Iron Carrier (50mg/mL) Cerium Carrier (1mg/mL) 1.25M Ca(NO₃)₂ 3.2M (NH₄)₂HPO₄ $2M AI(NO_3)_3$ 10% (w:w) TiCl₃ ²⁴²Pu (or ²³⁶Pu if meas. Np), ²⁴³Am and ²³²U tracers Oxalic acid/Ammonium oxalate Sulfamic Acid Ascorbic Acid Sodium Nitrite Denatured Ethanol

Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Analytical Balance Alpha Spectrometry System Vacuum Pump

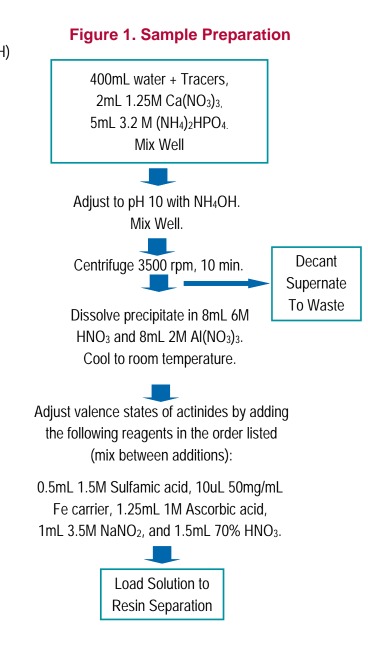
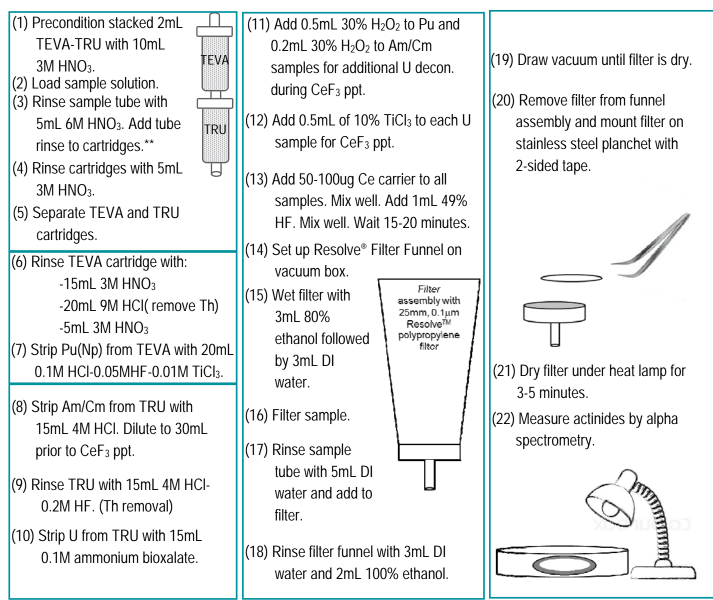


Figure 2. Actinide Separation on TEVA - TRU*



*Strontium may also be measured by adding a 2mL Sr Resin Cartridge below DGA and following the separation scheme in Eichrom application note AN-1411, "Rapid Determination of Sr in Emergency Water Samples."

**Adding 50uL of 30% H_2O_2 to the 6M HNO₃ tube rinse can help improve U recoveries and decontamination in the Pu/Np fraction.

References

1) Sherrod L. Maxwell, Brian K. Culligan, "Rapid separation method for emergency water and urine samples," *J. Radioanal. Nucl. Chem., 279(3), 901-907* (2009).

eichrom[®] Ra

Rapid Determination of ⁹⁰Sr in up to 40 Liter Seawater Samples

AN-1414-10

Summary of Method Yttrium-90, the daughter product of ⁹⁰Sr decay, is separated and concentrated from up to 40L samples of seawater. A ferric hydroxide precipitate enhanced with 10mg of lanthanum and 1mg of yttrium concentrates ⁹⁰Y, while rejecting much of the salt content of the seawater sample. A second precipitation with lanthanum fluoride removes additional matrix ions. Yttrium is separated from potentially interfering radionuclides in the sample, including rare earths such as ¹³⁸La and ^{139/144}Ce, using a 2mL cartridge of Eichrom DGA Resin. ⁹⁰Y is measured on a low background gas flow proportional counter following cerium fluoride microprecipitation onto an Eichrom Resolve[®] Filter. Chemical yield of stable yttrium is determined by ICP-MS or ICP-AES. Average chemical recovery of yttrium is 84 \pm 7% for 40L samples. Measured values of ⁹⁰Sr(⁹⁰Y) agree to within 5% of reference values, with two hour count times. The minimum detectable activity for ⁹⁰Sr for 40L samples with a two hour count time is 0.35mBq/L. The average time to complete the method is 8 hours. While standard methods targeting Sr are limited by the ~8mg/L native Sr content in seawater, targeting ⁹⁰Y directly allows for the efficient processing of very large seawater samples to

achieve very low minimum detectible activities. However, interference by the fission product 91 Y ($t_{1/2} = 58.51$ days) precludes application of this method for the measurement of 90 Sr(90 Y) immediately following a radiological incident involving the release of un-aged nuclear fuel or fission products.

Reagents

DGA Resin, Normal 2mL Cartridges (Eichrom DN-R50-S)Nitric Acid (70%)Hydrochloric Acid (37%)Hydrofluoric Acid (49%) or Sodium FluorideAmmonium Hydroxide (listed as 28% NH3 or 56% NH4OH)Deionized WaterIron Carrier (50mg/mL Fe, as ferric nitrate)Yttrium and Cerium Carriers (1mg/mL)Lanthanum Carrier (10mg/mL)1.25M Ca(NO3)22M Al(NO3)3%Sr standard

Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 500mL Centrifuge Tubes Centrifuge Gas Flow Proportional Counter Analytical Balance Vacuum Pump Heat Lamp

Figure 1. Sample Preparation

Up to 40L Sample of Seawater. Acidify to pH 2 with 37% HCl. Add 1mg Yttrium carrier.

Add 10mg La carrier. Add 50mg Fe carrier per liter of sample. Mix Well.

Adjust to pH 9 with 56% NH₄OH. Mix. Allow precipitate to settle. Decant supernate until ~2L remains.

Transfer remaining supernate and precipitate to 500mL centrifuge tubes. Centrifuge 3000rpm for 10 minutes. Decant supernate. Repeat until entire sample centrifuged.

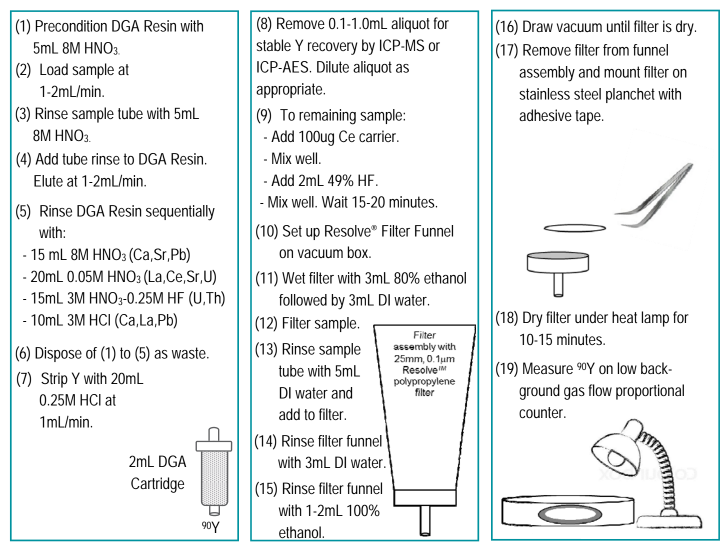
> Wash precipitate with 100mL water. Centrifuge. Decant supernate.

Dissolve precipitate in 100mL 1.5M HCl. Add 75mg Ca and 50mL 49% HF. Mix. Wait 15 minutes. Centrifuge. Decant supernate.

Dissolve precipitate in 10mL 3M HNO₃-0.25M Boric acid, 10mL 70% HNO₃, and 10mL 2M Al(NO₃)₃.

Load Solution for Sr separation.

Figure 2. Yttrium Separation on DGA and CeF₃ Microcprecipitation



Method Performance 10-40L Spike Seawater Samples

Sample	% Recovery	⁹⁰ Sr (mBq/L)	⁹⁰ Sr (mBq/L)	
Volume, L	Y carrier	Reference	Measured	% Bias
10	85.5	296	310	4.7
20	89.2	28.2	28.1	-0.4
30	72.3	18.8	18.5	-1.6
40	87.6	14.1	13.7	-2.8
40	86.5	14.1	13.9	-1.4

MDA for 40L sample = 0.35 mBq/L for 2 hour count time

MDA for 40L sample = 0.20 mBq/L for 8 hour count time

References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchinson, Robin C. Utsey, Daniel R. McAlister, "Rapid determination of ⁹⁰Sr in seawater samples," *J. Radioanal. Nucl. Chem.*, 303, 709-717 (2015).

Rapid Determination of ²¹⁰Po in Water Samples

AN-1415-10

Summary of Method A method for the measurement of ²¹⁰Po in terrestrial water samples is described, offering significant advantages in detection limit, processing time, and resistance to chemical and radiochemical interferences over standard methods where polonium is determined following spontaneous deposition onto metal planchets. ²¹⁰Po is concentrated from up to 1L samples of ground water or 2L samples of drinking water using a calcium phosphate precipitate. ²¹⁰Po is then separated from matrix ions and potentially interfering radionuclides using a 2mL cartridge of Eichrom DGA Resin. ²¹⁰Po is measured using alpha spectrometry following bismuth phosphate microprecipitation onto an Eichrom Resolve® Filter. Chemical recoveries of polonium, determined with a ²⁰⁹Po tracer, were typically 80-90%. ²¹⁰Po measurements typically agreed to reference values to within 3-5%. A single operator can prepare batches of 12-24 samples for alpha counting in 3-4 hours. Alpha spectrometry count times will vary depending on desired detection limit and data quality objectives. Polonium determination may also be integrated into methods for the determination of actinides (Eichrom Application Note AN-1416).

Reagents

DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Nitric Acid (70%) Hydrochloric Acid (37%) Ammonium Hydroxide (listed as 28% NH₃ or 56% NH₄OH) Hydrogen Peroxide (30%) Deionized Water 1.25M Ca(NO₃)₂ 3.2M (NH₄)₂HPO₄ ²⁰⁹Po tracer Bi standard solution (1mg/mL) Denatured Ethanol

Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Alpha Spectrometry System Analytical Balance Vacuum Pump Stainless steel planchets (1.25 inch) with adhesive tape Heat Lamp

Figure 1. Sample Preparation

1-2L Water Sample. Add ²⁰⁹Po tracer. Add 1-2mL of 30% H₂O₂.

Add 1mL 1.25M Ca(NO₃)₃ and 3mL 3.2M (NH₄)₂HPO₄. Mix Well.

Adjust to pH 9 with 56% NH₄OH. Mix. Allow precipitate to settle. Decant supernate to <200mL.

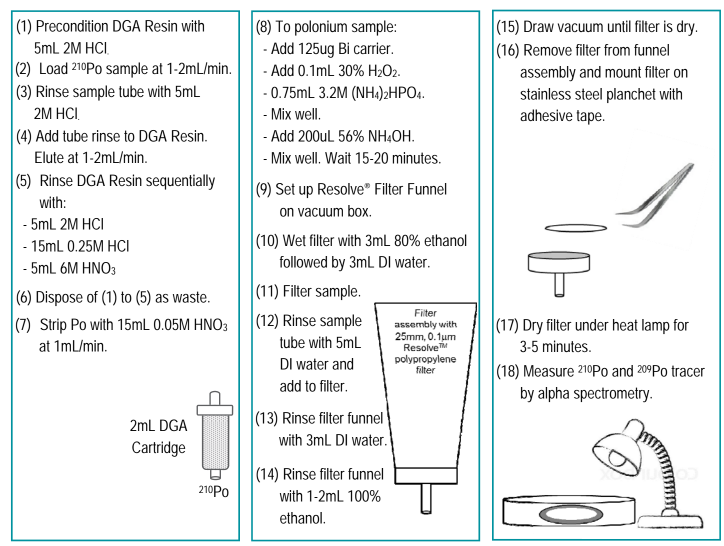
Transfer remaining supernate and precipitate to 250mL centrifuge tubes. Centrifuge 3500rpm for 10 minutes. Decant supernate.

Dissolve precipitate in 10mL 9M HCI.

Transfer to 50mL centrifuge tube. Rinse 250mL tube with 10mL 2M HCI. Transfer tube rinse to same 50mL centrifuge tube.

Load Solution for Po separation.

Figure 2. Polonium Separation on DGA and BiPO₄ Microprecipitation



Method Performance ²¹⁰Po in Water

	Volume		% Recovery	²¹⁰ Po (mBq/L)	²¹⁰ Po (mBq/L)	
Sample	mL	Replicates	²⁰⁹ Po tracer	Reference	Measured	% Bias
Ground Water	200	6	87.4 <u>+</u> 5.8	316	308 <u>+</u> 5	-2.5
Ground Water	200	7	82.3 <u>+</u> 3.9	1262	1289 <u>+</u> 6	2.1
Ground Water	1000	6	85.0 <u>+</u> 8.2	63.3	61.5 <u>+</u> 5.1	-2.8
Drinking Water	2000	4	80.0 <u>+</u> 9.6	63.3	61.1 <u>+</u> 6.2	-3.5

6-12 hour count time

References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchinson, Robin C. Utsey, Daniel R. McAlister, "Rapid determination of ²¹⁰Po in water samples," *J. Radioanal. Nucl. Chem.*, 298(3), 1977-1989 (2014).

Rapid Determination of Actinides and ²¹⁰Po in Water

AN-1416-10

Summary of Method A method for the measurement of ²¹⁰Po and actinides in terrestrial water samples is described, offering significant advantages in detection limit, processing time, and resistance to chemical and radiochemical interferences over standard methods where polonium is determined following spontaneous deposition onto metal planchets. ²¹⁰Po and actinides are concentrated from up to 1L samples of ground water or 2L samples of drinking water using a calcium phosphate precipitate. ²¹⁰Po and actinides are then separated from matrix ions and potentially interfering radionuclides using stacked 2mL cartridge of Eichrom TRU and DGA Resin. ²¹⁰Po and actinides are measured using alpha spectrometry following bismuth phosphate and cerium fluoride microprecipitation, respectively, onto Eichrom Resolve[®] Filters. Tracer recoveries averaged 81.5 \pm 2.6% for ²⁰⁹Po, 93.4 \pm 6.8% for ²⁴²Pu, 100.2 \pm 6.9% for ²⁴³Am and 96.6 \pm 2.5 for ²³²U. Measured values typically agreed to within 3-5% of reference values. A single operator can prepare batches of 12-24 samples for alpha counting in 4-6 hours. Alpha spectrometry count times will vary depending on desired detection limit and data quality objectives.

Reagents

TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Ammonium Hydroxide (Listed as 28% NH₃ or 56% NH₄OH) ²⁰⁹Po, ²³²U, ²⁴³Am, ²⁴²Pu tracers Bi and Ce carriers (1mg/mL) Hydrochloric Acid (37%) Nitric Acid (70%) Hydrofluoric Acid (49%) Hydrogen Peroxide (30%) **Deionized Water** 3.2M (NH₄)₂HPO₄ 1.25M Ca(NO₃)₂ $2M AI(NO_3)_3$ 10% (w:w) TiCl₃ Denatured Ethanol Oxalic acid/Ammonium Oxalate

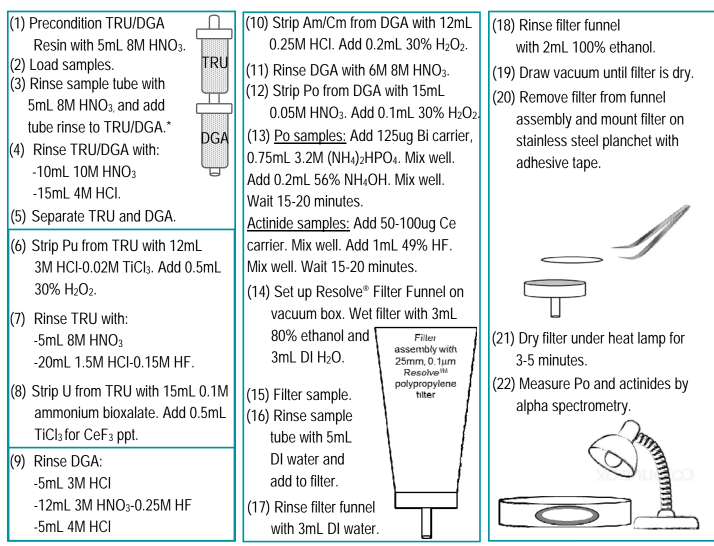
Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Alpha Spectrometry System Analytical Balance Vacuum Pump Heat Lamp Stainless steel planchets (1.25 inch) with adhesive tape

Figure 1. Sample Preparation

1-2L Water Sample. Add tracers. Add 1-2mL of 30% H₂O₂.
Add 1mL 1.25M Ca(NO₃)₃ and 3mL 3.2M (NH₄)₂HPO₄. Mix Well.
Adjust to pH 9 with NH₄OH. Mix. Alow precipitate to settle. Decant supernate to <200mL.
Transfer remaining supernate and precipitate to 250mL centrifuge tubes. Centrifuge 3500rpm for 10 minutes. Decant supernate.
Dissolve precipitate in 10mL 8M HNO₃, 3mL 2M Al(NO₃)₃, and 100uL 30% H₂O₂.
Load Solution for resin separation.

Figure 2. TRU/DGA Separation and Source Preparation



*Adding 50uL 30% H₂O₂ to the tube rinse can improve U recoveries and decontamination in Pu(Np) fractions.

Method Performance ²¹⁰ Po and Actinides in Water							
		% Recovery	Analyte (mBq/L)	Analyte (mBq/L)			
 Analyte	Tracer	of tracer	Reference	Measured	% Bias		
²¹⁰ Po	²⁰⁹ Po	81.5 <u>+</u> 2.6	1584	1660 <u>+</u> 3	4.8		
²³⁸ Pu	²⁴² Pu	93.4 <u>+</u> 6.8	370	381 <u>+</u> 4	3.0		
²⁴¹ Am	²⁴³ Am	100.2 <u>+</u> 6.9	370	381 <u>+</u> 3	3.0		
²⁴⁴ Cm	²⁴³ Am	100.2 <u>+</u> 6.9	328	328 <u>+</u> 4	0.1		
 ²³⁸ U	²³² U	96.6 <u>+</u> 2.5	655	627 <u>+</u> 4	-4.4		

200mL ground water samples, 6 replicates

8-16 hour count time

References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchinson, Robin C. Utsey, Daniel R. McAlister, "Rapid determination of ²¹⁰Po in water samples," *J. Radioanal. Nucl. Chem.*, 298(3), 1977-1989 (2014).

Rapid Determination of ^{226/228}Ra in Water Samples

AN-1417-11

Summary of Method Ra isotopes are separated and measured from 1.0-1.5 liter samples of terrestrial waters. Radium is concentrated from samples on MnO₂ Resin. After a >36 hour ingrowth period for ²²⁸Ac from ²²⁸Ra, radium and ²²⁸Ac are separated from matrix ions and potentially interfering radionuclides using stacked 2mL cartridges of Eichrom LN and DGA Resins. ²²⁸Ac is prepared for gas flow proportional counting using a cerium fluoride microprecipitation onto Eichrom Resolve[®] Filters. ²²⁶Ra is prepared for alpha spectrometry using a barium sulfate microprecipitation onto Eichrom Resolve[®] Filters. Chemical yield of radium is determined by adding a ¹³³Ba tracer. A single operator can process batches of 12-24 samples in 4-5 hours. Results for ²²⁶Ra and ²²⁸Ra can be obtained in 48 hours, including >36 hour ingrowth time for ²²⁸Ac. Results for ²²⁶Ra and ²²⁸Ra in spiked river and ground water samples typically agreed to within 5% of reference values.

Reagents

MnO₂ Resin (Eichrom MN-B100-A) LN Resin (Eichrom LN-R50-S) DGA Resin, Normal 2mL Cartridges (Eichrom DN-R50-S) Ammonium Hydroxide (listed as 28% NH₃ or 56% NH₄OH) ¹³³Ba Tracer Barium and Cerium Carriers (1mg/mL) Nitric Acid (70%) Hydrofluoric Acid (50%) Hydrogen Peroxide (30%) 1.25M Ca(NO₃)₂ Denatured Ethanol Deionized Water

Equipment

Plastic Chromatography Column (Eichrom AC-50E-5M) Column Extension Funnel (Eichrom AC-20X-20M) Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filter in Disposable Funnel (Eichrom RF-DF-25-25PP01) 50mL Centrifuge Tubes Stainless Steel Planchets with Adhesive Tape Alpha Spectroscopy System Gamma Spectroscopy System (if ¹³³Ba tracer used) Low Background Gas Flow Proportional Counter 150mL Glass beakers Vacuum Pump Hot Plate Heat Lamp

Figure 1. Sample Preparation

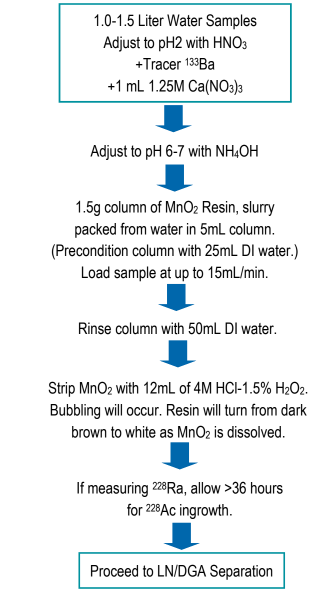
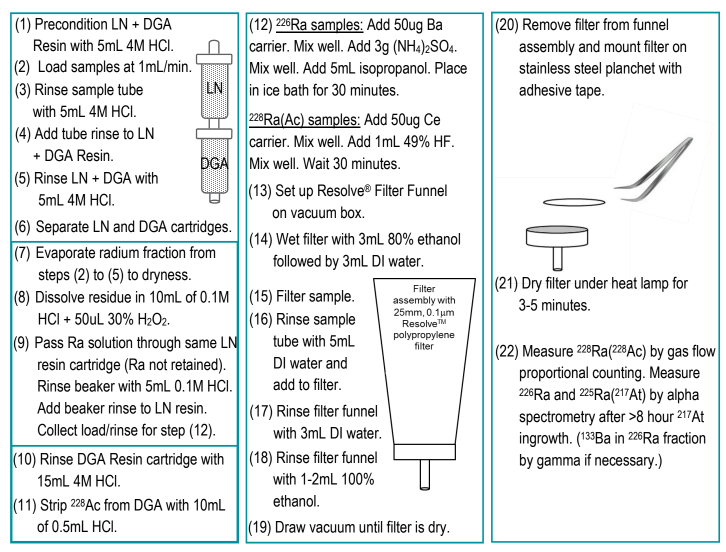


Figure 2. LN-DGA Separation and Alpha Source Preparation



Method Performance ^{226/228} Ra in Water						
	Volume		¹³³ Ba Tracer	% Recovery	% Recovery	
Sample	Liters	Replicates	% Recovery	²²⁶ Ra	²²⁸ Ra	
River Water	1.5	3	101 <u>+</u> 5	103 <u>+</u> 1	103 <u>+</u> 7	
Ground Water	1.0	5	95 <u>+</u> 4	104 <u>+</u> 1	102 <u>+</u> 8	

1040pCi ¹³³Ba, 5.0pCi ²²⁶Ra, 20pCi ²²⁸Ra

References

1) Sherrod L. Maxwell, "Rapid Method for ²²⁶Ra and ²²⁸Ra in Water Samples," *J. Radioanal. Nucl. Chem.*, 270(3), 651-655 (2006).

AN-1418-10

Rapid Determination of ²²⁶Ra in Water Samples

Summary of Method ²²⁶Ra is separated from up to 1 liter water samples and measured by alpha spectrometry. Radium is precipitated from samples with calcium carbonate. The calcium carbonate precipitate is dissolved in hydrochloric acid, and cation exchange chromatography is used to purify radium and barium from matrix ions. Eichrom DGA Resin is used to remove other alpha emitting nuclides from radium. Samples are prepared for radium measurement by alpha spectrometry using barium sulfate micro-precipitation method onto Eichrom[®] Resolve Filters. Sample preparation, including alpha spectrometry source preparation, for batches of 12-24 samples can be completed by a single operator in as little as 3-4 hours. Yields can be traced with ¹³³Ba by gamma spectrometry or ²²⁵Ra(²²⁹Th) by alpha spectrometry. If tracing with ²²⁵Ra, >8 hours of ingrowth time for the alpha emitting ²¹⁷At daughter of ²²⁵Ra is required prior to measurement by alpha spectrometry.

Reagents

Cation Exchange Resin (Eichrom C8-B500-F-H) DGA Resin, Normal 2mL Cartridges (Eichrom DN-R50-S) Ammonium Hydroxide (listed as 28% NH₃ or 56% NH₄OH) ¹³³Ba or ²²⁵Ra(²²⁹Th) Tracer

Nitric Acid (70%) 1.25M Ca(NO₃)₂ Barium Carrier (1mg/mL) Ammonium Sulfate Denatured Ethanol H₂O₂ (30%)

Equipment

Hydrochloric Acid (37%) 2M Na₂CO₃ Isopropyl Alcohol Ascorbic Acid Deionized Water

Figure 1. Sample Preparation

Water sample +Tracer ¹³³Ba or ²²⁵Ra(²²⁹Th) +10mL 56% NH₄OH Mix well.

Add 3mL 1.25M Ca(NO₃)₂ and 10mL 2M Na₂CO₃. Mix well.*

*When minimizing reagent blank is important, a calcium phosphate ppt. (AN1401) should be used.

Plastic Chromatography Column (Eichrom AC-50E-5M) Column Extension Funnel (Eichrom AC-20X-20M) Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filter in Disposable Funnel (Eichrom RF-DF-25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Stainless Steel Planchets with adhesive tape Hotplate Alpha Spectrometry System Gamma Spectrometry System (if ¹³³Ba tracer used) 150mL Glass beakers Vacuum Pump Heat Lamp

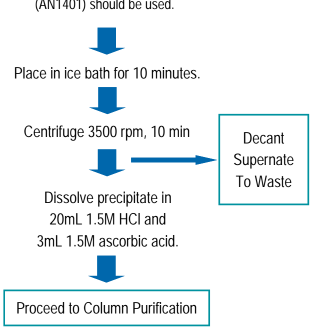
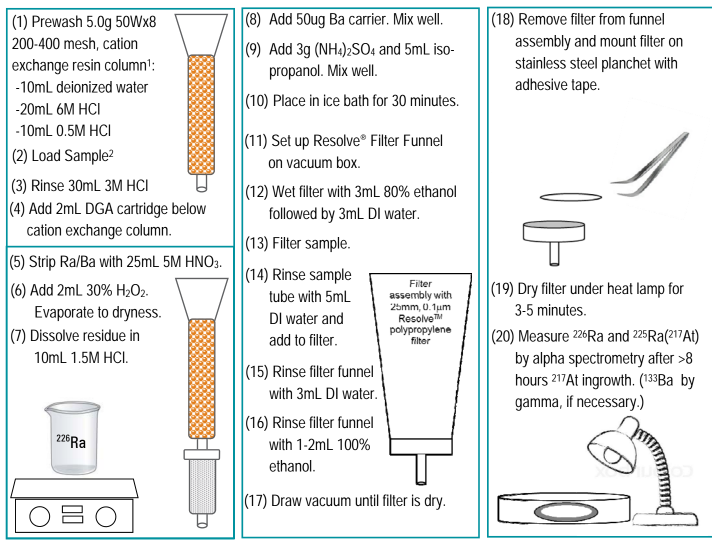


Figure 2. Column Purification and Alpha Source Preparation



¹If using ¹³³Ba tracer, 3.0g of cation exchange resin and proportionally smaller rinse volumes may be used. ²If tracing with ²²⁹Th, a 20mL 1M HCI-1M H₃PO₄ rinse following the sample load can improve purity of final ²²⁶Ra fraction.

~~~

|        | Method Performance <sup>226</sup> Ra in Water |                                                                                         |                   |       |  |  |  |
|--------|-----------------------------------------------|-----------------------------------------------------------------------------------------|-------------------|-------|--|--|--|
|        | <sup>225</sup> Ra( <sup>217</sup> At)         | <sup>225</sup> Ra( <sup>217</sup> At) <sup>226</sup> Ra(mBq/L) <sup>226</sup> Ra(mBq/L) |                   |       |  |  |  |
| Sample | % Yield*                                      | Reference                                                                               | Measured          | %Bias |  |  |  |
| 1      | 84.8                                          | 73.8                                                                                    | 69.6              | -5.7  |  |  |  |
| 2      | 87.3                                          | 73.8                                                                                    | 75.7              | 2.6   |  |  |  |
| 3      | 86.2                                          | 73.8                                                                                    | 71.3              | -3.4  |  |  |  |
| 4      | 98.7                                          | 73.8                                                                                    | 66.9              | -9.3  |  |  |  |
| AVG    | 89 <u>+</u> 6                                 | 73.8                                                                                    | 70.9 <u>+</u> 3.7 | -3.9  |  |  |  |

\*<sup>225</sup>Ra tracer is added as <sup>229</sup>Th in equilibrium with its daughters and measured by its alpha emitting <sup>217</sup>At daughter (7.066MeV) after
 >8 hr ingrowth.

### References

1) Sherrod L. Maxwell, Brian K. Culligan, "Rapid Determination of <sup>226</sup>Ra in Environmental Samples," *J. Radioanal. Nucl. Chem.*, 293(1), 149-155 (2012).

# Rapid Determination of <sup>226</sup>Ra in Concrete and Brick

#### AN-1419-10

**Summary of Method** <sup>226</sup>Ra is separated from 1 gram samples of concrete and brick and measured by alpha spectrometry. Samples are fused with sodium hydroxide at 600°C. The fusion cake is dissolved in water, and radium is precipitated from samples with calcium carbonate. The calcium carbonate precipitate is dissolved in hydrochloric acid, and cation exchange chromatography is used to purify radium and barium from matrix ions. Barium is removed from samples using Eichrom Sr Resin. Eichrom DGA Resin is used to separate other alpha emitting nuclides from radium. Samples are prepared for radium measurement by alpha spectrometry via barium sulfate micro-precipitation onto Eichrom<sup>®</sup> Resolve Filters. Sample preparation including alpha spectrometry source preparation for batches of 12 samples can be completed by a single operator in as little as 6 hours, with 85-90% yield of Radium. Yields are traced with <sup>225</sup>Ra(<sup>229</sup>Th) by alpha spectrometry. At least 8 hours of ingrowth time for the alpha emitting <sup>217</sup>At daughter of <sup>225</sup>Ra is required prior to measurement by alpha spectrometry.

#### Reagents

Cation Exchange Resin (Eichrom C8-B500-F-H) Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) DGA Resin, Normal 2mL Cartridges (Eichrom DN-R50-S) Nitric Acid (70%) Hydrochloric Acid (37%) <sup>225</sup>Ra(<sup>229</sup>Th) Tracer **Deionized Water** 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> 2M Na<sub>2</sub>CO<sub>3</sub> Barium Carrier (1mg/mL) Isopropyl Alcohol Ammonium Sulfate Sodium Hydroxide **Denatured Ethanol** Ascorbic Acid  $H_2O_2(30\%)$ 

#### Equipment

Plastic Chromatography Column (Eichrom AC-50E-5M) Column Extension Funnel (Eichrom AC-20X-20M) Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Resolve Filter in Disposable Funnel (Eichrom RF-DF-25-25PP01) Yellow Outer Tips (Eichrom AC-1000-OT) 50mL and 250mL Centrifuge Tubes Centrifuge Stainless Steel Planchets with adhesive tape Hotplate Alpha Spectrometry System 150mL Glass beakers Vacuum Pump 250mL Zirconium Crucible w/ lid Muffle Furnace Heat Lamp

#### Figure 1. Sample Preparation

1g finely milled concrete or brick +Tracer <sup>225</sup>Ra(<sup>229</sup>Th) +10g NaOH in Zr crucible.

Fuse at 600°C in muffle furnace for 15 minutes.

Remove from furnace. Cool 10 minutes. Dissolve fusion cake with 100mL DI water. Transfer to 250mL centrifuge tube. Add 10mL 37% HCI. Dilute to 150mL.

Add 0.5mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> and 10mL 2M Na<sub>2</sub>CO<sub>3</sub>. Mix well.

Place in ice bath for 10 minutes. Centrifuge 3500 rpm, 10 min

Decant

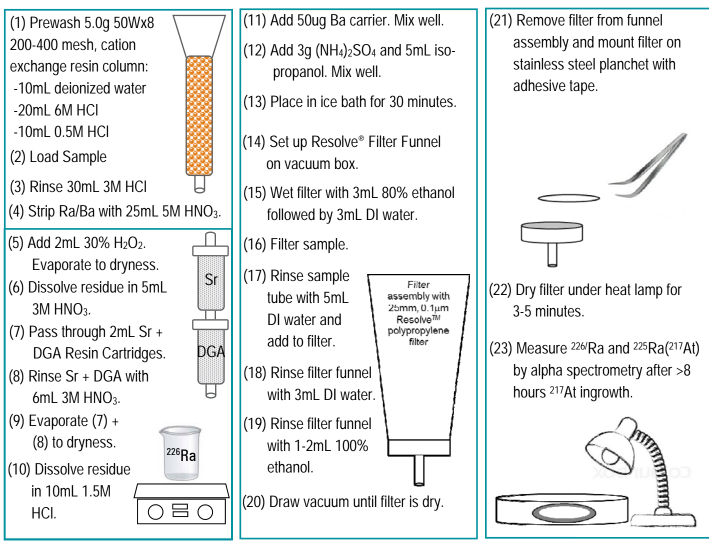
Supernate

To Waste

Dissolve precipitate in 20mL 1.5M HCl and 3mL 1.5M ascorbic acid.

Proceed to Column Purification

# Figure 2. Column Purification and Alpha Source Preparation



<sup>1</sup>If using <sup>133</sup>Ba tracer, 3.0g of cation exchange resin and proportionally smaller rinse volumes may be used. <sup>2</sup>If tracing with <sup>229</sup>Th, a 20mL 1M HCI-1M H<sub>3</sub>PO<sub>4</sub> rinse following the sample load can improve purity of final <sup>226</sup>Ra fraction.

| Method Performance <sup>226</sup> Ra in Concrete and Brick |                                       |                                                                                                        |                                                                                                                                                 |                                                                                                                                                                                                       |  |  |
|------------------------------------------------------------|---------------------------------------|--------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--|--|
|                                                            | <sup>225</sup> Ra( <sup>217</sup> At) | <sup>226</sup> Ra(mBq/g)                                                                               | <sup>226</sup> Ra(mBq/g)                                                                                                                        |                                                                                                                                                                                                       |  |  |
| Replicates                                                 | % Yield*                              | Reference                                                                                              | Measured                                                                                                                                        | %Bias                                                                                                                                                                                                 |  |  |
| 6                                                          | 85 <u>+</u> 7                         | 184.5                                                                                                  | 181 <u>+</u> 4                                                                                                                                  | -1.9                                                                                                                                                                                                  |  |  |
| 6                                                          | 87 <u>+</u> 7                         | 73.8                                                                                                   | 77.8 <u>+</u> 4.6                                                                                                                               | 5.4                                                                                                                                                                                                   |  |  |
|                                                            | Replicates<br>6                       | <sup>225</sup> Ra( <sup>217</sup> At)           Replicates         % Yield*           6         85 ± 7 | 225Ra( <sup>217</sup> At)         226Ra(mBq/g)           Replicates         % Yield*         Reference           6         85 ± 7         184.5 | 225Ra( <sup>217</sup> At)         226Ra(mBq/g)         226Ra(mBq/g)           Replicates         % Yield*         Reference         Measured           6         85 ± 7         184.5         181 ± 4 |  |  |

\*<sup>225</sup>Ra tracer is added as <sup>229</sup>Th in equilibrium with its daughters and measured by its alpha emitting <sup>217</sup>At daughter (7.066MeV) after >8 hr ingrowth.

# References

1) Sherrod L. Maxwell, Brian K. Culligan, "Rapid Determination of <sup>226</sup>Ra in Environmental Samples," *J. Radioanal. Nucl. Chem., 293(1), 149-155* (2012).

# Rapid Determination of <sup>226</sup>Ra in Glass Fiber Air Filters

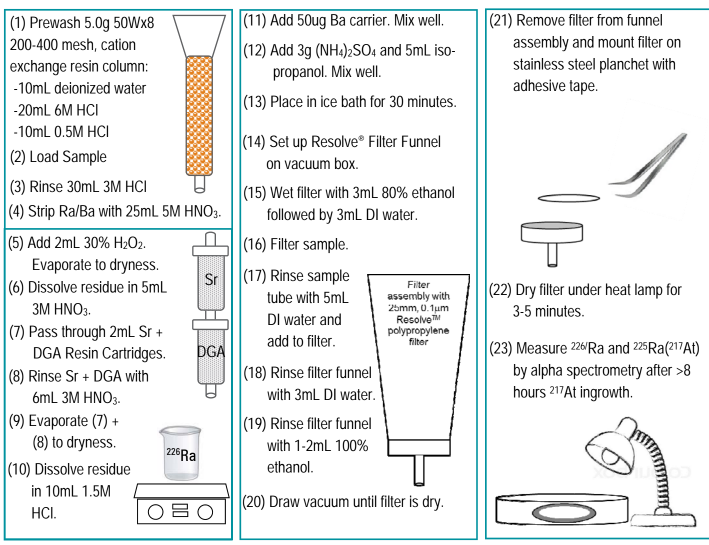
#### AN-1420-10

**Summary of Method** <sup>226</sup>Ra is separated from 47mm glass fiber air filters and measured by alpha spectrometry. Samples are fused with sodium hydroxide at 600°C. The fusion cake is dissolved in water, and radium is precipitated from samples with calcium carbonate. The calcium carbonate precipitate is dissolved in hydrochloric acid, and cation exchange chromatography is used to purify radium and barium from matrix ions. Barium is removed from samples using Eichrom Sr Resin. Eichrom DGA Resin is used to separate radium from other alpha emitting nuclides. Samples are prepared for alpha spectrometry by barium sulfate micro-precipitation onto Eichrom<sup>®</sup> Resolve Filters. Sample preparation, including alpha spectrometry source preparation, for batches of 12 samples can be completed by a single operator in as little as 6 hours, with 85-90% yield of Radium. Yields are traced with <sup>225</sup>Ra(<sup>229</sup>Th) by alpha spectrometry. At least 8 hours of ingrowth time for the alpha emitting <sup>217</sup>At daughter of <sup>225</sup>Ra is required prior to measurement by alpha spectrometry.

#### Reagents

Cation Exchange Resin (Eichrom C8-B500-F-H) Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) Figure 1. Sample Preparation DGA Resin, Normal 2mL Cartridges (Eichrom DN-R5S) Nitric Acid (70%) Hydrochloric Acid (37%) 47mm glass fiber air filter **Deionized Water** <sup>225</sup>Ra(<sup>229</sup>Th) Tracer +Tracer 225Ra(229Th) 1.25M Ca(NO<sub>3</sub>)<sub>2</sub>  $2M Na_2CO_3$ +10g NaOH in Zr crucible. Barium Carrier (1mg/mL) Isopropyl Alcohol Denatured Fthanol Ammonium Sulfate Fuse at 600°C in muffle furnace for 15 minutes. Ascorbic Acid Sodium Hydroxide  $H_2O_2(30\%)$ Remove from furnace. Cool 10 minutes. Equipment Dissolve fusion cake with 100mL DI water. Plastic Chromatography Column (Eichrom AC-50E-5M) Transfer to 250mL centrifuge tube. Add Column Extension Funnel (Eichrom AC-20X-20M) 10mL 37% HCI. Dilute to 150mL. Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Add 0.5mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> and Yellow Outer Tips (Eichrom AR-1000-OT) 10mL 2M Na<sub>2</sub>CO<sub>3</sub>. Mix well. Resolve Filter in Disposable Funnel (Eichrom RF-DF-25-25PP01) 50mL and 250mL Centrifuge Tubes Place in ice bath for 10 minutes. Centrifuge Stainless Steel Planchets with adhesive tape Centrifuge 3500 rpm, 10 min Decant Hotplate Supernate Alpha Spectrometry System 150mL Glass beakers To Waste Dissolve precipitate in Vacuum Pump 20mL 1.5M HCI and 250mL Zirconium Crucible w/ lid 3mL 1.5M ascorbic acid. Muffle Furnace Heat Lamp Proceed to Column Purification

# Figure 2. Column Purification and Alpha Source Preparation



<sup>1</sup>If using <sup>133</sup>Ba tracer, 3.0g of cation exchange resin and proportionally smaller rinse volumes may be used. <sup>2</sup>If tracing with <sup>229</sup>Th, a 20mL 1M HCI-1M H<sub>3</sub>PO<sub>4</sub> rinse following the sample load can improve purity of final <sup>226</sup>Ra fraction.

| I      | Method Performance <sup>226</sup> Ra in 47mm Glass Fiber Air Filter |                               |                               |       |  |  |
|--------|---------------------------------------------------------------------|-------------------------------|-------------------------------|-------|--|--|
|        | <sup>225</sup> Ra( <sup>217</sup> At)                               | <sup>226</sup> Ra(mBq/filter) | <sup>226</sup> Ra(mBq/filter) |       |  |  |
| Sample | % Yield*                                                            | Reference                     | Measured                      | %Bias |  |  |
| 1      | 80.7                                                                | 73.8                          | 70.5                          | -4.5  |  |  |
| 2      | 79.9                                                                | 73.8                          | 80.8                          | 9.5   |  |  |
| 3      | 78.6                                                                | 73.8                          | 77.0                          | 4.3   |  |  |
| 4      | 73.0                                                                | 73.8                          | 79.5                          | 7.7   |  |  |
| 5      | 71.5                                                                | 73.8                          | 77.7                          | 5.3   |  |  |
| AVG    | 77 <u>+</u> 4                                                       | 73.8                          | 77 <u>+</u> 4                 | 4.3   |  |  |

| Method Perform                        | ance <sup>226</sup> Ra in 47mn | n Glass Fiber Air Filter      |
|---------------------------------------|--------------------------------|-------------------------------|
| <sup>225</sup> Ra( <sup>217</sup> At) | <sup>226</sup> Ra(mBɑ/filter)  | <sup>226</sup> Ra(mBg/filter) |

\*<sup>225</sup>Ra tracer is added as <sup>229</sup>Th in equilibrium with its daughters and measured by its alpha emitting <sup>217</sup>At daughter (7.066MeV) after >8 hr ingrowth.

### References

1) Sherrod L. Maxwell, Brian K. Culligan, "Rapid Determination of <sup>226</sup>Ra in Environmental Samples," J. Radioanal. Nucl. Chem., 293(1), 149-155 (2012).

# Rapid Determination of <sup>226</sup>Ra in 1g Soil Samples

#### AN-1421-10

**Summary of Method** <sup>226</sup>Ra is separated from 1 gram samples of soil and measured by alpha spectrometry. Samples are fused with sodium hydroxide at 600°C. The fusion cake is dissolved in water, and radium is precipitated from samples with calcium carbonate. The calcium carbonate precipitate is dissolved in hydrochloric acid, and cation exchange chromatography is used to purify radium and barium from matrix ions. Barium is removed from samples using Eichrom Sr Resin. Eichrom DGA Resin is used to reomve other alpha emitting nuclides from radium. Samples are prepared for alpha spectrometry by barium sulfate micro-precipitation onto Eichrom<sup>®</sup> Resolve Filters. Sample preparation, including alpha spectrometry source preparation, for batches of 12-24 samples can be completed by a single operator in as little as 6 hours. Yields are traced with <sup>225</sup>Ra(<sup>229</sup>Th) by alpha spectrometry. At least 8 hours of ingrowth time for the alpha emitting <sup>217</sup>At daughter of <sup>225</sup>Ra is required prior to measurement by alpha spectrometry.

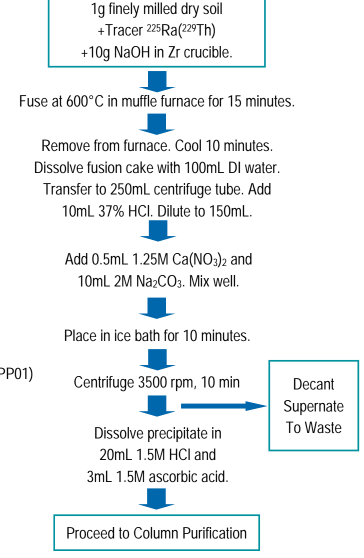
#### Reagents

Cation Exchange Resin (Eichrom C8-B500-F-H) Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) DGA Resin, Normal 2mL Cartridges (Eichrom DN-R50-S) Hydrochloric Acid (37%) Nitric Acid (70%) **Deionized Water** <sup>225</sup>Ra(<sup>229</sup>Th) Tracer 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> 2M Na<sub>2</sub>CO<sub>3</sub> Barium Carrier (1mg/mL) Isopropyl Alcohol Ammonium Sulfate **Denatured Ethanol** Ascorbic Acid Sodium Hydroxide  $H_2O_2(30\%)$ 

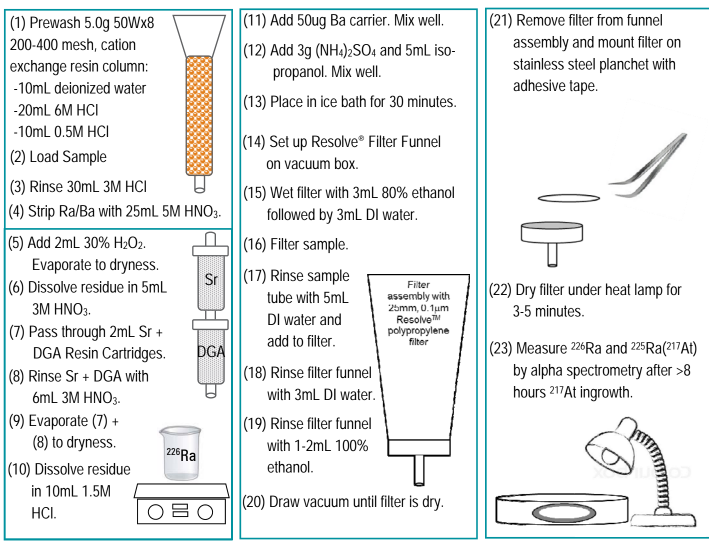
#### Equipment

Plastic Chromatography Column (Eichrom AC-50E-5M) Column Extension Funnel (Eichrom AC-20X-20M) Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filter in Disposable Funnel (Eichrom RF-DF-25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Stainless Steel Planchets with adhesive tape Hotplate Alpha Spectrometry System 150mL Glass beakers Vacuum Pump 250mL Zirconium Crucible w/ lid Muffle Furnace

#### Figure 1. Sample Preparation



# Figure 2. Column Purification and Alpha Source Preparation



<sup>1</sup>If using <sup>133</sup>Ba tracer, 3.0g of cation exchange resin and proportionally smaller rinse volumes may be used. <sup>2</sup>If tracing with <sup>229</sup>Th, a 20mL 1M HCI-1M H<sub>3</sub>PO<sub>4</sub> rinse following the sample load can improve purity of final <sup>226</sup>Ra fraction.

| Method Performance <sup>226</sup> Ra in 1g Soil Samples |                                       |                                                                                         |                |       |  |  |  |
|---------------------------------------------------------|---------------------------------------|-----------------------------------------------------------------------------------------|----------------|-------|--|--|--|
|                                                         | <sup>225</sup> Ra( <sup>217</sup> At) | <sup>225</sup> Ra( <sup>217</sup> At) <sup>226</sup> Ra(mBq/g) <sup>226</sup> Ra(mBq/g) |                |       |  |  |  |
| Sample                                                  | % Yield*                              | Reference                                                                               | Measured       | %Bias |  |  |  |
| 1                                                       | 75.2                                  | 184.5                                                                                   | 185.9          | 0.8   |  |  |  |
| 2                                                       | 77.9                                  | 184.5                                                                                   | 192.0          | 4.1   |  |  |  |
| 3                                                       | 74.8                                  | 184.5                                                                                   | 176.9          | -4.1  |  |  |  |
| 4                                                       | 73.3                                  | 184.5                                                                                   | 184.7          | 0.1   |  |  |  |
| AVG                                                     | 75 <u>+</u> 2                         | 184.5                                                                                   | 185 <u>+</u> 6 | 0.3   |  |  |  |

 $^{*225}$ Ra tracer is added as  $^{229}$ Th in equilibrium with its daughters and measured by its alpha emitting  $^{217}$ At daughter (7.066MeV) after >8 hr ingrowth.

# References

1) Sherrod L. Maxwell, Brian K. Culligan, "Rapid Determination of <sup>226</sup>Ra in Environmental Samples," *J. Radioanal. Nucl. Chem.*, 293(1), 149-155 (2012).

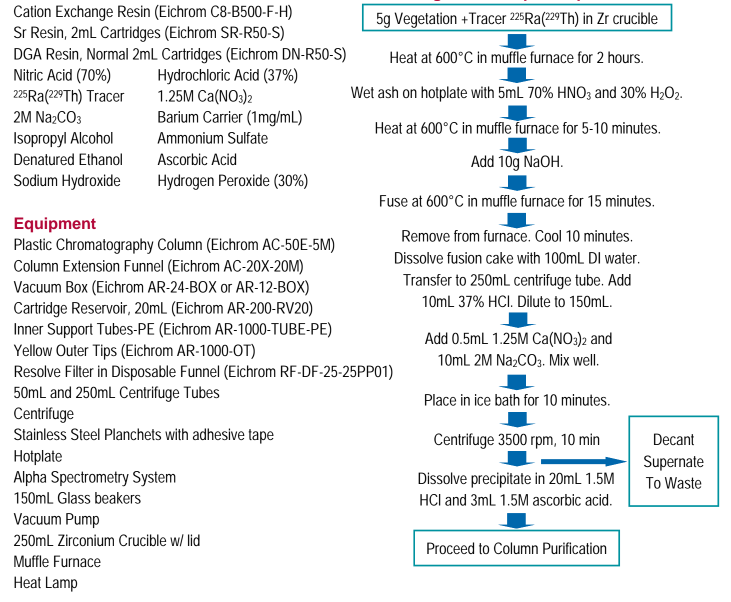
AN-1422-10

# Rapid Determination of <sup>226</sup>Ra in 5g Vegetation Samples

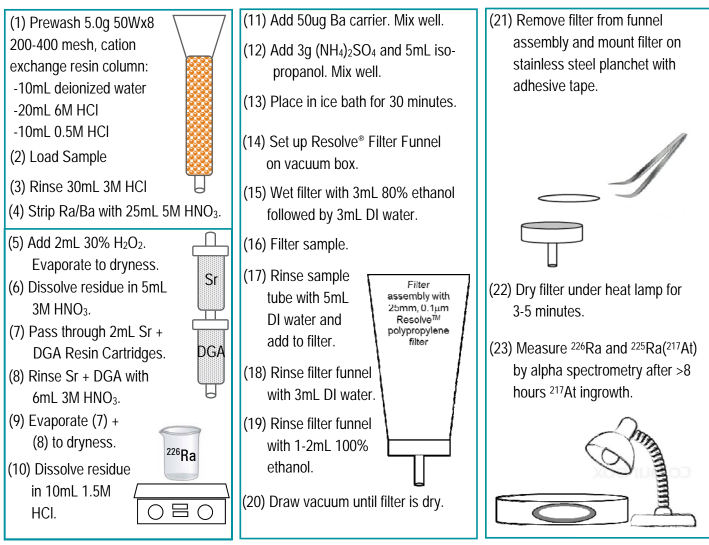
Figure 1. Sample Preparation

**Summary of Method** <sup>226</sup>Ra is separated from 5 gram samples of vegetation and measured by alpha spectrometry. Samples are fused with sodium hydroxide at 600°C. The fusion cake is dissolved in water, and radium is precipitated from samples with calcium carbonate. The calcium carbonate precipitate is dissolved in hydrochloric acid, and cation exchange chromatography is used to purify radium and barium from matrix ions. Barium is removed from samples using Eichrom Sr Resin. Eichrom DGA Resin is used to separate other alpha emitting nuclides from radium. Samples are prepared for alpha spectrometry by barium sulfate micro-precipitation onto Eichrom<sup>®</sup> Resolve Filters. Sample preparation, including alpha spectrometry source preparation, for batches of 12 samples can be completed by a single operator in as little as 6 hours. Yields are traced with <sup>225</sup>Ra(<sup>229</sup>Th) by alpha spectrometry. At least 8 hours of ingrowth time for the alpha emitting <sup>217</sup>At daughter of <sup>225</sup>Ra is required prior to measurement by alpha spectrometry.

#### Reagents



# Figure 2. Column Purification and Alpha Source Preparation



<sup>1</sup>If using <sup>133</sup>Ba tracer, 3.0g of cation exchange resin and proportionally smaller rinse volumes may be used. <sup>2</sup>If tracing with <sup>229</sup>Th, a 20mL 1M HCI-1M H<sub>3</sub>PO<sub>4</sub> rinse following the sample load can improve purity of final <sup>226</sup>Ra fraction.

| Method Performance <sup>226</sup> Ra in 5g Vegetation Samples |                                       |                          |                          |       |
|---------------------------------------------------------------|---------------------------------------|--------------------------|--------------------------|-------|
|                                                               | <sup>225</sup> Ra( <sup>217</sup> At) | <sup>226</sup> Ra(mBq/g) | <sup>226</sup> Ra(mBq/g) |       |
| Sample**                                                      | % Yield*                              | Reference                | Measured                 | %Bias |
| 1                                                             | 91.5                                  | 73.8                     | 70.8                     | -4.1  |
| 2                                                             | 88.3                                  | 73.8                     | 73.8                     | 0.0   |
| 3                                                             | 93.1                                  | 73.8                     | 69.8                     | -5.4  |
| 4                                                             | 82.2                                  | 73.8                     | 68.5                     | -7.2  |
| 5                                                             | 80.2                                  | 73.8                     | 81.4                     | 10.3  |
| AVG                                                           | 87 <u>+</u> 6                         | 73.8                     | 73 <u>+</u> 5            | -1.1  |

 $^{*225}$ Ra tracer is added as  $^{229}$ Th in equilibrium with its daughters and measured by its alpha emitting  $^{217}$ At daughter (7.066MeV) after >8 hours ingrowth.

\*\*5 grams of blank hay matrix spiked with <sup>226</sup>Ra

## References

1) Sherrod L. Maxwell, Brian K. Culligan, "Rapid Determination of <sup>226</sup>Ra in Environmental Samples," *J. Radioanal. Nucl. Chem.*, 293(1), 149-155 (2012).

# Rapid Determination of Pu, Np, and U in 1-8L Seawater Samples

#### AN-1423-10

**Summary of Method** Plutonium, Neptunium, and Uranium are separated and concentrated from up to 8L samples of seawater with a hydrous titanium oxide precipitation, enhanced with 5mg of lanthanum and 125mg of ferric iron. A second precipitation with lanthanum fluoride removes additional matrix ions, and Uranium and Pu+Np are separated from potentially interfering radionuclides in the sample using stacked 2mL cartridges of Eichrom TEVA and TRU Resins. Isotopic U and Pu+Np are measured by alpha spectrometry following cerium fluoride microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Chemical yields are determined by recovery of <sup>232</sup>U and <sup>242</sup>Pu (or <sup>236</sup>Pu if measuring <sup>237</sup>Np) tracers. Recoveries of <sup>232</sup>U average 95  $\pm$  6%, while <sup>236</sup>Pu average 90  $\pm$  9%. Measured values of <sup>238</sup>U, <sup>239</sup>Pu, and <sup>237</sup>Np typically agree to within 10% of reference value. A single operator can process batches of 12 samples through alpha source preparation in 6-8 hours. Alpha spectrometry count times will vary depending on desired detection limit and data quality objectives.

#### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) Nitric Acid (70%) Hydrochloric Acid (37%) Hydrofluoric Acid (49%) or Sodium Fluoride Ammonium Hydroxide (listed as 28% NH<sub>3</sub> or 56% NH<sub>4</sub>OH) Iron Carrier (50mg/mL Fe, as ferric nitrate) Lanthanum and Cerium Carriers (1mg/mL) <sup>232</sup>U and <sup>242</sup>Pu(or <sup>236</sup>Pu if meas. <sup>237</sup>Np) tracers Oxalic acid/Ammonium Oxalate Deionized Water  $H_2O_2(30\%)$ 10% (w:w) TiCl<sub>3</sub>  $2M AI(NO_3)_3$ Boric acid Sulfamic Acid NaNO<sub>2</sub> Ascorbic Acid Denatured Ethanol 1.25M Ca(NO<sub>3</sub>)<sub>2</sub>

### Equipment

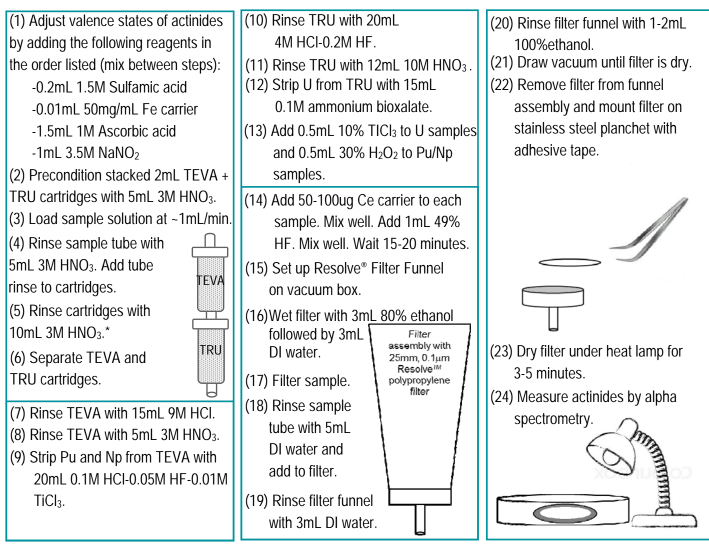
Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250-500mL Centrifuge Tubes Centrifuge Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Analytical Balance Vacuum Pump Heat Lamp

### Figure 1. Sample Preparation

1-8L Sample of Seawater. Acidify to pH 2 with 70% HCI. Add tracers. Mix well. Add 5mg La, 125mg Fe and 10mL 10% TiCl<sub>3</sub>. Mix well. Add 5mL 56% NH<sub>4</sub>OH. Mix well. Allow precipitate to settle. Decant supernate to <1L. Transfer remaining supernate and precipitate to 250-500mL centrifuge tubes. Centrifuge 3000rpm for 10 minutes. Decant supernate. Repeat until all sample processed. Partially dissolve precipitate in 80mL 1M HCI. Some solids will remain. Add 1mg La, 25mg Ca, 5mL 10% TiCl<sub>3</sub>. Mix. Add 25mL 49% HF. Mix. Centrifuge. Decant Supernate. Dissolve precipitate with 5mL 3M HNO<sub>3</sub>-0.25M boric acid, 7mL 2M AI(NO<sub>3</sub>)<sub>3</sub> and 8mL 7M HNO<sub>3</sub>.

Load solution to valence adjustment and TEVA-TRU separation.

# Figure 2. TEVA-TRU Separation and Alpha Source Preparation



\*Adding 50uL of 30% H<sub>2</sub>O<sub>2</sub> to tube rinse can improve Uranium recoveries and decontamination in Pu(Np) fractions.

|                   |           |            |                   | % Tracer      | Analyte(mBq/L) | Analyte(mBq/L)    |        |
|-------------------|-----------|------------|-------------------|---------------|----------------|-------------------|--------|
| Analyte           | Volume, L | Replicates | Tracer            | Recovery      | Reference      | Measured          | % Bias |
| <sup>239</sup> Pu | 2         | 5          | <sup>236</sup> Pu | 91 <u>+</u> 9 | 33.8           | 32.6 <u>+</u> 1.4 | -3.6   |
| <sup>239</sup> Pu | 4         | 1          | <sup>236</sup> Pu | 86            | 16.9           | 16.2              | -4.1   |
| <sup>239</sup> Pu | 8         | 2          | <sup>236</sup> Pu | 87 <u>+</u> 3 | 27.8           | 27.6 <u>+</u> 0.5 | -0.7   |
| <sup>237</sup> Np | 2         | 5          | <sup>236</sup> Pu | 91 <u>+</u> 9 | 17.4           | 17.7 <u>+</u> 1.5 | 1.7    |
| <sup>237</sup> Np | 4         | 1          | <sup>236</sup> Pu | 86            | 8.7            | 7.2               | -17    |
| <sup>237</sup> Np | 8         | 2          | <sup>236</sup> Pu | 87 <u>+</u> 3 | 4.4            | 4.2 <u>+</u> 0.4  | -4.5   |
| <sup>238</sup> U  | 2         | 5          | <sup>232</sup> U  | 99 <u>+</u> 2 | 51.8           | 49.3 <u>+</u> 1.5 | -4.8   |
| <sup>238</sup> U  | 4         | 1          | <sup>232</sup> U  | 86            | 25.9           | 25.0              | -3.6   |
| <sup>238</sup> U  | 8         | 2          | <sup>232</sup> U  | 92 <u>+</u> 5 | 96.3           | 94 <u>+</u> 3     | -2.4   |

Method Performance Pu, Np and U from Seawater

16 hour count times

#### References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchinson, Robin C. Utsey, Daniel R. McAlister, "Rapid determination of actinides in seawater samples," *J. Radioanal. Nucl. Chem., 300(3), 1175-1189* (2014).

# Rapid Determination of Pu, Am, and Cm in 80L Seawater Samples

#### AN-1424-10

**Summary of Method** Plutonium, Americium, and Curium are separated and concentrated from up to 80L samples of seawater with a hydrous titanium oxide precipitation, enhanced with lanthanum and ferric iron. A second precipitation with lanthanum fluoride removes additional matrix ions, and Pu and Am+Cm are separated from potentially interfering radionuclides in the sample using stacked 2mL cartridges of Eichrom TEVA and DGA Resins. Isotopic Pu and Am+Cm are measured by alpha spectrometry following cerium fluoride microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Chemical yields are determined by recovery of <sup>243</sup>Am and <sup>242</sup>Pu tracers. Recoveries of <sup>243</sup>Am average 94  $\pm$  3%, while <sup>242</sup>Pu average 86  $\pm$  4%. Measured values of <sup>239</sup>Pu, <sup>241</sup>Am, and <sup>244</sup>Cm typically agree to within 10% of reference values. A single operator can process batches of 12 samples through alpha source preparation in 6-8 hours. Alpha spectrometry count times will vary depending on desired detection limit and data quality objectives.

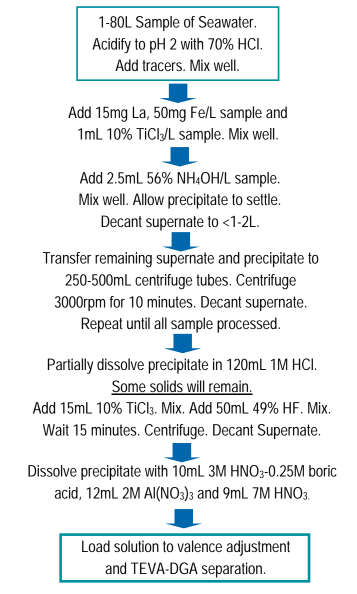
#### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Ammonium Hydroxide (listed as 28% NH<sub>3</sub> or 56% NH<sub>4</sub>OH) Nitric Acid (70%) Hydrochloric Acid (37%) Hydrofluoric Acid (49%) or Sodium Fluoride **Deionized Water** Iron Carrier (50mg/mL Fe, as ferric nitrate) Lanthanum and Cerium Carriers (1mg/mL) <sup>243</sup>Am and <sup>242</sup>Pu tracers 10% (w:w) TiCl<sub>3</sub>  $H_2O_2(30\%)$  $2M AI(NO_3)_3$ Boric acid Sulfamic Acid Ascorbic Acid NaNO<sub>2</sub> **Denatured Ethanol** 

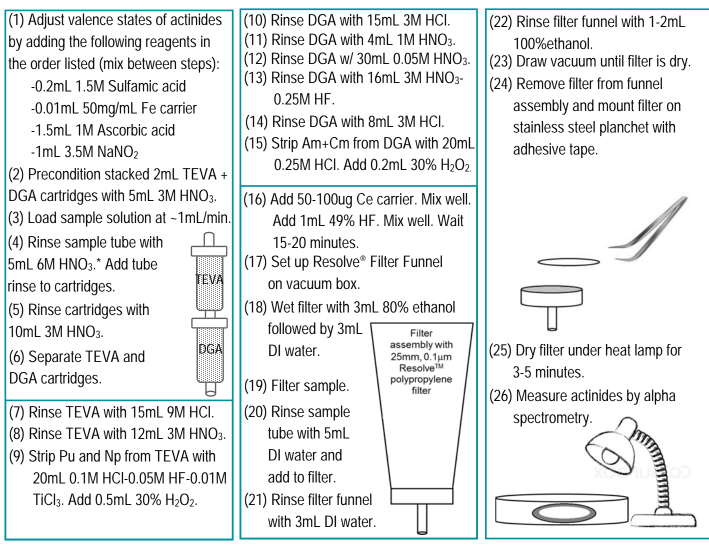
#### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250-500mL Centrifuge Tubes Centrifuge Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Analytical Balance Vacuum Pump Heat Lamp

### Figure 1. Sample Preparation



# Figure 2. TEVA-DGA Separation and Alpha Source Preparation



\*Adding 50uL of 30% H<sub>2</sub>O<sub>2</sub> to the tube rinse can improve Uranium recoveries and decontamination in Pu(Np) fractions.

|                   |           | Method P   | erforman          | ce Pu, Am a       | nd Cm from Seaw | ater               |        |
|-------------------|-----------|------------|-------------------|-------------------|-----------------|--------------------|--------|
|                   |           |            |                   | % Tracer          | Analyte(mBq/L)  | Analyte(mBq/L)     |        |
| Analyte           | Volume, L | Replicates | Tracer            | Recovery          | Reference       | Measured           | % Bias |
| <sup>239</sup> Pu | 16        | 2          | <sup>242</sup> Pu | 90 <u>+</u> 1     | 4.22            | 4.67 <u>+</u> 0.05 | 11     |
| <sup>239</sup> Pu | 25        | 2          | <sup>242</sup> Pu | 84.6 <u>+</u> 0.2 | 3.22            | 3.3 <u>+</u> 0.1   | 2.5    |
| <sup>239</sup> Pu | 40        | 2          | <sup>242</sup> Pu | 86 <u>+</u> 2     | 0.81            | 0.82 <u>+</u> 0.02 | 1.2    |
| <sup>239</sup> Pu | 80        | 2          | <sup>242</sup> Pu | 85 <u>+</u> 5     | 0.40            | 0.37 <u>+</u> 0.01 | -7.5   |
| <sup>241</sup> Am | 16        | 2          | <sup>243</sup> Am | 95 <u>+</u> 4     | 3.31            | 3.1 <u>+</u> 0.1   | -6.3   |
| <sup>241</sup> Am | 25        | 2          | <sup>243</sup> Am | 93.1 <u>+</u> 0.1 | 2.12            | 1.9 <u>+</u> 0.1   | -10    |
| <sup>241</sup> Am | 40        | 2          | <sup>243</sup> Am | 96 <u>+</u> 2     | 0.53            | 0.51 <u>+</u> 0.02 | -3.8   |
| <sup>241</sup> Am | 80        | 2          | <sup>243</sup> Am | 93 <u>+</u> 4     | 0.27            | 0.25 <u>+</u> 0.01 | -7.4   |
| <sup>244</sup> Cm | 16        | 2          | <sup>243</sup> Am | 95 + 4            | 2.16            | 2.1 + 0.2          | -2.8   |
| <sup>244</sup> Cm | 25        | 2          | <sup>243</sup> Am | 93.1 <u>+</u> 0.1 | 1.35            | 1.3 <u>+</u> 0.1   | -3.7   |
| <sup>244</sup> Cm | 40        | 2          | <sup>243</sup> Am | 96 <u>+</u> 2     | 0.85            | 0.78 <u>+</u> 0.04 | -8.2   |
| <sup>244</sup> Cm | 80        | 2          | <sup>243</sup> Am | 93 <u>+</u> 4     | 0.42            | 0.41 <u>+</u> 0.01 | -2.3   |

| ethod Performance | Pu, Am | and Cm | from | Seawater |  |
|-------------------|--------|--------|------|----------|--|

16 hour count times

# References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchinson, Robin C. Utsey, Daniel R. McAlister, "Rapid determination of actinides in seawater samples," J. Radioanal. Nucl. Chem., 300(3), 1175-1189 (2014).

# Rapid Determination of Actinides in 10g Emergency Food Samples

#### AN-1425-10

**Summary of Method** U, Pu, Np, Am and Cm are separated and concentrated from 10 gram food samples. Samples are muffled at 600°C in zirconium crucibles 2 hours to destroy organic content. The residue is wet ashed with HNO<sub>3</sub>-H<sub>2</sub>O<sub>2</sub> and then fused with 15g NaOH at 600°C for ten minutes. The fusion cakes are dissolved in water, transferred to 250mL centrifuge tubes and precipitated sequentially with hydrous titanium oxide and lanthanum fluoride to facilitate matrix removal. Actinides are separated on stacked 2mL cartridges of Eichrom TEVA, TRU and DGA resins. Actinides are measured by alpha spectrometry following CeF<sub>3</sub> microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Chemical yields of tracers ranged from 93-98% for <sup>236</sup>Pu, 85-93% for <sup>243</sup>Am, and 78-89% for <sup>232</sup>U. Measured values typically agreed to within 10% of reference values. Sample preparation for batches of 12 samples can be completed by a single operator in <8 hours. Alpha spectrometry count times will depend on detection limit and data quality objectives.

#### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) **Deionized Water** 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu (or <sup>236</sup>Pu if meas. Np), <sup>243</sup>Am and <sup>232</sup>U tracers Oxalic acid/Ammonium oxalate La and Ce carriers (1mg/mL) 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>  $2M AI(NO_3)_3$ 10% (w:w) TiCl<sub>3</sub> HNO<sub>3</sub> (70%) HCI (37%) NaOH HF (49%) or NaF Boric acid  $H_2O_2$  (30%) NaNO<sub>2</sub> Denatured ethanol Sulfamic Acid Ascorbic Acid

#### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Heat Lamp Muffle Furnace Hot Plate Analytical Balance 250mL Zirconium crucibles with zirconium lids Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Vacuum Pump

#### Figure 1. Sample Preparation

10g Food sample + tracers in zirconium crucible . Muffle at 600°C for 2 hours. Wet ash on hotplate with 5mL 70% HNO<sub>3</sub> and 5mL 30% H<sub>2</sub>O<sub>2</sub>. Fuse samples with 15g NaOH at 600°C for 10minutes. Dissolve fusion cake with H<sub>2</sub>O. Transfer to 250mL c-tube. Add 10mL 3M HNO<sub>3</sub> to crucible. Heat to dissolve residue. Transfer to same 25mL c-tube. Add 125mg Fe and 5mg La to c-tube. Dilute to 180mL. Add 4mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub>, 5mL 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>, 5mL 10% TiCl<sub>3</sub>. Mix. Cool in ice bath for 10min. Centrifuge at 3500rpm. Decant Supernate. Partially dissolve precipitate in 60mL 1M HCI. Some solids will remain. Dilute to 170mL. Add 1mg La, 1mL 1.25M Ca(NO<sub>3</sub>)<sub>3</sub>, 3mL 10% TiCl<sub>3</sub>. Mix. Add 20mL 49% HF. Centrifuge at 3500rpm. Decant Supernate. Dissolve precipitate in 5mL 3M HNO<sub>3</sub>-0.25M Boric acid, 7mL 70% HNO<sub>3</sub>, and 7mL 2M AI(NO<sub>3</sub>)<sub>3</sub>. Fix valence states. Mix between each addition of: 0.5mL 1.5M sulfamic acid, 10uL 50mg/mL Fe, 1.5mL 1M ascorbic acid, 1mL 3.5M NaNO<sub>2</sub>, 1.5mL 70% HNO<sub>3</sub>.

# Figure 2. Actinide Separation on TEVA - TRU - DGA\* and Source Preparation

|                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         | -                                                                                                                                                                                                                                                                               |
|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <ul> <li>(1) Precondition stacked 2mL<br/>TEVA, TRU, DGA cartridges<br/>with 10mL 3M HNO<sub>3</sub>.</li> <li>(2) Load sample solution.</li> <li>(3) Rinse sample tube<br/>with 5mL 3M HNO<sub>3</sub>. Add<br/>tube rinse to cartridges.</li> <li>(4) Rinse cartridges with<br/>10mL 3M HNO<sub>3</sub>.*</li> <li>(5) Separate TEVA, TRU,<br/>and DGA cartridges.</li> <li>(6) Rinse TEVA cartridge with:<br/>-10mL 3M HNO<sub>3</sub><br/>-20mL 9M HCI</li> </ul> | <ul> <li>(12) Rinse DGA cartridge sequentially</li> <li>with: -5mL 3M HCI</li> <li>-3mL 1M HNO3</li> <li>-15mL 0.05M HNO3</li> <li>(13) Strip Am and Cm from DGA with</li> <li>10mL 0.25M HCI.</li> <li>(14) Rinse TRU cartridge with 15mL</li> <li>4M HCI-0.2M HF-0.002M TiCl3 .</li> <li>(15) Strip U from TRU with 15mL of</li> <li>0.1M ammonium bioxalate.</li> <li>(16) Add 0.5mL 10% TiCl3 to U samples,</li> <li>0.5mL 30% H2O2 to Pu, and 0.2mL</li> <li>30% H2O2 to Am/Cm samples.</li> </ul> | <ul> <li>(22) Rinse filter<br/>funnel with 3mL<br/>DI water and 2mL<br/>100%ethanol.</li> <li>(23) Draw vacuum<br/>until filter is dry.</li> <li>(24) Remove filter from<br/>funnel assembly and<br/>mount filter on stainless<br/>steel planchet with 2-sided tape.</li> </ul> |
| -5mL 3M HNO <sub>3</sub><br>(7) Strip Pu (and Np) from TEVA<br>cartridge with 20mL 0.1M HCI-<br>0.05MHF-0.01M TiCI <sub>3</sub> .                                                                                                                                                                                                                                                                                                                                     | (17) Add 50-100ug Ce carrier to all<br>samples. Mix well. Add 1mL 49% HF.<br>Mix well. Wait 15-20 minutes.                                                                                                                                                                                                                                                                                                                                                                                              |                                                                                                                                                                                                                                                                                 |
| <ul> <li>(8) Rinse DGA cartridge with 10mL</li> <li>0.1M HNO<sub>3</sub>.</li> <li>(9) Place TRU cartridge above DGA.</li> <li>(10) Strip Am/Cm from TRU onto<br/>DGA with 15mL 3M HCI .</li> <li>(11) Separate TRU cartridge from<br/>DGA cartridge.</li> </ul>                                                                                                                                                                                                      | <ul> <li>(18) Set up Resolve<sup>®</sup> Filter Funnel on vacuum box.</li> <li>(19) Wet filter with 3mL 80% ethanol followed by 3mL DI water.</li> <li>(20) Filter sample.</li> <li>(21) Rinse sample tube with 5mL DI water and add to filter.</li> </ul>                                                                                                                                                                                                                                              | <ul> <li>(25) Dry filter under heat lamp for 3-5 minutes.</li> <li>(26) Measure actinides by alpha spectrometry.</li> </ul>                                                                                                                                                     |

\*Adding 50uL 30% H<sub>2</sub>O<sub>2</sub> to the tube rinse can improve Uranium recoveries and decontamination in Pu(Np) fractions.

| Method Performance Actinides in 10 Gram | n Food Samples (16 hour count times) |
|-----------------------------------------|--------------------------------------|
|-----------------------------------------|--------------------------------------|

|           |            |                   |                   |                   | Analyte   | Analyte          |        |        | •          |                   |                   |                   | Analyte   | Analyte          |        |
|-----------|------------|-------------------|-------------------|-------------------|-----------|------------------|--------|--------|------------|-------------------|-------------------|-------------------|-----------|------------------|--------|
|           |            |                   |                   | % Tracer          | (mBq/g)   | (mBq/g)          |        |        |            |                   |                   | % Tracer          | (mBq/g)   | (mBq/g)          |        |
| Sample    | Replicates | Analyte           | Tracer            | Recovery          | Reference | Measured         | % Bias | Sample | Replicates | Analyte           | Tracer            | Recovery          | Reference | Measured         | % Bias |
| Baby Food | 5          | <sup>238</sup> Pu | <sup>236</sup> Pu | 93.5 <u>+</u> 7.5 | 2.9       | 2.9 <u>+</u> 0.1 | -0.7   | Apples | 5          | <sup>238</sup> Pu | <sup>236</sup> Pu | 98 <u>+</u> 12    | 2.9       | 2.9 <u>+</u> 0.1 | -0.5   |
|           | 5          | <sup>239</sup> Pu | <sup>236</sup> Pu | 93.5 <u>+</u> 7.5 | 3.6       | 3.3 <u>+</u> 0.4 | -7.9   |        | 5          | <sup>239</sup> Pu | <sup>236</sup> Pu | 98 <u>+</u> 12    | 3.6       | 3.6 <u>+</u> 0.4 | -0.9   |
|           | 5          | <sup>237</sup> Np | <sup>236</sup> Pu | 93.5 <u>+</u> 7.5 | 3.7       | 3.4 <u>+</u> 0.2 | -8.1   |        | 5          | <sup>237</sup> Np | <sup>236</sup> Pu | 98 <u>+</u> 12    | 3.7       | 3.3 <u>+</u> 0.1 | -11.5  |
|           | 5          | <sup>241</sup> Am | <sup>243</sup> Am | 84.6 + 6.3        | 5.1       | 5.0 <u>+</u> 0.1 | -3.5   |        | 5          | <sup>241</sup> Am | <sup>243</sup> Am | 93.4 <u>+</u> 8.5 | 5.1       | 4.9 <u>+</u> 0.3 | -2.8   |
|           | 5          | <sup>244</sup> Cm | <sup>243</sup> Am | 84.6 <u>+</u> 6.3 | 3.5       | 3.7 <u>+</u> 0.3 | 4.4    |        | 5          | <sup>244</sup> Cm | <sup>243</sup> Am | 93.4 <u>+</u> 8.5 | 3.5       | 3.7 <u>+</u> 0.5 | 6.3    |
|           | 5          | <sup>238</sup> U  | <sup>232</sup> U  | 78 <u>+</u> 10    | 5.7       | 5.6 <u>+</u> 0.4 | -1.5   |        | 5          | <sup>238</sup> U  | <sup>232</sup> U  | 89 <u>+</u> 10    | 5.7       | 5.6 <u>+</u> 0.3 | -1.2   |
|           | 5          | <sup>234</sup> U  | <sup>232</sup> U  | 78 <u>+</u> 10    | 5.9       | 5.9 <u>+</u> 0.2 | -0.3   |        | 5          | <sup>234</sup> U  | <sup>232</sup> U  | 89 <u>+</u> 10    | 5.9       | 5.5 <u>+</u> 0.4 | -6.4   |

### References

1) Sherrod L. Maxwell, Brian K. Culligan, Angel Kelsy-Wall, Patrick J. Shaw, "Rapid separation of actinides and in emergency food samples," *J. Radioanal. Nucl. Chem.*, 292(1), 339-347 (2011).

# **Cichrom**<sup>•</sup> Rapid Determination of Actinides in 100g Emergency Food Samples

**Summary of Method** U, Pu, Np, Am and Cm are separated and concentrated from 100gram food samples. Samples are muffled at 600°C in zirconium crucibles 2 hours to destroy organic content. The residue is wet ashed with HNO<sub>3</sub>-H<sub>2</sub>O<sub>2</sub> and then fused with 15g NaOH at 600°C for ten minutes. The fusion cakes are dissolved in water, transferred to 250mL centrifuge tubes and precipitated sequentially with hydrous titanium oxide and lanthanum fluoride to facilitate matrix removal. Actinides are separated on stacked 2mL cartridges of Eichrom TEVA, TRU and DGA resins. Actinides are measured by alpha spectrometry following CeF<sub>3</sub> microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Chemical yields of tracers ranged from 93-98% for <sup>236</sup>Pu, 85-93% for <sup>243</sup>Am, and 78-89% for <sup>232</sup>U. Measured values typically agreed to within 10% of reference values. Sample preparation for batches of 12 samples can be completed by a single operator in <16hours. Alpha spectrometry count times will depend on detection limit and data quality objectives.

#### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu (or <sup>236</sup>Pu if meas. Np), <sup>243</sup>Am and <sup>232</sup>U tracers Oxalic acid/Ammonium oxalate La and Ce carriers (1mg/mL)

Deionized Water $1.25M Ca(NO_3)_2$  $3.2M (NH_4)_2HPO_4$  $2M Al(NO_3)_3$  $10\% (w:w) TiCl_3$  $HNO_3 (70\%)$ HCI (37%)NaOHHF (49%) or NaFBoric acid $H_2O_2 (30\%)$  $NaNO_2$ Denatured ethanolSulfamic AcidAscorbic AcidAightarrow and an analysis

### Equipment

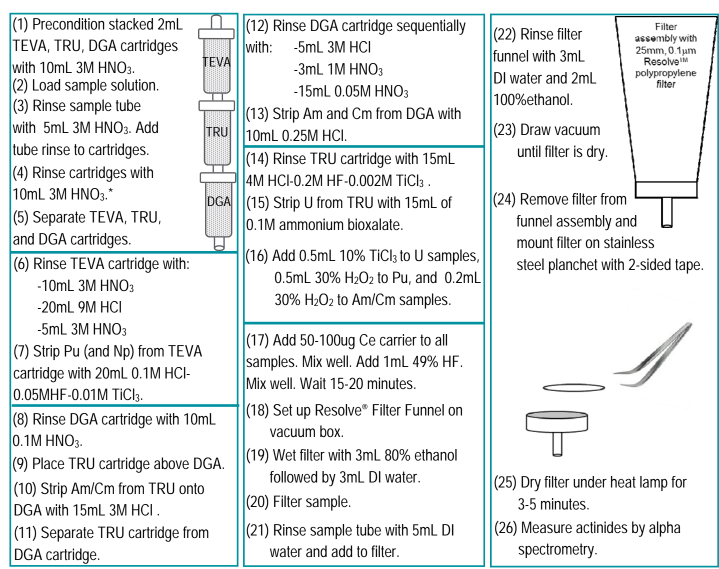
Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes 250mL Zirconium crucibles with zirconium lids Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Centrifuge Muffle Furnace Analytical Balance 1L Glass Beakers Vacuum Pump Heat Lamp

### Figure 1. Sample Preparation

100g Food sample + tracers in 1L glass beaker Muffle at 550°C for 12 hours. Transfer ash to Zr-crucible. Rinse beaker with 10mL 70% HNO<sub>3</sub> and 10mL 30% H<sub>2</sub>O<sub>2</sub>. Transfer to crucible. Wet ash. Fuse samples with 15g NaOH at 600°C for 10minutes. Dissolve fusion cake with H<sub>2</sub>O. Transfer to 250mL c-tube. Add 10mL 3M HNO<sub>3</sub> to crucible. Heat to dissolve residue. Add to same c-tube. Add 125mg Fe and 5mg La to c-tube. Dilute to 180mL. Add 4mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub>, 5mL 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>, 5mL 10% TiCl<sub>3</sub>. Mix. Cool in ice bath for 10min. Centrifuge at 3500rpm. Decant Supernate. Partillay dissolve precipitate in 60mL 1M HCI. Some solids will remain. Dilute to 170mL. Add 1mg La, 1mL 1.25M Ca(NO<sub>3</sub>)<sub>3</sub>, 3mL 10% TiCl<sub>3</sub>. Mix. Add 20mL 49% HF. Centrifuge at 3500rpm. Decant Supernate. Dissolve precipitate in 5mL 3M HNO<sub>3</sub>-0.25M Boric acid, 7mL 70% HNO<sub>3</sub>, and 7mL 2M AI(NO<sub>3</sub>)<sub>3</sub>. Fix valence states. Mix between each addition of: 0.5mL

1.5M sulfamic acid, 10uL 50mg/mL Fe, 1.5mL 1M ascorbic acid, 1mL 3.5M NaNO<sub>2</sub>, 1.5mL 70% HNO<sub>3</sub>.

## Figure 2. Actinide Separation on TEVA - TRU - DGA\* and Source Preparation



\*Adding 50uL 30% H<sub>2</sub>O<sub>2</sub> to tube rinse can improve Uranium recoveries and decontamination in Pu(Np) fractions.

#### Method Performance 100g Apple Samples (16 hr count times)

|        |            |                   |                   |               | Analyte   | Analyte            |        |
|--------|------------|-------------------|-------------------|---------------|-----------|--------------------|--------|
|        |            |                   |                   | % Tracer      | (mBq/g)   | (mBq/g)            |        |
| Sample | Replicates |                   | Tracer            | Recovery      | Reference | Measured           | % Bias |
| Apples | 5          | <sup>238</sup> Pu | <sup>236</sup> Pu | 78 <u>+</u> 8 | 0.29      | 0.30 <u>+</u> 0.02 | 3.1    |
|        | 5          | <sup>239</sup> Pu | <sup>236</sup> Pu | 78 <u>+</u> 8 | 0.36      | 0.37 <u>+</u> 0.05 | 4.0    |
|        | 5          | <sup>237</sup> Np | <sup>236</sup> Pu | 78 <u>+</u> 8 | 0.37      | 0.36 <u>+</u> 0.02 | -3.3   |
|        | 5          | <sup>241</sup> Am | <sup>243</sup> Am | 76 <u>+</u> 3 | 0.25      | 0.25 + 0.02        | -2.3   |
|        | 5          | <sup>244</sup> Cm | <sup>243</sup> Am | 76 <u>+</u> 3 | 0.35      | 0.41 <u>+</u> 0.03 | 16     |
|        | 5          | <sup>238</sup> U  | <sup>232</sup> U  | 71 <u>+</u> 5 | 0.57      | 0.56 + 0.04        | -1.4   |
|        | 5          | <sup>234</sup> U  | <sup>232</sup> U  | 71 <u>+</u> 5 | 0.59      | 0.58 <u>+</u> 0.05 | -2.7   |

### References

1) Sherrod L. Maxwell, Brian K. Culligan, Angel Kelsy-Wall, Patrick J. Shaw, "Rapid separation of actinides and in emergency food samples," *J. Radioanal. Nucl. Chem.*, 292(1), 339-347 (2011).

# Rapid Determination of Plutonium in Large Rice Samples

#### AN-1427-10

**Summary of Method** Plutonium is separated and measured from up to 1.5kg rice samples. Rice samples are muffled and wet ashed to reduce volume and destroy organic content. The residue is then fused with sodium hydroxide. Precipitation steps remove additional matrix and prepare plutonium for separation on Eichrom TEVA resin. Plutonium is measured by alpha spectrometry following cerium fluoride microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Plutonium recovery through the method, determined using <sup>242</sup>Pu tracer, was  $87 \pm 4\%$  for 1kg samples. Measured values for <sup>239</sup>Pu and <sup>238</sup>Pu agreed within 6% of reference values, even when refractory <sup>239</sup>Pu was present in the sample. Sample preparation can be completed in less than 48 hours.

#### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S)

Iron carrier (50mg/mL Fe, as ferric iron nitrate)

La carrier (10mg/mL) Ce carrier (1mg/mL) Deionized Water 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> 10% (w:w) TiCl<sub>3</sub> HCI (37%) HF (49%) or NaF H<sub>2</sub>O<sub>2</sub> (30%) Denatured ethanol Ascorbic Acid

1.25M Ca(NO<sub>3</sub>)<sub>2</sub> 2M AI(NO<sub>3</sub>)<sub>3</sub> HNO<sub>3</sub> (70%) NaOH Boric acid NaNO<sub>2</sub> Sulfamic Acid <sup>242</sup>Pu tracer

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes 250mL Zirconium crucibles with zirconium lids Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Centrifuge Heat Lamp Muffle Furnace Hot Plate Analytical Balance Vacuum Pump 600mL glass beakers

#### **Figure 1. Sample Preparation**

Rice Sample + tracers in 600mL glass beaker(s). Multiple beakers may be needed for large samples.

Muffle for 5 hours at 350°C, then 550°C for 12 hours.

Carefully wet ash with enough 1:1 (v:v) 70% HNO<sub>3</sub>:30%  $H_2O_2$  to cover sample.

Heat to dryness. Muffle 550°C for 6-12 hours.

Transfer residue to Zr crucible. Rinse beaker with 70% HNO<sub>3</sub>. Add rinse to crucible. Wet ash with 1:1 70% HNO<sub>3</sub>:30% H<sub>2</sub>O<sub>2</sub>. Heat to dryness.

Repeat wet ash until no black char remains (violet residue common).

Fuse samples with 15g NaOH at 600°C for 20-30 minutes.

Dissolve fusion cake with H<sub>2</sub>O. Transfer to 250mL c-tube.

Add 10mL 3M HNO $_3$  to crucible. Heat to dissolve residue. Transfer to same 25mL c-tube.

Add 125mg Fe and 10mg La to c-tube. Dilute to 180mL.

Add 2mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub>, 1mL 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>, 6mL 10% TiCl<sub>3</sub>. Mix. Cool in ice bath for 10min.

Centrifuge at 3500rpm. Decant Supernate.

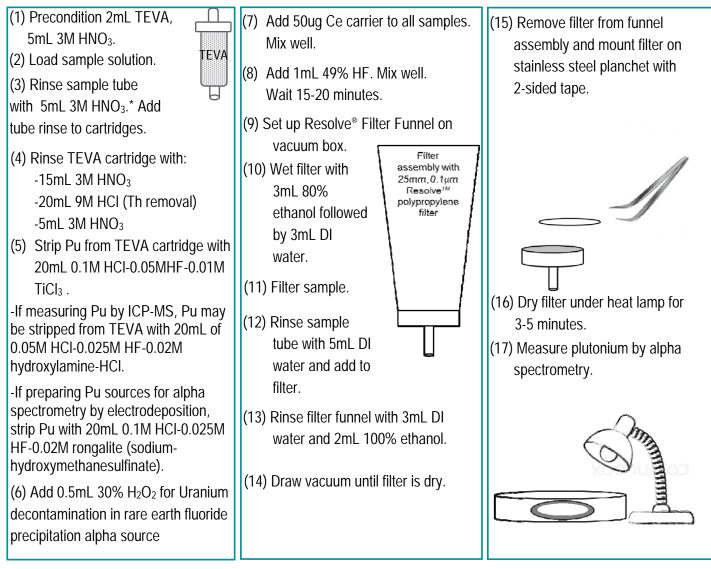
Partially dissolve precipitate in 100mL 1.5M HCl. Some solids will remain. Dilute to 170mL. Add 5mL 10% TiCl<sub>3</sub> and 22mL 49% HF. Mix.

Centrifuge at 3500rpm. Decant Supernate.

Dissolve precipitate in 5mL 3M HNO<sub>3</sub>-0.25M Boric acid, 7mL 6M HNO<sub>3</sub>, and 7mL 2M Al(NO<sub>3</sub>)<sub>3</sub>. Cool to room temperature.

Fix valence states. Mix between each addition of: 0.5mL 1.5M sulfamic acid, 40uL 50mg/mL Fe, 1.5mL 1M ascorbic acid, 1mL 3.5M NaNO<sub>2</sub>.

# Figure 2. Plutonium Separation on TEVA Resin and Source Preparation



\*Adding 50uL of 30%  $H_2O_2$  to tube rinse can improve Uranium decontamination.

#### Method Performance

|             |            | <sup>242</sup> Pu |                   |                   |                   |                   |                   |                   |
|-------------|------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|
|             |            | Tracer            | Reference         | e (mBq/kg)        | Measured          | (mBq/kg)          | % E               | Bias              |
| Sample (kg) | Replicates | % Yield           | <sup>239</sup> Pu | <sup>238</sup> Pu | <sup>239</sup> Pu | <sup>238</sup> Pu | <sup>239</sup> Pu | <sup>238</sup> Pu |
| 1.0         | 8          | 87 <u>+</u> 4     | 12.5              | 10.6              | 11.8 <u>+</u> 1.0 | 10.5 <u>+</u> 0.7 | -5.6              | -0.7              |

MDA for 1 kg sample, 30hours count time, 0.37uBq/kg

## References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchinson, "Rapid fusion method for determination of plutonium isotopes in large rice samples," *J. Radioanal. Nucl. Chem.*, 298(2), 1367-1374 (2013).

# Rapid Determination of Actinides in Fecal Samples

#### AN-1428-10

**Summary of Method** Actinides are separated and measured from fecal samples. Fecal samples are muffled and wet ashed prior to fusion with sodium hydroxide. Sequential precipitation steps remove sample matrix prior to actinide separation on 2mL cartridges of Eichrom TEVA, TRU and DGA resins. Actinides are measured by alpha spectrometry following cerium fluoride microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Samples can be prepared for measurement in less than 24 hours.

#### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu (or <sup>236</sup>Pu if meas. Np), <sup>243</sup>Am and <sup>232</sup>U tracers Oxalic acid/Ammonium oxalate

Ce carrier (1mg/mL)

1.25M Ca(NO<sub>3</sub>)<sub>2</sub>

 $2M AI(NO_3)_3$ 

HNO<sub>3</sub> (70%)

NaOH

NaNO<sub>2</sub>

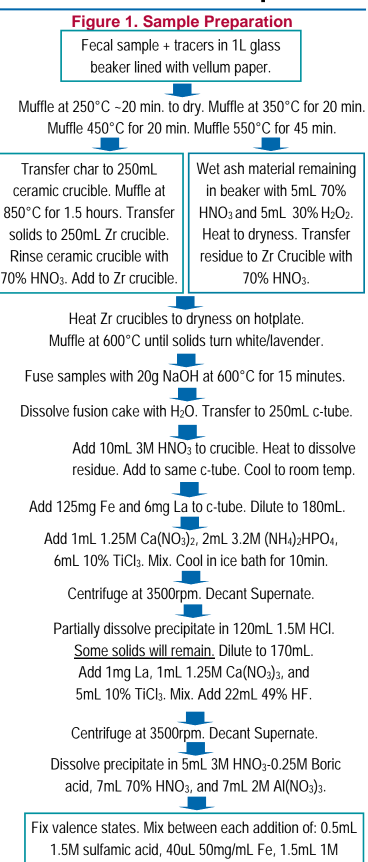
Boric acid

Sulfamic Acid

La carrier (10mg/mL) Deionized Water 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> 10% (w:w) TiCl<sub>3</sub> HCI (37%) HF (49%) or NaF H<sub>2</sub>O<sub>2</sub> (30%) Denatured ethanol Ascorbic Acid

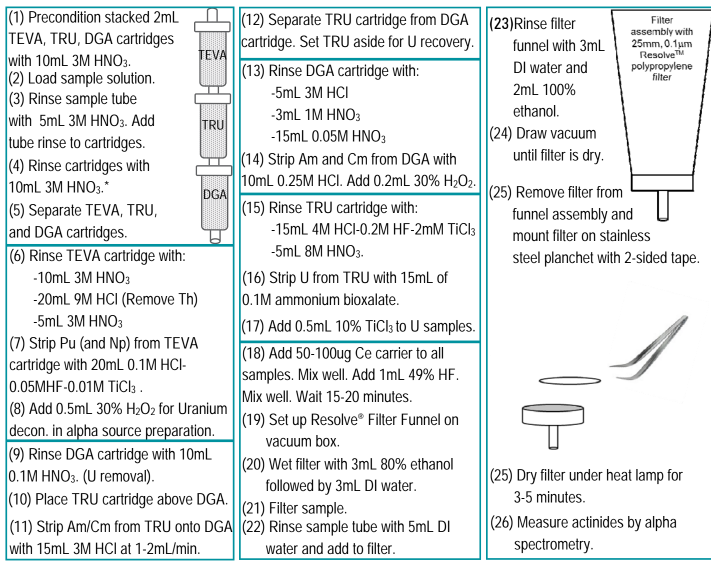
Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes 250mL Ceramic crucibles Hot Plate 250mL Zirconium crucibles with zirconium lids Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Vellum paper Centrifuge Muffle Furnace Analytical Balance **1L Glass Beakers** Vacuum Pump Heat Lamp



ascorbic acid, 1mL 3.5M NaNO<sub>2</sub>, 1.5mL 70% HNO<sub>3</sub>.

## Figure 2. Actinide Separation on TEVA - TRU - DGA\* and Source Preparation



\*Adding 50uL 30% H<sub>2</sub>O<sub>2</sub> can improve Uranium recoveries and decontamination in Pu(Np) fractions.

Method Performance

| Analyte               | Samples | Tracer            | % Tracer<br>Recovery | Reference<br>(Bq/sample) | Measurement<br>(Bq/sample) | % Bias      |
|-----------------------|---------|-------------------|----------------------|--------------------------|----------------------------|-------------|
| <sup>239/240</sup> Pu | 5       | <sup>242</sup> Pu | 95 <u>+</u> 9        | 0.085 - 0.204            | 0.081 - 0.198              | -11 to -1.5 |
| <sup>238</sup> Pu     | 5       | <sup>242</sup> Pu | 95 <u>+</u> 9        | 0.066 - 0.156            | 0.071 - 0.146              | -5.3 to 3.0 |
| <sup>241</sup> Am     | 5       | <sup>243</sup> Am | 83 <u>+</u> 4        | 0.199 - 0.476            | 0.201 - 0.464              | -11 to 1.0  |
| <sup>238</sup> U      | 5       | <sup>232</sup> U  | 63 <u>+</u> 7        | 0.226 - 0.541            | 0.196 - 0.592              | -9.0 to 2.9 |
| <sup>234</sup> U      | 5       | <sup>232</sup> U  | 63 <u>+</u> 7        | 0.218 - 0.521            | 0.206 - 0.536              | -13 to 9.4  |

6 hour count time

### References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchinson, Ronie B. Spencer "Rapid fusion method for determination of actinides in fecal samples," *J. Radioanal. Nucl. Chem.*, 298(3), 1533-1542 (2013).

# **Rapid Determination of Actinides in Asphalt Samples**

#### AN-1429-10

Summary of Method Actinides are separated and measured from 1g samples of asphalt. Asphalt samples are fused in zirconium crucibles with sodium hydroxide. Sequential precipitations remove matrix prior to separation of actinides on 2mL cartridges of Eichrom TRU and DGA resins. Actinides are measured by alpha spectrometry following cerium fluoride microprecipitation onto Eichrom Resolve® Filters. Chemical recoveries averaged 91±6%, 84+12%, and 86+7%, respectively, for <sup>242</sup>Pu, <sup>243</sup>Am and <sup>232</sup>U tracers. Measured values typically agreed to within 2-6% of reference values. Batches of 12 samples can be prepared for measurement in as little as 4 hours.

#### Reagents

TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu (or <sup>236</sup>Pu if meas. Np), <sup>243</sup>Am and <sup>232</sup>U tracers Oxalic acid/Ammonium oxalate

La carrier (10mg/mL) **Deionized Water** 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> 10% (w:w) TiCl<sub>3</sub> HCI (37%) HF (49%) or NaF  $H_2O_2$  (30%) Denatured ethanol Ascorbic Acid

Ce carrier (1mg/mL) 1.25M Ca(NO<sub>3</sub>)<sub>2</sub>  $2M AI(NO_3)_3$ HNO<sub>3</sub> (70%) NaOH Boric acid NaNO<sub>2</sub> Sulfamic Acid

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Muffle Furnace **Analytical Balance** 250mL Zirconium crucibles with zirconium lids Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Vacuum Pump Heat Lamp

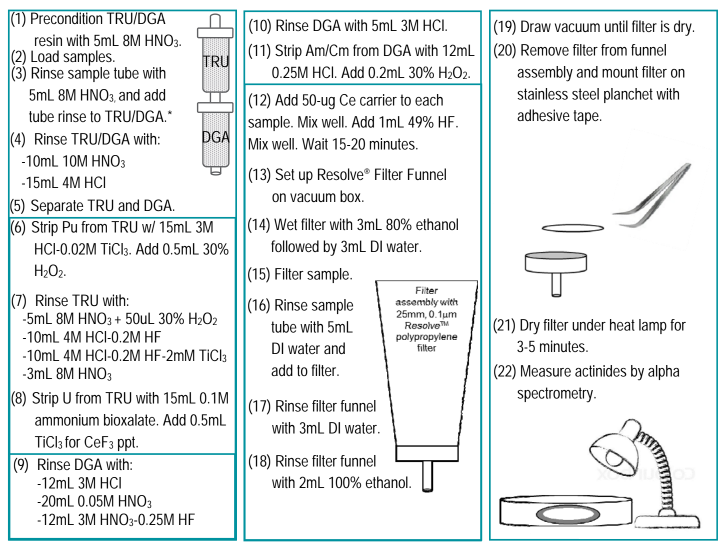
# 1q finely ground asphalt + tracers in 250mL Zr-crucible Fuse samples with 15g NaOH at 600°C for 15 minutes. Add 10mL 3M HNO<sub>3</sub> to crucible. Heat to dissolve residue. Add to same c-tube. Add 125mg Fe and 5mg La to c-tube. Dilute to 180mL. Add 2mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub>, 5mL 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>, 10mL 10% TiCl<sub>3</sub>. Mix. Cool in ice bath for 10min. Centrifuge at 3500rpm. Decant Supernate. Partially dissolve precipitate in 60-80mL 1.5M HCI.\* Dilute to 170mL with 0.01M HCI. Add 1mg La and 5mL 10% TiCl<sub>3</sub>. Mix. Add 20mL 49% HF. Mix. Wait 10 min. \*The entire precipitate will not dissolve in HCI. Dissolution will be completed with the HF addition. Centrifuge at 3500rpm. Decant Supernate. Dissolve precipitate in 5mL 3M HNO<sub>3</sub>-0.25M Boric acid, 7mL 70% HNO<sub>3</sub>, and 7mL 2M AI(NO<sub>3</sub>)<sub>3</sub>.

Fix valence states. Mix between each addition of: 0.5mL 1.5M sulfamic acid, 10uL 50mg/mL Fe, 1.5mL 1M ascorbic acid, 1mL 3.5M NaNO<sub>2</sub>, 1.5mL 70% HNO<sub>3</sub>.

### Figure 1. Sample Preparation

Dissolve fusion cake with H<sub>2</sub>O. Transfer to 250mL c-tube.

### Figure 2. Actinide Separation on TRU/DGA and Source Preparation



\*Adding 50uL of 30% H<sub>2</sub>O<sub>2</sub> to the tube rinse can help improve U recoveries and decontamination in Pu(Np) fractions.

| Analyte           | Replicates | Tracer            | % Tracer<br>Recovery | Analyte<br>Reference<br>(mBq/g) | Analyte<br>Measured<br>(mBq/g) | % Bias |
|-------------------|------------|-------------------|----------------------|---------------------------------|--------------------------------|--------|
| <sup>239</sup> Pu | 8          | <sup>242</sup> Pu | 91 + 6               | 39.2                            | 40 + 2                         | 2.0    |
| <sup>241</sup> Am | 8          | <sup>243</sup> Am | 84 <u>+</u> 13       | 24.4                            | 23 + 3                         | -5.7   |
| <sup>244</sup> Cm | 8          | <sup>243</sup> Am | 84 <u>+</u> 13       | 35.5                            | 37 <u>+</u> 5                  | 4.2    |
| <sup>238</sup> U  | 8          | <sup>232</sup> U  | 86 <u>+</u> 7        | 73.6                            | 72 <u>+</u> 8                  | -2.1   |
| <sup>234</sup> U  | 8          | <sup>232</sup> U  | 86 <u>+</u> 7        | 73.6                            | 72 <u>+</u> 9                  | -2.1   |

### References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchinson, "Rapid determination of actinides and in asphalt samples," *J. Radioanal. Nucl. Chem.*, 299(3), 1891-1901 (2014).

# Rapid Determination of Actinides in Soil Samples

### AN-1430-10

**Summary of Method** Actinides are separated and measured from 1-2g samples of soil. Soil samples are fused in zirconium crucibles with sodium hydroxide. Sequential precipitations remove matrix prior to separation of actinides on 2mL cartridges of Eichrom TRU and DGA resins. Actinides are measured by alpha spectrometry following cerium fluoride microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Chemical recoveries averaged  $97\pm9\%$ ,  $96\pm7\%$ , and  $91\pm4\%$ , respectively, for <sup>242</sup>Pu, <sup>243</sup>Am and <sup>232</sup>U tracers. Measured values typically agreed to within 3% of reference values. Batches of 12 samples can be prepared for measurement in as little as 4 hours.

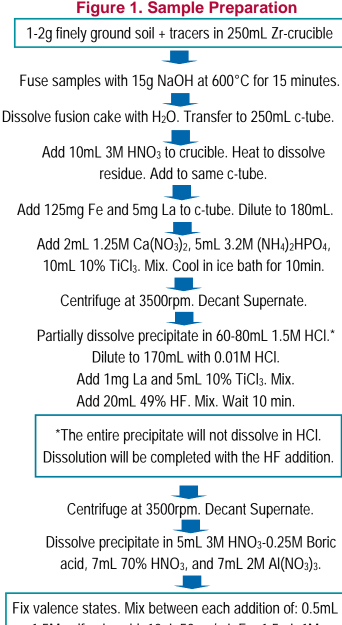
#### Reagents

TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu (or <sup>236</sup>Pu if meas. Np), <sup>243</sup>Am and <sup>232</sup>U tracers Oxalic acid/Ammonium oxalate

La carrier (10mg/mL) Deionized Water 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> 10% (w:w) TiCl<sub>3</sub> HCI (37%) HF (49%) or NaF H<sub>2</sub>O<sub>2</sub> (30%) Denatured ethanol Ascorbic Acid Ce carrier (1mg/mL) 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> 2M Al(NO<sub>3</sub>)<sub>3</sub> HNO<sub>3</sub> (70%) NaOH Boric acid NaNO<sub>2</sub> Sulfamic Acid

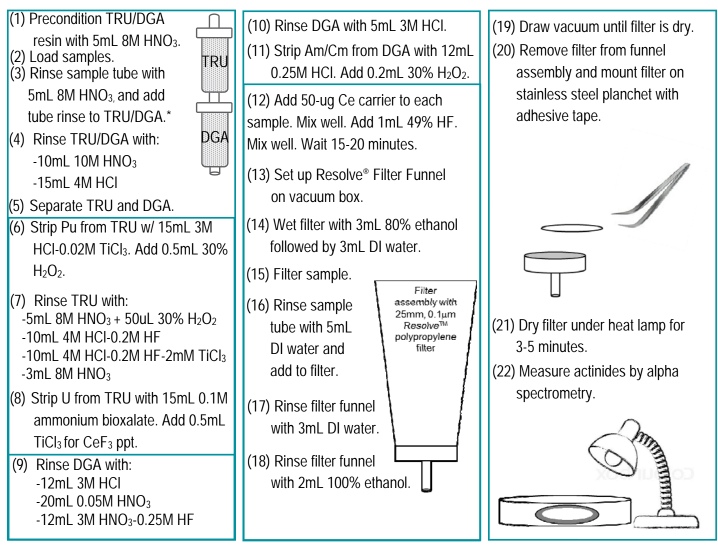
### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Muffle Furnace Analytical Balance 250mL Zirconium crucibles with zirconium lids Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Vacuum Pump Heat Lamp



1.5M sulfamic acid, 10uL 50mg/mL Fe, 1.5mL 1M ascorbic acid, 1mL 3.5M NaNO<sub>2</sub>, 1.5mL 70% HNO<sub>3</sub>.

### Figure 2. Actinide Separation on TRU/DGA and Source Preparation



\*Adding 50uL of 30% H<sub>2</sub>O<sub>2</sub> to tube rinse can improve U recoveries and decontamination in Pu(Np) fractions.

#### Method Performance

|                   |            |                   |               | Analyte   | Analyte        |        |
|-------------------|------------|-------------------|---------------|-----------|----------------|--------|
|                   |            |                   | % Tracer      | Reference | Measured       |        |
| Analyte           | Replicates | Tracer            | Recovery      | (mBq/g)   | (mBq/g)        | % Bias |
| <sup>239</sup> Pu | 7          | <sup>242</sup> Pu | 97 <u>+</u> 9 | 98.0      | 95 <u>+</u> 3  | -3.1   |
| <sup>241</sup> Am | 7          | <sup>243</sup> Am | 96 <u>+</u> 7 | 61.1      | 59 <u>+</u> 4  | -3.4   |
| <sup>238</sup> U  | 7          | <sup>232</sup> U  | 91 <u>+</u> 4 | 184       | 183 <u>+</u> 6 | -0.5   |

16 hour counts

### References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchinson, "Rapid determination of actinides and in asphalt samples," *J. Radioanal. Nucl. Chem.*, 299(3), 1891-1901 (2014).

# Rapid Determination of Pu, Np, Am and Cm in 100g Soil Samples

#### AN-1431-10

**Summary of Method** Pu(Np) and Am-Cm are separated and concentrated from 100-200 gram soil samples. Samples are muffled at 550°C to destroy organic content and wet ashed and leached with HNO<sub>3</sub> and HCI. The filtered leachates are evaporated to dryness and fused with NaOH in Zr crucibles. Sequential precipitations facilitate matrix removal. Actinides are separated on stacked 2mL cartridges of Eichrom TEVA, TRU, and DGA resins. Native rare earths from the samples are removed from Am-Cm using TEVA Resin and ammonium thiocyanate. Actinides are measured by alpha spectrometry following CeF<sub>3</sub> microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Chemical yields of tracers ranged

from 93-98% for <sup>236</sup>Pu and 85-93% for <sup>243</sup>Am. Measured values typically agreed to within 10% of reference values. Sample preparation for batches of 12 samples can be completed by a single operator in <8 hours.

#### Figure 1. Sample Preparation

Dry 100-200g soil at 110°C. Blend and Size. Aliquot sample to 1L glass beaker. Add tracers. Muffle at 550°C for 4 hours. Cool. Add 75mL 70% HNO<sub>3</sub> and 25mL 37% HCl. Heat to dryness on hotplate. Add 75mL 70% HNO<sub>3</sub>. Warm sample. Transfer solids and liquid to 250mL centrifuge tube. Add 25mL 70% HNO<sub>3</sub> to beaker. Warm beaker. Transfer solids and liquid to 250mL centrifuge tube. Centrifuge 3500 rpm, 10 min. Filter leachate through 25mm 0.45um filter. Transfer leachate to 600mL beaker. Add 25mL 70% HNO<sub>3</sub> to solids. Mix. Centrifuge. Filter leachate into same 600mL beaker. Repeat once. Add 25mL warm 4M HCI to solids. Mix. Centrifuge. Filter leachate into same 600mL beaker. Repeat once. Evaporate supernate in 600mL beaker to dryness. Wet ash with 15mL 70% HNO<sub>3</sub> to dryness. Repeat once. Transfer solids to 250mL Zr crucibles. Rinse beakers with 70% HNO<sub>3</sub>. Transfer to same crucible. Heat crucibles to dryness. Add 20g NaOH to each crucible. Fuse at 600°C for 20min. Dissolve fusion cakes with water. Transfer to 250mL c-tubes. Rinse crucibles with 10mL 3M HNO<sub>3</sub>. Heat to dissolve residue. Transfer to same c-tube. Continue to precipitation steps.

#### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu (or <sup>236</sup>Pu if meas. Np), and <sup>243</sup>Am tracers

La carrier (10mg/mL) Deionized Water 10% (w:w) TiCl<sub>3</sub> HCI (37%) HF (49%) or NaF H<sub>2</sub>O<sub>2</sub> (30%) Denatured ethanol Ascorbic Acid Formic Acid Ce carrier (1mg/mL) 2M AI(NO<sub>3</sub>)<sub>3</sub> HNO<sub>3</sub> (70%) NaOH Boric acid NaNO<sub>2</sub> Sulfamic Acid Ammonium Thiocyanate

#### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 1L and 600mL Glass beakers 250mL Zirconium crucibles with lids Stainless Steel Planchets with adhesive tape Alpha Spectrometry System 50mL and 250mL Centrifuge Tubes 25mm 0.45um filters Centrifuge Heat Lamp Muffle Furnace Hot Plate Analytical Balance Vacuum Pump

# Figure 2. Actinide Separation on TEVA - TRU - DGA and Source Preparation

| <ul> <li>Dilute samples to 180mL.</li> <li>Add 7mg La and 20mL 10% TiCl3.</li> <li>Mix. Cool to room temperature.</li> <li>Centrifuge 3500 rpm. 5min.</li> <li>Decant supernate</li> <li>Partially dissolve in 60mL 1.5M HCl.</li> <li>Solids will remain. Dilute to 170mL.</li> <li>Add 22mL 49% HF. Mix.</li> <li>Centrifuge 3500 rpm. 5min.</li> <li>Decant supernate</li> <li>Add 22mL 49% HF. Mix.</li> <li>Centrifuge 3500 rpm. 5min.</li> <li>Decant supernate</li> <li>Of Strip Pu from TEVA with 20mL</li> <li>(1) Strip Pu from TEVA with 20mL</li> <li>(1) Strip Pu from TEVA with 15mL 4M HCl.</li> <li>(2) Mad 50ug Ce carrier to all samples. Add 0.5mL 30% H<sub>2</sub>O<sub>2</sub> to Pu samples. Mix. Add 1mL 49%</li> <li>(1) Strip Pu from TEVA with 20mL</li> <li>(1) Strip Pu from TEVA with 20mL</li> <li>(1) Strip Pu from TEVA with 15mL 4M HCl.</li> <li>(2) Wet filter with 3mL 80% ethanol followed by 3mL DI water.</li> <li>(2) Wet filter with 3mL 80% ethanol followed by 3mL DI water.</li> <li>(2) Add 20mL 70% HNO3.</li> <li>(2) Wet filter sample.</li> <li>(2) Add 20mL 70% HNO3.</li> <li>(2) Wet filter funnel with 5mL DI water and add to filter.</li> <li>(2) Add 2mL 70% HNO3. +20L 10%</li> <li>(2) Add 2mL 70% HNO3. +20L 10%</li> <li>(2) Add 2mL 70% HNO3.</li> <li>(2) Add 2mL 70% HNO3. +20L 10%</li> <li>(2) Add 2mL 70% HNO3. +20L 10%</li> <li>(2) Add 2mL 70% HNO3. +20L 30% H<sub>2</sub>O2.</li> <li>(2) Add 2mL 70% HNO3. +20L 30% H<sub>2</sub>O2.</li> <li>(2) Add 2mL 70% HNO3. +20L 30% H<sub>2</sub>O2.</li> <li>(2) Risse filter from funnel assembly and mount filter on stainless steel planchet with</li> <li>2-sided tape.</li> <li>(2) Precondition 2mL TEVA. TRU,</li> <li>(3) Rinse TEVA w/ 10mL 1.5M</li> <li>(4) NH<sub>4</sub>SCN-0.1M Formic acid.</li> <li>(1) Precondition 2mL TEVA, TRU,</li> <li>(2) Load Sample.</li> <li>(3) Rinse c-tube with 5mL 6M HNO3.</li> <li>(4) Sing Am/Cm from TEVA with</li> </ul>                                                                                                                                                                                                                                                                                                                                                                                | rigare zi / tetimae eepai                                    |                                          |                                                   |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------|------------------------------------------|---------------------------------------------------|
| <ul> <li>Add /mg La and 20mL 10% rIL(s.<br/>Mix. Cool to room temperature.<br/>Centrifuge 3500 rpm. 5min.<br/>Decant supernate</li> <li>Partially dissolve in 60mL 1.5M HCI.<br/>Solids will remain. Dilute to 170mL.</li> <li>Add 22mL 49% HF. Mix.<br/>Add 22mL 49% HF. Mix.<br/>Centrifuge 3500 rpm. 5min.<br/>Decant supernate</li> <li>Centrifuge 3500 rpm. 5min.<br/>Decant supernate</li> <li>Cost and form. 20% HF. Mix.</li> <li>Centrifuge 3500 rpm. 5min.<br/>Decant supernate</li> <li>Cost and form. 20% HF. Mix.</li> <li>Centrifuge 3500 rpm. 5min.<br/>Decant supernate</li> <li>Cost and form. 20% HF. Mix.</li> <li>Cost and form. 20% HF. Mix.</li> <li>Cost asmples to room temp. Fix<br/>valence by adding: (mix between steps)</li> <li>Cool samples to room temp. Fix</li> <li>MI NH<sub>4</sub>SCN-0.1M Formic acid.</li> <li>MI Sinse TEVA with 10mL 1.5M</li> <li>M</li></ul> | · ·                                                          | , , , , , , , , , , , , , , , , , , ,    | (20) Add 50ug Ce carrier to all                   |
| Mix. Cool to room temperature.<br>Centrifuge 3500 rpm. 5min.<br>Decant supernate(6) Rinse TEVA with:<br>-10mL 6M HNO3<br>.20mL 9M HCl (Th removal)<br>-5mL 3M HNO3samples and 0.2mL 30% H <sub>2</sub> O <sub>2</sub> to<br>Am/Cm samples. Mix. Add 1mL 49%<br>HF. Mix. Wait 15-20 minutes.Partially dissolve in 60mL 1.5M HCl.<br>Solids will remain.<br>Ditate to 170mL.(7) Strip Pu from TEVA with 20mL<br>0.1M HCl-0.05M HF-0.03M TiCls.(2) Wet filter vith 3mL 80% ethanol<br>followed by 3mL DI water.Add 22mL 49% HF. Mix.<br>Add 22mL 49% HF. Mix.<br>Decant supernate(8) Rinse TEV-DGA with 15mL 4M HCl.<br>0.1M HCl-0.05M HF-0.03M TiCls.(2) Wet filter with 3mL 80% ethanol<br>followed by 3mL DI water.Decant supernate<br>Dissolve solids in 5mL 3M HNO3-<br>0.25M Boric acid, 6mL 7M HNO3, and<br>7.5mL 2M Al(NO3).(3) Rinse TEVA with 20mL<br>0.1M HCl-0.05M HFO3.(2) Rinse sample tube with 5mL DI<br>water and add to filter.Cool samples to room temp. Fix<br>valence by adding: (mix between steps)<br>0.5mL 1.5M sulfamic acid<br>-40uL 50mg/mL Fe carrier<br>-1.5mL 1M ascorbic acid (Wait 3 min)<br>-11mL 3.5M NANO2(1) Precondition 2mL TEVA, TRU,<br>DGA cartridge with 10mL 8M HNO3.(1) Strip Am/Cm beaker with 5mL<br>4M NH <sub>4</sub> SCN-0.1M Formic acid.<br>(1) Precondition 2mL TEVA, TRU,<br>DGA cartridge with 10mL 8M HNO3.(1) Strip Am/Cm torm TEVA with<br>4M NH <sub>4</sub> SCN-0.1M Formic acid.<br>(1) Precondition 2mL TEVA, TRU,<br>DGA cartridge with 10mL 8M HNO3.(3) Rinse TEVA w/10mL 1.5M<br>NH <sub>4</sub> SCN-0.1M Formic acid.<br>(1) Strip Am/Cm torm TEVA with<br>4M NH <sub>4</sub> SCN-0.1M Formic acid.<br>(1) Precondition 2mL TEVA, TRU,<br>DGA cartridge with 10mL 8M HNO3.(3) Rinse TEVA w/10mL 1.5M<br>NH <sub>4</sub> SCN-0.1M Formic acid.<br>(1) Precondition 2mL TEVA, TRU,<br>DGA cartridge with 10mL 8M HNO3.(3) Rinse TEVA w/10mL 1.5M<br>NH <sub>4</sub> SCN-0.1M Formic acid.<                                                                                                                                                                                                                                                                                                                                                                                                                        | ů, s                                                         | (5) Separate TEVA from TRU-DGA.          |                                                   |
| Centrifuge 3500 rpm. 5min.<br>Decant supernate-10mL 6M HNO3<br>.10mL 3M HNO3Am/Cm samples. Mix. Add 1mL 49%<br>HF. Mix.<br>.20mL 9M HCl (Th removal)<br>.5mL 3M HNO3Partially dissolve in 60mL 1.5M HCl.<br>Solids will remain. Dilute to 170mL.<br>Add 22mL 49% HF. Mix.<br>Add 22mL 49% HF. Mix.<br>Centrifuge 3500 rpm. 5min.<br>Decant supernate(7) Strip Pu from TEVA with 20mL<br>0.1M HCl-0.05M HF-0.03M TiCl3.(2) Wet filter with 3mL 80% ethanol<br>followed by 3mL DI water.<br>(2) Wet filter with 3mL 80% ethanol<br>followed by 3mL DI water.<br>(2) Wet filter sample.Decant supernate<br>Dissolve solids in 5mL 3M HNO3-<br>0.25M Boric acid, 6mL 7M HNO3, and<br>7.5mL 2M AI(NO3). Warming samples<br>can improve dissolution.(1) Rinse DGA w/ 20mL 0.05M HNO3-<br>(1) Strip Am/Cm w 10mL 0.25M HCl.<br>(1) Strip Am/Cm. Evaporate to dryness<br>(14) Dissolve An/Cm. Evaporate to dryness<br>(14) Dissolve An/Cm. Evaporate to dryness<br>(15) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3-<br>(10) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3-<br>(1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3-<br>(2) Load Sample10mL 6M HNO3<br>(19) Strip Am/Cm from TEVA with<br>(19) Strip Am/Cm from TEVA with<br>(19) Strip Am/Cm from TEVA with<br>(19) Strip Am/Cm from TEVA withAm/Cm samples.<br>(2) Measure actinides by alpha<br>spectrometry.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 | Mix. Cool to room temperature.                               |                                          |                                                   |
| Decant supernateInfinition of the second                                                                                                                                                                                                                                                                         | Centrifuge 3500 rpm 5min                                     |                                          |                                                   |
| Partially dissolve in 60mL 1.5M HCI.<br>Solids will remain. Dilute to 170mL.SmL 3M HNO3(21) Set up Resolve* Filter Funnel on<br>vacuum box.Add 2mg La and 10mL 30% H <sub>2</sub> O2. Mix.<br>Add 22mL 49% HF. Mix.<br>Centrifuge 3500 rpm. 5min.<br>Decant supernate<br>Dissolve solids in 5mL 3M HNO3-<br>0.25M Boric acid, 6mL 7M HNO3, and<br>7.5mL 2M Al(NO3)3. Warming samples<br>can improve dissolution.(3) Rinse TRU-DGA with 15mL 4M HCI.<br>(9) Discard TRU cartridge.<br>(10) Rinse DGA w/ 20mL 0.05M HNO3.<br>(11) Strip Am/Cm w/ 10mL 0.25M HCI.<br>(12) Add 2mL 70% HNO3 + 50uL 10%<br>H2SO4 to Am/Cm. Evaporate to dryness<br>(13) Ash to dryness with 3mL 70%<br>(14) Dissolve Am/Cm in 5mL<br>(14) Dissolve Am/Cm in 5mL<br>(14) Dissolve Am/Cm in 5mL<br>(14) Dissolve Am/Cm in 5mL<br>(15) Precondition 2mL TEVA with 5mL<br>(15) Precondition 2mL TEVA with 5mL<br>(16) Load Am/Cm on TEVA.<br>(16) Load Am/Cm on TEVA.(21) Set up Resolve* Filter Funnel on<br>vacuum box.<br>(22) Wet filter with 3mL 80% ethanol<br>followed by 3mL DI water.<br>(23) Filter sample.<br>(24) Rinse sample tube with 5mL DI<br>water and add to filter.<br>(25) Rinse filter funnel with 3mL DI<br>water and 2mL 100% ethanol.<br>(26) Draw vacuum until filter is dry.<br>(27) Remove filter from funnel<br>assembly and mount filter on<br>stainless steel planchet with<br>2-sided tape.<br>(28) Dry filter under heat lamp for<br>3-5 minutes.0.5mL 1.5M sulfamic acid<br>(10) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8m HNO3.<br>(2) Load Sample.<br>(3) Rinse c-tube with 5mL 6M HNO3.*<br>(3) Rinse c-tube with 5mL 6M HNO3.*(10) Strip Am/Cm from TEVA with<br>(19) Strip Am/Cm from TEVA with(21) Set up Resolve* Filter Funnel on<br>vacuum box.<br>(23) Pitter under heat lamp for<br>3-5 minutes.<br>(29) Measure actinides by alpha<br>spectrometry.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |                                                              |                                          | HF. Mix. Wait 15-20 minutes.                      |
| Partially dissolve in 60mL 1.5M HCl.<br>Solids will remain, Dilute to 170mL.(7) Strip Pu from TEVA with 20mL<br>0.1M HCl-0.05M HF-0.03M TiCl3.Vacuum box.Add 2mg La and 10mL 30% H <sub>2</sub> O2. Mix.<br>Add 22mL 49% HF. Mix.<br>Decant supernate(7) Strip Pu from TEVA with 20mL<br>0.1M HCl-0.05M HF-0.03M TiCl3.(2) Wet filter with 3mL 80% ethanol<br>followed by 3mL DI water.Decant supernate(10) Rinse DGA w/ 20mL 0.05M HNO3<br>(11) Strip Am/Cm w/ 10mL 0.25M HCl.<br>(12) Add 2mL 70% HNO3 + 50uL 10%<br>H2SO4 to Am/Cm. Evaporate to dryness<br>(13) Ash to dryness with 3mL 70%<br>(13) Ash to dryness with 3mL 70%<br>(14) Dissolve Am/Cm in 5mL<br>can improve dissolution.(26) Draw vacuum until filter is dry.<br>(27) Remove filter from funnel<br>assembly and mount filter on<br>stainless steel planchet with<br>2-sided tape.Cool samples to room temp. Fix<br>valence by adding: (mix between steps)<br>(.5mL 1.5M sulfamic acid<br>-40uL 50mg/mL Fe carrier<br>(15) Precondition 2mL TEVA with 5mL<br>1.5mL 1M ascorbic acid (Wait 3 min)<br>(11) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.(18) Rinse TEVA w/ 10mL 1.5M<br>NH4SCN-0.1M Formic acid.<br>(19) Strip Am/Cm from TEVA with(29) Measure actinides by alpha<br>spectrometry.(10) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.(18) Rinse TEVA w/ 10mL 1.5M<br>NH4SCN-0.1M Formic acid.<br>(19) Strip Am/Cm from TEVA with(29) Measure actinides by alpha<br>spectrometry.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      | Decant supernate                                             | · · · ·                                  | (21) Set up Resolve <sup>®</sup> Filter Funnel on |
| Solids will remain.Dilute to 170mL.0.1M HCI-0.05M HF-0.03M TiCl3.(22) Wet filter with 3mL 80% ethanol<br>followed by 3mL DI water.Add 2mg La and 10mL 30% H2O2. Mix.<br>Add 22mL 49% HF. Mix.<br>Decant supernate(8) Rinse TRU-DGA with 15mL 4M HCI.<br>(9) Discard TRU cartridge.<br>(10) Rinse DGA w/ 20mL 0.05M HNO3.<br>(11) Strip Am/Cm w/ 10mL 0.25M HCI.<br>(12) Add 2mL 70% HNO3 + 50uL 10%<br>H2SO4 to Am/Cm. Evaporate to dryness.(24) Rinse sample tube with 5mL DI<br>water and add to filter.<br>(25) Rinse filter funnel with 3mL DI<br>water and 2mL 100% ethanol.0.25M Boric acid, 6mL 7M HNO3.<br>(0.55M Boric acid, 6mL 7M HNO3., and<br>7.5mL 2M Al(NO3)3. Warming samples<br>can improve dissolution.(13) Ash to dryness with 3mL 70%<br>HNO3 + 2mL 30% H2O2.<br>(14) Dissolve Am/Cm in 5mL<br>4M NH4SCN-0.1M Formic acid.(26) Draw vacuum until filter is dry.<br>(27) Remove filter from funnel<br>assembly and mount filter on<br>stainless steel planchet with<br>2-sided tape.0.5mL 1.5M sulfamic acid<br>-40uL 50mg/mL Fe carrier<br>(1) Precondition 2mL TEVA. TRU,<br>DGA cartridges with 10mL 8M HNO3.(13) Rinse TEVA w/ 10mL 1.5M<br>NH4SCN-0.1M Formic acid.(29) Measure actinides by alpha<br>spectrometry.(1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.(18) Rinse TEVA w/ 10mL 1.5M<br>NH4SCN-0.1M Formic acid.(29) Measure actinides by alpha<br>spectrometry.(1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.(19) Strip Am/Cm from TEVA with(29) Measure actinides by alpha<br>spectrometry.(1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.(19) Strip Am/Cm from TEVA with(19) Strip Am/Cm from TEVA with                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | Partially dissolve in 60mL 1.5M HCI.                         | -                                        | vacuum box.                                       |
| Add 2mg La and 10mL 30% H2O2. Mix.<br>Add 22mL 49% HF. Mix.<br>Centrifuge 3500 rpm. 5min.<br>Decant supernate6.0. mit ref closed with 15mL 4M HCl.<br>(9) Discard TRU cartridge.<br>(10) Rinse DGA w/ 20mL 0.05M HNO3.<br>(11) Strip Am/Cm w/ 10mL 0.25M HCl.<br>(12) Add 2mL 70% HNO3 + 50uL 10%<br>H2SO4 to Am/Cm. Evaporate to dryness<br>(13) Ash to dryness with 3mL 70%<br>(14) Dissolve Am/Cm is 5mL<br>(13) Ash to dryness with 3mL 70%<br>(14) Dissolve Am/Cm. Evaporate to dryness<br>(13) Ash to dryness with 3mL 70%<br>(14) Dissolve Am/Cm. Evaporate to dryness<br>(13) Ash to dryness with 3mL 70%<br>(14) Dissolve Am/Cm. 5mL<br>(15) Precondition 2mL TEVA with 5mL<br>(15) Precondition 2mL TEVA with 5mL<br>(16) Load Am/Cm on TEVA.<br>(17) Rinse Am/Cm beaker with 5mL<br>(16) Load Am/Cm on TEVA.followed by 3mL DI water.<br>(24) Rinse sample tube with 5mL DI<br>water and add to filter.<br>(25) Rinse filter from funnel<br>assembly and mount filter on<br>stainless steel planchet with<br>2-sided tape.<br>(28) Dry filter under heat lamp for<br>3-5 minutes.0.5mL 1.5M sulfamic acid<br>-40UL 50mg/mL Fe carrier<br>-1.5mL 1M ascorbic acid (Wait 3 min)<br>-1mL 3.5M NaNO2(18) Rinse TEVA w/ 10mL 1.5M<br>NH4SCN-0.1M Formic acid.<br>(19) Rinse Am/Cm beaker with 5mL<br>(19) Strip Am/Cm from TEVA with(29) Measure actinides by alpha<br>spectrometry.(11) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.<br>(2) Load Sample.<br>(3) Rinse c-tube with 5mL 6M HNO3.*(18) Rinse TEVA w/ 10mL 1.5M<br>NH4SCN-0.1M Formic acid.<br>(19) Strip Am/Cm from TEVA withvalue actinides by alpha<br>spectrometry.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                | Solids will remain. Dilute to 170mL.                         |                                          | (22) Wet filter with 3ml_80% ethanol              |
| Add 22mL 49% HF. Mix.<br>Centrifuge 3500 rpm. 5min.<br>Decant supernate(9) Discard TRU cartridge.<br>(10) Rinse DGA w/ 20mL 0.05M HNO3.<br>(11) Strip Am/Cm w/ 10mL 0.25M HCI.<br>(12) Add 2mL 70% HNO3 + 50uL 10%<br>H2SO4 to Am/Cm. Evaporate to dryness.<br>(13) Ash to dryness with 3mL 70%<br>H2SO4 to Am/Cm. Evaporate to dryness.<br>(13) Ash to dryness with 3mL 70%<br>HNO3 + 2mL 30% H2O2.<br>(14) Dissolve Am/Cm in 5mL<br>assembly and mount filter on<br>stainless steel planchet with<br>2-Sided tape.<br>(25) Mass steel planchet with<br>2-sided tape.(26) Draw vacuum until filter is dry.<br>(26) Draw vacuum until filter on<br>stainless steel planchet with<br>2-sided tape.0.5mL 1.5M sulfamic acid<br>-40uL 50mg/mL Fe carrier<br>-1.5mL 1M ascorbic acid (Wait 3 min)<br>-1mL 3.5M NaNO2(15) Precondition 2mL TEVA.<br>(17) Rinse Am/Cm beaker with 5mL<br>(17) Rinse Am/Cm beaker with 5mL<br>(18) Rinse TEVA w/ 10mL 1.5M<br>M4sCN-0.1M Formic acid.<br>(19) Strip Am/Cm from TEVA with(29) Measure actinides by alpha<br>spectrometry.(1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.<br>(2) Load Sample.<br>(3) Rinse c-tube with 5mL 6M HNO3.**(19) Strip Am/Cm from TEVA with(29) Measure actinides by alpha<br>spectrometry.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | Add 2mg La and 10mL 30% H <sub>2</sub> O <sub>2</sub> . Mix. |                                          |                                                   |
| Centrifuge 3500 rpm. 5min.<br>Decant supernate(10) Rinse DGA w/ 20mL 0.05M HNO3.<br>(11) Strip Am/Cm w/ 10mL 0.25M HCI.<br>(12) Add 2mL 70% HNO3 + 50uL 10%(24) Rinse sample tube with 5mL DI<br>water and add to filter.<br>(25) Rinse filter funnel with 3mL DI<br>water and 2mL 100% ethanol.Dissolve solids in 5mL 3M HNO3-<br>0.25M Boric acid, 6mL 7M HNO3, and<br>7.5mL 2M Al(NO3)3. Warming samples<br>can improve dissolution.(13) Ash to dryness with 3mL 70%<br>HNO3 + 2mL 30% H <sub>2</sub> O2.<br>(14) Dissolve Am/Cm in 5mL<br>4M NH4SCN-0.1M Formic acid.(26) Draw vacuum until filter is dry.<br>(27) Remove filter from funnel<br>assembly and mount filter on<br>stainless steel planchet withCool samples to room temp. Fix<br>valence by adding: (mix between steps)<br>-0.5mL 1.5M sulfamic acid<br>-40uL 50mg/mL Fe carrier<br>-1.5mL 1M ascorbic acid (Wait 3 min)<br>-1mL 3.5M NaNO2(15) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.<br>(10) Rinse TEVA w/ 10mL 1.5M<br>NH4SCN-0.1M Formic acid.<br>(16) Load Sample.<br>(3) Rinse c-tube with 5mL 6M HNO3.*(13) Strip Am/Cm from TEVA with<br>Formic acid.<br>(14) Distrip Am/Cm from TEVA with(24) Rinse sample tube with 5mL DI<br>water and add to filter.<br>(25) Rinse filter funnel with 3mL DI<br>water and 2mL 100% ethanol.<br>(26) Draw vacuum until filter is dry.<br>(27) Remove filter from funnel<br>assembly and mount filter on<br>stainless steel planchet with<br>2-sided tape.<br>(28) Dry filter under heat lamp for<br>3-5 minutes.<br>(29) Measure actinides by alpha<br>spectrometry.(10) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.<br>(2) Load Sample.<br>(3) Rinse c-tube with 5mL 6M HNO3.**(19) Strip Am/Cm from TEVA with                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   | Add 22mL 49% HF. Mix.                                        | . ,                                      | (23) Filter sample.                               |
| Celliningle 3000 rpm. shift.(11) Strip Am/Cm w/ 10mL 0.25M HCL.water and add to filter.Decant supernate(12) Add 2mL 70% HNO3 + 50uL 10%(25) Rinse filter funnel with 3mL DIDissolve solids in 5mL 3M HNO3-H2SO4 to Am/Cm. Evaporate to drynesswater and 2mL 100% ethanol.0.25M Boric acid, 6mL 7M HNO3, and(13) Ash to dryness with 3mL 70%(26) Draw vacuum until filter is dry.7.5mL 2M Al(NO3)3. Warming samples(14) Dissolve Am/Cm in 5mLassembly and mount filter oncoal samples to room temp. Fix4M NH4SCN-0.1M Formic acid.(27) Remove filter from funnelvalence by adding: (mix between steps)(15) Precondition 2mL TEVA with 5mL2-sided tape0.5mL 1.5M sulfamic acid4M NH4SCN-0.1M Formic acid.(28) Dry filter under heat lamp for-40uL 50mg/mL Fe carrier(16) Load Am/Cm on TEVA.(29) Measure actinides by alpha-1.5mL 1M ascorbic acid (Wait 3 min)(17) Rinse Am/Cm beaker with 5mL(29) Measure actinides by alpha-1mL 3.5M NaNO218) Rinse TEVA w/ 10mL 1.5Mspectrometry.(2) Load Sample.(19) Strip Am/Cm from TEVA withLint 4.5CN-0.1M Formic acid.(3) Rinse c-tube with 5mL 6M HNO3.*(19) Strip Am/Cm from TEVA withLint 4.5CN-0.1M Formic acid.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |                                                              |                                          |                                                   |
| Decarit superate(12) Add 2mL 70% HNO3 + 50uL 10%(25) Rinse filter funnel with 3mL DIDissolve solids in 5mL 3M HNO3-(12) Add 2mL 70% HNO3 + 50uL 10%(26) Rinse filter funnel with 3mL DI0.25M Boric acid, 6mL 7M HNO3, and(13) Ash to dryness with 3mL 70%(26) Draw vacuum until filter is dry.7.5mL 2M Al(NO3)3. Warning samples<br>can improve dissolution.(14) Dissolve Am/Cm in 5mL(27) Remove filter from funnelCool samples to room temp. Fix4M NH4SCN-0.1M Formic acid.stainless steel planchet with0.5mL 1.5M sulfamic acid4M NH4SCN-0.1M Formic acid.2-sided tape0.5mL 1.5M sulfamic acid(16) Load Am/Cm on TEVA.3-5 minutes1.5mL 1M ascorbic acid (Wait 3 min)(17) Rinse Am/Cm beaker with 5mL(29) Measure actinides by alpha.1mL 3.5M NANO2(18) Rinse TEVA w/ 10mL 1.5Mspectrometry.(1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.(18) Rinse TEVA w/ 10mL 1.5Mspectrometry.(2) Load Sample.(19) Strip Am/Cm from TEVA withspectrometry.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | <b>3</b> 1                                                   | · ,                                      |                                                   |
| Dissolve solids in 5mL 3M HNO3-<br>0.25M Boric acid, 6mL 7M HNO3, and<br>7.5mL 2M Al(NO3)3. Warming samples<br>can improve dissolution.H2SO4 to Am/Cm. Evaporate to dryness.<br>(13) Ash to dryness with 3mL 70%<br>HNO3 + 2mL 30% H2O2.<br>(14) Dissolve Am/Cm in 5mL<br>4M NH4SCN-0.1M Formic acid.<br>(15) Precondition 2mL TEVA with 5mL<br>(16) Load Am/Cm on TEVA.<br>(17) Rinse Am/Cm beaker with 5mL<br>(18) Rinse TEVA w/ 10mL 1.5M<br>(2) Load Sample.water and 2mL 100% ethanol.<br>(26) Draw vacuum until filter is dry.<br>(27) Remove filter from funnel<br>assembly and mount filter on<br>stainless steel planchet with<br>2-sided tape.0.5mL 1.5M sulfamic acid<br>-40uL 50mg/mL Fe carrier<br>-1.5mL 1M ascorbic acid (Wait 3 min)<br>-1mL 3.5M NANO2(16) Load Am/Cm on TEVA.<br>(17) Rinse Am/Cm beaker with 5mL<br>(17) Rinse Am/Cm beaker with 5mL<br>(18) Rinse TEVA w/ 10mL 1.5M<br>NH4SCN-0.1M Formic acid.<br>(19) Strip Am/Cm from TEVA with(29) Measure actinides by alpha<br>spectrometry.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | Decant supernate                                             | <u> </u>                                 | (25) Rinse filter funnel with 3mL DI              |
| <ul> <li>0.25M Boric acid, 6mL 7M HNO3, and<br/>7.5mL 2M Al(NO3)3. Warming samples<br/>can improve dissolution.</li> <li>Cool samples to room temp. Fix<br/>valence by adding: (mix between steps)</li> <li>0.5mL 1.5M sulfamic acid</li> <li>4M NH4SCN-0.1M Formic acid.</li> <li>(15) Precondition 2mL TEVA with 5mL</li> <li>(16) Load Am/Cm on TEVA.</li> <li>(17) Rinse Am/Cm beaker with 5mL</li> <li>(17) Rinse Am/Cm beaker with 5mL</li> <li>(17) Rinse TEVA w/ 10mL 1.5M</li> <li>(29) Measure actinides by alpha<br/>spectrometry.</li> <li>(19) Strip Am/Cm from TEVA with</li> </ul>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             | Dissolve solids in 5mL 3M HNO <sub>3</sub> -                 |                                          |                                                   |
| 7.5mL 2M Al(NO <sub>3</sub> ) <sub>3</sub> . Warming samples<br>can improve dissolution.HNO <sub>3</sub> + 2mL 30% H <sub>2</sub> O <sub>2</sub> .<br>(14) Dissolve Am/Cm in 5mL(27) Remove filter from funnel<br>assembly and mount filter on<br>stainless steel planchet with<br>2-sided tape.Cool samples to room temp. Fix<br>valence by adding: (mix between steps)<br>-0.5mL 1.5M sulfamic acid<br>-40uL 50mg/mL Fe carrier<br>-1.5mL 1M ascorbic acid (Wait 3 min)<br>-1mL 3.5M NaNO24M NH <sub>4</sub> SCN-0.1M Formic acid.<br>(16) Load Am/Cm on TEVA.<br>(17) Rinse Am/Cm beaker with 5mL<br>                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      | 0.25M Boric acid, 6mL 7M HNO <sub>3</sub> , and              | 1 5                                      | (26) Draw vacuum until filter is dry.             |
| can improve dissolution.(14) Dissolve Am/Cm in 5mLassembly and mount filter onCool samples to room temp. Fix4M NH4SCN-0.1M Formic acid.stainless steel planchet withvalence by adding: (mix between steps)(15) Precondition 2mL TEVA with 5mL2-sided tape0.5mL 1.5M sulfamic acid4M NH4SCN-0.1M Formic acid.(28) Dry filter under heat lamp for-40uL 50mg/mL Fe carrier(16) Load Am/Cm on TEVA.3-5 minutes1.5mL 1M ascorbic acid (Wait 3 min)(17) Rinse Am/Cm beaker with 5mL(29) Measure actinides by alpha-1mL 3.5M NaNO24M NH4SCN-0.1M Formic acid.spectrometry.(1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.(18) Rinse TEVA w/ 10mL 1.5Mspectrometry.(2) Load Sample.NH4SCN-0.1M Formic acid.H4SCN-0.1M Formic acid.spectrometry.(3) Rinse c-tube with 5mL 6M HNO3.*(19) Strip Am/Cm from TEVA withstate actinides by alpha                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | 7.5mL 2M AI(NO <sub>3</sub> ) <sub>3</sub> . Warming samples | • •                                      |                                                   |
| valence by adding: (mix between steps)(15) Precondition 2mL TEVA with 5mL2-sided tape0.5mL 1.5M sulfamic acid4M NH4SCN-0.1M Formic acid.(28) Dry filter under heat lamp for-40uL 50mg/mL Fe carrier(16) Load Am/Cm on TEVA.3-5 minutes1.5mL 1M ascorbic acid (Wait 3 min)(17) Rinse Am/Cm beaker with 5mL(29) Measure actinides by alpha-1mL 3.5M NaNO24M NH4SCN-0.1M Formic acid.spectrometry.(1) Precondition 2mL TEVA, TRU,Add to TEVA.spectrometry.DGA cartridges with 10mL 8M HNO3.(18) Rinse TEVA w/ 10mL 1.5MH4SCN-0.1M Formic acid.(2) Load Sample.NH4SCN-0.1M Formic acid.Image: 10 Strip Am/Cm from TEVA with                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | can improve dissolution.                                     | (14) Dissolve Am/Cm in 5mL               |                                                   |
| valence by adding: (mix between steps)(15) Precondition 2mL TEVA with 5mL2-sided tape0.5mL 1.5M sulfamic acid4M NH4SCN-0.1M Formic acid.(28) Dry filter under heat lamp for-40uL 50mg/mL Fe carrier(16) Load Am/Cm on TEVA.3-5 minutes1.5mL 1M ascorbic acid (Wait 3 min)(17) Rinse Am/Cm beaker with 5mL(29) Measure actinides by alpha-1mL 3.5M NaNO24M NH4SCN-0.1M Formic acid.spectrometry.(1) Precondition 2mL TEVA, TRU,Add to TEVA.spectrometry.DGA cartridges with 10mL 8M HNO3.(18) Rinse TEVA w/10mL 1.5MLHascon 2mL 4ML(2) Load Sample.NH4SCN-0.1M Formic acid.state actinities acting and the state acting ac                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | Cool samples to room temp. Fix                               | 4M NH <sub>4</sub> SCN-0.1M Formic acid. | stainless steel planchet with                     |
| -0.5mL 1.5M sulfamic acid4M NH4SCN-0.1M Formic acid.(28) Dry filter under heat lamp for<br>3-5 minutes40uL 50mg/mL Fe carrier(16) Load Am/Cm on TEVA.3-5 minutes1.5mL 1M ascorbic acid (Wait 3 min)<br>-1mL 3.5M NaNO2(17) Rinse Am/Cm beaker with 5mL(29) Measure actinides by alpha<br>spectrometry.(1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.Add to TEVA.(29) Measure actinides by alpha<br>spectrometry.(2) Load Sample.(18) Rinse TEVA w/ 10mL 1.5M<br>NH4SCN-0.1M Formic acid.H4SCN-0.1M Formic acid.(3) Rinse c-tube with 5mL 6M HNO3.*(19) Strip Am/Cm from TEVA withImage: Complement of the spectrometry with the spectrom teva with term teva with teva w                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |                                                              | (15) Precondition 2mL TEVA with 5mL      | 2-sided tape.                                     |
| -1.5mL 1M ascorbic acid (Wait 3 min)<br>-1mL 3.5M NaNO2(17) Rinse Am/Cm beaker with 5mL<br>4M NH4SCN-0.1M Formic acid.(29) Measure actinides by alpha<br>spectrometry.(1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.Add to TEVA.spectrometry.(2) Load Sample.<br>(3) Rinse c-tube with 5mL 6M HNO3.*(19) Strip Am/Cm from TEVA withspectrometry.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | J U I I I                                                    | 4M NH <sub>4</sub> SCN-0.1M Formic acid. | (28) Dry filter under heat lamp for               |
| -1mL 3.5M NaNO24M NH4SCN-0.1M Formic acid.spectrometry.(1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.Add to TEVA.spectrometry.(2) Load Sample.<br>(3) Rinse c-tube with 5mL 6M HNO3.*(19) Strip Am/Cm from TEVA withspectrometry.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | -40uL 50mg/mL Fe carrier                                     | (16) Load Am/Cm on TEVA.                 | 3-5 minutes.                                      |
| (1) Precondition 2mL TEVA, TRU,<br>DGA cartridges with 10mL 8M HNO3.Add to TEVA.(2) Load Sample.(18) Rinse TEVA w/ 10mL 1.5M<br>NH4SCN-0.1M Formic acid.(3) Rinse c-tube with 5mL 6M HNO3.*(19) Strip Am/Cm from TEVA with                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | -1.5mL 1M ascorbic acid (Wait 3 min)                         | (17) Rinse Am/Cm beaker with 5mL         | (29) Measure actinides by alpha                   |
| (1) Freedration 2012 FEVA, Free,DGA cartridges with 10mL 8M HNO3.(18) Rinse TEVA w/ 10mL 1.5M(2) Load Sample.NH₄SCN-0.1M Formic acid.(3) Rinse c-tube with 5mL 6M HNO3.*(19) Strip Am/Cm from TEVA with                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | -1mL 3.5M NaNO <sub>2</sub>                                  | 4M NH₄SCN-0.1M Formic acid.              | spectrometry.                                     |
| (2) Load Sample.NH₄SCN-0.1M Formic acid.(3) Rinse c-tube with 5mL 6M HNO3.*(19) Strip Am/Cm from TEVA with                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    | (1) Precondition 2mL TEVA, TRU,                              | Add to TEVA.                             |                                                   |
| (3) Rinse c-tube with 5mL 6M HNO <sub>3</sub> .* (19) Strip Am/Cm from TEVA with                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              | DGA cartridges with 10mL 8M HNO <sub>3</sub> .               | (18) Rinse TEVA w/ 10mL 1.5M             |                                                   |
|                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               | -                                                            | NH <sub>4</sub> SCN-0.1M Formic acid.    |                                                   |
|                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               | (3) Rinse c-tube with 5mL 6M HNO <sub>3</sub> .*             | (19) Strip Am/Cm from TEVA with          |                                                   |
| Add to stacked cartridges. 20mL 1M HCI.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | Add to stacked cartridges.                                   | 20mL 1M HCI.                             |                                                   |

\*Adding 50uL of 30% H<sub>2</sub>O<sub>2</sub> to tube rinse can help improve U decontamination.

|   |          |            | Method Pe         | rformance         |                   |                   |
|---|----------|------------|-------------------|-------------------|-------------------|-------------------|
|   |          |            | <sup>242</sup> Pu | <sup>238</sup> Pu | <sup>243</sup> Am | <sup>241</sup> Am |
|   | Sample   |            | Tracer            | Measured          | Tracer            | Measured          |
|   | Size (g) | Replicates | % Recovery        | % Bias            | % Recovery        | % Bias            |
|   | 100      | 3          | 86 <u>+</u> 7     | -3.0              | 94 <u>+</u> 4     | -10               |
|   | 100      | 3          | 81 <u>+</u> 15    | -6.0              | 80 <u>+</u> 5     | -13               |
|   | 200      | 2          | 82 <u>+</u> 1     | 2.0               | 93 <u>+</u> 5     | -19               |
| 2 | 200      | 3          | 80 <u>+</u> 8     | -5.0              | 93 <u>+</u> 5     | -18               |

## References

1) Sherrod L. Maxwell, "Rapid method for determination of plutonium, americium, and curium in large soil samples," *J. Radioanal. Nucl. Chem.*, 275(2), 395-402 (2008).

# **Cichrom**<sup>\*</sup> Rapid Determination of Actinides in 1g Concrete and Brick Samples

**Summary of Method** U, Pu, Np, Am and Cm are separated and concentrated from 1 gram samples of concrete and brick. Samples are fused with NaOH at 600°C in zirconium crucibles. The fusion cakes are dissolved in water, transferred to 250mL centrifuge tubes and precipitated sequentially with iron-hydrous titanium oxide and lanthanum fluoride to facilitate matrix removal. Actinides are separated on stacked 2mL cartridges of Eichrom TEVA, TRU and DGA resins. Actinides are measured by alpha spectrometry following CeF<sub>3</sub> microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Chemical yields of tracers ranged from 79-98% for <sup>236</sup>Pu, 77-90% for <sup>243</sup>Am, and 72-81% for <sup>232</sup>U. Measured values typically agreed to within 10% of reference values. Sample preparation for batches of 12 samples can be completed by a single operator in <8 hours. Alpha spectrometry count times will depend on detection limit and data quality objectives.

#### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu (or <sup>236</sup>Pu if meas. Np), <sup>243</sup>Am and <sup>232</sup>U tracers Oxalic acid/Ammonium oxalate

La carrier (10mg/mL) Deionized Water 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> 10% (w:w) TiCl<sub>3</sub> HCI (37%) HF (49%) or NaF H<sub>2</sub>O<sub>2</sub> (30%) Denatured ethanol Ascorbic Acid

Ce carrier (1mg/mL) 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> 2M AI(NO<sub>3</sub>)<sub>3</sub> HNO<sub>3</sub> (70%) NaOH Boric acid NaNO<sub>2</sub> Sulfamic Acid

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Heat Lamp Muffle Furnace Hot Plate Analytical Balance 250mL Zirconium crucibles with zirconium lids Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Vacuum Pump

### Figure 1. Sample Preparation

| 1 | g milled Concrete or Brick + tracers in zirconium crucible.                                                                                                                    |
|---|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|   | Fuse samples with 15g NaOH at 600°C for 15 minutes.                                                                                                                            |
| [ | Dissolve fusion cake with $H_2O$ . Transfer to 250mL c-tube.                                                                                                                   |
| ŀ | Add 10mL 3M HNO <sub>3</sub> to crucible. Heat to dissolve residue.<br>Transfer to same 25mL c-tube.                                                                           |
|   | Add 125mg Fe and 5mg La to c-tube. Dilute to 180mL.                                                                                                                            |
|   | Add 2mL 1.25M Ca(NO <sub>3</sub> ) <sub>2</sub> , 3mL 3.2M (NH <sub>4</sub> ) <sub>2</sub> HPO <sub>4</sub> ,<br>5mL 10% TiCl <sub>3</sub> . Mix. Cool in ice bath for 10 min. |
|   | Centrifuge at 3500rpm. Decant Supernate.                                                                                                                                       |
|   | Partially dissolve precipitate in 60mL 1.5M HCl.                                                                                                                               |
|   | Some solids will remain. Dilute to 170mL.                                                                                                                                      |
|   | Add 1mg La, and 3mL 10% TiCl <sub>3</sub> . Mix.                                                                                                                               |
|   | Add 20mL 49% HF. Cool in ice bath for 10 min.                                                                                                                                  |
|   | Centrifuge at 3500rpm. Decant Supernate.                                                                                                                                       |
|   | Dissolve precipitate in 5mL 3M HNO <sub>3</sub> -0.25M Boric                                                                                                                   |
|   | acid, 7mL 70% HNO <sub>3</sub> , and 7mL 2M Al(NO <sub>3</sub> ) <sub>3</sub> .                                                                                                |
|   | Warming samples can help complete dissolution.                                                                                                                                 |
|   | Cool samples to room temperature.                                                                                                                                              |
|   | Fix valence states. Mix between each addition of: 0.5mL<br>1.5M sulfamic acid, 10uL 50mg/mL Fe, 1.5mL 1M                                                                       |
|   | ascorbic acid, 1mL 3.5M NaNO <sub>2</sub> , 1.5mL 70% HNO <sub>3</sub> .                                                                                                       |

# Figure 2. Actinide Separation on TEVA - TRU - DGA\* and Source Preparation

| •                                                                                                                                                                                                                                                                      |                                                                                                                                                                                                                                                                                                              | · · · · · · · · · · · · · · · · · · ·                                                                                                               |
|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------|
| (1) Precondition stacked 2mL                                                                                                                                                                                                                                           | (12) Separate TRU cartridge from DGA cartridge. Set TRU aside for U recovery.                                                                                                                                                                                                                                | (23) Rinse sample tube with 5mL DI water and add to filter.                                                                                         |
| <ul> <li>with 10mL 3M HNO<sub>3</sub>.</li> <li>(2) Load sample solution.</li> <li>(3) Rinse sample tube</li> <li>with 5mL 3M HNO<sub>3</sub>.* Add</li> <li>tube rinse to cartridges.</li> <li>(4) Rinse cartridges with</li> <li>10mL 3M HNO<sub>3</sub>.</li> </ul> | <ul> <li>(13) Rinse DGA cartridge sequentially</li> <li>with: -5mL 4M HCI</li> <li>-5mL 1M HNO3</li> <li>-15mL 0.05M HNO3</li> <li>(14) Strip Am and Cm from DGA with</li> <li>10mL 0.25M HCI. Add 0.2mL 30% H<sub>2</sub>O<sub>2</sub>.</li> </ul>                                                          | <ul> <li>(24) Rinse filter<br/>funnel with 3mL<br/>DI water and 2mL<br/>100% ethanol.</li> <li>(25) Draw vacuum<br/>until filter is dry.</li> </ul> |
| <ul> <li>(5) Separate TEVA, TRU,<br/>and DGA cartridges.</li> <li>(6) Rinse TEVA cartridge with:<br/>-10mL 3M HNO<sub>3</sub></li> <li>-20mL 9M HCI<br/>-5mL 3M HNO<sub>3</sub></li> <li>(7) Strip Pu (and Np) from TEVA</li> </ul>                                    | <ul> <li>(15) Rinse TRU cartridge with 15mL</li> <li>4M HCI-0.2M HF-0.002M TiCl<sub>3</sub>.</li> <li>(16) Rinse TRU cartridge with 10mL</li> <li>8M HNO<sub>3</sub>.</li> <li>(17) Strip U from TRU with 15mL of</li> <li>0.1M ammonium bioxalate.</li> <li>(18) Add 0.5mL 10% TiCl<sub>3</sub>.</li> </ul> | (26) Remove filter from<br>funnel assembly and<br>mount filter on stainless<br>steel planchet with 2-sided tape.                                    |
| cartridge with 20mL 0.1M HCI-<br>0.05MHF-0.01M TiCl <sub>3</sub> .<br>(8) Add 0.5mL 30% H <sub>2</sub> O <sub>2</sub> for Uranium<br>decon. in alpha source preparation.<br>(9) Rinse DGA cartridge with 10mL                                                          | <ul> <li>(19) Add 50ug Ce carrier to all samples.</li> <li>Mix well. Add 1mL 49% HF. Mix well.</li> <li>Wait 15-20 minutes.</li> <li>(20) Set up Resolve<sup>®</sup> Filter Funnel on</li> </ul>                                                                                                             |                                                                                                                                                     |
| 0.1M HNO <sub>3</sub> .<br>(10) Place TRU cartridge above DGA.<br>(11) Strip Am/Cm from TRU onto<br>DGA with 15mL 4M HCI.                                                                                                                                              | vacuum box.<br>(21) Wet filter with 3mL 80% ethanol<br>followed by 3mL DI water.<br>(22) Filter sample.                                                                                                                                                                                                      | <ul><li>(27) Dry filter under heat lamp for<br/>3-5 minutes.</li><li>(28) Measure actinides by alpha<br/>spectrometry.</li></ul>                    |

\*Adding 50uL 30% H<sub>2</sub>O<sub>2</sub> to the tube rinse can improve Uranium recoveries and decontamination in the Pu/Np fraction.

|                   | Method Performance |                   |               |           |               |        |  |  |  |
|-------------------|--------------------|-------------------|---------------|-----------|---------------|--------|--|--|--|
|                   |                    |                   |               | Analyte   | Analyte       |        |  |  |  |
|                   |                    |                   | Tracer        | Reference | Measured      |        |  |  |  |
| Analyte           | Replicates         | Tracer            | % Yield       | (mBq/g)   | (mBq/g)       | % Bias |  |  |  |
| <sup>239</sup> Pu | 5                  | <sup>236</sup> Pu | 90 <u>+</u> 7 | 18.0      | 18 <u>+</u> 2 | 0.0    |  |  |  |
| <sup>238</sup> Pu | 5                  | <sup>236</sup> Pu | 90 <u>+</u> 7 | 14.8      | 15 <u>+</u> 2 | 1.4    |  |  |  |
| <sup>237</sup> Np | 5                  | <sup>236</sup> Pu | 90 <u>+</u> 7 | 37.0      | 33 <u>+</u> 1 | -11    |  |  |  |
| <sup>241</sup> Am | 5                  | <sup>243</sup> Am | 85 <u>+</u> 6 | 25.4      | 24 <u>+</u> 1 | -5.5   |  |  |  |
| <sup>244</sup> Cm | 5                  | <sup>243</sup> Am | 85 <u>+</u> 6 | 35.0      | 35 <u>+</u> 2 | 0.0    |  |  |  |
| <sup>238</sup> U  | 5                  | <sup>232</sup> U  | 77 <u>+</u> 3 | 29.6      | 31 <u>+</u> 3 | 4.7    |  |  |  |
| <sup>234</sup> U  | 5                  | <sup>232</sup> U  | 77 <u>+</u> 3 | 28.4      | 26 <u>+</u> 4 | -8.5   |  |  |  |

### References

1) Sherrod L. Maxwell, Brian K. Culligan, Angel Kelsey-Wall, Patrick J. Shaw, "Rapid radiochemical method for determination of actinides in emergency concrete and brick samples," *Analytica Chimica Acta*, 701(1), 112-118 (2011).

# **Rapid Determination of Actinides** eichrom in Emergency Air Filter Samples

#### AN-1433-10

**Summary of Method** U, Pu, Np, Am and Cm are separated and concentrated from air filters. Samples are digested in Teflon beakers once with HNO<sub>3</sub>-H<sub>2</sub>O<sub>2</sub>-HF and then several times with HNO<sub>3</sub>-H<sub>2</sub>O<sub>2</sub>. After evaporating to dryness from HNO<sub>3</sub>-H<sub>3</sub>BO<sub>3</sub> to complex any residual fluoride, actinides are valence adjusted and separated on stacked 2mL cartridges of Eichrom TEVA and TRU resins. Actinides are measured by alpha spectrometry following CeF<sub>3</sub> microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Chemical yields of tracers averaged from 94+12% for <sup>242</sup>Pu, 87+6% for <sup>243</sup>Am, and 67+32% for <sup>232</sup>U. Poor <sup>232</sup>U recoveries in some samples were traced to insufficient mass of Ce carrier in the source preparation step. Recovery of <sup>232</sup>U improved upon increasing to 100ug of Ce carrier. Measured values typically agreed to within 10% of reference values. Sample preparation for batches of 12 samples can be completed by a single operator in <8 hours. Alpha spectrometry count times will depend on detection limit and data quality objectives.

#### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu (or <sup>236</sup>Pu if meas. Np), <sup>243</sup>Am and <sup>232</sup>U tracers Oxalic acid/Ammonium oxalate

Ce carrier (1mg/mL) Deionized water 10% (w:w) TiCl<sub>3</sub>

 $2M AI(NO_3)_3$ HNO<sub>3</sub> (70%) HF (49%) or NaF  $H_2O_2$  (30%) Denatured ethanol Ascorbic Acid

### Equipment

Sulfamic Acid

HCI (37%)

Boric acid

NaNO<sub>2</sub>

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL Centrifuge Tubes Centrifuge Heat Lamp Hot Plate **Analytical Balance** 250mL Teflon beakers Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Vacuum Pump

# Figure 1. Sample Preparation Air Filter + tracers in Teflon beaker. Wet ash to dryness with 3mL 70% HNO<sub>3</sub>, 3mL 30% H<sub>2</sub>O<sub>2</sub> and 5mL 49% HF. Wet ash to dryness with 3mL 70% HNO<sub>3</sub> and 3mL 30% H<sub>2</sub>O<sub>2</sub>. Wet ash to dryness with 3mL 70% HNO<sub>3</sub> and 3mL 30% H<sub>2</sub>O<sub>2</sub>.

Wet ash to dryness with 3mL 70% HNO<sub>3</sub> and 3mL 30% H<sub>2</sub>O<sub>2</sub>.

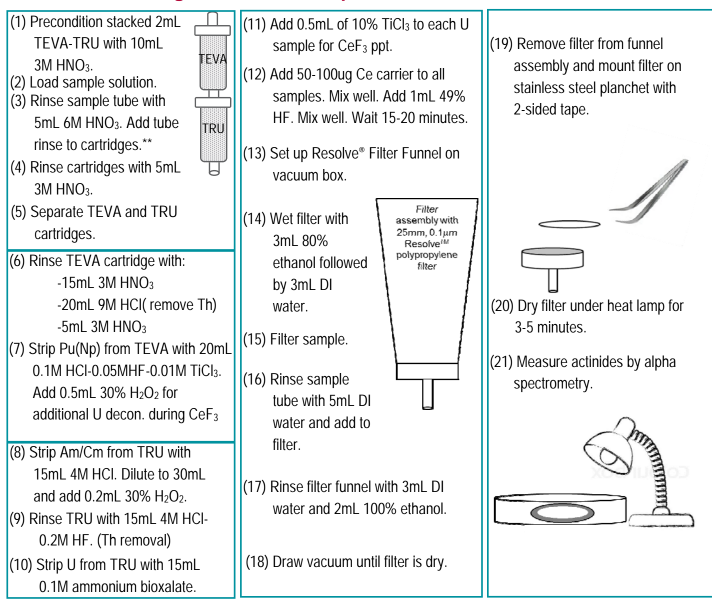
Add 3mL 3M HNO<sub>3</sub> –0.25M H<sub>3</sub>BO<sub>3</sub>. Evaporate to dryness.

Dissolve residue in 6mL 6M HNO<sub>3</sub> and 6mL 2M AI(NO<sub>3</sub>)<sub>3</sub>.

Cool samples to room temperature.

Fix valence states. Mix between each addition of: 0.5mL 1.5M sulfamic acid, 10uL 50mg/mL Fe, 1.5mL 1M ascorbic acid, 1mL 3.5M NaNO<sub>2</sub>.

# Figure 2. Actinide Separation on TEVA - TRU\*



\*89/90Sr can also be measured by placing a 2mL Sr Resin cartridge below DGA and following the separation scheme in application note AN-1434

\*\*Adding 50uL of 30% H<sub>2</sub>O<sub>2</sub> to the 6M HNO<sub>3</sub> tube rinse can help improve U recoveries and decontamination in the Pu/Np fraction.

## References

1) Sherrod L. Maxwell, Brian K. Culligan, Gary W. Noyes, "Rapid separation method for actinides in emergency air filter samples," *Applied Radiation and Isotopes*, 68(12), 2125-2131 (2010).

# Rapid Determination of Sr in Emergency Air Filter Samples

#### AN-1434-10

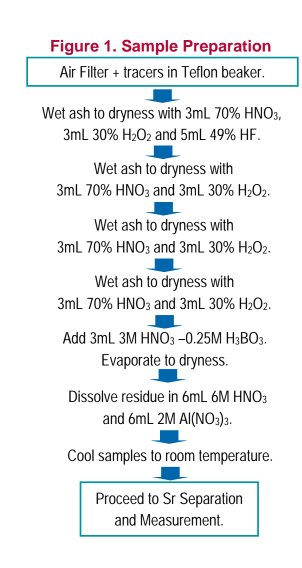
**Summary of Method** Strontium is separated and concentrated from air filters. Samples are digested in Teflon beakers once with HNO<sub>3</sub>-H<sub>2</sub>O<sub>2</sub>-HF and then several times with HNO<sub>3</sub>-H<sub>2</sub>O<sub>2</sub>. After evaporating to dryness from HNO<sub>3</sub>-H<sub>3</sub>BO<sub>3</sub> to complex any residual fluoride, strontium is separated on a 2mL cartridges of Eichrom Sr resin. Radiostrontium is measured by low background gas flow proportional counting or liquid scintillation counting. Chemical yield of strontium, which averaged 86±5%, is determined by gravimetric recovery of stable strontium carrier or ICP-AES measurement. <sup>90</sup>Sr measurements agreed to within 10% of reference values. <sup>89</sup>Sr and <sup>90</sup>Sr activities can be determined by Cerenkov counting or by subsequent <sup>90</sup>Y ingrowth, separation and measurement. Sample preparation for batches of 12 samples can be completed by a single operator in <8 hours.

#### Reagents

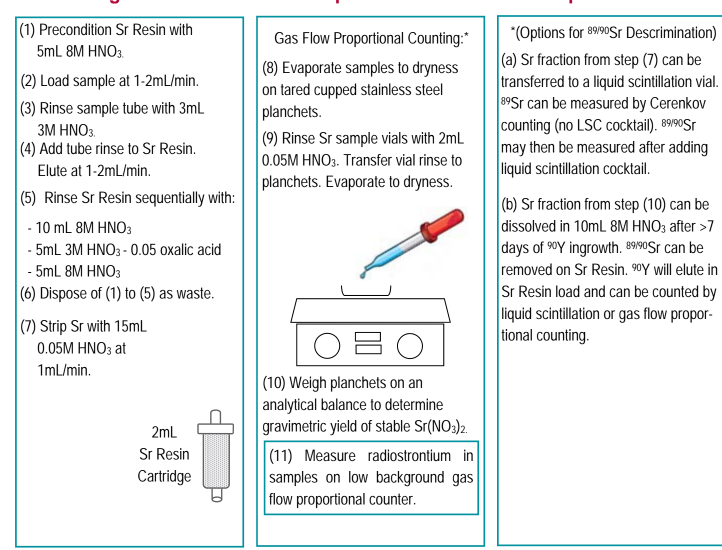
Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) Oxalic acid/Ammonium oxalate Sr carrier (10mg/mL) Deionized Water 2M Al(NO<sub>3</sub>)<sub>3</sub> HNO<sub>3</sub> (70%) HF (49%) or NaF Boric acid H<sub>2</sub>O<sub>2</sub> (30%)

#### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Hot Plate Analytical Balance 250mL Teflon beakers Cupped Stainless Steel Planchets (~5mL volume) Low background gas flow proportional counter Vacuum Pump



### Figure 2. Load Solution Preparation and Strontium Separation



\*Actinides also be measured by placing 2mL cartridges of TEVA, TRU and DGA resin above Sr Resin and following the separation scheme in application note AN-1433.

### References

1) Sherrod L. Maxwell, Brian K. Culligan, Gary W. Noyes, "Rapid separation method for actinides in emergency air filter samples," *Applied Radiation and Isotopes*, 68(12), 2125-2131 (2010).

AN-1435-10

# Rapid Determination of Np/Pu in 20-50g Soil Samples

# **Summary of Method** Plutonium and Neptunium are separated and concentrated from 20-50 gram soil samples. Samples are leached with HNO<sub>3</sub> and HCI. The leachates are evaporated to dryness, and sequential precipitations with Fe/Ti-hydroxide and LaF<sub>3</sub> facilitate matrix removal. Pu-Np are separated on 2mL cartridges of Eichrom TEVA resin. Pu-Np are measured by alpha spectrometry following CeF<sub>3</sub> microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Chemical yields of the <sup>236</sup>Pu tracer ranged from 82-96%. Measured values typically agreed to within 10% of reference values. Sample preparation for batches of 12 samples can be completed by a single operator in <8 hours.

#### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>236</sup>Pu tracer NH<sub>4</sub>OH (28% NH<sub>3</sub> or 56% NH<sub>4</sub>OH) La carrier (10mg/mL) Ce carrier (1mg/mL) **Deionized Water** 2M AI(NO<sub>3</sub>)<sub>3</sub> HNO<sub>3</sub> (70%) 10% (w:w) TiCl<sub>3</sub> HF (49%) or NaF HCI (37%) Boric acid  $H_2O_2$  (30%) NaNO<sub>2</sub> Denatured ethanol Sulfamic Acid Ascorbic Acid

### Figure 1. Sample Preparation

Dry 20-50g soil at 110°C. Blend and Size. Aliquot sample to 600mL glass beaker. Add <sup>236</sup>Pu. Add 1.5mL 70% HNO<sub>3</sub> and 0.5mL 37% HCl per gram of sample. Heat to 80°C on hotplate. Transfer liquid to 250mL centrifuge tube. Add 20mL 70% HNO<sub>3</sub> to beaker. Warm beaker. Transfer liquid to same 250mL centrifuge tube. Repeat once. Centrifuge 3500 rpm, 10 min. Transfer leachate to 600mL beaker. Evaporate to dryness. Dissolve residue in 20mL 1M HCI. Warm if necessary. Dilute samples to 180mL. Add 5mg La, 125mg of Fe, and 20mL 10% TiCl<sub>3</sub>. Mix. Add 25mL 56% NH<sub>4</sub>OH. Mix. Centrifuge 3500 rpm. 5min. Decant supernate Partially dissolve in 60mL 1.5M HCI. Solids will remain. Dilute to 170mL. Add 3mg La and 20mL 10% TiCl<sub>3</sub>. Mix.

> Add 22mL 49% HF. Mix. Place in ice bath for 10min.

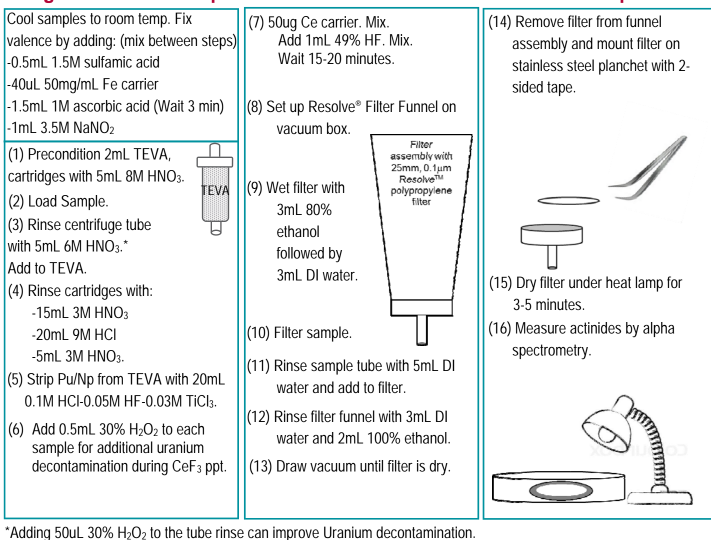
Centrifuge 3500 rpm. 5min. Decant supernate.

Dissolve solids in 6mL 3M HNO<sub>3</sub>-0.25M H<sub>3</sub>BO<sub>3</sub>, 8.5mL 7M HNO<sub>3</sub>, and 8mL 2M Al(NO<sub>3</sub>)<sub>3</sub>. Warming samples can improve dissolution.

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 600mL Glass beakers Stainless Steel Planchets with adhesive tape Alpha Spectrometry System 50mL and 250mL Centrifuge Tubes Centrifuge Heat Lamp Hot Plate Analytical Balance Vacuum Pump

### Figure 2. Actinide Separation on TEVA - TRU - DGA and Source Preparation



|           |            | <sup>236</sup> Pu | <sup>239</sup> Pu | <sup>239</sup> Pu |                   | <sup>238</sup> Pu | <sup>238</sup> Pu |                   | <sup>237</sup> Np | <sup>237</sup> Np |                   |
|-----------|------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|-------------------|
|           |            | Tracer            | Reference         | Measured          | <sup>239</sup> Pu | Reference         | Measured          | <sup>238</sup> Pu | Reference         | Measured          | <sup>237</sup> Np |
| Sample, g | replicates | % Yield           | (mBq)             | (mBq)             | %Bias             |                   | (mBq)             | %Bias             |                   | (mBq)             | %Bias             |
| 20        | 6          | 89 <u>+</u> 6     | 116.3             | 118 <u>+</u> 7    | 1.5               | 63.2              | 67 <u>+</u> 4     | 6.0               | 37.0              | 39 <u>+</u> 4     | 5.4               |
| 20        | 6          | 96 <u>+</u> 7     | 1.69              | 2.1 <u>+</u> 0.4  | 24                | 25.3              | 25 <u>+</u> 2     | -1.2              | 37.0              | 35 <u>+</u> 2     | -5.4              |
| 30        | 6          | 82 <u>+</u> 6     | 116.3             | 121 <u>+</u> 5    | 4.0               | 63.2              | 68 <u>+</u> 5     | 7.6               | 37.0              | 39 <u>+</u> 4     | 5.4               |
| 50        | 6          | 88 <u>+</u> 5     | 116.3             | 114 <u>+</u> 3    | -2.0              | 63.2              | 64 <u>+</u> 2     | 1.3               | 37.0              | 21 <u>+</u> 11    | -43               |

### References

1) Sherrod L. Maxwell, Brian K. Culligan, Gary W. Noyes, "Rapid separation method for <sup>237</sup>Np and Pu isotopes in large soil samples," *Applied Radiation and Isotopes*, 69(7), 917-925 (2011).

### **Rapid Determination of Np/Pu** in 20-75g Soil Samples (ICP-MS)

#### AN-1436-10

**Summary of Method** Plutonium and Neptunium are separated and concentrated from 20-75 gram soil samples. Samples are leached with HNO<sub>3</sub> and HCI. The leachates are evaporated to dryness, and sequential precipitations with Fe/Ti-hydroxide and LaF<sub>3</sub> facilitate matrix removal. Pu-Np are separated on 2mL cartridges of Eichrom TEVA and DGA resins. Pu-Np are measured by ICP-MS. Chemical yields of the <sup>242</sup>Pu tracer were 87+4%, 75+6%, and 70+3% for 20, 50 and 75g samples, respectively. Measured values for <sup>239</sup>Pu agreed to within 1% of reference values, while <sup>237</sup>Np agreed to within 15%. Decontamination factors of >106 were achieved for Pu over U (238U-H can interfere with the measurement of <sup>239</sup>Pu by ICP-MS). Sample preparation for batches of 12 sam-**Figure 1. Sample Preparation** 

ples can be completed by a single operator in <8 hours.

### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) DGA Resin, Normal, 2mL Cartridges (Eichrom DN-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu tracer La carrier (10mg/mL) **Deionized Water**  $2M AI(NO_3)_3$ 10% (w:w) TiCl<sub>3</sub> HNO<sub>3</sub> (70%) NH<sub>4</sub>OH (28% HN<sub>3</sub> or 56% NH<sub>4</sub>OH) HCI (37%) HF (49%) or NaF Boric acid NaNO<sub>2</sub> Sulfamic Acid Ascorbic Acid Hydroxylamine Hydrochloride

Dry 20-75g soil at 110°C. Blend and Size.

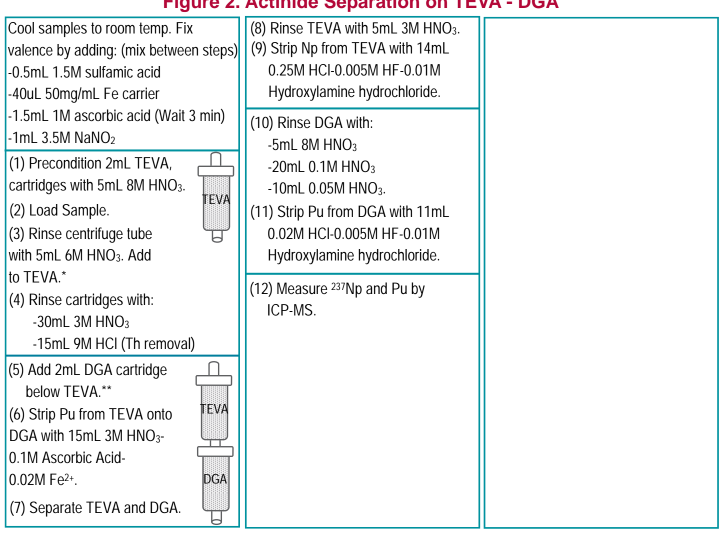
Aliguot sample to 600mL glass beaker. Add <sup>242</sup>Pu. Add 1.5mL 70% HNO<sub>3</sub> and 0.5mL 37% HCl per gram of sample. Heat to 80°C on hotplate. Transfer liquid to 250mL centrifuge tube. Add 20mL 70% HNO<sub>3</sub> to beaker. Warm beaker. Transfer liquid to same 250mL centrifuge tube. Repeat once. Centrifuge 3500 rpm, 10 min. Transfer leachate to 600mL beaker. Evaporate to dryness. Dissolve residue in 20mL 1M HCI. Warm if necessary. Dilute samples to 180mL. Add 5mg La, 125mg of Fe, and 20mL 10% TiCl<sub>3</sub>. Mix. Add 25mL 56% NH<sub>4</sub>OH. Mix. Centrifuge 3500 rpm. 5min. Decant supernate Partially dissolve in 60mL 1.5M HCI. Solids will remain. Dilute to 170mL. Add 3mg La and 20mL 10% TiCl<sub>3</sub>. Mix. Add 22mL 49% HF. Mix. Place in ice bath for 10min. Centrifuge 3500 rpm. 5min. Decant supernate. Dissolve solids in 6mL 3M HNO<sub>3</sub>-0.25M H<sub>3</sub>BO<sub>3</sub>, 8.5mL 7M HNO<sub>3</sub>, and 8mL 2M AI(NO<sub>3</sub>)<sub>3</sub>. Warming

samples can improve dissolution.

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 600mL Glass beakers 50mL and 250mL Centrifuge Tubes **ICP-MS** system Centrifuge Hot Plate Analytical Balance Vacuum Pump

### Figure 2. Actinide Separation on TEVA - DGA



\*Adding 50uL of 30% H<sub>2</sub>O<sub>2</sub> to the 6M HNO<sub>3</sub> tube rinse can further improve uranium decontamination factors.

\*\*Placing a 1mL UTEVA cartridge between TEVA and DGA can provide additional decontamination from uranium.

### References

- 1) Sherrod L. Maxwell, Brian K. Culligan, Gary W. Noyes, "Rapid separation method for <sup>237</sup>Np and Pu isotopes in large soil samples," Applied Radiation and Isotopes, 69(7), 917-925 (2011).
- 2) Sherrod L. Maxwell, Brian K. Culligan, Vernon D. Jones, Sheldon T. Nichols, Gary W. Noyes, Maureen A. Bernard, "Rapid Determination of <sup>237</sup>Np and Plutonium Isotopes by ICP-MS and Alpha Spectrometry," Health Physics, 101(2), 180-186 (2011).

# Rapid Determination of Actinides in Urine by ICP-MS + Alpha Spec.

#### AN-1437-10

**Summary of Method** Actinides are separated and concentrated from 100mL urine samples. Actinides are concentrated from urine samples using a calcium phosphate precipitation. Pu, Np, Am-Cm, and U are separated on 2mL cartridges of Eichrom TEVA, TRU and DGA resins. Pu-Np are measured by ICP-MS. Measured values for <sup>239</sup>Pu and <sup>237</sup>Np agreed to within 1-2% of reference values, while <sup>241</sup>Am and <sup>244</sup>Cm agreed to within 2-3%. Decontamination factors of >10<sup>6</sup> were achieved for Pu over U (<sup>238</sup>U-H can interfere with the measurement of <sup>239</sup>Pu by ICP-MS). Sample preparation for batches of 12 samples can be completed by a single operator in <8 hours.

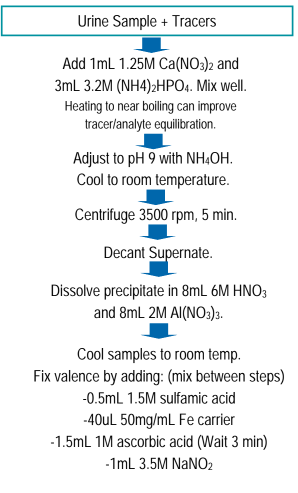
### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) DGA Resin, Normal, 2mL Cartridges (Eichrom DN-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) 242Pu (ICP-MS) or 236Pu (alpha) tracer <sup>233</sup>U (ICP-MS) or U<sup>232</sup> (alpha) tracer <sup>243</sup>Am tracer Ce carrier (1mg/mL) 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> **Deionized Water** 2M AI(NO<sub>3</sub>)<sub>3</sub> HNO<sub>3</sub> (70%) HCI (37%) NH<sub>4</sub>OH HF (49%) or NaF NaNO<sub>2</sub> Denatured ethanol Sulfamic Acid Ascorbic Acid Oxalic acid/Ammonium oxalate Hydroxylamine Hydrochloride

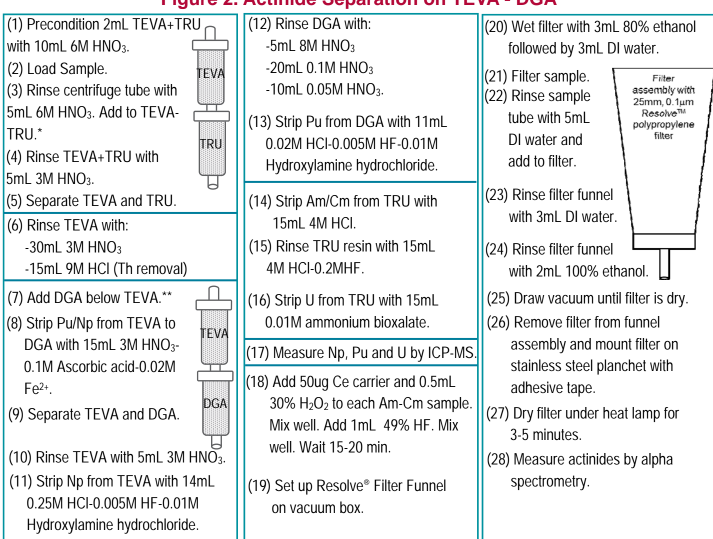
### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 600mL Glass beakers Stainless Steel Planchets with adhesive tape Alpha Spectrometry System ICP-MS System 50mL and 250mL Centrifuge Tubes Centrifuge Heat Lamp Hot Plate Analytical Balance Vacuum Pump

### Figure 1. Sample Preparation



### Figure 2. Actinide Separation on TEVA - DGA



\* Adding 50uL of 30% H<sub>2</sub>O<sub>2</sub> to the 6M HNO<sub>3</sub> tube rinse can help improve U recoveries and decontamination in the Np and Pu fractions.

\*\* Adding a 1mL UTEVA cartridge between TEVA and DGA can help improve uranium decontamination.

### References

- 1) Sherrod L. Maxwell, Vernon D. Jones, "Rapid determination of Actinides in urine by ICP-MS and alpha spectrometry: A hybrid approach," *Talanta,* 80(1), 143-150 (2009).
- Sherrod L. Maxwell, Brian K. Culligan, Vernon D. Jones, Sheldon T. Nichols, Gary W. Noyes, Maureen A. Bernard, "Rapid Determination of <sup>237</sup>Np and Plutonium Isotopes by ICP-MS and Alpha Spectrometry," *Health Physics*, 101(2), 180-186 (2011).

### Rapid Determination of Np/Pu in Water Samples by ICP-MS

### AN-1438-10

**Summary of Method** Plutonium and Neptunium are separated and concentrated from 200mL water samples. Pu and Np are concentrated from the water sample using a calcium phosphate precipitation. Pu-Np are separated on 2mL cartridges of Eichrom TEVA and DGA resins. Pu-Np are measured by ICP-MS. Measured values for <sup>239</sup>Pu, <sup>242</sup>Pu, and <sup>237</sup>Np agreed to within 1-4% of reference values, while <sup>237</sup>Np agreed to within 15%. Decontamination factors of >10<sup>6</sup> were achieved for Pu over U (<sup>238</sup>U-H can interfere with the measurement of <sup>239</sup>Pu by ICP-MS). Sample preparation for batches of 12 samples can be completed by a single operator in <4 hours.

### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) DGA Resin, Normal, 2mL Cartridges (Eichrom DN-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu tracer 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> **Deionized Water**  $2M AI(NO_3)_3$ HNO<sub>3</sub> (70%) HCI (37%) NH<sub>4</sub>OH HF (49%) or NaF NaNO<sub>2</sub> Sulfamic Acid Ascorbic Acid Hydroxylamine Hydrochloride

### Figure 1. Sample Preparation

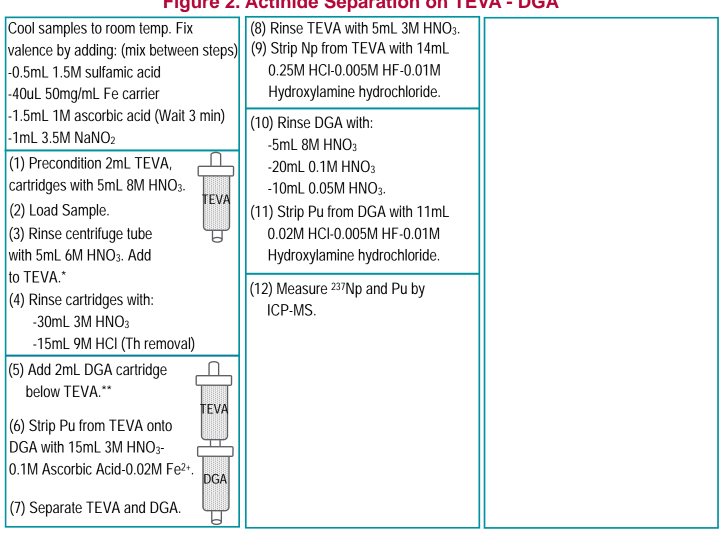
Water Sample + <sup>242</sup>Pu Tracer Add 1mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> and 3mL 3.2M (NH4)<sub>2</sub>HPO<sub>4</sub>. Mix well. Heating to near boiling can improve tracer/analyte equilibration. Adjust to pH 9 with NH<sub>4</sub>OH. Cool to room temperature. Centrifuge 3500 rpm, 5 min. Decant Supernate. Dissolve precipitate in 8mL 6M HNO<sub>3</sub>

and 8mL 2M AI(NO<sub>3</sub>)<sub>3</sub>.

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 600mL Glass beakers 50mL and 250mL Centrifuge Tubes Centrifuge Hot Plate Analytical Balance Vacuum Pump ICP-MS System

### Figure 2. Actinide Separation on TEVA - DGA



\* Adding 50uL of 30%  $H_2O_2$  to 6M HNO<sub>3</sub> tube rinse can help improve U decontamination.

\*\* Adding a 1mL UTEVA cartridge between TEVA and DGA can provide additional uranium decontamination.

### References

- 1) Sherrod L. Maxwell, Brian K. Culligan, Vernon D. Jones, Sheldon T. Nichols, Gary W. Noyes, "Rapid determination of <sup>237</sup>Np and Pu isotopes in water by ICP-MS and alpha spectrometry," J. Radioanal. Nucl. Chem., 287(1), 223-230 (2011).
- 2) Sherrod L. Maxwell, Brian K. Culligan, Vernon D. Jones, Sheldon T. Nichols, Gary W. Noyes, Maureen A. Bernard, "Rapid Determination of <sup>237</sup>Np and Plutonium Isotopes by ICP-MS and Alpha Spectrometry," Health Physics, 101(2), 180-186 (2011).

### Determination of <sup>227</sup>Ac in Geological Samples

### AN-1601-10

**Summary of Method** Soil or rock samples are pulverized to <1mm and dissolved, either by acid digestion or sodium hydroxide fusion. <sup>227</sup>Ac is separated from matrix ions using a ferric hydroxide precipitation step. Following dissolution in 4M HCl, <sup>227</sup>Ac is separated from radiometric impurities using a 2mL cartridge of DGA, Normal resin. <sup>227</sup>Ac is prepared for measurement using a CeF<sub>3</sub> microprecipitation onto Resolve<sup>(R)</sup> Filters. An <sup>225</sup>Ac(<sup>229</sup>Th) tracer is used to

measure chemical recovery of actinium. After a 30 minute ingrowth time, the <sup>225</sup>Ac tracer yield is measured via alpha spectrometry using the <sup>221</sup>Fr and <sup>217</sup>At daughters of <sup>225</sup>Ac. <sup>227</sup>Ac is measured via its <sup>227</sup>Th and <sup>223</sup>Ra daughters after a longer period of ingrowth (30-90 days). Ac yields are typically 70-90%. MDA for <sup>227</sup>Ac was 0.05Bq/kg for 3 day count times after 90 days ingrowth period.

### Reagents

DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Iron Carrier (50mg/mL Fe, as ferric nitrate) Cerium Carrier (10mg/mL) <sup>229</sup>Th(<sup>225</sup>Ac) tracer Hydrofluoric Acid (49%) or Sodium Fluoride Boric acid HNO<sub>3</sub> (70%) HCI (37%) NaOH Deionized Water H<sub>2</sub>O<sub>2</sub> (30%) Optional for additional Th/U removal: TRU Resin, 2mL cartridges (Eichrom TR-R50-S)

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Alpha Spectrometry System Ball mill grinder or equivalent Centrifuge Vacuum Pump Heat Lamp Analytical Balance

### **Fusion Option**

250mL Zirconium crucibles with zirconium lids Muffle Furnace

### **Digestion Option**

Hot Plate Teflon Beakers

### **Sample Preparation**

0.25-50g Soil or Rock

Pulverize to <1mm.

Aliquot Sample. Add <sup>229</sup>Th(<sup>225</sup>Ac) tracer.

### **Acid Digestion Option**

Digest in Teflon beaker on hotplate with 2:1 conc. HNO<sub>3</sub>:HF to near dryness.

Digest in Teflon beaker on hotplate with conc. HNO<sub>3</sub> + Boric Acid.

Dissolve Residue in 4M HCl + 0.25M Boric acid. If solids remain. Repeat digestion. Proceed to ferric hydroxide precipitation.

### **Fusion Option**

In Zr crucible. Add 10-15g NaOH

Muffle at 600°C for 15-30 minutes.

Cool. Dissolve fusion cake with 50mL water. Heat as necessary. Rinse crucible with 50mL 4M HCl. Proceed to ferric hydroxide precipitation.

### Ferric Hydroxide Precipitation

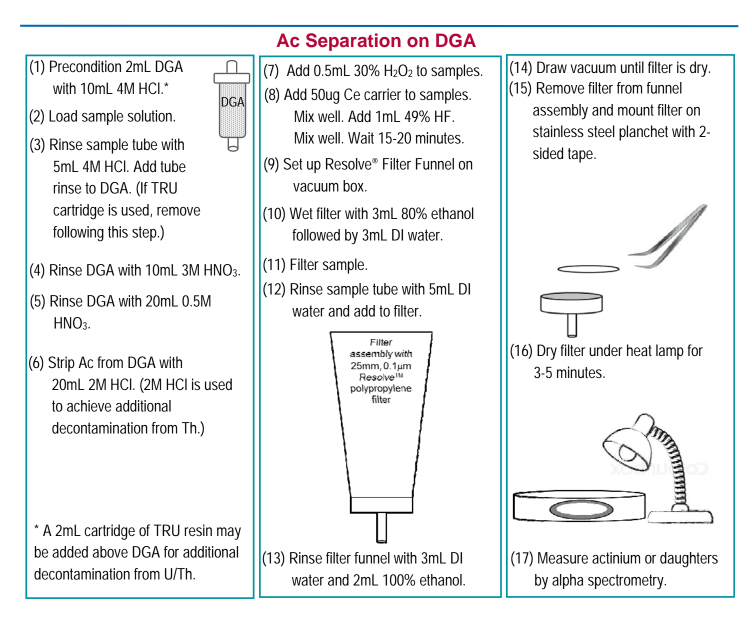
Transfer sample to 250mL centrifuge tube.

Dilute to 150mL with water. Add 25mg Fe carrier. Mix well.

Centrifuge 2500 rpm for 10 minutes. Decant Supernate.

Rinse ppt with 50mL water. Centrifuge. Decant Supernate.

Dissolve precipitate with 10mL conc. HCI. Dilute to 30mL.



### **Method Performance**

|          | <sup>227</sup> Ac    | <sup>227</sup> Ac |        |          |
|----------|----------------------|-------------------|--------|----------|
| Rock     | Measured             | Reference         |        | Tracer   |
| Standard | Bq/kg                | Bq/kg             | % Bias | Recovery |
| BCR-2    | 0.955 <u>+</u> 0.083 | 0.967             | -1.2   | 83       |
| BHVO-1   | 0.299 <u>+</u> 0.017 | 0.283             | 5.7    | 71       |
| HK-018   | 0.965 <u>+</u> 0.009 | 0.948             | 1.8    | 86       |
| HK-019   | 0.962 <u>+</u> 0.073 | 0.966             | -0.4   | 91       |
| HK-021   | 0.559 <u>+</u> 0.055 | 0.572             | -2.3   | 80       |
| HK-022   | 0.887 <u>+</u> 0.080 | 0.862             | 2.9    | 68       |
| SAV B6   | 0.677 <u>+</u> 0.067 | 0.680             | -0.4   | 66       |

### References

1) H. Dulaiova, K.W.W. Sims, M.A. Charette, J. Prytulak, J.S. Blusztajn "A new method for the determination of actinium-227 in geological samples," *J. Radioanal. Nucl. Chem., 296, 279-283* (2013).

### Determination of <sup>227</sup>Ac in Water Samples

### AN-1602-10

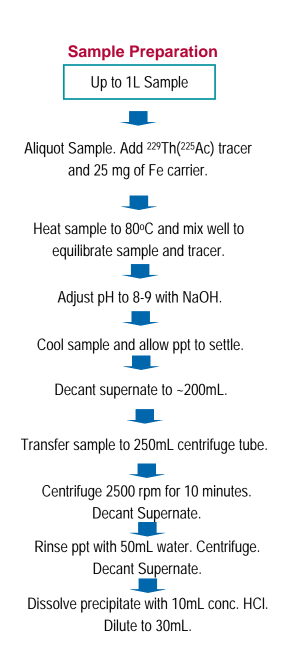
**Summary of Method** <sup>227</sup>Ac is preconcentrated from up to 1L of water sample using a ferric hydroxide precipitation. Following dissolution in 4M HCl, <sup>227</sup>Ac is separated from radiometric impurities using a 2mL cartridge of DGA, Normal resin. <sup>227</sup>Ac is prepared for measurement using a CeF<sub>3</sub> microprecipitation onto Resolve<sup>(R)</sup> Filters. An <sup>225</sup>Ac(<sup>229</sup>Th) tracer is used to measure chemical recovery of actinium. After a 30 minute ingrowth time, the <sup>225</sup>Ac tracer yield is measured via alpha spectrometry. <sup>227</sup>Ac is measured via its <sup>227</sup>Th and <sup>223</sup>Ra daughters after a longer period of ingrowth (30-90 days). Actinium yields are typically 70-90%. MDA for <sup>227</sup>Ac was 0.05Bq/L for 3 day count times after 90 days ingrowth period.

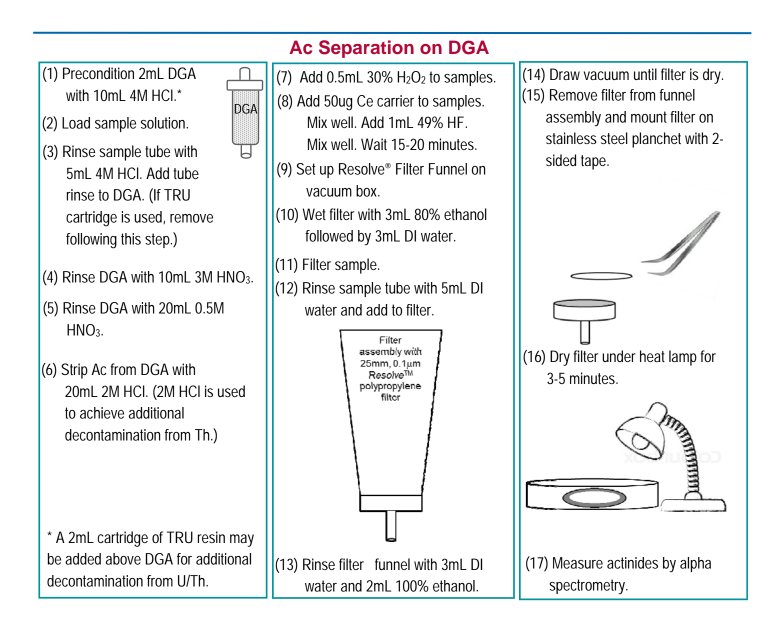
### Reagents

DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Iron Carrier (50mg/mL Fe, as ferric nitrate) Cerium Carrier (10mg/mL) <sup>229</sup>Th(<sup>225</sup>Ac) tracer Hydrofluoric Acid (49%) or Sodium Fluoride Nitric Acid (70%) Hydrochloric Acid (37%) Sodium Hydroxide Deionized Water H<sub>2</sub>O<sub>2</sub> (30%) Optional for additional Th/U removal: TRU Resin, 2mL cartridges (Eichrom TR-R50-S)

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Alpha Spectrometry System Centrifuge Vacuum Pump Heat Lamp Analytical Balance Hot Plate 1L Glass beakers pH meter or pH strips or pH indicator (pH 8-9)





| Method Performance |                   |                    |        |                    |  |  |
|--------------------|-------------------|--------------------|--------|--------------------|--|--|
|                    | <sup>227</sup> Ac | <sup>227</sup> Ac  |        |                    |  |  |
| Water<br>Standard  | Measured<br>Bq/kg | Reference<br>Bq/kg | % Bias | Tracer<br>Recovery |  |  |
| IAEA               |                   |                    |        |                    |  |  |

### References

1) H. Dulaiova, K.W.W. Sims, M.A. Charette, J. Prytulak, J.S. Blusztajn "A new method for the determination of actinium-227 in geological samples," *J. Radioanal. Nucl. Chem., 296, 279-283* (2013).

### **Rapid Determination of Actinides** eichrom in Limestone and Marble

### AN-1603-10

**Summary of Method** Actinides are separated and concentrated from 1.5 gram samples of limestone or marble. Samples are finely ground and fused in a zirconium crucible for 15 minutes at 600°C with 15 grams of sodium hydroxide. The fusion cake is dissolved in water, and actinides are concentrated and separated from the matrix using a calcium phosphate precipitate enhanced with iron. A secondary precipitation with calcium fluoride removes additional matrix (including silicates) and decreases the volume of precipitate. The calcium fluoride precipitate is dissolved with nitric acid-boric acid-aluminum nitrate to form the load solution. Analytes are separated from remaining matrix and potentially interfering radionuclides using stacked TEVA, TRU, and DGA Resin cartridges. Actinides are measured by alpha spectrometry after CeF<sub>3</sub> microprecipitation onto Resolve<sup>(R)</sup> Filters. Simultaneous separation of radiostrontium can be achieved by using the separation method in AN-1604.

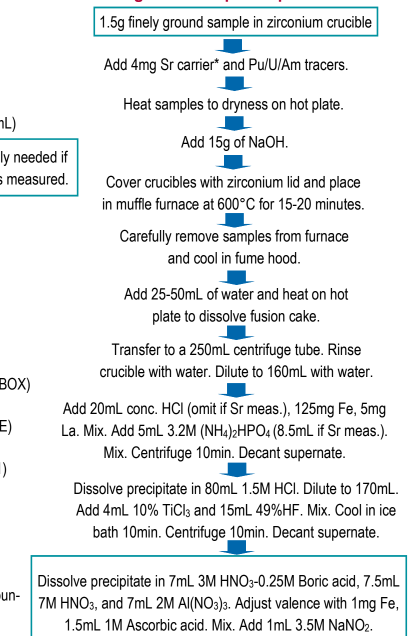
### Reagents

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S)\* Sr Resin, 1mL Cartridges (Eichrom SR1ML-R50-S)\* TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) DGA Resin, 2mL Cartridges (Eichrom DN-R50-S) Strontium\*, Lanthanum and Cerium Carriers (10mg/mL) Iron Carrier (50mg/mL Fe, as ferric nitrate) \*Only needed if <sup>242</sup>Pu (or <sup>236</sup>Pu if Np is measured) tracer Sr is measured. <sup>243</sup>Am and <sup>232</sup>U tracers 10% TiCl<sub>3</sub> <sup>90</sup>Sr standard\* HF(49%) 30% H<sub>2</sub>O<sub>2</sub> Nitric Acid (70%) Hydrochloric Acid (37%) **Deionized Water** 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>  $2M AI(NO_3)_3$ Oxalic acid Boric acid Sodium Hydroxide Ascorbic acid NaNO<sub>2</sub> Sulfamic acid

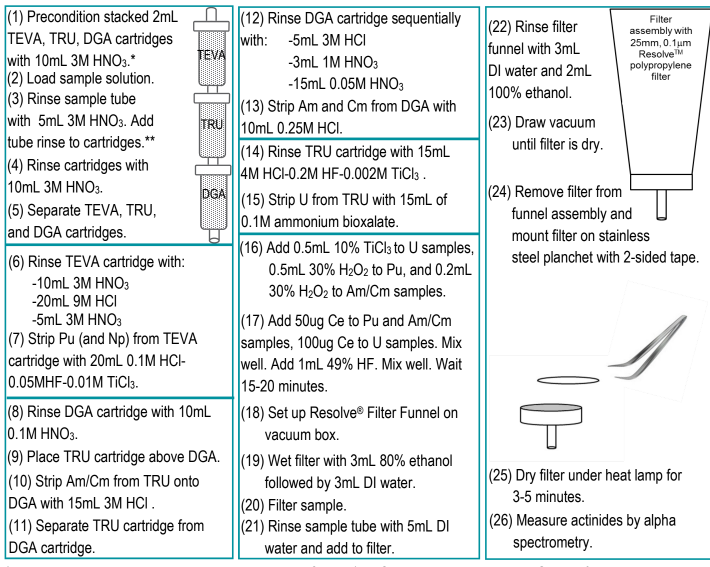
### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Cupped Stainless Steel Planchets (~5mL volume)\* 250mL Zirconium crucibles with zirconium lids Alpha Spectrometry System Stainless Steel planchets with two sided tape Centrifuge Gas Flow Proportional Counter\* Muffle Furnace Hot Plate/Heat Lamp

### Figure 1. Sample Preparation



### Figure 2. Actinide Separation on TEVA - TRU - DGA\* and Source Preparation



\*Radiostrontium may also be measured by adding a 2mL + 1mL Sr Resin cartridge below DGA and following separation scheme in Eichrom Application note AN-1604-10.

\*\*Adding 50uL 30% H<sub>2</sub>O<sub>2</sub> to the tube rinse can improve Uranium recoveries and decontamination in Pu(Np) fractions. Method Performance

| Sample         | replicates | analyte               | tracer            | % tracer<br>recovery | mBq/g<br>reference | mBq/g<br>measured | % bias     |
|----------------|------------|-----------------------|-------------------|----------------------|--------------------|-------------------|------------|
| 1.5g limestone | 6          | <sup>239/240</sup> Pu | <sup>242</sup> Pu | 100 <u>+</u> 5       | 29.4               | 30 <u>+</u> 2     | <u>+</u> 5 |
| 1.5g limestone | 6          | <sup>239/240</sup> Pu | <sup>236</sup> Pu | 93 <u>+</u> 6        | 23.0               | 24 <u>+</u> 1     | <u>+</u> 5 |
| 1.5g limestone | 6          | <sup>238</sup> Pu     | <sup>236</sup> Pu | 93 <u>+</u> 6        | 28.8               | 29 <u>+</u> 2     | <u>+</u> 5 |
| 1.5g limestone | 6          | <sup>237</sup> Np     | <sup>236</sup> Pu | 93 <u>+</u> 6        | 37.0               | 39 <u>+</u> 3     | <u>+</u> 7 |
| 1.5 g marble   | 4          | <sup>239/240</sup> Pu | <sup>242</sup> Pu | 96 <u>+</u> 3        | 29.4               | 30 <u>+</u> 2     | <u>+</u> 6 |
| 1.5 g marble   | 4          | <sup>241</sup> Am     | <sup>243</sup> Am | 89 <u>+</u> 4        | 29.1               | 29 <u>+</u> 1     | <u>+</u> 3 |
| 1.5 g marble   | 4          | <sup>244</sup> Cm     | <sup>243</sup> Am | 89 <u>+</u> 4        | 34.8               | 35 <u>+</u> 3     | <u>+</u> 6 |
| 1.5 g marble   | 7          | <sup>238</sup> U      | <sup>232</sup> U  | 93 <u>+</u> 6        | 50.2               | 48 <u>+</u> 1     | <u>+</u> 4 |

### References

1) Maxwell, Culligan, Hutchinson, Utsey, Sudowe, McAlister, "Rapid Method to Determine Actinides and Sr-89/90 in Limestone and Marble Samples," *J. Radioanal. Nucl. Chem.* 310, 377-388 (2016).

### Rapid Determination of <sup>89/90</sup>Sr in Limestone and Marble

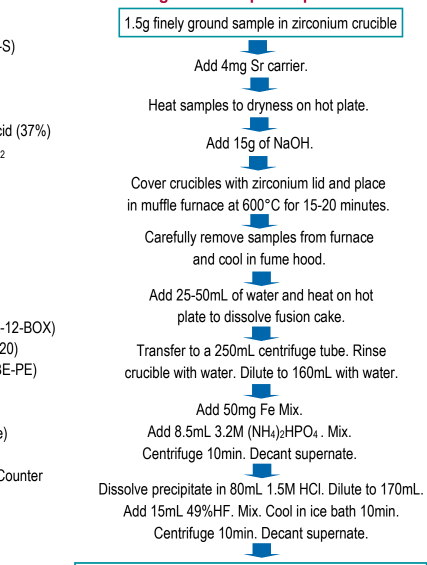
#### AN-1604-10

**Summary of Method** Strontium is separated and concentrated from 1.5 gram samples of limestone or marble. Samples are finely ground and fused in a zirconium crucible for 15 minutes at 600°C with 15 grams of sodium hydroxide. The fusion cake is dissolved in water, and strontium is concentrated and separated from the matrix using a calcium phosphate precipitate enhanced with iron. A secondary precipitation with calcium fluoride removes additional matrix (including silicates) and decreases the volume of precipitate. The calcium fluoride precipitate is dissolved with nitric acid-boric acid-aluminum nitrate to form the load solution. Strontium is separated from remaining matrix and potentially interfering radionuclides using stacked 2mL + 1mL Sr Resin cartridges. Radiostrontium is measured by gas flow proportional counting or liquid scintillation counting. Chemical yields of strontium are determined by gravimetric yield or by ICP-AES. Batches of 12-24 samples can be prepared for analysis in less than 8 hours. Simultaneous separation of actinides can be achieved by using the separation method in AN-1603.

### Reagents

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S) Sr Resin, 1mL Cartridges (Eichrom SR1ML-R50-S) Strontium Carrier (10mg/mL) Iron Carrier (50mg/mL Fe, as ferric nitrate) <sup>90</sup>Sr standard HF(49%) Nitric Acid (70%) Hydrochloric Acid (37%) Deionized Water 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> 2M Al(NO<sub>3</sub>)<sub>3</sub> Oxalic acid Boric acid Sodium Hydroxide

### Figure 1. Sample Preparation

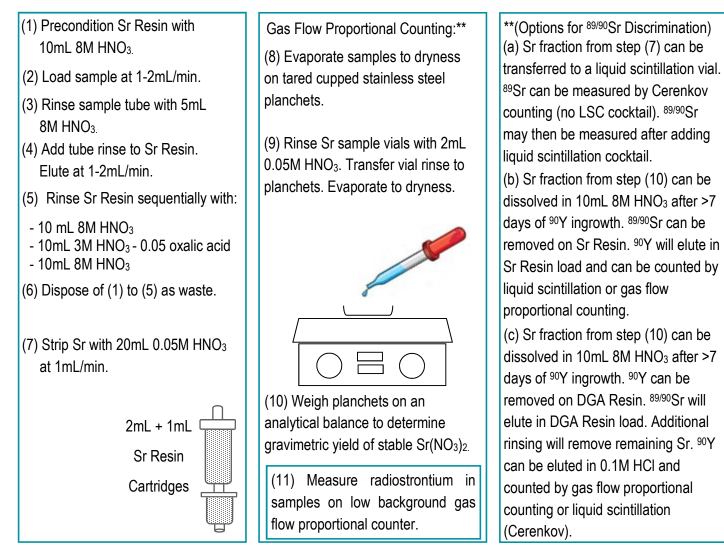


Dissolve precipitate in 7mL 3M HNO<sub>3</sub>-0.25M Boric acid, 7.5mL 7M HNO<sub>3</sub>, and 7mL 2M Al(NO<sub>3</sub>)<sub>3</sub>. Warm as needed.

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50mL and 250mL Centrifuge Tubes Cupped Stainless Steel Planchets (~5mL volume) 250mL Zirconium crucibles with zirconium lids Centrifuge Gas Flow Proportional Counter Muffle Furnace Hot Plate/Heat Lamp Analytical Balance Vacuum Pump

### Figure 2. Strontium Resin Separation (Optional <sup>90</sup>Y Ingrowth)\*



\*Actinides may also be measured by adding 2mL TEVA, TRU and DGA Resin cartridges above Sr Resin and following separation scheme in Eichrom Application note AN-1603.

\*\*Additional discussion of <sup>89/90</sup>Sr separation and measurement options can be found in Eichrom Application Note AN-1624-10.

### Method Performance

|        | % Sr tracer       | <sup>90</sup> Sr Bq/g | <sup>90</sup> Sr Bq/g |        |
|--------|-------------------|-----------------------|-----------------------|--------|
| Sample | recovery          | reference             | measured              | % bias |
| 1      | 84.1              | 1.415                 | 1.41                  | -0.1   |
| 2      | 84.8              | 1.415                 | 1.42                  | 0.4    |
| 3      | 84.8              | 1.415                 | 1.38                  | -2.7   |
| AVG    | 84.6 <u>+</u> 0.4 |                       | 1.40 <u>+</u> 0.02    |        |

### References

1) Maxwell, Culligan, Hutchinson, Utsey, Sudowe, McAlister, "Rapid Method to Determine Actinides and Sr-89/90 in Limestone and Marble Samples," *J. Radioanal. Nucl. Chem.* 310, 377-388 (2016).

### Rapid Determination of <sup>89/90</sup>Sr in 5g Concrete Samples

### AN-1605-10

**Summary of Method** Strontium is separated and concentrated from 5 gram concrete samples. Samples are finely ground and fused in a zirconium crucible for 30 minutes at 600°C with 30 grams of sodium hydroxide. The fusion cake is dissolved in water, and strontium is concentrated and separated from the matrix using a calcium phosphate precipitate enhanced with iron. A secondary precipitation with calcium fluoride removes additional matrix (including silicates) and decreases the volume of precipitate. The calcium fluoride precipitate is dissolved with nitric acid-boric acid -aluminum nitrate to form the load solution. Analytes are separated from remaining matrix and potentially interfering radionuclides using two stacked 2mL Sr Resin cartridges. Radiostrontium is measured by gas flow proportional counting or liquid scintillation counting. Chemical yields of strontium are determined by gravimetric yield or by ICP-AES. Batches of 12-24 samples can be prepared for analysis in less than 8 hours. For aged samples, where the shorter lived <sup>89</sup>Sr (t<sub>1/2</sub> = 50.55 days) is unlikely to be present, <sup>90</sup>Sr can be determined from the direct separation of its <sup>90</sup>Y daughter from up to 10g concrete samples, using Eichrom Application Note AN-1606.

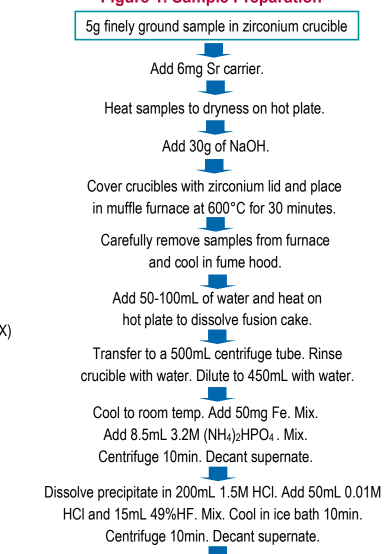
### Reagents

Sr Resin, 2mL Cartridges (Eichrom SR-R50-S)Strontium Carrier (10mg/mL)Iron Carrier (50mg/mL Fe, as ferric nitrate)90Sr standardHF(49%)30% H2O2Nitric Acid (70%)Hydrochloric Acid (37%)Deionized Water1.25M Ca(NO3)23.2M (NH4)2HPO42M Al(NO3)3Oxalic acidBoric acidSodium Hydroxide

### Equipment

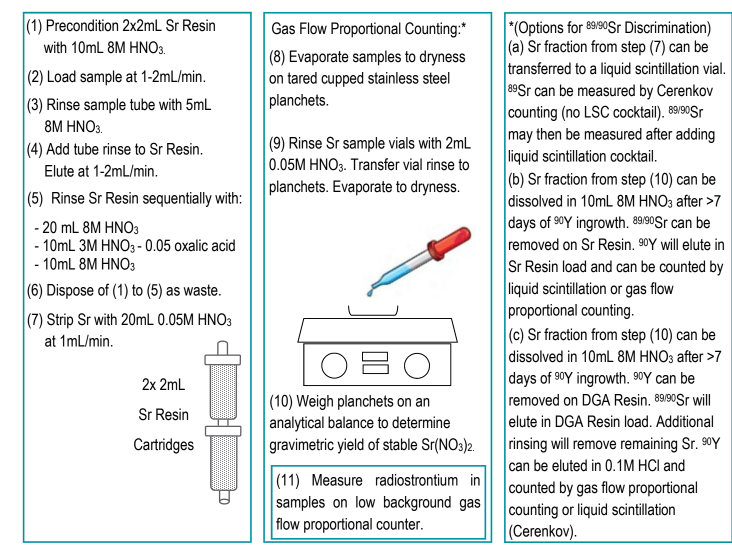
Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50mL and 500mL Centrifuge Tubes Cupped Stainless Steel Planchets (~5mL volume) 250mL Zirconium crucibles with zirconium lids Centrifuge Gas Flow Proportional Counter Muffle Furnace Hot Plate/Heat Lamp Analytical Balance Vacuum Pump

### Figure 1. Sample Preparation



Dissolve precipitate in 7mL 3M HNO<sub>3</sub>-0.25M Boric acid, 7mL conc. HNO<sub>3</sub>, 7mL 8M HNO<sub>3</sub> and 7mL 2M Al(NO<sub>3</sub>)<sub>3</sub>. Warm as needed.

### Figure 2. Strontium Resin Separation (Optional <sup>90</sup>Y Ingrowth)



\*Additional discussion of <sup>89/90</sup>Sr separation and measurement options can be found in Eichrom Application Note AN-1624-10.

### Method Performance (5g gram Concrete, Sr Resin Method)

|   |        | % Sr tracer   | <sup>90</sup> Sr Bq/g | <sup>90</sup> Sr Bq/g |        |
|---|--------|---------------|-----------------------|-----------------------|--------|
| ę | Sample | recovery      | reference             | measured              | % bias |
|   | 1      | 78.5          | 1.416                 | 1.51                  | 6.6    |
|   | 2      | 77.8          | 1.416                 | 1.35                  | -4.6   |
|   | 3      | 80.5          | 1.416                 | 1.42                  | 0.2    |
|   | 4      | 62.2          | 1.416                 | 1.49                  | 5.2    |
|   | AVG    | 75 <u>+</u> 8 |                       | 1.44 <u>+</u> 0.07    |        |

### References

1) Maxwell, Culligan, Hutchinson, Utsey, Sudowe, McAlister, "Rapid Method to Determine 89/90Sr in Large Concrete Samples," *J. Radioanal. Nucl. Chem.* 310, 399-411 (2016).

### Rapid Determination of <sup>90</sup>Sr in 10g Concrete Samples

#### AN-1606-10

**Summary of Method** <sup>90</sup>Sr is determined by the direct separation of its daughter <sup>90</sup>Y from 10 gram concrete samples. Samples are finely ground and fused in a zirconium crucible for 30 minutes at 600°C with 40 grams of sodium hydroxide. The fusion cake is dissolved in water, and strontium is concentrated and separated from the matrix using a ferric hydroxide precipitate. A secondary precipitation with Y/Ca-fluoride removes additional matrix (including silicates) and decreases the volume of precipitate. The Y/Ca-fluoride precipitate is dissolved with nitric acid-boric acid-aluminum nitrate to form the load solution. <sup>90</sup>Y is separated from remaining matrix and potentially interfering radionuclides using stacked 2mL TRU and DGA Resin cartridges. <sup>90</sup>Y is measured by gas flow proportional counting following microprecipitation onto Resolve<sup>®</sup> Filters. Chemical yields are determined by ICP-AES analysis. Batches of 12-24 samples can be prepared for analysis in less than 8 hours. This method is only suitable for aged samples, where the shorter lived <sup>89</sup>Sr (t<sub>1/2</sub> = 50.55 days) and fission products such as <sup>91</sup>Y are unlikely to be present. For samples not

meeting this criterion, <sup>89/90</sup>Sr can be determined from up to 5g concrete samples, using Eichrom Application Note AN-1605.

### Reagents

DGA Resin, 2mL Cartridges (Eichrom DN-R50-S)TRU Resin, 2mL Cartridges (Eichrom TR-R50-S)Yttrium Carrier (10mg/mL)Iron Carrier (50mg/mL Fe, as ferric nitrate)<sup>90</sup>Sr standardHF(49%)Nitric Acid (70%)Sodium HydroxideHydrochloric Acid (37%)Deionized Water1.25M Ca(NO<sub>3</sub>)23.2M (NH<sub>4</sub>)2HPO<sub>4</sub>2M Al(NO<sub>3</sub>)3Oxalic acid

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters with funnel (Eichrom RF-DF25-25PP01) 50mL and 500mL Centrifuge Tubes Stainless Steel Planchets with two sided tape 250mL Zirconium crucibles with zirconium lids Centrifuge Gas Flow Proportional Counter Muffle Furnace Hot Plate/Heat Lamp Analytical Balance Vacuum Pump

### Figure 1. Sample Preparation

10g finely ground sample in zirconium crucible

Add 2mg Y carrier. Heat samples to dryness on hot plate.

Add 40g of NaOH. Cover crucibles with zirconium lid and place in muffle furnace at 600°C for 30 minutes.

Carefully remove samples from furnace and cool in fume hood.

Add 50-100mL of water and heat on hot plate to dissolve fusion cake.

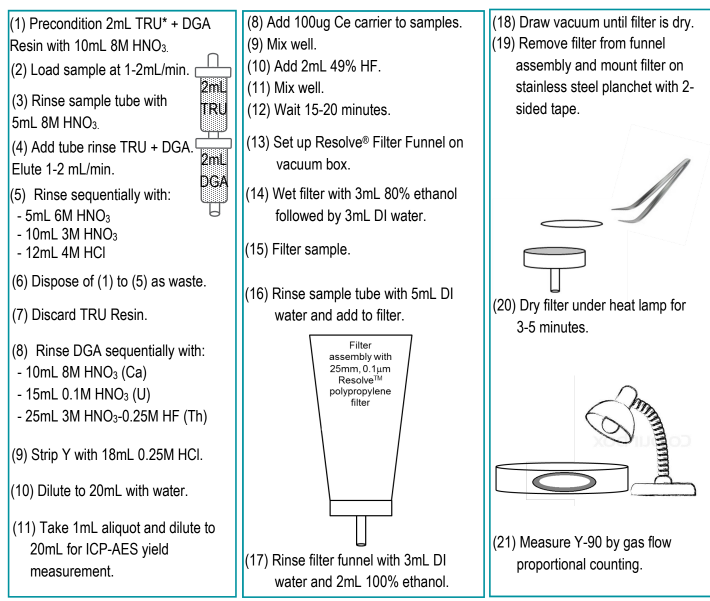
Transfer to a 500mL centrifuge tube. Rinse crucible with water. Dilute to 450mL with water. Cool to room temp. Add 125mg Fe Mix. Centrifuge 10min. Decant supernate.

Rinse precipitate with 150mL pH ~9 NaOH. Centrifuge. Decant Supernate. Repeat.

Dissolve precipitate in 200mL 1.5M HCI. Add 50mL 0.01M HCI and 15mL 49%HF. Mix. Cool in ice bath 10min. Centrifuge 10min. Decant supernate.

Dissolve precipitate in 7mL 3M HNO<sub>3</sub>-0.25M Boric acid, 7mL conc. HNO<sub>3</sub>, 7mL 8M HNO<sub>3</sub> and 7mL 2M Al(NO<sub>3</sub>)<sub>3</sub>. Warm as needed.

### Figure 2. TRU-DGA Separation and Gas Flow Proportional Counting



\*TRU Resin improves decontamination factors for U, Th and Bi isotopes, which could interfere with the measurement of <sup>90</sup>Y by gas flow proportional counting.

### Method Performance (10g gram Concrete, TRU/DGA Resin Method)

|        | % Y tracer    | <sup>90</sup> Sr Bq/g | <sup>90</sup> Sr Bq/g |        |
|--------|---------------|-----------------------|-----------------------|--------|
| Sample | recovery      | reference             | measured              | % bias |
| 1      | 81.7          | 0.0327                | 0.031                 | -5.4   |
| 2      | 83.3          | 0.0327                | 0.033                 | 1.2    |
| 3      | 83.7          | 0.0327                | 0.031                 | -5.0   |
| 4      | 86.3          | 0.0327                | 0.033                 | -0.6   |
| AVG    | 84 <u>+</u> 2 |                       | 0.032 <u>+</u> 0.001  |        |

### References

1) Maxwell, Culligan, Hutchinson, Utsey, Sudowe, McAlister, "Rapid Method to Determine 89/90Sr in Large Concrete Samples," *J. Radioanal. Nucl. Chem.* 310, 399-411 (2016).

AN-1607-10

### Rapid Determination of Np/Pu/Am in 10-20g Soil Samples

**Summary of Method** Plutonium, Neptunium, and Americium are separated and concentrated from 10 gram soil samples. Samples are fused in zirconium crucibles with 40g NaOH to facilitate complete dissolution. Actinides are separated from matrix using an iron/titanium hydroxide precipitate. A second precipitate with Ca/La-fluoride is used to remove additional matrix, particularly silicates, and decrease the volume of precipitate. Actinides are separated from potential radiometric impurities using 2mL cartridges of TEVA and DGA Resins. Am/Cm fractions may require additional purification using TEVA-NH<sub>4</sub>SCN to remove native rare earths which can degrade alpha spectra. Pu/Np and Am/Cm are

prepared for alpha spectrometry measurement via CeF<sub>3</sub> microprecipitation onto Resolve<sup>®</sup> Filters. To further lower detection limits, two 10g soil aliquots can be fused separately, combined following the Fe/Ti hydroxide Aliq precipitate, and then processed through the remaining steps of the method.

### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) DGA Resin, Normal, 2mL Cartridges (Eichrom DN-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu (or <sup>236</sup>Pu if Np Measured) tracer <sup>243</sup>Am tracer La carrier (10mg/mL) Ce Carrier (10mg/mL) 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> **Deionized Water**  $2M AI(NO_3)_3$ 10% (w:w) TiCl<sub>3</sub> HNO<sub>3</sub> (70%) HCI (37%) NaOH HF (49%) or NaF Boric acid NaNO<sub>2</sub> Sulfamic Acid Ascorbic Acid 30% H<sub>2</sub>O<sub>2</sub> NH<sub>4</sub>SCN (rare earth separation)

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters with funnel (Eichrom RF-DF25-25PP01) 250mL Zirconium Crucibles with lids 50mL and 500mL Centrifuge Tubes Alpha Spectrometry System Stainless Steel Planchets with double sided tape Centrifuge Hot Plate Analytical Balance Vacuum Pump Muffle Furnace

### Figure 1. Sample Preparation

Dry soil at 110°C. Blend and Crush to fine powder.

Aliquot 10g sample to Zr crucible. Add <sup>242</sup>Pu(<sup>236</sup>Pu)/<sup>243</sup>Am.

Dry on Hotplate. Place in Muffle furnace at 250°C. Ramp to 600°C. Heat 2 hrs to destroy organics.

Remove from furnace. Add 40g NaOH.

Fuse at 600°C for 30 minutes.

Transfer to 500mL centrifuge tube with water. Add 125mg Fe, 100mg Ca, 10mg La, 10mL 10% TiCl<sub>3</sub>. Dilute to 450 mL. Mix. Cool to room temp. in ice bath.

Centrifuge 10 min. Discard Supernate.

Rinse ppt with 150 mL pH ~9 NaOH. Centrifuge 10min. Decant Supernate.

Partially dissolve in 200mL 1.5M HCl. Solids will remain. Dilute to 250mL. Add 6mL 10% TiCl<sub>3</sub> and 40mL 49% HF. Mix. Place in ice bath for 10min.

Centrifuge 10min. Decant supernate.

Dissolve solids in 10mL 3M HNO<sub>3</sub>-0.25M H<sub>3</sub>BO<sub>3</sub>, 6mL 7M HNO<sub>3</sub>, 8mL 2M Al(NO<sub>3</sub>)<sub>3</sub> and 3mL 3M HNO<sub>3</sub>. Warming samples can improve dissolution.

Cool to room temperature. Adjust valence with 1mg Fe, 1.5mL 1M Ascorbic acid. Mix. Add 1mL 3.5M NaNO<sub>2</sub>.

### Figure 2. Actinide Separation on TEVA - DGA

| Figure 2. Actinide Separation on TEVA - DGA                                                                                                                                                                                                                                                                                                                                                                                           |                                                                                                                                                                                                                                                                                                                                                                  |                                                                                                                                                                                                                                                                                  |  |  |  |  |  |  |  |
|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--|--|--|--|--|--|--|
| <ul> <li>(1) Precondition 2mL TEVA + DGA<br/>cartridges with 5mL 8M HNO<sub>3</sub>.</li> <li>(2) Load Sample.</li> <li>(3) Rinse centrifuge tube<br/>with 5mL 6M HNO<sub>3</sub>. Add<br/>to TEVA + DGA.*</li> <li>(4) Rinse TEVA + DGA with<br/>10mL 3M HNO<sub>3</sub>.</li> <li>(5) Separate TEVA and DGA.</li> <li>(6) Rinse TEVA with<br/>-15mL 3M HNO<sub>3</sub><br/>-20mL 9M HCI (Th)<br/>-5mL 3M HNO<sub>3</sub></li> </ul> | Optional: Rare Earth Removal Steps(10) Add 3mL 3M HNO3 and 3mL 30%H2O2 to Am/Cm fraction.(11) Wet ash on hotplate to dryness.(12) Dissolve in 10mL 1.5M NH4SCN.(13) Precondition 2mL TEVA with5mL 1.5M NH4SCN.(14) Load Am/Cm fraction.(15) Rinse beaker with 5mL 1.5MNH4SCN. Add to TEVA.(16) Rinse TEVA with 5mL 1.5MNH4SCN.(17) Strip Am/Cm with 20mL 1M HCI. | <ul> <li>(23) Rinse filter<br/>funnel with 3mL<br/>DI water and 2mL<br/>100% ethanol.</li> <li>(24) Draw vacuum<br/>until filter is dry.</li> <li>(25) Remove filter from<br/>funnel assembly and<br/>mount filter on stainless<br/>steel planchet with 2-sided tape.</li> </ul> |  |  |  |  |  |  |  |
| (7) Strip Pu/Np from TEVA with 20mL<br>0.1M HCI-0.05M HF-0.01M TiCI <sub>3</sub> .                                                                                                                                                                                                                                                                                                                                                    | (18) Add 50ug Ce carrier and 0.5mL $30\%$ H <sub>2</sub> O <sub>2</sub> to all samples. Mix well. Add 1mL 49% HF. Mix well. Wait 15-20 minutes.                                                                                                                                                                                                                  |                                                                                                                                                                                                                                                                                  |  |  |  |  |  |  |  |
| <ul> <li>(8) Rinse DGA with:</li> <li>-10mL 3M HCI (Ca)</li> <li>-3mL 1M HNO<sub>3</sub></li> <li>-15mL 0.1M HNO<sub>3</sub> (La, Ca)</li> <li>-25mL 3M HNO<sub>3</sub>-0.25M HF (Th)</li> <li>-5mL 4M HCI</li> <li>(9) Strip Am/Cm with 12mL 0.25M HCI.</li> </ul>                                                                                                                                                                   | <ul> <li>(19) Set up Resolve<sup>®</sup> Filter Funnel on vacuum box.</li> <li>(20) Wet filter with 3mL 80% ethanol followed by 3mL DI water.</li> <li>(21) Filter sample.</li> <li>(22) Rinse sample tube with 5mL DI water and add to filter.</li> </ul>                                                                                                       | <ul> <li>(26) Dry filter under heat lamp for 3-5 minutes.</li> <li>(27) Measure actinides by alpha spectrometry.</li> </ul>                                                                                                                                                      |  |  |  |  |  |  |  |

\*Adding 50uL of 30%  $H_2O_2$  to the 6M HNO<sub>3</sub> tube rinse can further improve uranium decontamination factors.

### Method Performance

|          |            |                       |                   | % tracer      | mBq/g     | mBq/g              |            |
|----------|------------|-----------------------|-------------------|---------------|-----------|--------------------|------------|
| Sample   | replicates | analyte               | tracer            | recovery      | reference | measured           | % bias     |
| 10g Soil | 10         | <sup>239/240</sup> Pu | <sup>236</sup> Pu | 85 <u>+</u> 8 | 3.43      | 3.41 <u>+</u> 0.22 | <u>+</u> 5 |
| 10g Soil | 6          | <sup>237</sup> Np     | <sup>236</sup> Pu | 82 <u>+</u> 4 | 3.99      | 4.19 <u>+</u> 0.16 | <u>+</u> 6 |
| 10g Soil | 11         | <sup>241</sup> Am     | <sup>243</sup> Am | 89 <u>+</u> 4 | 2.14      | 2.07 <u>+</u> 0.16 | <u>+</u> 6 |

### References

1) Maxwell, Culligan, Hutchinson, McAlister, "Rapid Fusion Method for the Determination of Pu, Np, and Am in Large Soil Samples," *J. Radioanal. Nucl. Chem.* 305 : 599-608 (2015).

### **Rapid Fusion Method for** Refractory Th, U, and Pu in Soils

#### AN-1608-10

**Summary of Method** U, Th and Pu are separated and concentrated from 1-2 gram soil samples. Samples are fused with NaOH at 600°C in zirconium crucibles. The fusion cakes are dissolved in water, transferred to 250mL centrifuge tubes and precipitated sequentially with Fe/Ti-hydroxide and lanthanum fluoride to facilitate matrix removal. U, Th, and Pu are separated on stacked 2mL cartridges of Eichrom TEVA and TRU resins. U, Th, and Pu are measured by alpha spectrometry following CeF<sub>3</sub> microprecipitation onto Eichrom Resolve<sup>®</sup> Filters. Batches of 12-24 samples can be prepared for alpha spectrometry in less than 8 hours. Method ruggedness has been demonstrated with successful analysis of high fired refractory material from MAPEP 30 soil standards. For one gram soil samples and 16 hour count times, MDA for this method is ~500uBq/q.

### Reagents

TEVA Resin, 2mL Cartridges (Eichrom TE-R50-S) TRU Resin, 2mL Cartridges (Eichrom TR-R50-S) Iron carrier (50mg/mL Fe, as ferric iron nitrate) <sup>242</sup>Pu, <sup>232</sup>U and <sup>229</sup>Th tracers Oxalic acid/Ammonium oxalate La carrier (10mg/mL) Ce carrier (1mg/mL) 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> **Deionized Water** 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub> 2M AI(NO<sub>3</sub>)<sub>3</sub> 10% (w:w) TiCl<sub>3</sub> HNO<sub>3</sub> (70%) HCI (37%) NaOH HF (49%) or NaF Boric acid

 $H_2O_2$  (30%) Denatured ethanol NaNO<sub>2</sub> Ascorbic Acid

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50mL and 250mL Centrifuge Tubes Centrifuge Heat Lamp Muffle Furnace Hot Plate Analytical Balance 250mL Zirconium crucibles with zirconium lids Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Vacuum Pump

### **Figure 1. Sample Preparation**

1-2g finely ground soil + tracers in zirconium crucible.

Place in Furnace at 250°C. Ramp to 600°C. Heat 30 min.

Cool. Add 15g NaOH. Fuse at 600°C for 15-20 minutes.

Dissolve fusion cake with 2x 50mL H<sub>2</sub>O. Transfer to 250mL c-tube.

Add 10mL 3M HNO<sub>3</sub> to crucible. Heat to dissolve residue. Transfer to same 250mL c-tube.

Add 150mg Fe and 5mg La to c-tube. Dilute to 180mL.

Add 2mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub>, 3mL 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>, 10mL 10% TiCl<sub>3</sub>. Mix. Cool in ice bath for 10 min.

Centrifuge 10min. Decant Supernate.

Partially dissolve precipitate in 80mL 1.5M HCI. Some solids will remain. Dilute to 170mL. Add 1mg La, 0.5mL 1.25M Ca, and 6mL 10% TiCl<sub>3</sub>. Mix. Add 25mL 49% HF. Cool in ice bath for 10 min.

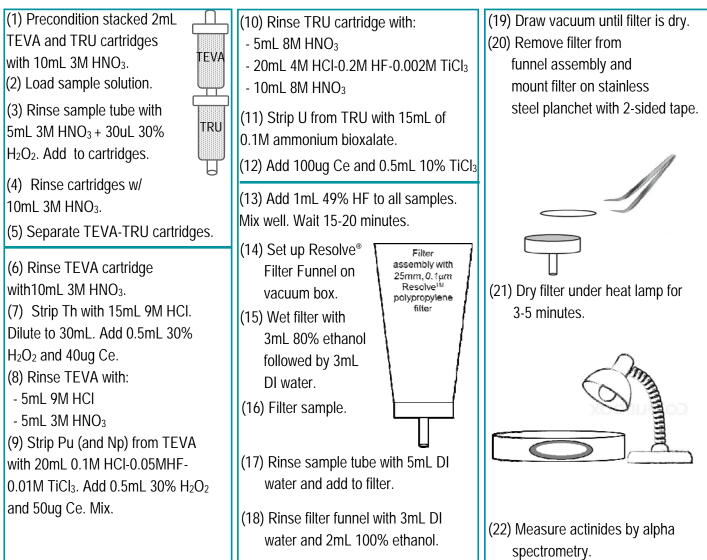
Centrifuge 10min. Decant Supernate.

Dissolve precipitate in 10mL 3M HNO<sub>3</sub>-0.25M Boric acid, 6mL 7M HNO<sub>3</sub>, and 8.5mL 2M AI(NO<sub>3</sub>)<sub>3</sub>. Warming samples can help complete dissolution.

Cool samples to room temperature.

Fix valence states. Mix between each addition of: 20uL 50mg/mL Fe 1.5mL 1M ascorbic acid 1mL 3.5M NaNO<sub>2</sub>.

### Figure 2. Actinide Separation on TEVA - TRU - DGA\* and Source Preparation



### Method Performance

|         |            |                   |                   | % tracer      | mBq/g     | mBq/g         |            |
|---------|------------|-------------------|-------------------|---------------|-----------|---------------|------------|
| Sample  | replicates | analyte           | tracer            | recovery      | reference | measured      | % bias     |
| 1g Soil | 12         | <sup>238</sup> U  | <sup>232</sup> U  | 86 <u>+</u> 8 | 83.0      | 85 <u>+</u> 3 | <u>+</u> 3 |
| 1g Soil | 12         | <sup>234</sup> U  | <sup>232</sup> U  | 86 <u>+</u> 8 | 81.0      | 80 <u>+</u> 2 | <u>+</u> 2 |
| 1g Soil | 12         | <sup>228</sup> Th | <sup>229</sup> Th | 91 <u>+</u> 6 | 51.1      | 50 <u>+</u> 2 | <u>+</u> 4 |
| 1g Soil | 12         | <sup>230</sup> Th | <sup>229</sup> Th | 91 <u>+</u> 6 | 96.2      | 98 <u>+</u> 6 | <u>+</u> 5 |
| 1g Soil | 12         | <sup>232</sup> Th | <sup>229</sup> Th | 91 <u>+</u> 6 | 48.8      | 50 <u>+</u> 3 | <u>+</u> 6 |
| 1g Soil | 3          | <sup>239</sup> Pu | <sup>242</sup> Pu | 91 <u>+</u> 6 | 76.8      | 79 <u>+</u> 3 | <u>+</u> 4 |
| 1g Soil | 3          | <sup>239</sup> Pu | <sup>242</sup> Pu | 91 <u>+</u> 6 | 96.0      | 98 <u>+</u> 5 | <u>+</u> 4 |

### References

1) Maxwell, Hutchinson, McAlister, "Rapid Fusion Method for the Determination of Refractory Thorium and Uranium Isotopes in Soil Samples" *Analytica Chimica Acta*, 701(1), 112-118 (2015).

### Measurement of Tritium in Water

#### AN-1609-10

**Summary of Method** Tritium is measured in 5-10mL aliquots of water using liquid scintillation counting. An Eichrom Tritium column is used to remove potentially interfering nuclides and matrix which can cause quench in the liquid scintillation cocktail. Sample size will be limited by the amount of sample that can be effectively mixed with the liquid scintillation cocktail (typically 5-10mL) and the salt content of the sample which can impact the separation of difficult to remove nuclides, such as isotopes of Cs. For samples which this method is not adequate, distillation methods, such as ASTM D4107 are recommended.

### Reagents

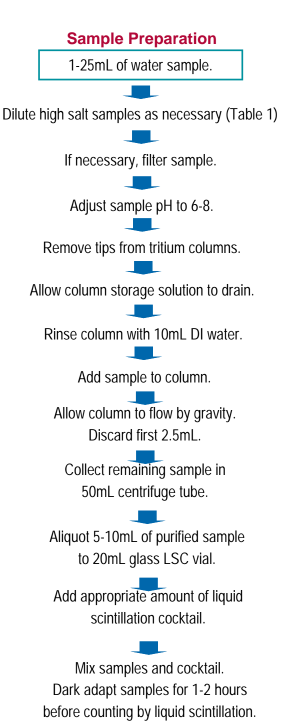
Tritium Column (Eichrom H3-C50-A) Deionized Water Liquid Scintillation Cocktail Nitromethane (Or other quench agent) <sup>3</sup>H Standard (To measure LSC quench curve) HCI (for sample pH adjustment) NaOH or NH₄OH (for sample pH adjustment)

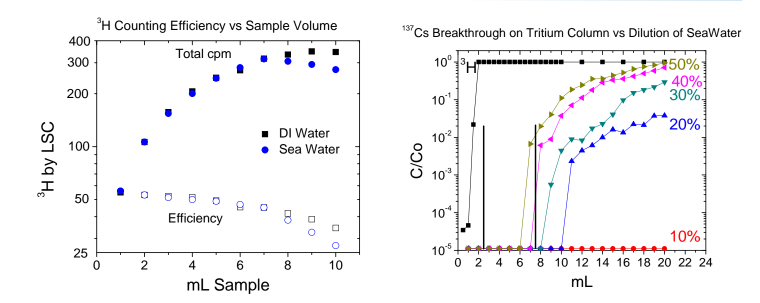
### Equipment

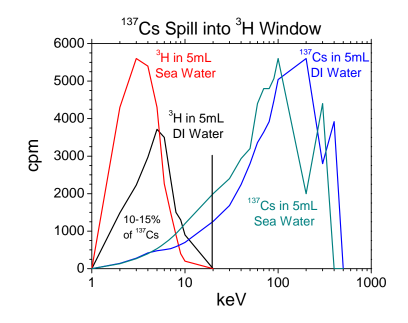
Column Rack (Eichrom AC-103) Extension Funnels (Eichrom AC-120) Centrifuge Tubes - 50mL 20mL glass liquid scintillation tubes Liquid scintillation counter Calibrated pipets and disposable tips pH meter or pH strips Analytical balance

| Table 1. Sam | ple Processing | ı on Tritium | Column    |
|--------------|----------------|--------------|-----------|
|              |                |              | oolallill |

|            | Тар   | Ground | Sea        |
|------------|-------|--------|------------|
|            | Water | Water  | Water      |
| Sample mL  | 15    | 15     | 3          |
| Dilution   | none  | none   | 3mL sample |
| Dilution   | none  | none   | to 10mL    |
| Discard mL | 2.5   | 2.5    | 2.5        |
| Collect mL | 12.5  | 12.5   | 5.0        |
| mL to LSC  | 10.0  | 10.0   | 5.0        |







### References

1) Eichrom Method H3W02. "Tritium in water," http://www.eichrom.com/eichrom/radiochem/methods/eichrom/

### Measurement of 59/63Ni in Water

### AN-1610-20

**Summary of Method** Nickel-59/63 is separated and measured from up to 500mL aliquots of water. Samples are preconcentrated by evaporation or ferric hydroxide precipitation, dissolved in 1M HCl, buffered with ammonium citrate and adjusted to pH 8-9 with ammonium hydroxide. Citrate complexes Fe(III), preventing precipitation at pH 8-9. Nickel is loaded onto 2mL cartridges of Nickel Resin. Yields can be improved by adding a 2mL cartridge of prefilter resin below the Nickel cartridge to minimize losses of the Ni-DMG complex. Nickel is recovered in 3M HNO<sub>3</sub> and measured by liquid scintillation counting. Chemical recovery of nickel is determined by ICP-AES measurement of 1-2mg of stable nickel cartrier.

### Reagents

Nickel Resin Cartridges (Eichrom NI-R50-S) Prefilter Resin Cartridges (Eichrom PF-R50-L) Anion Exchange Resin Cartridges (Eichrom A8-R50-M-CI)\* Deionized Water Ammonium Citrate Ammonium Hydroxide Sodium Hydroxide HCI HNO<sub>3</sub> Iron(III) Carrier (10mg/mL) Nickel Carrier (10mg/mL)\* Phenolphthalein pH indicator Liquid Scintillation Cocktail

### Equipment

Vacuum Box (Eichrom AR-12-BOX or AR-24-BOX) Vacuum Box Inner Liner (Eichrom AR-12-LINER or AR-24-LINER) Yellow Outer Tips (Eichrom AR-1000-OT) Inner Support Tube (Eichrom AR-1000-TUBE-PE) Cartridge Reservoirs (Eichrom AR-200-RV20) Centrifuge Tubes - 50mL and 250mL 20mL glass liquid scintillation tubes Liquid scintillation counter Calibrated pipets and disposable tips Appropriately Sized Glass Beakers ICP-AES system for Ni chemical yield measurement Analytical balance Vacuum Pump Centrifuge Hotplate

### **Sample Preparation**

Up to 500mL of water sample in glass beaker.

Add 1-2 mg of Ni carrier\*

Evaporate to dryness or proceed to ferric hydroxide precipitation steps below.

### **Ferric Hydroxide Precipitation**

Add 2mg of Fe(III) carrier and pH indicator

Heat sample to 80°C

Adjust to pH 8-9 with NaOH.

Mix sample and allow to cool to room temperature.

Allow ppt to settle and decant supernate to <200mL.

Transfer to 250mL centrifuge tube. Rinse beaker with water to ensure complete transfer of ppt.

Centrifuge 10min. Decant supernate to waste.

\*1mg of Co and carrier may also be added to improve decontamination from cobalt isotopes. For samples with very high <sup>58/60</sup>Co content, additional separation of cobalt on anion exchange resin may be required. (See Next Page)

### \*Optional Co/Fe Separation on Anion Exchange Resin

- 1) Following addition of Ni, Fe, Co carriers and sample evaporation or preconcentration be Fe(OH)<sub>3</sub> ppt.
- 2) Dissolve sample residue or Fe(OH)<sub>3</sub> ppt in 10mL 10M HCl.
- 3) Precondition 2mL anion exchange cartridge with 5mL 10M HCl.
- 4) Load sample in 10M HCl on anion exchange resin (1mL/min. Fe/Co retained). Collect Ni eluate in glass beaker.
- 5) Rinse cartridge with 10mL 10M HCI. Collect Ni eluate in glass beaker.
- 6) Carefully evaporate eluate from steps 4-5 to dryness.

### **Load Solution Preparation**

- 1) Dissolve ppt/residue in 5-10mL 1M HCI.
- 2) Add 1-2mL of 1M ammonium citrate.
- 3) Add pH indicator.
- 4) Adjust to pH 8-9 with ammonium hydroxide.
- 5) If ppt. forms, add additional ammonium citrate.



### **Nickel Separation**

- 1) Set up vacuum box with Nickel cartridges.\*\*\*
- 2) Precondition with 5mL 0.2M ammonium citrate.
- 3) Load samples on Nickel/Prefilter Resin.
- Rinse Nickel/Prefilter Resin with 20mL 0.2M ammonium citrate.
- 5) Strip Ni with 10-15mL of 3M HNO<sub>3</sub>.
- 6) Take aliquots for ICP-AES and Liquid Scintillation.

\*\*\* optional: prefilter cartridges below Ni to improve yield.

### References

1) Eichrom Method NIW01VBS. "Nickel-59/63 in water," http://www.eichrom.com/eichrom/radiochem/methods/eichrom/

AN-1611-10

### Measurement of <sup>55</sup>Fe in Water (TEVA Separation)

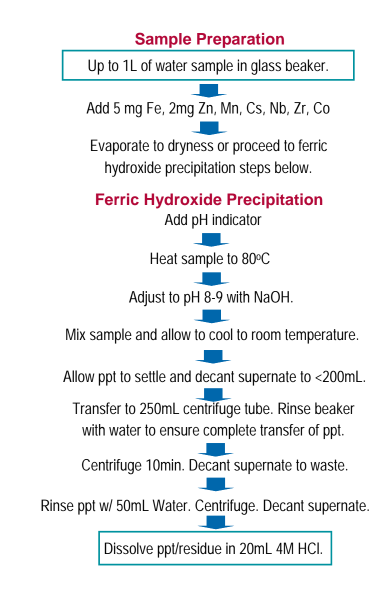
**Summary of Method** <sup>55</sup>Fe is separated and measured from up to 1L aliquots of water. Samples are preconcentrated by evaporation or ferric hydroxide precipitation, dissolved in 4M HCl and loaded onto 2mL cartridges of TEVA Resin. Hold back carriers, 2mg each of Zn, Mn, Cs, Nb, Zr, and Co are added to improve separation from radionuclides of these elements. An iron phosphate precipitate at pH 2.8-3.2 is used to prepare samples for liquid scintillation counting and remove remaining traces of Zn, which can co-elute with iron from TEVA resin. Chemical recovery of iron is determined by ICP-AES measurement of 5mg of stable iron carrier. <sup>55</sup>Fe may also be determined using TRU resin, AN-1612 from nitrate media. AN-1612 allows <sup>55</sup>Fe incorporation into standard TEVA-TRU actinide separations methods, but is limited to 2mg Fe per sample for a 2mL TRU resin cartridge.

### Reagents

TEVA Resin Cartridges (Eichrom TE-R50-S) Deionized Water Sodium Hydroxide HCI HNO<sub>3</sub> H<sub>3</sub>PO<sub>4</sub> LSC Cocktail Fe, Zn, Mn, Cs, Nb, Zr, Co carriers (10mg/mL) Phenolphthalein pH indicator <sup>55</sup>Fe standard Nitromethane or other LSC quench agent

### Equipment

Vacuum Box (Eichrom AR-12-BOX or AR-24-BOX) Vacuum Box Inner Liner (Eichrom AR-12-LINER or AR-24-LINER) Yellow Outer Tips (Eichrom AR-1000-OT) Inner Support Tube (Eichrom AR-1000-TUBE-PE) Cartridge Reservoirs (Eichrom AR-200-RV20) Centrifuge Tubes - 50mL and 250mL 20mL glass liquid scintillation tubes Liquid scintillation counter Calibrated pipets and disposable tips Appropriately Sized Glass Beakers ICP-AES system for Fe chemical yield measurement Analytical balance Vacuum Pump Centrifuge Hotplate

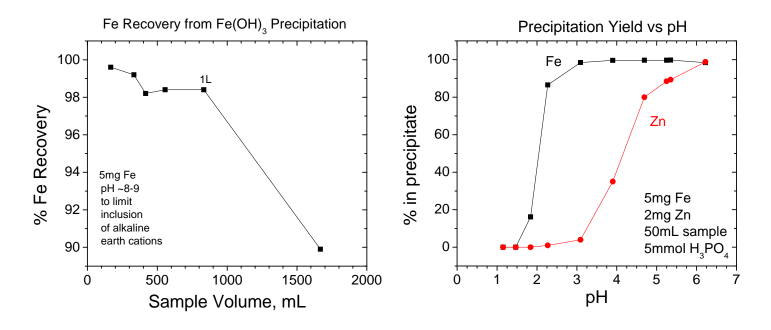


### **Iron Separation**

- 1) Set up vacuum box with TEVA cartridges.
- 2) Precondition with 5mL 4M HCl.
- 3) Load samples on TEVA Resin.
- 4) Rinse tube with 5mL 4M HCI. Add to TEVA.
- 5) Rinse TEVA with 10mL 4M HCl.
- 6) Strip Fe from TEVA with 20mL 0.1M HNO<sub>3</sub>.
- 7) Add 5mL 1M H<sub>3</sub>PO<sub>4</sub>. Mix.
- 8) Adjust to pH 2.8-3.2 with NaOH/H<sub>3</sub>PO<sub>4</sub>. Mix.

9) Centrifuge. Decant Supernate.

- 10) Wash ppt with 50mL H $_2$ O. Centrifuge. Decant Supernate.
- 11) Dissolve ppt with minimal 6M HCl.
- 12) Transfer to 10mL volumetric flask. Dilute to 10mL.
- 13) Take 0.1-0.2 mL, dilute to 10mL for ICP-AES Fe yield.
- 14) Transfer balance of sample to 20mL glass LSC vial.
- 15) Add 6 drops  $H_3PO4$ . Evap. on hotplate to ~0.5mL.
- 16) Add 1mL  $H_2O$ . Cool. Add 15mL LSC cocktail. Mix.



### Method Performance

|        |           | %Rec   |          |                  |      |           |
|--------|-----------|--------|----------|------------------|------|-----------|
|        |           | 2mg Fe | Fe-55    | Fe-55            |      |           |
| Method | Replicate | tracer | raw %rec | Tracer corrected | Bias | Impurity* |
| TEVA   | 1         | 95.8   | 89.2     | 93.1             | -6.9 | <0.5%     |
|        | 2         | 94.4   | 89.7     | 95.0             | 5.0  |           |
|        | 3         | 97.6   | 87.2     | 89.4             | 10.6 |           |
|        | 4         | 95.3   | 88.2     | 92.6             | 7.4  |           |
|        | 5         | 83.9   | 79.8     | 95.1             | 4.9  |           |
|        | 6         | 89.1   | 89.6     | 100.5            | -0.5 |           |
|        | 7         | 80.6   | 86.4     | 107.2            | -7.2 |           |
|        | AVG       | 91.0   | 87.2     | 96.1             |      |           |
|        | SD        | 6.6    | 3.5      | 5.9              |      |           |
|        |           |        |          |                  |      |           |

### References

1) ASTM Method D4922. "Standard Test Method for Determination of Radioactive Iron in Water."

0/ Do

AN-1612-10

### Measurement of <sup>55</sup>Fe in Water (TRU Separation)

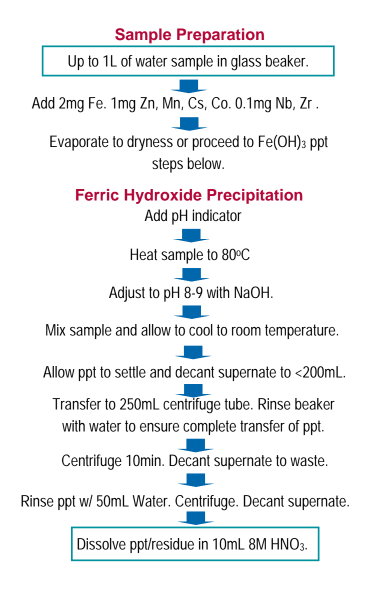
**Summary of Method** <sup>55</sup>Fe is separated and measured from up to 500mL aliquots of water. Samples are preconcentrated by evaporation or ferric hydroxide precipitation and purified on 2mL cartridges of TRU Resin. Holdback carriers, 0.1-1mg each of Zn, Mn, Cs, Nb, Zr, and Co are added to improve separation from these nuclides of these elements. An iron phosphate precipitate at is used to prepare samples for liquid scintillation counting. Chemical recovery of iron is determined by ICP-AES measurement of 2mg of stable iron carrier. <sup>55</sup>Fe may also be determined from chloride media using TEVA resin (Eichrom AN-1611). AN-1612 provides higher Zn decontamination and can be incorporated into TEVA-TRU actinide separations, but is limited to 2mg total Fe per 2mL cartridge. AN-1611 can process 5-6mg of Fe, but is less rugged for Zn decontamination.

### Reagents

TRU Resin Cartridges (Eichrom TE-R50-S) Deionized Water Sodium Hydroxide HCI HNO<sub>3</sub> H<sub>3</sub>PO<sub>4</sub> LSC Cocktail Fe, Zn, Mn, Cs, Nb, Zr, Co carriers (10mg/mL) Phenolphthalein pH indicator <sup>55</sup>Fe standard Nitromethane or other LSC quench agent

### Equipment

Vacuum Box (Eichrom AR-12-BOX or AR-24-BOX) Vacuum Box Inner Liner (Eichrom AR-12-LINER or AR-24-LINER) Yellow Outer Tips (Eichrom AR-1000-OT) Inner Support Tube (Eichrom AR-1000-TUBE-PE) Cartridge Reservoirs (Eichrom AR-200-RV20) Centrifuge Tubes - 50mL and 250mL 20mL glass liquid scintillation tubes Liquid scintillation counter Calibrated pipets and disposable tips Appropriately Sized Glass Beakers ICP-AES system for Fe chemical yield measurement Analytical balance Vacuum Pump Centrifuge Hotplate

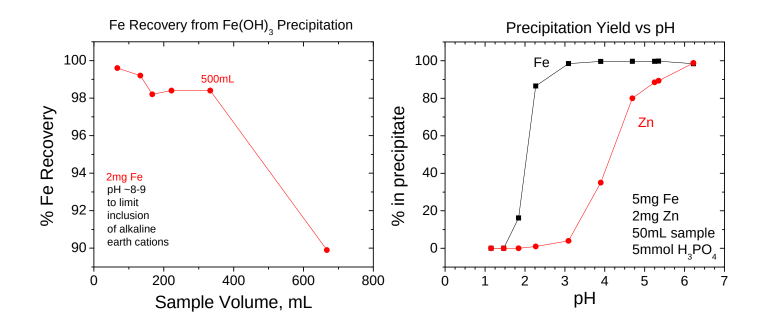


### **Iron Separation**

- 1) Set up vacuum box with TRU cartridges.
- 2) Precondition with 5mL 8M HNO<sub>3</sub>.
- 3) Load samples on TRU Resin.
- 4) Rinse tube with  $5mL 8M HNO_3$ . Add to TRU.
- 5) Rinse TRU with 10mL 8M HNO<sub>3</sub>.
- 6) Strip Fe from TRU with 15mL 2M HNO<sub>3.</sub>
- 7) Add 5mL 1M  $H_3PO_4$ . Mix.
- 8) Adjust to pH 2.8-3.2 with NaOH. Mix.

9) Centrifuge. Decant Supernate.

- 10) Wash ppt with 50mL  $H_2O$ . Centrifuge. Decant Supernate.
- 11) Dissolve ppt with minimal 6M HCl.
- 12) Transfer to 10mL volumetric flask. Dilute to 10mL.
- 13) Take 0.1-0.2 mL, dilute to 10mL for ICP-AES Fe yield.
- 14) Transfer balance of sample to 20mL glass LSC vial.
- 15) Add 6 drops  $H_3PO4$ . Evap. on hotplate to ~0.5mL.
- 16) Add 1mL H<sub>2</sub>O. Cool. Add 15mL LSC cocktail. Mix.



### Method Performance

|           | %Rec                                     |                                                                                                                                                                                                                            |                                                                                                                                                                                                                                                                                                                                                      |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
|-----------|------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
|           | 2mg Fe                                   | Fe-55                                                                                                                                                                                                                      | Fe-55                                                                                                                                                                                                                                                                                                                                                |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| Replicate | tracer                                   | raw %rec                                                                                                                                                                                                                   | Tracer corrected                                                                                                                                                                                                                                                                                                                                     | Bias                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       | Impurity*                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
| 1         | 90.6                                     | 93.1                                                                                                                                                                                                                       | 102.8                                                                                                                                                                                                                                                                                                                                                | 2.8                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        | <0.5%                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| 2         | 90.0                                     | 92.3                                                                                                                                                                                                                       | 102.5                                                                                                                                                                                                                                                                                                                                                | 2.5                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 3         | 94.8                                     | 92.4                                                                                                                                                                                                                       | 97.5                                                                                                                                                                                                                                                                                                                                                 | -2.5                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 4         | 89.5                                     | 94.0                                                                                                                                                                                                                       | 105.0                                                                                                                                                                                                                                                                                                                                                | 5.0                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 5         | 95.8                                     | 94.3                                                                                                                                                                                                                       | 98.5                                                                                                                                                                                                                                                                                                                                                 | -1.5                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 6         | 95.8                                     | 92.8                                                                                                                                                                                                                       | 96.9                                                                                                                                                                                                                                                                                                                                                 | -3.1                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| AVG       | 92.8                                     | 93.2                                                                                                                                                                                                                       | 100.5                                                                                                                                                                                                                                                                                                                                                |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| SD        | 3.0                                      | 0.8                                                                                                                                                                                                                        | 3.3                                                                                                                                                                                                                                                                                                                                                  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
|           | 1<br>2<br>3<br>4<br>5<br>6<br><b>AVG</b> | 2mg Fe           Replicate         tracer           1         90.6           2         90.0           3         94.8           4         89.5           5         95.8           6         95.8           AVG         92.8 | 2mg Fe         Fe-55           Replicate         tracer         raw %rec           1         90.6         93.1           2         90.0         92.3           3         94.8         92.4           4         89.5         94.0           5         95.8         94.3           6         95.8         92.8           AVG         92.8         93.2 | 2mg Fe         Fe-55         Fe-55           Replicate         tracer         raw %rec         Tracer corrected           1         90.6         93.1         102.8           2         90.0         92.3         102.5           3         94.8         92.4         97.5           4         89.5         94.0         105.0           5         95.8         94.3         98.5           6         95.8         92.8         96.9           AVG         92.8         93.2         100.5 | 2mg Fe         Fe-55         Fe-55           Replicate         tracer         raw %rec         Tracer corrected         Bias           1         90.6         93.1         102.8         2.8           2         90.0         92.3         102.5         2.5           3         94.8         92.4         97.5         -2.5           4         89.5         94.0         105.0         5.0           5         95.8         94.3         98.5         -1.5           6         95.8         92.8         96.9         -3.1           AVG         92.8         93.2         100.5         -3.1 |

### References

1) Eichrom Method FEW01VBS. "Iron-55 in water," http://www.eichrom.com/eichrom/radiochem/methods/eichrom/

### 68Ga Generator

#### AN-1613-10

**Summary of Method** <sup>68</sup>Ga is a positron emitting radionuclide which has garnered interest for use in positron emission tomography (PET).  $^{68}$ Ga ( $t_{1/2}$  = 68 min) can be readily isolated from its parent  $^{68}$ Ge ( $t_{1/2}$  = 271 days), which is produced by cyclotron irradiation of gallium or zinc target material. Classic <sup>68</sup>Ga generators consist of <sup>68</sup>Ge adsorbed onto an inorganic exchanger, such as Al<sub>2</sub>O<sub>3</sub>, SnO<sub>2</sub> or TiO<sub>2</sub>. The <sup>68</sup>Ga is then periodically eluted with 0.1-1.0M HCl or dilute EDTA. These generators are simple and robust, yielding 60-80% of 68Ga with minimal 68Ge breakthrough over many elutions. However, the classic generator can be limited by the relatively large volume of solution needed to elute the <sup>68</sup>Ga and by metal ion impurities arising from the inorganic substrate. An alternative generator system has been developed, in which the 68Ge source material is stored in dilute HCI. 68Ga is then selectively retained on cation exchange resin, while the <sup>68</sup>Ge is remains in solution for future use. A small amount of rinsing of the cation exchange column, completes the 68Ge source recovery. 68Ga is then stripped from the cation exchange resin using a small volume of 4M HCI and adsorbed on a second cartridge of UTEVA resin. A small volume of rinse with 4M HCI provides additional decontamination from 68Ge, and 68Ga is recovered in a small volume of dilute HCI (0.05-0.5M HCI). The chemistry is robust and scalable. The separation has been demonstrated using 0.5 - 2mL columns/cartridges. Typical decay corrected yields of 68Ga are 95 + 1% in 2-5mL of 0.1M HCl, with <10-7% 68Ge impurity. Stable metal ion impurities are typically in the low parts per billion range. Operation of the generator has also been demonstrated with the Northstar Medical Radioistope automated generator system.

#### Reagents

UTEVA Resin Cartridges (Eichrom UT-R50-S) Cation Exchange 2mL Cartridges (Eichrom C8-R50-H) <sup>68</sup>Ge Source\* Deionized Water HCI

<sup>\*</sup>Germanium chloride is relatively volatile and can be spread through the air. Care should be taken to minimize contamination of personnel and work spaces. Use of sealed systems for steps during separation is recommended.

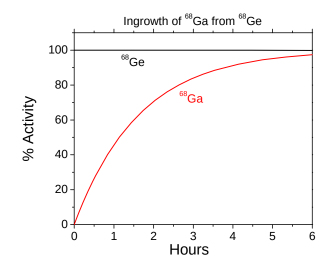
#### Equipment

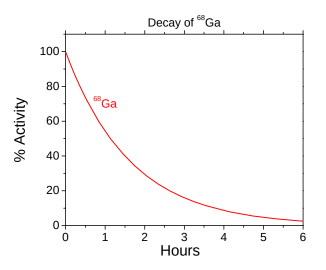
Glass vials for storage of <sup>68</sup>Ge source.

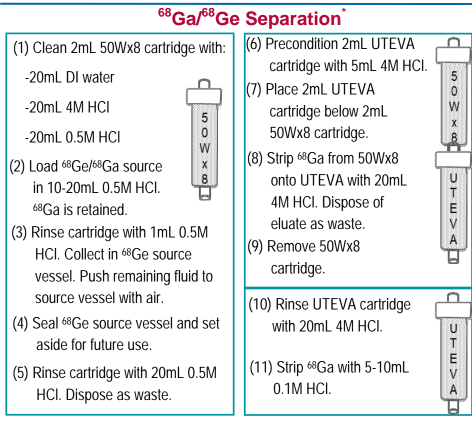
Glass or plastic vials/bottles for collection <sup>68</sup>Ga product and waste.

5, 10 or 20mL plastic luer lock syringes

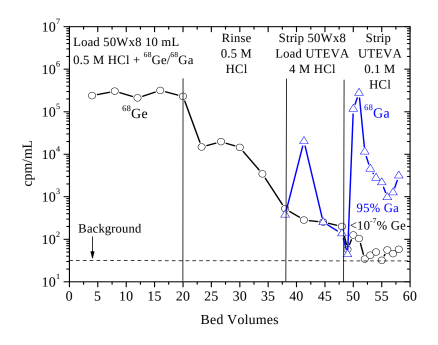
Gamma spectroscopy system for measurement of <sup>68</sup>Ga. (The electron capture of <sup>68</sup>Ge can be measured by liquid scintillation or <sup>68</sup>Ge can be determined after decay/ ingrowth of <sup>68</sup>Ga using the 511keV emission following positron annihilation.)

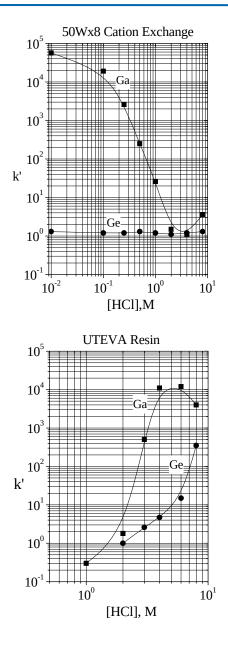






<sup>\*</sup>The separation may also be performed using 0.5mL or 1mL columns/ cartridges and proportionally scaled eluate volumes to improve method speed and reduce losses from <sup>68</sup>Ga decay during separation.





### References

1) McAlister and Horwitz, "Automated Two Column Generator Systems for Medical Radionuclides," *Applied Radiation and Isotopes*, 67:1985-1991 (2009).

### <sup>225</sup>Ac/<sup>225</sup>Ra Generator

#### AN-1614-11

**Summary of Method** A method for the preparation of <sup>225</sup>Ac (t<sub>1/2</sub> = 10 days) and <sup>225</sup>Ra (t<sub>1/2</sub> = 14.8 days) from <sup>229</sup>Th (t<sub>1/2</sub> = 7340 years) source material is presented. The method employs 2mL cartridges of UTEVA and DGA resins to obtain high purity <sup>225</sup>Ac in small volumes of eluate while preserving valuable <sup>229</sup>Th source material. The method is meant for <sup>225</sup>Ac tracer production from <sup>229</sup>Th containing 5-10mg or less of total Th. For separations from larger masses of Th see the Eichrom website bibliography for other options (Recent Advances in the Recovery and Purification of Actinium Isotopes, Horwitz and McAlister, National Meeting of the American Chemical Society, 2009). The source material, containing <sup>229</sup>Th, <sup>225</sup>Ac, <sup>225</sup>Ra and other daughter nuclides in 4M HNO<sub>3</sub>, is loaded onto stacked 2mL cartridges of UTEVA and DGA resins. <sup>229</sup>Th is retained on UTEVA, while <sup>225</sup>Ac is retained on DGA and <sup>225</sup>Ra passes through both cartridges. <sup>225</sup>Ac is recovered from DGA with a small volume of 2.0M HCl. The <sup>229</sup>Th source is recovered from UTEVA with a small volume of 0.5M HCl. Following a suitable ingrowth period, the <sup>229</sup>Th can be acidified to 4M HNO<sub>3</sub> and used to produce additional <sup>225</sup>Ac and <sup>225</sup>Ra. The <sup>229</sup>Th is preserved nearly indefinitely and continuously purified from chemical and radiologic impurities run to run.

#### Reagents

UTEVA Resin Cartridges (Eichrom UT-R50-S) DGA Resin Cartridges (Eichrom DN-R50-S) <sup>229</sup>Th Source Deionized Water HCI HNO<sub>3</sub>

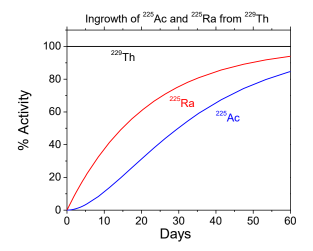
### Equipment

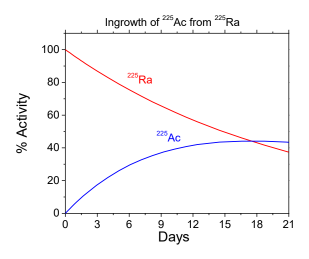
Glass vials for storage of <sup>229</sup>Th source.

Glass or plastic vials/bottles for collection of <sup>225</sup>Ac. <sup>225</sup>Ra and waste.

5, 10 or 20mL plastic luer lock syringes

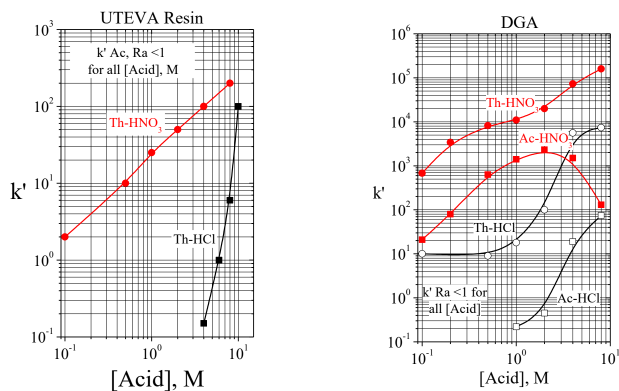
Gamma spectrometry system and/or alpha spectrometry for measurement of <sup>225</sup>Ac(<sup>221</sup>Fr), <sup>225</sup>Ra and <sup>229</sup>Th.





#### <sup>225</sup>Ac/<sup>225</sup>Ra/<sup>229</sup>Th Separation (6) Rinse DGA with 10mL 8M HCl. (1) Precondition stacked 2mL $\square$ cartridges of UTEVA and U T (7) Strip <sup>225</sup>Ac with 10mL 2M DGA with 10mL 4M HNO<sub>3</sub>. Ē V HCI. (Traces of <sup>229</sup>Th that D (2) Acidify <sup>229</sup>Th eluate from may have broken through G А previous separation with A UTEVA will be retained on 5mL HNO<sub>3</sub>. (If new <sup>229</sup>Th DGA.) source, dilute to 20mL with D G 4M HNO<sub>3</sub>.) (8) Place DGA (from which А <sup>225</sup>Ac has been stripped) D (3) Load <sup>229</sup>Th and daughters G above the UTEVA in 20mL 4M HNO<sub>3</sub>. Collect А cartridge. and save eluate containing <sup>225</sup>Ra.\* (9) Strip <sup>229</sup>Th from DGA-(4) Rinse UTEVA/DGA with 10mL UTEVA cartridges with U 4M HNO<sub>3</sub>. Collect <sup>225</sup>Ra.\* Т 15mL 0.5M HCI. Save E (5) Separate UTEVA and DGA <sup>229</sup>Th for future use. V cartridges. A

\*225Ra can used directly as a tracer or as a source of additional <sup>225</sup>Ra.



### References

1) McAlister and Horwitz, "Chromatographic Generator Systems for the actinides and natural decay series elements," *Radiochimica Acta*, 99:1-9 (2011).

### 90Y Generator

#### AN-1615-11

**Summary of Method** A method for the preparation of <sup>90</sup>Y ( $t_{1/2} = 64.1$  hours) from <sup>90</sup>Sr ( $t_{1/2} = 28.6$  years) source material is presented. The method employs 2mL cartridges of Sr and DGA resins to obtain high purity <sup>90</sup>Y in small volumes of eluate while preserving valuable <sup>90</sup>Sr source material. The source material, containing <sup>90</sup>Sr/<sup>90</sup>Y, in 4M HNO<sub>3</sub>, is loaded onto stacked 2mL cartridges of Sr and DGA resins. <sup>90</sup>Sr is retained on Sr Resin, while <sup>90</sup>Y is retained on DGA. The <sup>90</sup>Sr source is recovered from Sr Resin with a small volume of 0.1M HCI. Following a suitable ingrowth period, the <sup>90</sup>Sr can be acidified to 4M HNO<sub>3</sub> and used to produce additional <sup>90</sup>Y. The <sup>90</sup>Sr is preserved nearly indefinitely and continuously purified from chemical and radiologic impurities run to run. <sup>90</sup>Y is recovered from DGA resin with 0.1M HCI. For applications where <sup>90</sup>Y must be recovered in minimal volumes, DGA, Branched may be used in place of DGA, Normal.

### Reagents

Sr Resin Cartridges (Eichrom SR-R50-S) DGA, Normal Resin Cartridges (Eichrom DN-R50-S) or DGA, Branched Resin Cartridges (Eichrom DB-R50-S) Liquid Scintillation Cocktail <sup>90</sup>Sr Source Deionized Water HCI HNO<sub>3</sub>

### Equipment

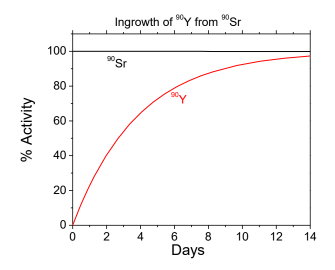
Glass vials for storage of <sup>90</sup>Sr source.

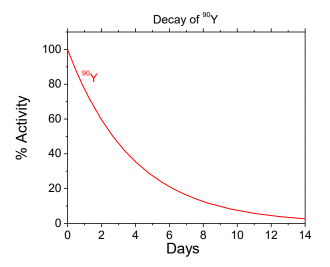
Glass or plastic vials/bottles for collection of <sup>90</sup>Y and waste.

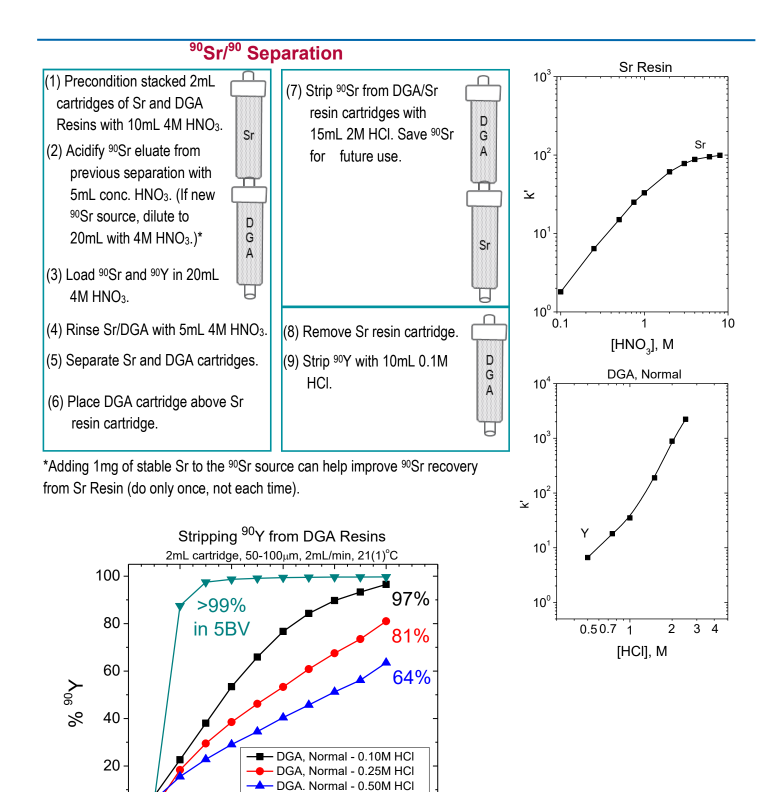
5, 10 or 20mL plastic luer lock syringes

Liquid Scintillation system for measurement of  ${}^{90}\text{Sr}\,\text{and}\,{}^{90}\text{Y}.^{*}$ 

<sup>\*90</sup>Y may also be measured by Cerenkov counting without the addition of scintillation cocktail.







### References

0

0

1

2

3

**Bed Volumes** 

1) McAlister and Horwitz, "Chromatographic Generator Systems for the actinides and natural decay series elements," *Radiochimica Acta*, 99:1-9 (2011).

5

6

DGA, Branched - 0.10M HCI

## <sup>210</sup>Po/<sup>210</sup>Bi Generator

#### AN-1616a-10

**Summary of Method** A method for the preparation of <sup>210</sup>Po ( $t_{1/2} = 138.4$  days) and <sup>210</sup>Bi ( $t_{1/2} = 5.013$  days) from <sup>210</sup>Pb ( $t_{1/2} = 22.26$  years) source material is presented. The method employs 2mL cartridges of UTEVA and Sr resins to obtain high purity <sup>210</sup>Po and <sup>210</sup>Bi in small volumes of eluate while preserving valuable <sup>210</sup>Pb source material. The source material, containing <sup>210</sup>Pb/<sup>210</sup>Bi/<sup>210</sup>Bi in 2.67M HCl, is loaded onto stacked 2mL cartridges of UTEVA and Sr resins. <sup>210</sup>Po is retained on UTEVA Resin, while <sup>210</sup>Pb is retained on Sr Resin and <sup>210</sup>Bi is not retained. The <sup>210</sup>Pb source is recovered from Sr Resin with a small volume of 8M HCl. Following a suitable ingrowth period, the <sup>210</sup>Pb can be diluted to 2.67M HCl and used to produce additional <sup>210</sup>Po and <sup>210</sup>Bi. The <sup>210</sup>Pb is preserved nearly indefinitely and continuously purified from chemical and radiologic impurities run to run. <sup>210</sup>Po is recovered from UTEVA resin with 6M HNO<sub>3</sub>.

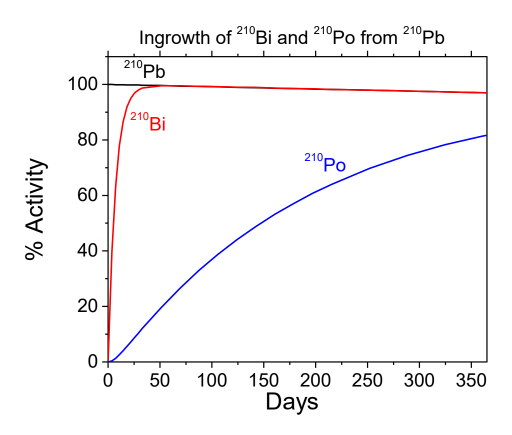
#### Reagents

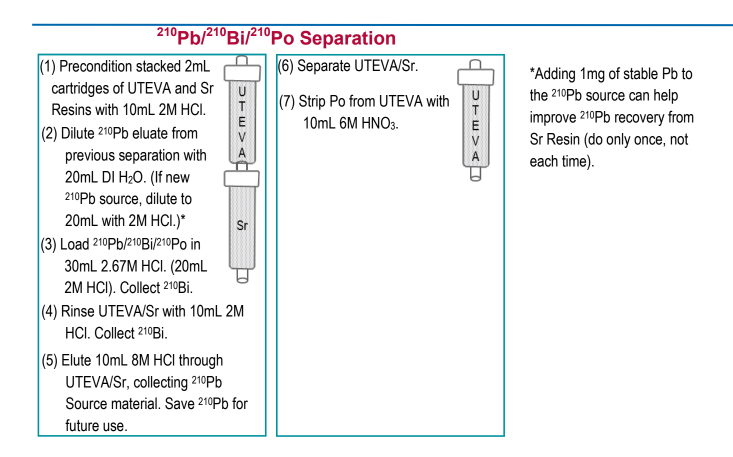
Sr Resin Cartridges (Eichrom SR-R50-S) UTEVA Cartridges (Eichrom UT-R50-S) Liquid Scintillation Cocktail <sup>210</sup>Pb Source Deionized Water HCI HNO<sub>3</sub>

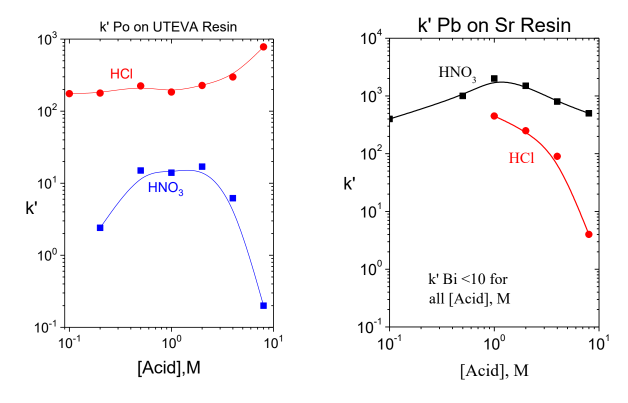
#### Equipment

Glass vials for storage of <sup>210</sup>Pb source. Glass or plastic vials/bottles for collection of <sup>210</sup>Po, <sup>210</sup>Bi and waste. 10, 20 or 30mL plastic luer lock syringes Liquid Scintillation System for measurement of <sup>210</sup>Bi and <sup>210</sup>Po.

Gamma Spectrometry System for measurement of <sup>210</sup>Pb.







### References

1) McAlister and Horwitz, "Chromatographic Generator Systems for the actinides and natural decay series elements," *Radiochimica Acta*, 99:1-9 (2011).

## <sup>210</sup>Po Generator

#### AN-1616b-11

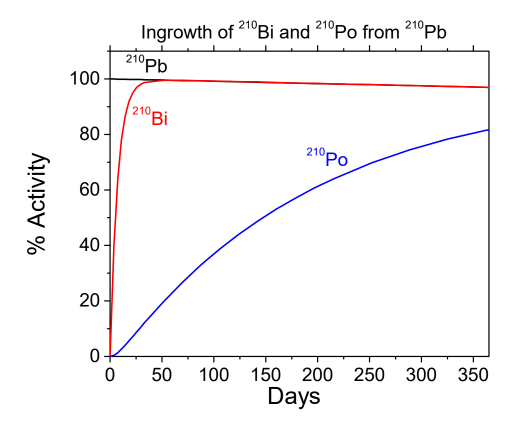
**Summary of Method** A method for the preparation of <sup>210</sup>Po (t<sub>1/2</sub> = 138.4 days) from <sup>210</sup>Pb (t<sub>1/2</sub> = 22.26 years) source material is presented. The method employs 2mL cartridges of DGA and Sr resins to obtain high purity <sup>210</sup>Po in small volumes of eluate while preserving valuable <sup>210</sup>Pb source material. The source material, containing <sup>210</sup>Pb/<sup>210</sup>Bi/<sup>210</sup>Bi in 2.67M HCl, is loaded onto stacked 2mL cartridges of DGA and Sr resins. <sup>210</sup>Po and <sup>210</sup>Bi are retained on DGA Resin, while <sup>210</sup>Pb is retained on Sr Resin. The <sup>210</sup>Pb source is recovered from Sr Resin with a small volume of 8M HCl. Following a suitable ingrowth period, the <sup>210</sup>Pb can be diluted to 2.67M HCl and used to produce additional <sup>210</sup>Po. The <sup>210</sup>Pb is preserved nearly indefinitely and continuously purified from chemical and radiologic impurities run to run. <sup>210</sup>Po is recovered from DGA resin with 0.05M HN0<sub>3</sub>, but should be acidified to 1M HNO<sub>3</sub> to prevent loss of Po to glass vials. The <sup>210</sup>Bi will remain on the DGA resin during the Po elution, and can be recovered with 10mL of 0.05M ammonium bioxalate. The DGA/Sr Resin chemistry is an improvement over the UTEVA/Sr Resin chemistry previously described (AN-1616a), which required 6M HNO<sub>3</sub> to recover the <sup>210</sup>Po.

#### Reagents

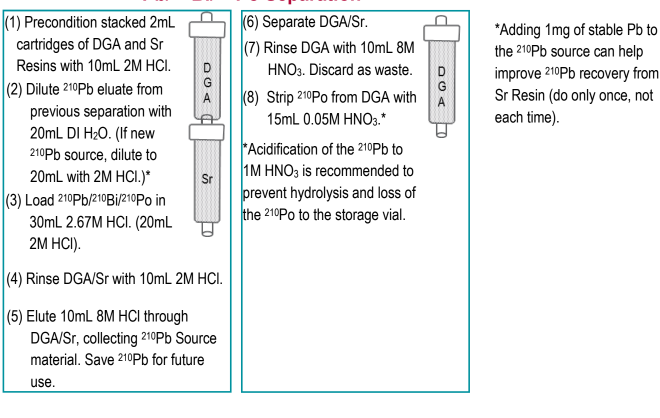
Sr Resin Cartridges (Eichrom SR-R50-S) DGA, Normal Cartridges (Eichrom DN-R50-S) Liquid Scintillation Cocktail <sup>210</sup>Pb Source Deionized Water HCI HNO<sub>3</sub>

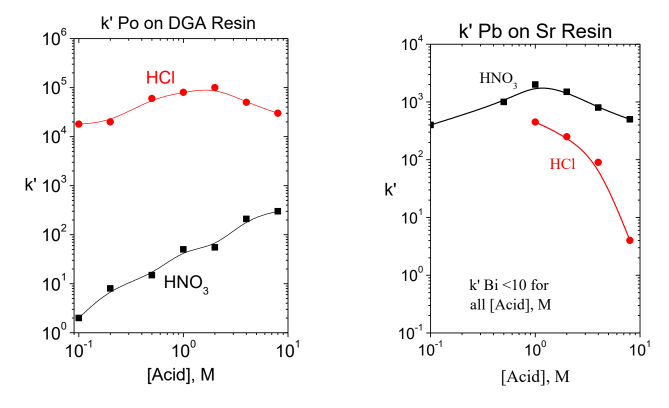
#### Equipment

Glass vials for storage of <sup>210</sup>Pb source. Glass or plastic vials/bottles for collection of <sup>210</sup>Po, <sup>210</sup>Bi and waste. 10, 20 or 30mL plastic luer lock syringes Liquid Scintillation System for measurement of <sup>210</sup>Bi and <sup>210</sup>Po. Gamma Spectrometry System for measurement of <sup>210</sup>Pb.



## <sup>210</sup>Pb/<sup>210</sup>Bi/<sup>210</sup>Po Separation





### References

1) McAlister and Horwitz, "Chromatographic Generator Systems for the actinides and natural decay series elements," *Radiochimica Acta*, 99:1-9 (2011).

## <sup>227</sup>Th/<sup>223</sup>Ra Generator

#### AN-1617-10

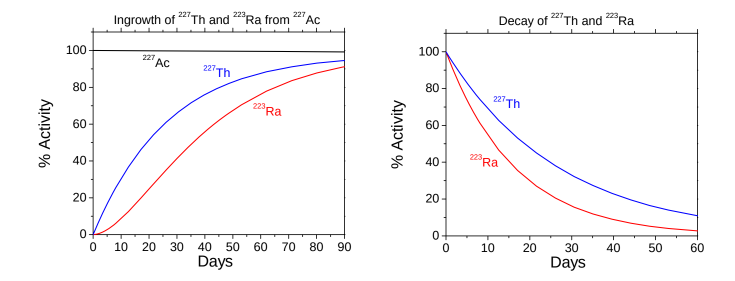
**Summary of Method** A method for the preparation of <sup>227</sup>Th ( $t_{1/2} = 18.72$  days) and <sup>223</sup>Ra ( $t_{1/2} = 11.43$  days) from <sup>227</sup>Ac ( $t_{1/2} = 21.77$  years) source material is presented. The method employs 2mL cartridges of UTEVA and DGA resins to obtain high purity <sup>227</sup>Th and <sup>223</sup>Ra in small volumes of eluate while preserving valuable <sup>227</sup>Ac source material. The source material, containing <sup>227</sup>Ac/<sup>227</sup>Th/<sup>223</sup>Ra in 4M HNO<sub>3</sub>, is loaded onto stacked 2mL cartridges of UTEVA and DGA resins. <sup>227</sup>Th is retained on UTEVA Resin, while <sup>227</sup>Ac is retained on DGA Resin and <sup>223</sup>Ra is not retained. The <sup>227</sup>Ac source is recovered from DGA Resin with a small volume of 0.1M HCI. Following a suitable ingrowth period, the <sup>227</sup>Ac can be acidified to 4M HNO<sub>3</sub> and used to produce additional <sup>227</sup>Th and <sup>223</sup>Ra. The <sup>227</sup>Ac is preserved nearly indefinitely and continuously purified from chemical and radiologic impurities run to run. <sup>227</sup>Th is recovered from UTEVA resin with 0.5M HCI.

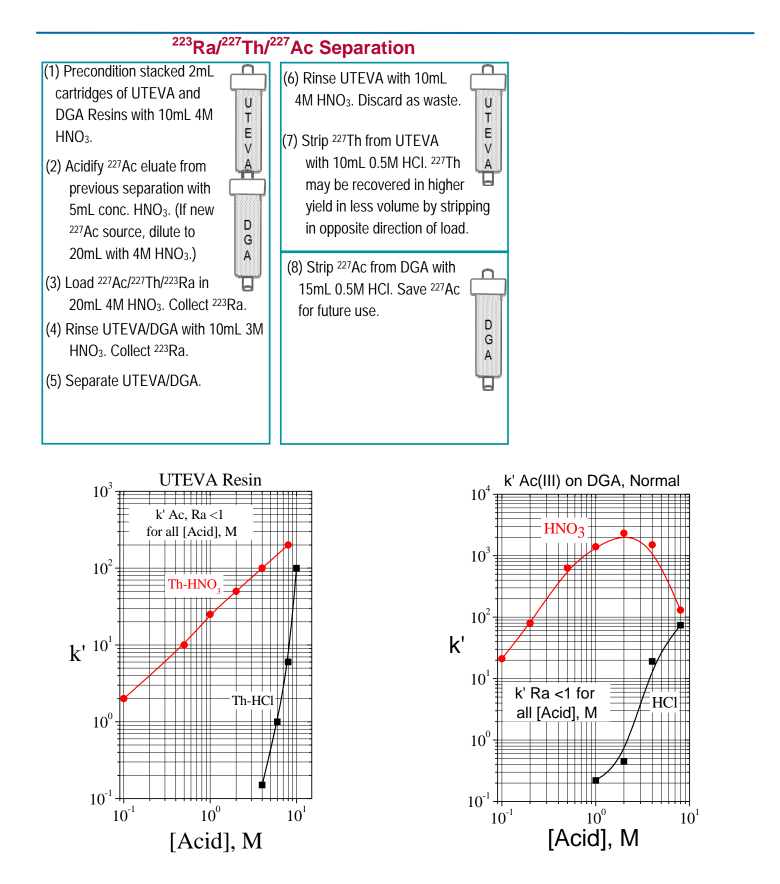
#### Reagents

UTEVA Cartridges (Eichrom UT-R50-S) DGA, Normal Cartridges (Eichrom DN-R50-S) <sup>227</sup>Ac Source Deionized Water HCI HNO<sub>3</sub>

#### Equipment

Glass vials for storage of <sup>227</sup>Ac source. Glass or plastic vials/bottles for collection of <sup>223</sup>Ra, <sup>227</sup>Th and waste. 10, 20 or 30mL plastic luer lock syringes Gamma Spectrometry System for measurement of <sup>227</sup>Th and <sup>223</sup>Ra.





### References

1) McAlister and Horwitz, "Chromatographic Generator Systems for the actinides and natural decay series elements," *Radiochimica Acta*, 99:1-9 (2011).

## <sup>228</sup>Th/<sup>231</sup>Th Generator

#### AN-1618-10

**Summary of Method** A method for the preparation of <sup>228</sup>Th ( $t_{1/2} = 1.913$  years) from <sup>232</sup>U ( $t_{1/2} = 72$  years) source material or <sup>231</sup>Th ( $t_{1/2} = 25.52$  hours) from <sup>235</sup>U ( $t_{1/2} = 7.04E8$  years) is presented. The method employs 2mL cartridges of TEVA and UTEVA resins to obtain high purity <sup>228</sup>Th or <sup>231</sup>Th in small volumes of eluate while preserving valuable <sup>232</sup>U or <sup>235</sup>U source material. The source material in 4M HNO<sub>3</sub>, is loaded onto stacked 2mL cartridges of TEVA and UTEVA resins. <sup>228</sup>Th or <sup>231</sup>Th is retained on TEVA Resin, while <sup>232</sup>U or <sup>235</sup>U is retained on UTEVA Resin. The <sup>232</sup>U or <sup>235</sup>U source is recovered from UTEVA Resin with a small volume of 1M HCI. Following a suitable ingrowth period, the <sup>232</sup>U or <sup>235</sup>U can be acidified to 4M HNO<sub>3</sub> and used to produce additional <sup>228</sup>Th or <sup>231</sup>Th. The <sup>232</sup>U or <sup>235</sup>U is preserved nearly indefinitely and continuously purified from chemical and radiologic impurities run to run. <sup>228</sup>Th or <sup>231</sup>Th is recovered from TEVA

#### Reagents

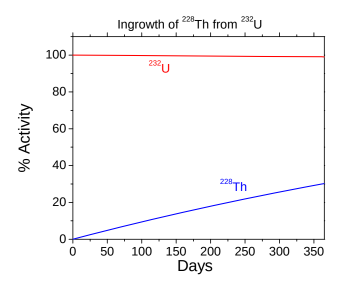
UTEVA Cartridges (Eichrom UT-R50-S) TEVA Cartridges (Eichrom TE-R50-S) <sup>232</sup>U or <sup>235</sup>U Source Deionized Water HCI HNO<sub>3</sub>

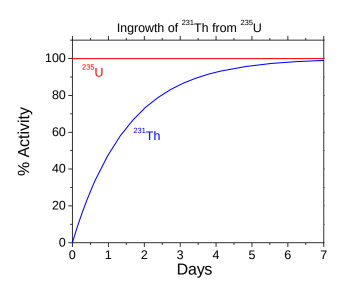
#### Equipment

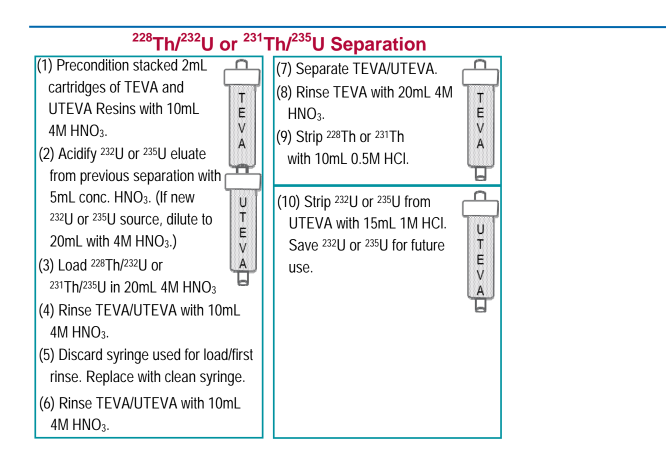
Glass vials for storage of <sup>232</sup>U or <sup>235</sup>U source. Glass or plastic vials/bottles for collection of <sup>228</sup>Th or <sup>231</sup>Th and waste.

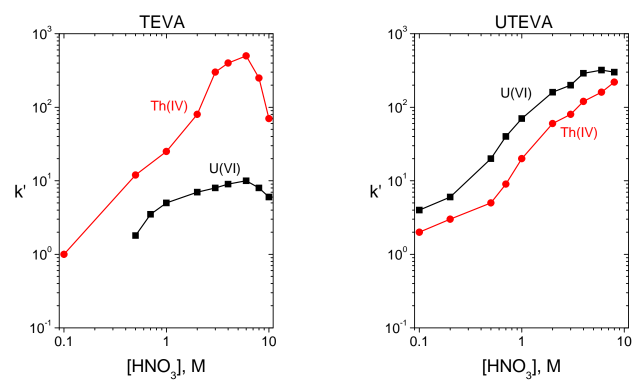
10, 20 or 30mL plastic luer lock syringes

Gamma Spectrometry System and/or Alpha Spectrometry System for measurement of <sup>228</sup>Th and <sup>232</sup>U or <sup>231</sup>Th and <sup>235</sup>U.









### References

1) McAlister and Horwitz, "Chromatographic Generator Systems for the actinides and natural decay series elements," *Radiochimica Acta*, 99:1-9 (2011).

## <sup>239</sup>Np Generator

#### AN-1619-10

**Summary of Method** A method for the preparation of <sup>239</sup>Np (t<sub>1/2</sub> = 2.355 days) from <sup>243</sup>Am (t<sub>1/2</sub> = 7380 years) source material is presented. The method employs 2mL cartridges of UTEVA and DGA resins to obtain high purity <sup>239</sup>Np in small volumes of eluate, while preserving valuable <sup>243</sup>Am material. The source material is adjusted to 4M HNO<sub>3</sub>, treated with iron, sulfamic acid and ascorbic acid to fix the Np(IV) oxidation state, and loaded onto stacked 2mL cartridges of UTEVA and DGA resins. <sup>239</sup>Np is retained on UTEVA Resin, while <sup>243</sup>Am is retained on DGA Resin. The <sup>243</sup>Am source is recovered from DGA Resin with a small volume of 0.5M HCI. Following a suitable ingrowth period, the <sup>243</sup>Am can be acidified to 4M HNO<sub>3</sub> and used to produce additional <sup>239</sup>Np. The <sup>243</sup>Am is preserved nearly indefinitely and continuously purified from chemical and radiologic impurities run to run. <sup>239</sup>Np is recovered from UTEVA resin with 0.5M HCI.

#### Reagents

UTEVA Cartridges (Eichrom UT-R50-S) DGA Cartridges (Eichrom DN-R50-S) <sup>243</sup>Am Source Deionized Water HCI HNO<sub>3</sub> Sulfamic Acid Fe carrier (10mg/mL) Ascorbic Acid

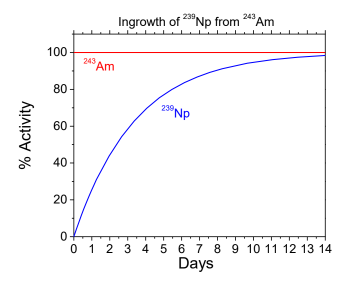
#### Equipment

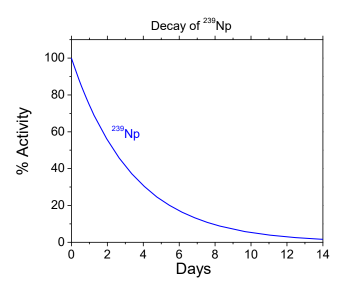
Glass vials for storage of <sup>243</sup>Am.

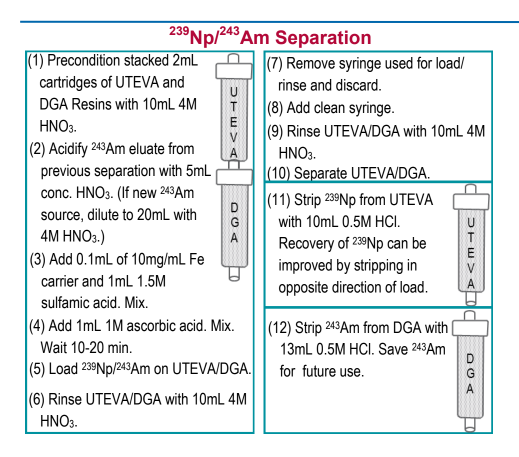
Glass or plastic vials/bottles for collection of <sup>239</sup>Np and waste.

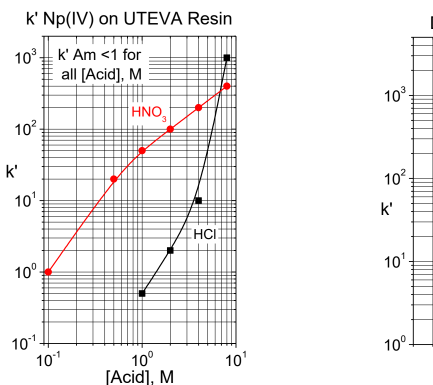
10, 20 or 30mL plastic luer lock syringes

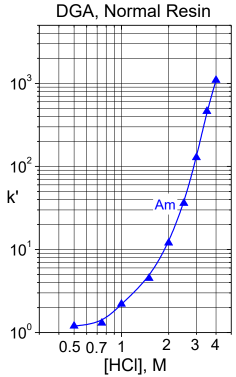
Gamma Spectrometry System for measurement of <sup>239</sup>Np and <sup>243</sup>Am.











### References

1) McAlister and Horwitz, "Chromatographic Generator Systems for the actinides and natural decay series elements," *Radiochimica Acta*, 99:1-9 (2011).

eichrom<sup>\*</sup> 1955 University Lane Lisle, IL 60532, USA +1 (630)963-0320 www.eichrom.com

## <sup>224</sup>Ra/<sup>212</sup>Pb Generator

#### AN-1620-10

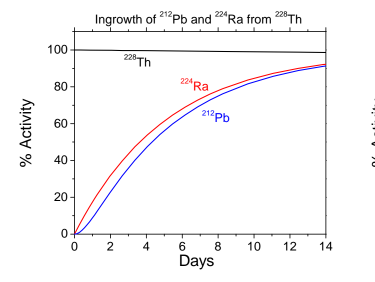
**Summary of Method** A method for the preparation of <sup>224</sup>Ra (t<sub>1/2</sub> = 3.62 days) and <sup>212</sup>Pb (t<sub>1/2</sub> = 10.64 hours) from <sup>228</sup>Th (t<sub>1/2</sub> = 1.913 years) source material is presented. The method employs 2mL cartridges of UTEVA and Sr resins to obtain high purity <sup>224</sup>Ra and <sup>212</sup>Pb in small volumes of eluate, while preserving valuable <sup>228</sup>Th material. The source material is adjusted to 4M HNO<sub>3</sub> and loaded onto stacked 2mL cartridges of UTEVA and Sr resins. <sup>228</sup>Th is retained on UTEVA Resin, while <sup>212</sup>Pb is retained on Sr Resin and <sup>224</sup>Ra is unretained. The <sup>228</sup>Th source is recovered from UTEVA Resin with a small volume of 0.5M HCl. Following a suitable ingrowth period, the <sup>228</sup>Th can be acidified to 4M HNO<sub>3</sub> and used to produce additional <sup>224</sup>Ra and <sup>212</sup>Pb. The <sup>228</sup>Th is preserved nearly completely and continuously purified from chemical and radiologic impurities run to run, allowing repeated use until radioactive decay depletes the <sup>228</sup>Th activity. <sup>212</sup>Pb may recovered from Sr resin with a variety of reagents, including 6-8M HCl, citrate, tartrate, acetate and bioxalate.

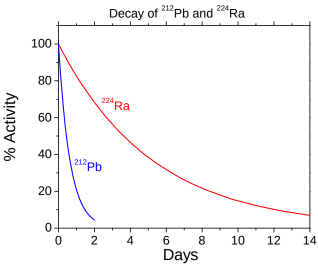
#### Reagents

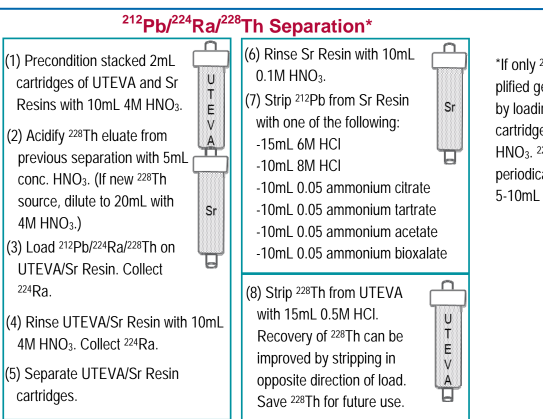
UTEVA Cartridges (Eichrom UT-R50-S) Sr Resin Cartridges (Eichrom SR-R50-S) <sup>228</sup>Th Source Deionized Water HCI HNO<sub>3</sub> <u>Option for <sup>224</sup>Ra only:</u> LN Resin cartridges (Eichrom LN-R50-S)

#### Equipment

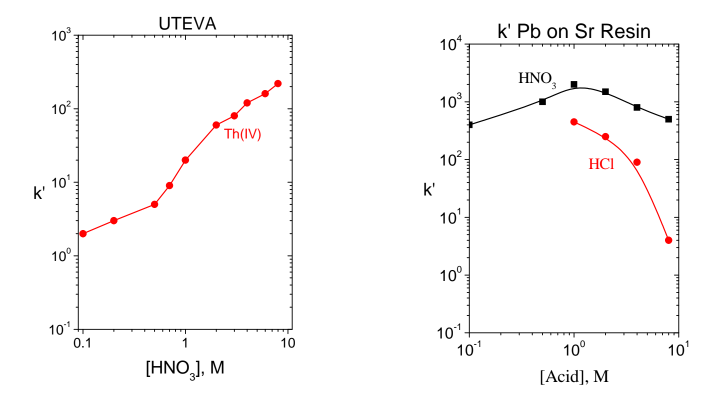
Glass vials for storage of <sup>228</sup>Th source.
Glass or plastic vials/bottles for collection of <sup>224</sup>Ra, <sup>212</sup>Pb and waste.
10, 20 or 30mL plastic luer lock syringes
Gamma Spectrometry System of alternative for measurement of <sup>228</sup>Th, <sup>224</sup>Ra, and <sup>212</sup>Pb.







\*If only <sup>224</sup>Ra is desired, a simplified generator can be made by loading <sup>228</sup>Th onto a 2mL cartridge of LN resin from 0.1M HNO<sub>3</sub>. <sup>224</sup>Ra can then be periodically milked using 5-10mL of 0.1M HNO<sub>3</sub> or HCI.



### References

1) McAlister and Horwitz, "Chromatographic Generator Systems for the actinides and natural decay series elements," *Radiochimica Acta*, 99:1-9 (2011).

## <sup>234</sup>Th Generator

#### AN-1621-10

**Summary of Method** A method for the preparation of  ${}^{234}$ Th ( $t_{1/2} = 24.1$  days) from natural or depleted Uranium ( $t_{1/2} = 4.47$ E9 years) source material is presented. The method utilizes extraction chromatography with a column of DGA, Normal resin and 2mL cartridges of DGA and UTEVA resins to obtain high purity  ${}^{234}$ Th in small volumes of eluate, while preserving  ${}^{238}$ U material. The source material is adjusted to 2M HNO<sub>3</sub> and loaded onto a column of DGA, Normal resin.  ${}^{234}$ Th is retained on DGA Resin from up to 0.2M uranium, while uranium is unretained. The uranium source is recovered and, following a suitable ingrowth period, can be used to produce additional  ${}^{234}$ Th.  ${}^{234}$ Th is stripped from the DGA resin column and further purified using 2mL cartridges of DGA and TEVA resins.

#### Reagents

TEVA Cartridges (Eichrom TE-R50-S) DGA Cartridges (Eichrom DN-R50-S) DGA, Normal Resin (Eichrom DN-B25-A) Natural or Depleted U Source Deionized Water Oxalic Acid Ammonium Oxalate HCI HNO<sub>3</sub>

#### Equipment

Glass/Plastic bottles for storage of Uranium source. Glass or plastic vials/bottles for collection of <sup>234</sup>Th and waste.

10, 20 or 30mL plastic luer lock syringes.

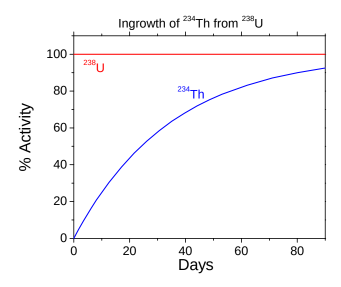
Gamma Spectrometry System or alternative for measurement of <sup>234</sup>Th.

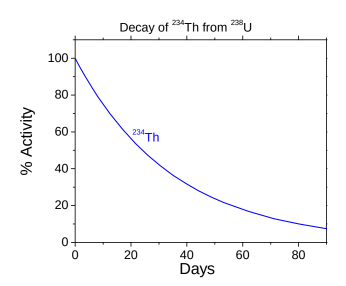
ICP-AES or alternative for measurement of U.

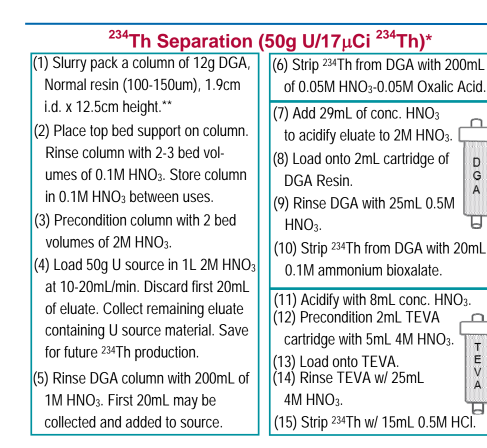
1.9cm i.d. glass or plastic column, minimum 15cm height, with 250mL-1L reservoir.

Glass wool or frit material for top bed support.

Peristaltic pump or alternative to increase flow rate.

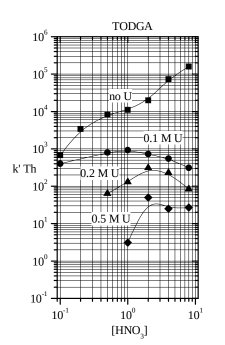


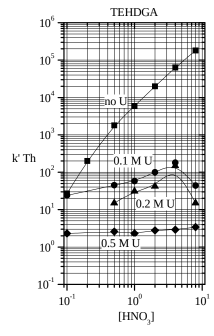


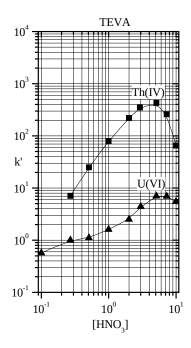


\*Separation is scalable. Simply adjust volumes of the initial DGA column and load solution to accommodate other source sizes.

\*\*DGA resin can be difficult to wet. Slurry the resin in 2x its volume of 1.0 -1.5M HNO<sub>3</sub> by gently swirling for 2-3 minutes (avoid vigorous shaking as this can incorporate air bubbles and cause resin to float). Centrifuge resin slurry for 5-10 minutes. Repeat until most of the resin sinks to the bottom of tube. Repeat swirling/ centrifugation, if needed. Use only well wetted resin to pack the column (omit floating resin). The column may be reused many times if stored in dilute acid between uses.







### References

1) E. P. Horwitz and D. R. McAlister, "The recovery of trace thorium from large quantities of uranium," *Solv. Extr. Ion Exch.*, 27, 474-488, (2009).

## Separation of <sup>89</sup>Zr From Y Target

#### AN-1622-10

**Summary of Method** A method for the separation of <sup>89</sup>Zr (t<sub>1/2</sub> = 78.43 hours) from yttrium target material is presented. The method employs 2mL cartridges of LN3 and Anion Exchange resins to obtain high purity <sup>89</sup>Zr in small volumes of eluate, while providing high separation factors from chemical and radiologic impurities. The primary separation of <sup>89</sup>Zr from the dissolved yttrium target can be performed in 2-8M HNO<sub>3</sub> or HCl using LN3 resin. <sup>89</sup>Zr is retained while yttrium passes through LN3. <sup>89</sup>Zr is recovered from LN3 with a small volume of 0.05M HCl-oxalic acid and directly loaded onto a 2mL cartridge of Anion Exchange resin. <sup>89</sup>Zr is retained while additional decontamination from yttrium and niobium is achieved. <sup>89</sup>Zr is then recovered in a small volume of 2-4M HCl. Average yield of Zr, separated from 500mg Y, was >90%, with >10<sup>6</sup> separation factor from Y and Nb.

#### Reagents

LN3 Cartridges (Eichrom L3-R50-S) Anion Exchange Cartridges (Eichrom A8-R50-M-Cl) Deionized Water Oxalic Acid Ammonium Oxalate HCl HNO<sub>3</sub>

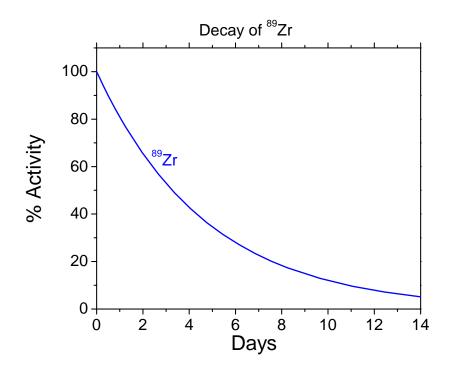
#### Equipment

Glass or plastic vials/bottles for collection of <sup>89</sup>Zr and waste.

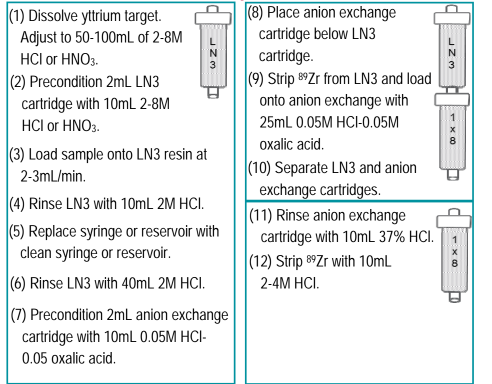
30mL and 60mL plastic luer lock syringes.

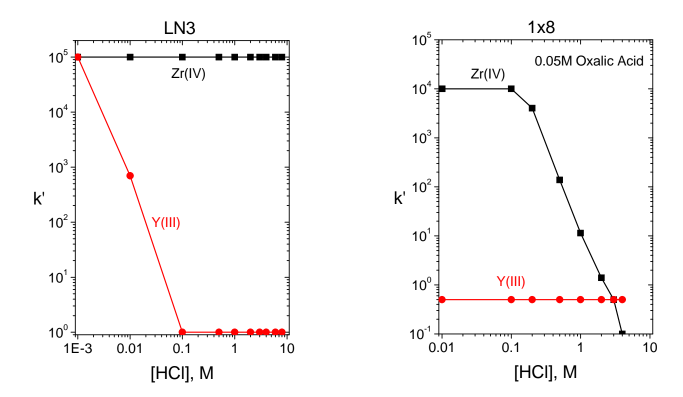
Gamma Spectrometry System or alternative for measurement of <sup>89</sup>Zr.

ICP-AES or alternative for measurement of Y.



## <sup>89</sup>Zr Separation





### References

1) E. P. Horwitz and D. R. McAlister, Unpublished data (2015 and 2016)

## Separation of <sup>86</sup>Y From Sr Target

#### AN-1623-10

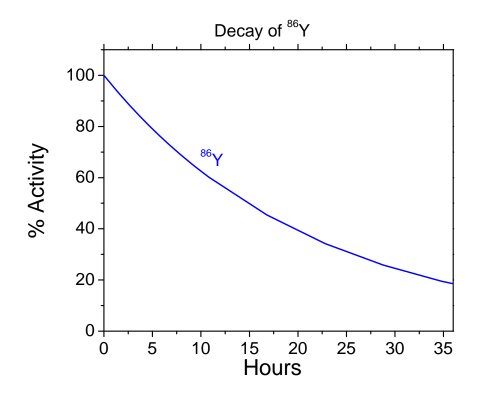
**Summary of Method** A method for the separation of <sup>86</sup>Y ( $t_{1/2} = 14.74$  hours) from strontium target material is presented. The method employs 2mL cartridges of DGA and LN resins to obtain high purity <sup>86</sup>Y in small volumes of eluate, while providing high separation factors from chemical and radiologic impurities. The primary separation of <sup>86</sup>Y from the dissolved yttrium target can be performed in 8M HNO<sub>3</sub> or HCl using DGA resin. <sup>86</sup>Y is retained while strontium passes through DGA. <sup>86</sup>Y is recovered from DGA with a small volume of 0.25M HCl and directly loaded onto a 2mL cartridge of LN resin. <sup>86</sup>Y is retained while additional decontamination from strontium is achieved. <sup>86</sup>Y is then stripped from LN resin onto a second 2mL cartridge of DGA resin using 8M HCl. <sup>86</sup>Y is then eluted from DGA using 10mL 0.1M HCl. DGA, Branched is used to allow stripping of <sup>86</sup>Y in a minimal volume of 0.1M HCl. Average yield of Y separation from 500mg of Sr was >95% with >10<sup>10</sup> separation factor from Sr.

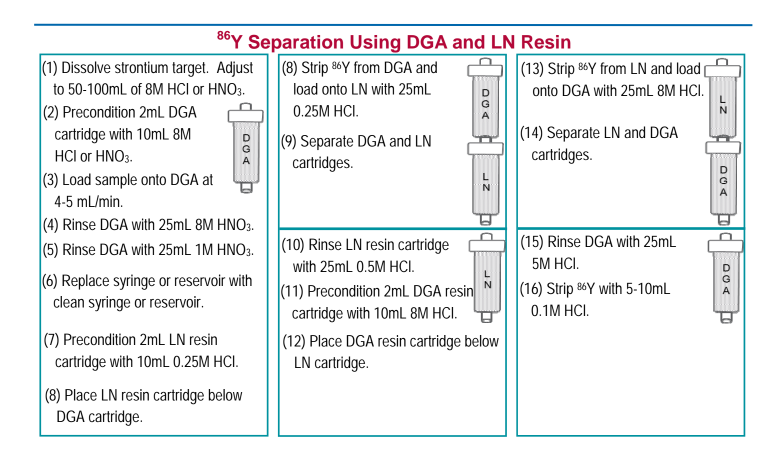
#### Reagents

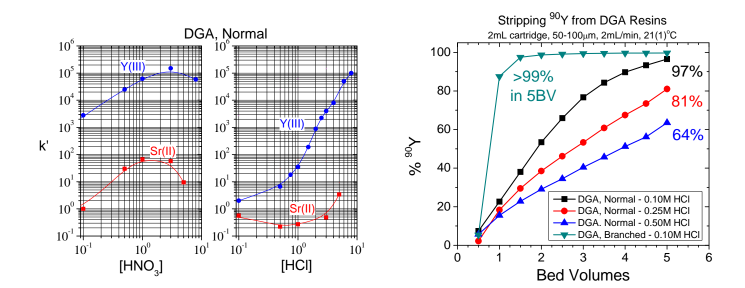
DGA, Branched Cartridges (Eichrom DB-R50-S) LN Resin Cartridges (Eichrom LN-R50-S) Deionized Water HCI HNO<sub>3</sub>

#### Equipment

Glass or plastic vials/bottles for collection of <sup>89</sup>Zr and waste. 30mL and 60mL plastic luer lock syringes. Gamma Spectrometry System or alternative for measurement of <sup>86</sup>Y. ICP-AES or alternative for measurement of Sr.







#### References

1) E. P. Horwitz and D. R. McAlister, Unpublished data (2015 and 2016).

## Options for <sup>89</sup>Sr and <sup>90</sup>Sr Determination

#### AN-1624-11

**Summary** There are many methods (Table 1) available for the measurement of radiostrontium (Table 2) from environmental, building materials, or biological samples. Typically, analysts are interested in the measurement of the fission products <sup>89</sup>Sr (decays to stable <sup>89</sup>Y) and <sup>90</sup>Sr (decays to  $\beta$ <sup>-</sup> emitting <sup>90</sup>Y). Stable strontium or <sup>85</sup>Sr may also be used as a chemical yield tracer. Methods typically begin with a concentration or matrix removal step, followed by the separation of strontium from interfering nuclides using Sr Resin (Figure 1A). Methods may also incorporate steps to discriminate between <sup>89</sup>Sr and <sup>90</sup>Sr, including multiple counts, nuclide selective counting techniques and ingrowth and secondary separations of <sup>90</sup>Y (daughter of <sup>90</sup>Sr). Measurement instrumentation includes low background gas flow proportional counting (GFPC), liquid scintillation (LSC), Cerenkov counting, and inductively coupled plasma-mass spectrometry (ICP-MS).

This application note will offer guidance in choosing an appropriate method for the determination of radiostrontium, taking into account process knowledge, available measurement equipment and data quality objectives (required detection limits, urgency of measurement, age of sample, and need for <sup>89/90</sup>Sr discrimination). The measurement methods are meant be to used in concert with the appropriate sample preparation method for the matrix being analyzed. For a more comprehensive treatment of sample preparation methods, see the application notes available at http://www.eichrom.com/eichrom/appnotes/applications/index.aspx. A detailed discussion of interferences for the various measurement techniques can also be found in references [2] and [5].

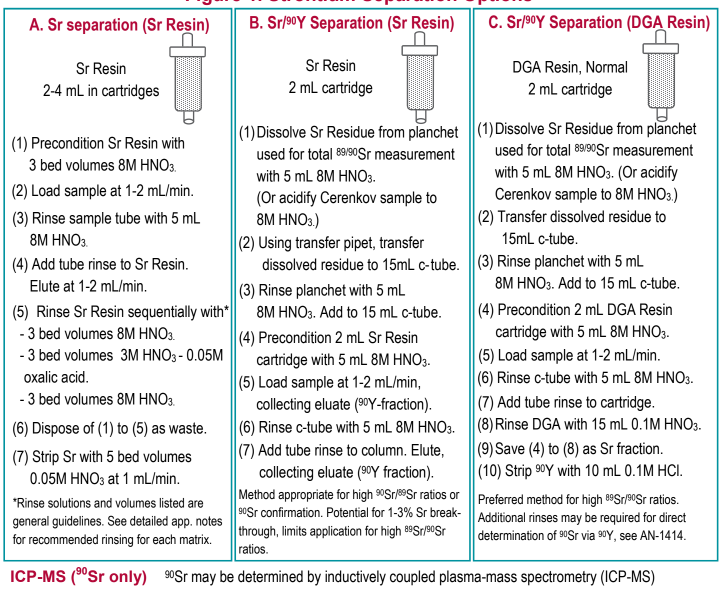
|                                                                 | Table 1. Summary of St Method Options |                                     |            |                                  |                             |      |
|-----------------------------------------------------------------|---------------------------------------|-------------------------------------|------------|----------------------------------|-----------------------------|------|
|                                                                 | Primary                               | Primary                             | Secondary  | Secondary                        | Sr Yield                    |      |
| Method                                                          | Separation                            | Measurement                         | Separation | Measurement                      | Monitor                     | Ref. |
| ICP-MS                                                          | 1A                                    | <sup>90</sup> Sr by ICP-MS          | None       | None                             | Stable Sr                   | 1    |
| Total <sup>89/90</sup> Sr                                       | 1A                                    | <sup>89/90</sup> Sr by GFPC/LSC     | None       | None                             | Stable Sr                   | 4,5  |
| <sup>90</sup> Y Direct                                          | 1C                                    | <sup>90</sup> Y by GFPC/LSC/Cerenko | v None     | None                             | Stable Sr                   | 6    |
| Two Count <sup>89</sup> Sr and <sup>90</sup> Sr                 | 1A                                    | <sup>89/90</sup> Sr by GFPC/LSC     | None       | <sup>89/90</sup> Sr by GFPC/LSC* | Stable Sr                   | 2    |
| Rapid <sup>89</sup> Sr and <sup>90</sup> Sr                     | 1A                                    | <sup>89</sup> Sr by Cerenkov        | None       | <sup>89/90</sup> Sr by LSC       | Stable Sr                   | 5    |
| Cerenkov <sup>89</sup> Sr and <sup>90</sup> Sr( <sup>90</sup> Y | ) 1A                                  | <sup>89</sup> Sr by Cerenkov        | 1B or 1C   | <sup>90</sup> Y by Cerenkov      | Stable Sr/ <sup>85</sup> Sr | 3    |
| Gas Flow Porportional                                           | 1A                                    | <sup>89/90</sup> Sr by GFPC         | 1B or 1C   | <sup>90</sup> Y by GFPC          | Stable Sr/ <sup>85</sup> Sr | 4    |
| 00 00                                                           |                                       | 00                                  | 00         |                                  |                             |      |

#### Table 1. Summary of <sup>89/90</sup>Sr Method Options

\*<sup>89</sup>Sr/<sup>90</sup>Sr descrimination by solving equations for <sup>89</sup>Sr decay and <sup>90</sup>Y ingrowth during time between 2 measurements

|                  | Table 2. Properties of Selected Nuclides |                       |                                                            |      |     |          |        |       |  |
|------------------|------------------------------------------|-----------------------|------------------------------------------------------------|------|-----|----------|--------|-------|--|
|                  | Decay Detector Suitable for Measuremen   |                       |                                                            |      |     | nt       |        |       |  |
| Nuclide          | Half-Life                                | Mode                  | Energy                                                     | GFPC | LSC | Cerenkov | MS/AES | Gamma |  |
| Stabel Sr        | <sup>84</sup> Sr (0.56%),                | <sup>86</sup> Sr(9.86 | %), <sup>87</sup> Sr (7.0%), <sup>88</sup> Sr (82.58%)     | No   | No  | No       | Yes    | No    |  |
| <sup>85</sup> Sr | 64.849 days                              | ε/γ                   | γ = 514 keV (96%)                                          | No   | Yes | No       | No     | Yes   |  |
| <sup>89</sup> Sr | 50.563<br>days                           | β <sup>-</sup>        | β <sub>max</sub> = 1500 keV<br>β <sub>mean</sub> = 587 keV | Yes  | Yes | Yes      | No     | No    |  |
| <sup>90</sup> Sr | 28.79 years                              | β <sup>-</sup>        | $\beta_{max}$ = 546 keV<br>$\beta_{mean}$ = 196 keV        | Yes  | Yes | No       | Yes    | No    |  |
| <sup>90</sup> Y  | 64 hours                                 | $\beta^-$             | β <sub>max</sub> = 2280 keV<br>β <sub>mean</sub> = 934 keV | Yes  | Yes | Yes      | No     | No    |  |

### Figure 1. Strontium Separation Options



following separation on Sr Resin (Figure 1A). However, due to the relatively short half-life of <sup>90</sup>Sr, **application to low level environmental analyses will be limited** by the available mass spectrometry technology and concentration chemistries. Recent publications suggest detection limits of ~1Bq/g are possible. However the achievable detection limit will depend on several factors, including sample type, sample size, and the age and type of MS instrument available [1]. **Radiometric techniques will allow lower detection limits** for <sup>90</sup>Sr and the ability to also determine <sup>89</sup>Sr.

Accurate measurement of strontium chemical yield as <sup>88</sup>Sr by ICP-MS has very little chance for impact by **isobaric interferences** (<sup>87</sup>Rb-H<sup>+</sup>, which should be effectively removed by Sr Resin Separation). <sup>90</sup>Zr is the only significant isobaric interference for <sup>90</sup>Sr measurement. Separation chemistries should be tailored to ensure complete removal of zirconium (rinsing with 3M HNO<sub>3</sub>-0.05M oxalic acid for Sr Resin separations). Determination of <sup>90</sup>Sr by ICP-MS **greatly simplifies the calculation of <sup>90</sup>Sr activity**, which can be complicated in radiometric detection techniques by the ingrowth of <sup>90</sup>Y and the decay of <sup>89</sup>Sr.

**ICP-MS** and **ICP-AES** (atomic emission spectrometry) are also very effective tools for the determination of **strontium chemical yield** when using radiometric detection methods for <sup>89</sup>Sr and <sup>90</sup>Sr. ICP-MS and ICP-AES will often give more precise chemical yield data than gravimetric techniques, while also allowing the use of **less Sr carrier** (<1 mg vs 4-10 mg stable Sr). Many environmental samples (soils, brines, sea water and some ground waters) may contain significant levels of stable strontium. Pre-anlysis of these samples by ICP-MS or ICP-AES for **native Sr** content may be necessary to adjust the amount of stable Sr yield monitor added and to ensure accurate measurement of the Sr yield throughout the separation process.

**Total** <sup>89/90</sup>**Sr** In instances where <sup>90</sup>**Sr is the only radiostrontium isotope** likely to be present or where total <sup>89</sup>Sr + <sup>90</sup>Sr determination is desired, radiostrontium may be determined by liquid scintillation counting or gas flow proportional counting following separation on Sr Resin (Figure 1A). **Confirmation of** <sup>90</sup>**Sr** activity will require **discrimination from** <sup>89</sup>**Sr** through two count methods and calculations [2], ingrowth and separation of <sup>90</sup>Y (Figure 1B or 1C) [4,5] or selective <sup>89</sup>Sr/<sup>90</sup>Y measurement(s) using Cerenkov counting [3].

<sup>90</sup>Sr by Direct Recovery of <sup>90</sup>Y In the analysis of debris from decommissioning of older facilities or other instances where <sup>90</sup>Sr is likely to be present in the absence of short-lived fission products (<sup>91</sup>Y, <sup>89</sup>Sr, etc.) and where <sup>90</sup>Y is in equilibrium with <sup>90</sup>Sr, <sup>90</sup>Sr may be determined by liquid scintillation, gas flow proportional counting, or Cerenkov counting following the direct separation of <sup>90</sup>Y on DGA Resin (Figure 1C) [6]. The very high retention of <sup>90</sup>Y on DGA resin allows recovery of <sup>90</sup>Y from difficult matrices, large sample volumes, and/or samples with high native Sr content that may prove problematic using the isolation of Sr on Sr Resin. However, the presence of fresh fission products, such as <sup>91</sup>Y and rare earth radionuclides, will cause significant positive bias in the <sup>90</sup>Sr determination performed using this method. Stable yttrium carrier yield can be measured by ICP-MS or ICP-AES.

**Two Count Method** Total <sup>89/90</sup>Sr can be measured in an **initial count** using GFPC or LSC following separation on Sr Resin (Figure 1A). Following a period of **ingrowth for** <sup>90</sup>Y, the samples can be **counted again**, and individual <sup>89</sup>Sr and <sup>90</sup>Sr activities calculated by **solving a series of equations** related to the decay of <sup>89</sup>Sr and the ingrowth of <sup>90</sup>Y (two count method) [2]. Ideally, the **initial count** is performed **as quickly as possible** following the separation on Sr Resin to minimize <sup>90</sup>Y ingrowth. The **second count** is ideally performed **after 1-2 weeks of** <sup>90</sup>Y **ingrowth**. Computational methods for resolving the <sup>89</sup>Sr and <sup>90</sup>Sr activities are outlined in Appendix B of reference [2].

**Cerenkov vs LSC** The beta emission of <sup>90</sup>Sr is below the threshold for efficient measurement by Cerenkov counting (LSC without addition of cocktail). <sup>89</sup>Sr may be determined by Cerenkov counting immediately following separation of radiostrontium on Sr Resin (Figure 1A). <sup>90</sup>Y will also be efficiently measured by Cerenkov counting.

Counting of <sup>89</sup>Sr by Cerenkov counting is **less efficient than by liquid scintillation** (LSC). However, the lower counting efficiency is partially offset by the **lower background** observed in Cerenkov counting. Furthermore, Cerenkov counting **eliminates interferences** from many low energy beta emitters that may cause bias in measurements made by LSC or GFPC. Additionally, Cerenkov counting **does not require the addition of cocktail**, offering cost savings in waste disposal and reagent costs. Cerenkov counting **does not suffer from quenching** which can occur in liquid scintillation.

Counting <sup>89</sup>Sr by Cerenkov or total <sup>89/90</sup>Sr by GFPC as a first measurement leaves open the option to separate <sup>90</sup>Y from the strontium fraction for further confirmation of <sup>90</sup>Sr or discrimination of <sup>89</sup>Sr/<sup>90</sup>Sr. Counting the strontium fraction by LSC limits any further separation on the sample due to the addition of the LSC cocktail, leaving the two count method as the only viable <sup>89</sup>Sr/<sup>90</sup>Sr discrimination option after LSC measurements of <sup>89/90</sup>Sr, unless the sample is split prior to LSC.

<sup>89</sup>Sr by Cerenkov, <sup>89/90</sup>Sr by LSC Counting (Rapid <sup>89/90</sup>Sr) <sup>89</sup>Sr is determined by Cerenkov counting immediately following separation of radiostrontium on Sr Resin (Figure 1A) [3]. After the addition of LSC cocktail, <sup>90</sup>Sr may then be determined by measuring total radiostrontium (<sup>89</sup>Sr + <sup>90</sup>Sr) by standard liquid scintillation counting and calculating the difference (total radiostrontium - <sup>89</sup>Sr), taking into account ingrowth of <sup>90</sup>Y and decay of <sup>89</sup>Sr during counting.

This approach may not be appropriate for samples that require long count times to meet data quality objectives or for samples with low ratios of <sup>89</sup>Sr/<sup>90</sup>Sr, due to interference from <sup>90</sup>Y ingrowth. However, this method can be used very effectively to rapidly determine <sup>89</sup>Sr and <sup>90</sup>Sr in samples containing relatively high amounts of radiostrontium in the immediate aftermath of a release of fresh fission products.

When utilizing Cerenkov and/or liquid scintillation counting, chemical yield of Sr throughout the separation process is most effectively measured via stable strontium carrier by taking a small aliquot of the separated strontium fraction for analysis by ICP-MS or ICP-AES. Recovery by <sup>85</sup>Sr gamma emission is not recommended, due to interference with the measurement of total radiostrontium by liquid scintillation counting.

**DGA vs Sr Resin for** <sup>90</sup>**Y Separation** For samples with expected high ratios of <sup>89</sup>Sr/<sup>90</sup>Sr, using DGA to selectively retain <sup>90</sup>Y (Figure 1C) is recommended, as the Sr can be effectively rinsed from the column, while <sup>90</sup>Y recovery remains strongly adsorbed (k' Y on DGA >>1000). In high <sup>89</sup>Sr/<sup>90</sup>Sr samples, small breakthrough of strontium from separation on Sr Resin (Figure 1B) into the <sup>90</sup>Y fraction (k' Sr only ~100 on Sr Resin) can lead to high bias of the <sup>90</sup>Sr measurement and a corresponding low bias of the <sup>89</sup>Sr.

<sup>89</sup>Sr by Cerenkov, <sup>90</sup>Y by Cerenkov Following Ingrowth and Separation <sup>89</sup>Sr is measured by Cerenkov counting following separation of strontium on Sr Resin (Figure 1A). <sup>90</sup>Sr may then be determined by waiting 1-14 days for <sup>90</sup>Y ingrowth, acidifying the Cerenkov counted sample with HNO<sub>3</sub>, separating the <sup>90</sup>Y using Sr Resin (Figure 1B) or DGA Resin (Figure 1C), and measuring <sup>90</sup>Y by Cerenkov counting.

This method is **not as rapid as the Cerenkov/LSC rapid** <sup>89/90</sup>Sr **method**, due to the <sup>90</sup>Y ingrowth period. However, due to the separation of pure <sup>90</sup>Y from strontium and other beta emitting nuclides with high decontamination factors, this method **may provide more accurate and sensitive measurements** for samples with **high** <sup>89</sup>Sr/<sup>90</sup>Sr **or** <sup>90</sup>Sr/<sup>89</sup>Sr ratios. The chemical yield of Sr may be measured via stable strontium carrier by taking a small aliquot of the separated strontium fraction for analysis by ICP-MS or ICP-AES or by <sup>85</sup>Sr gamma emission. Yttrium yields may also be tracked using stable Y and measurement by ICP-MS or ICP-AES.

<sup>89/90</sup>Sr by GFPC, <sup>90</sup>Y by GFPC Following Ingrowth and Separation Total <sup>89/90</sup>Sr can be measured effectively using low background gas flow proportional counting (GFPC) following separation of strontium on Sr Resin (Figure 1A) [4,5]. Prior to measurement by GFPC, purified strontium fractions are dried onto cupped planchets or precipitated as carbonates or phosphates and collected on filter paper (and dried). Immediate counts of the purified strontium fractions yield the total radiostrontium (<sup>89</sup>Sr + <sup>90</sup>Sr). Following the initial count for total radiostrontium and a period of <sup>90</sup>Y ingrowth, the dried sample can be dissolved from the planchet or filter using HNO<sub>3</sub>, and additional separations performed to isolate <sup>90</sup>Y from the strontium fraction using Sr Resin (Figure 1B) or DGA Resin (Figure 1C). <sup>90</sup>Y can then be measured by GFPC after evaporation on a stainless steel planchet or collection of an YF<sub>3</sub> precipitate on a filter. <sup>90</sup>Sr activity can be calculated using the measured <sup>90</sup>Y activity and the period of ingrowth from the initial separation. <sup>89</sup>Sr activity can be calculating by difference (total radiostrontium - <sup>90</sup>Sr).

When using GFPC, strontium chemical yield may be determined via stable strontium using ICP-MS, ICP-AES or gravimetric methods. Additionally, multiple drawer GFPC systems allow for the **simultaneous counting** of multiple samples, an option which is normally not available for Cerenkov or LSC.

#### References

1) Ohno, T., Hirono, M., Sakata, S., "Determination of Strontium 90 in environmental samples by triple quadrapole ICP-MS and its application to Fukashima soil samples." *J. Anal. At. Spectrom.* 33, 1081-1085 (2018).

2) EPA 402-R-10-001d, "Rapid Radiochemical Method for Total Radiostrontium (Sr-90) In Water for Environmental Remediation Following Homeland Security Events," October 2011.

3) Banavali, A. D. et al. "Strontium-89, 90 Analysis by Eichrom Column Chemistry and Cerenkov Counting". 38th Annual Conference on Bioassay Analytical and Environmental Radiochemistry. Santa Fe, NM. November 1992.

4) ASTM D5811-08, "Standard Test Method for Strontium-90 in Water."

5) IAEA Analytical Quality in Nuclear Applications Series No. 27. "Rapid Simultaneous Determination fo 89Sr and 90Sr in Milk: A Procedure Using Cerenkov and Scintillation Counting," IAEA/AQ/27 (2013). https://www-pub.iaea.org/MTCD/ Publications/PDF/IAEA-AQ-27\_web.pdf

6) Maxwell, S.L.; Culligan, B.K.; Hutchinson, J.B.; Utsey, R.C.; McAlister, D.R.; "Rapid Determination of <sup>90</sup>Sr in Seawater Samples," *J. Radional. Nucl. Chem.*, 303, 709-717 (2015).

## Measurement of <sup>36</sup>Cl and <sup>129</sup>l in Water

#### AN-1701-10

**Summary of Method** Chlorine-36 and Iodine-129 are separated and measured from up to 500mL aliquots of water. Samples are adjusted to 0.1-1.0M H<sub>2</sub>SO<sub>4</sub> and 0.1M SnSO<sub>4</sub> is added to ensure reduction of any oxidized species to chloride (Cl<sup>-</sup>) and Iodide (I<sup>-</sup>). The CL resin is prepared by Ioading with Ag<sup>+</sup> cations which will facilitate the retention of Cl<sup>-</sup> and I<sup>-</sup>. The sample is Ioaded onto the CL Resin, the column is rinsed with deionized water, and Cl<sup>-</sup> is recovered using 0.1M ammonium thiocyanate (NH<sub>4</sub>SCN). If iodine determination is also required, the resin is the rinsed with 0.1% NaOH and then I<sup>-</sup> is recovered using 0.35M sodium sulfide (Na<sub>2</sub>S). <sup>36</sup>Cl (t<sub>1/2</sub> = 3.01E5 years,  $\beta_{\text{mean}}^-$  = 251keV,  $\beta_{\text{max}}^-$  = 709.55keV, abundance = 98.1% ) and <sup>129</sup>I (t<sub>1/2</sub> = 1.57E7 years,  $\beta_{\text{mean}}^-$  = 40.03keV,  $\beta_{\text{max}}^-$  = 149.3keV, abundance = 100% ) can then be measured using liquid scintillation counting.

#### Reagents

CI Resin Bulk (CL-B50-A or CL-B50-S)\* -or-CI Resin Prepacked 2mL columns (CL-C50-A) Deionized Water NaOH \*If packing own columns for H<sub>2</sub>SO<sub>4</sub> gravity flow, then use CL-B50-A. SnSO<sub>4</sub> If using vacuum assisted flow, Na<sub>2</sub>S then use CL-B50-S. NH<sub>4</sub>SCN AgNO<sub>3</sub> (or Solution of 10mg/mL Ag<sup>+</sup>) Liquid Scintillation Cocktail

#### Equipment

Centrifuge tubes - 50mL Liquid scintillation vials, 20mL glass Liquid scintillation counter Calibrated pipets and disposable tips Appropriately sized glass beakers and flasks Analytical balance Filter funnel and paper **Optional:** Empty columns and frits for packing own columns

#### **Sample Preparation**

Up to 500mL of water sample in glass beaker.

Adjust to 0.1-1.0M H<sub>2</sub>SO<sub>4</sub>.

Add 1 mL of 0.1M SnSO<sub>4</sub> per 50mL of sample.

### CL Resin Preparation (Batch Mode)

## Reagent amounts may be adjusted proportionally to as needed.

Weigh 10 grams CL Resin into a 250mL flask.

Dissolve 0.65 grams AgNO\_3 in 100mL 1M  $H_2SO_4.$ 

Add Ag/H<sub>2</sub>SO<sub>4</sub> solution to resin and Mix for 2 hrs.

Filter CL Resin and rinse twice with 1M H<sub>2</sub>SO<sub>4</sub>.

Slurry resin in 100mL 0.1M H<sub>2</sub>SO<sub>4</sub> and pack into the appropriate sized column.

#### CL Resin Preparation (Column)

#### Reagent amounts may be adjusted

Slurry resin in 100mL 0.1M H<sub>2</sub>SO<sub>4</sub> and pack into the appropriate sized column or obtain prepacked 2mL column of CL Resin.

Rinse 2mL column with 10mL 1M  $H_2SO_4$ .

Dissolve 0.65 grams AgNO<sub>3</sub> in 100mL 1M H<sub>2</sub>SO<sub>4</sub>. Load 2 mL of solution onto each 2mL column. Wait 2hr for Ag to be completely absorbed.

Rinse 2mL column with 10mL 1M  $H_2SO_4$ .

Mix sample well.

#### **Chlorine/Iodine Separation**

- 1) Load Sample onto 2mL CL Resin column at 2mL/min.
- 2) Rinse 2mL CL Resin column with 10mL 0.1M  $H_2SO_4$ .
- 3) Rinse 2mL CL Resin column with 10mL deionized water.
- Strip 2mL CL Resin into 20mL LSC vial with 5mL 0.1M NH<sub>4</sub>SCN to recover chloride.
- 5) Rinse 2mL CL Resin column with 10mL 0.1% NaOH.
- Strip 2mL CL Resin column into a 2mL LSC vial with 5mL 0.35M Na<sub>2</sub>S.
- 7) Add 15 LSC cocktail to each LSC vial.
- 8) Count <sup>36</sup>Cl and <sup>129</sup>l samples by LSC.

| Decontamination Factors (DF)<br>for Chloride and Iodide Frations |                                     |       |  |  |  |
|------------------------------------------------------------------|-------------------------------------|-------|--|--|--|
| Analyte                                                          | lyte DF CI- fraction DF I- fraction |       |  |  |  |
| Ва                                                               | >1000                               | >600  |  |  |  |
| Cd                                                               | >6900                               | >7700 |  |  |  |
| Cu                                                               | >210                                | >190  |  |  |  |
| Mn                                                               | >210                                | >370  |  |  |  |
| Ni                                                               | >170                                | >320  |  |  |  |
| Pb                                                               | >300                                | >720  |  |  |  |
| U                                                                | >1900                               | >200  |  |  |  |
| <sup>60</sup> Co                                                 | >320                                | >320  |  |  |  |
| <sup>137</sup> Cs                                                | >150                                | >150  |  |  |  |
| <sup>90</sup> Sr/ <sup>90</sup> Y                                | >180                                | >160  |  |  |  |
| <sup>36</sup> Cl                                                 | N/A                                 | >160  |  |  |  |
| <sup>129</sup>                                                   | >420                                | N/A   |  |  |  |

| Retention of Cl and I                              |      |  |  |  |
|----------------------------------------------------|------|--|--|--|
| on CL Resin from 1M H <sub>2</sub> SO <sub>4</sub> |      |  |  |  |
| Analyte Dw                                         |      |  |  |  |
| <sup>36</sup> Cl                                   | 1600 |  |  |  |
| <sup>129</sup>                                     | 1980 |  |  |  |

Chloride capcity: 4 mg / 2mL column Iodide capcity: 15 mg / 2mL column

### References

1) A. Zulauf, S. Happel, M.B. Mokili, A. Bombard, H. Jungclas, "Characterization of an extraction chromatographic resin for the separation and determination of 36Cl and 129I." J. Radioanal. Nucl. Chem. 286(2), 539-546 (2010).

## Converting Methods from Gravity Columns to Cartridges

#### AN-1702-10

**Summary** Performing separations using Eichrom 2 mL pre-packed cartridges on a vacuum box system offers many advantages over gravity flow columns, including faster flow rates, improved chromatographic resolution, and the ability to stack multiple cartridges and measure multiple analytes from one sample aliquot. Converting separations methods from gravity flow columns to Eichrom cartridges is normally simple:



1) Obtain a 12-Hole or 24-Hole vacuum box, vacuum pump and tubing.



2) Add cartridges appropriate for the separation.3) Run separation procedure using the same solutions and volumes as the column method, adjusting the vacuum to achieve the optimal flow rate:

- 1 to 2 mL/min for sample load and rinses.
- 1 mL/min for stripping steps.

4) Drawing air through the cartridges for a short time (<5-10 minutes) between elution steps will not adversely impact the separation. Some samples will run faster than others.





5) Sample aliquots may be collected in individual 50 mL centrifuge tubes or collectively in the vacuum box liner.



6) Change cartridge reservoirs and inner and outer tips prior to elution of each analyte to improve purity.

7) The Eichrom vacuum box can also be used for CeF<sub>3</sub> microprecipitation source preparation using resolve filters.



Table 1. TEVA Resin Performance (2006-2017)\*

|                                    | 2 mL Cartridge    | 2 mL Column       |  |  |  |  |
|------------------------------------|-------------------|-------------------|--|--|--|--|
| Parameter                          | 50-100 μm         | 100-150 μm        |  |  |  |  |
| <sup>230</sup> Th % yield          | 99.3 <u>+</u> 0.5 | 99.1 <u>+</u> 0.7 |  |  |  |  |
| <sup>239</sup> Pu % yield          | 98.3 <u>+</u> 1.6 | 98.1 <u>+</u> 1.5 |  |  |  |  |
| <sup>239</sup> Pu % impurity in Th | 0.2 <u>+</u> 0.1  | 0.2 <u>+</u> 0.1  |  |  |  |  |
| <sup>230</sup> Th % impurity in Pu | 0.3 <u>+</u> 0.2  | 0.3 <u>+</u> 0.2  |  |  |  |  |

#### Table 2. TRU Resin Performance (2006-2017)\*

|                                    | 2 mL Cartridge    | 2 mL Column       |  |  |  |
|------------------------------------|-------------------|-------------------|--|--|--|
| Parameter                          | 50-100 μm         | 100-150 μm        |  |  |  |
| <sup>241</sup> Am % yield          | 99.4 <u>+</u> 0.4 | 99.4 <u>+</u> 0.6 |  |  |  |
| <sup>239</sup> Pu % yield          | 97.2 <u>+</u> 0.6 | 97.7 <u>+</u> 0.7 |  |  |  |
| <sup>239</sup> Pu % impurity in Am | 0.3 <u>+</u> 0.2  | 0.3 <u>+</u> 0.1  |  |  |  |
| <sup>241</sup> Am % impurity in Pu | 0.3 <u>+</u> 0.2  | 0.4 <u>+</u> 0.2  |  |  |  |
| *Eichrom Quality Method QA-0212    |                   |                   |  |  |  |

Table 3. UTEVA Resin Performance (2006-2017)\*

2 mL Cartridge

50-100 µm

99 + 3

99 + 4

0.2 <u>+</u> 0.1

0.1 + 0.1

2 mL Column

100-150 μm

99 + 3

99 + 4

0.1 + 0.1

0.1 + 0.1

\*Eichrom Quality Method QA-0213

Vacuum Box -12 Hole (AR-12-BOX) Includes:

- Rack for 50mL c-tubes (AR-12-RACK)
- Vacuum Gauge (AR-01-GAUGE-PVC)
- Vacuum Box Lid (AR-12-LID)
- White Inner Support Tubes (25)
- Yellow Outer Tips (25)
- Vacuum Box Manifold Plugs (50)
- Cartridge Reservoir, 10 mL (25)

Optional:

- Inner Liner (AR-12-LINER)
- Top Support (AR-12-TS)

Vacuum Box –24 Hole (AR-24-BOX) Includes:

- Rack for 50mL c-tubes (AR-24-RACK)
- Vacuum Gauge (AR-01-GAUGE-PVC)
- Vacuum Box Lid (AR-24-LID)
- White Inner Support Tubes (50)
- Yellow Outer Tips (50)
- Vacuum Box Manifold Plugs (50)
- Cartridge Reservoir, 10 mL (25) Optional:
- Inner Liner (AR-24-LINER)
- Top Support (AR-24-TS)

### Additional Equipment

- Vacuum Pump (Fisher no. 01-092-25 or equivalent)
- Tubing Tygon 1/4 in. I.D., 7/16 in. O.D. (Fisher no. 14-169-1K, or equivalent)
- White Inner Support Tubes (AR-1000-TUBE-PE)
- Yellow Outer Tips (AR-1000-OT)
- Stopcock, Polycarbonate (12) (AR-12-PC)
- 10 mL Cartridge Reservoir (200) (AR-200-RV10)
- 20 mL Cartridge Reservoir (200) (AR-200-RV20)

\*Eichrom Quality Method QA-0214

Parameter

<sup>230</sup>Th % yield

<sup>233</sup>U % yield

<sup>233</sup>U % impurity in Th

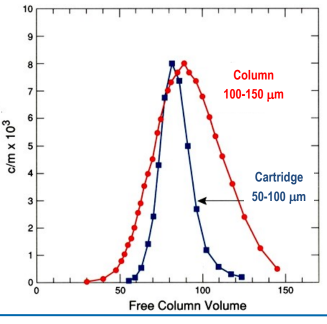
<sup>230</sup>Th % impurity in U

#### Table 4. Sr Resin Performance (2006-2017)\*

|                     | 2 mL Cartridge   | 2 mL Column      |
|---------------------|------------------|------------------|
| Parameter           | 50-100 μm        | 100-150 μm       |
| Sr % yield          | 93 <u>+</u> 3    | 95 <u>+</u> 3    |
| Ba % impurity in Sr | 0.2 <u>+</u> 0.1 | 0.2 <u>+</u> 0.1 |
| Ca % impurity in Sr | < 0.02           | < 0.02           |
| Y % impurity in Sr  | < 0.01           | < 0.01           |

\*Eichrom Quality Method QA-0215

#### Comparison of Elution Curves for Sr<sup>2+</sup> for Two Particle sizes of Sr Resin Elutrient 3.2 <u>M</u> HNO<sub>3</sub>, 23-24° C



#### +1 (630)963-0320

www.eichrom.com

# eichrom<sup>®</sup>

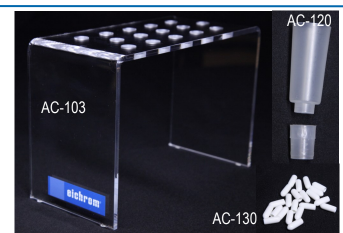
## **Packing Eichrom** 2 mL Columns

#### AN-1703-10

**Summary** Eichrom offers empty 2 mL columns and bulk resin for customers who wish to pack their own columns. This application note will offer advice on slurry packing columns that will exhibit favorable flow conditions and efficient separations. Some hydrophobic, difficult to wet resins may require additional treatment prior to slurry packing or may be dry packed.



Eichrom snap tip 2 mL Columns (AC-141-AL) come with the bottom frit inserted. Top frits and column top caps are also included.



2 mL Column Racks (AC-103), Column Tip Closures (AC-130), and 2-piece 25 mL Extension Funnels (AC-120) are also available.

The first step in slurry packing columns is to wet the resin by mixing with an appropriate aqueous phase. For most Eichrom resins, a solution of 0.05M HNO<sub>3</sub> is ideal for column packing. Add a volume of resin sufficient to pack your columns to 3-5x that volume of 0.05M HNO<sub>3</sub>. Mix the resin by vortexing, swirling or gently tumbling. Avoid vigorous shaking, which can lead to air bubbles that can degrade column flow and separation efficiency. If a portion of the resin floats on the surface of the 0.05M HNO<sub>3</sub>, centrifuge the sample. Repeat mixing and centrifuging as necessary to achieve a well wetted resin with minimal amount of floating material.

Some more hydrophobic resins will not wet well in 0.05M HNO<sub>3</sub>. Table 1 lists some difficult to wet resins and alternative matrices to facilitate wetting. The resins in Table 1 can be wetted by replacing the 0.05M HNO<sub>3</sub> with the alternative slurry matrix and following the steps above. Once DGA, LN2 and LN3 have been wetted with the alternative slurry matrix, centrifuge and decant the aqueous phase and replace with 0.05M HNO<sub>3</sub> for storage and column packing.

Wetted

resin.

slurry

| Table 1. Slurry Matrices for Difficult to wet Resins |                        |  |  |  |  |  |
|------------------------------------------------------|------------------------|--|--|--|--|--|
| Resin                                                | Slurry Matrix          |  |  |  |  |  |
| Prefilter                                            | 0.05M HNO <sub>3</sub> |  |  |  |  |  |
| Ni Resin                                             | 0.15M Ammonium Citrate |  |  |  |  |  |
| DGA, Normal                                          | 2M HNO <sub>3</sub>    |  |  |  |  |  |
| DGA, Branched                                        | 2M HNO <sub>3</sub>    |  |  |  |  |  |
| Cu Resin*                                            | 0.05 - 2M HCI          |  |  |  |  |  |
| LN2                                                  | 1M HNO <sub>3</sub>    |  |  |  |  |  |
| LN3                                                  | 2M HNO <sub>3</sub>    |  |  |  |  |  |
| *Cu Resin will float on the surface of the solution  |                        |  |  |  |  |  |
| even when wetted.                                    |                        |  |  |  |  |  |

Table 1 Slurry Matrices for Difficult to Wat Pasing

ideal for packing columns.

Floating or poorly wetted resin. difficult to slurry pack into columns.

The packing method is written assuming a 0.05M HNO<sub>3</sub> slurry matrix. For Ni Resin and Cu Resin, replace 0.05M HNO<sub>3</sub> with the appropriate alternative. Pre-packed Eichrom 2 mL columns contain 1.6 mL of resin. This method was written to replicate this fill volume.

As the resin is wetting in the appropriate matrix, add empty 2 mL columns to a column rack or other support. Add a small volume of 0.05M HNO<sub>3</sub> to soak the bottom frit and remove air bubbles. Soak until no can be seen escaping from the frits.



Add enough top frits for each column to a centrifuge tube with a small volume of 0.05M HNO<sub>3</sub>. Soak the frits to remove air bubbles. Swirl or vortex to mix, but avoid vigorous shaking.



Decant the 0.05M HNO<sub>3</sub> from the empty columns. Mix the slurry of resin and 0.05M HNO<sub>3</sub> to suspend the resin. Add the resin slurry to each column until the reservoir above the column is ~half full. Allow the resin to settle (~1 hr).



Full 2 mL columns should have a bed height of  $4.1 \pm 0.2$  cm. Add additional slurry to meet this height or remove excess resin using a plastic transfer pipet. Leave enough 0.05M HNO<sub>3</sub> above the packed bed to fill the column and a portion of the reservoir.

Place a pre-soaked frit into each column. Using a glass or plastic stir rod, push the frit to the top of the packed bed. Decant the 0.05M HNO<sub>3</sub> above the top frit and rinse away any residual resin from above the top frit using 0.05M HNO<sub>3</sub>.





If storing the columns for future use, fill the reservoir above the top frit ~half full with  $0.05M \text{ HNO}_3$  and place a top cap on each column. If using the column immediately, snap off the bottom tip, allow any excess  $0.05M \text{ HNO}_3$  to drain, and begin the column preconditioning step.

**Dry Packing Columns** Some difficult to wet resins can also be dry-packed into columns:

1) Place 2 mL columns with bottom frits in column rack.

- 2) Weigh  $0.65 \pm 0.05$  g of dry resin into each column.
- 3) Tap to settle the resin.

4) Place a top frit on each column and push the frit to the top of the resin bed.

5) Rinse away any excess resin above the top frit.

6) Add preconditioning solution to the column reservoir. Over pressure or vacuum may be required to initiate column flow.

7) Allow solution to drain through column.

**Cichrom** 1955 University Lane Lisle, IL 60532, USA +1 (630)963-0320 www.eichrom.com

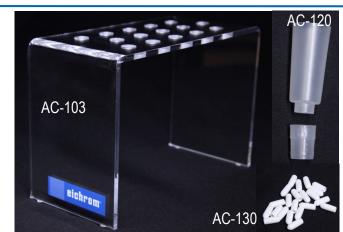
## Packing Eichrom 2 mL Columns

#### AN-1703-11

**Summary** Eichrom offers empty 2 mL columns and bulk resin for customers who wish to pack their own columns. This application note will offer advice on slurry packing columns that will exhibit favorable flow conditions and efficient separations. Some hydrophobic, difficult to wet resins may require additional treatment prior to slurry packing or may be dry packed.



Eichrom snap tip 2 mL Columns (AC-141-AL) come with the bottom frit inserted. Top frits and column top caps are also included.



2 mL Column Racks (AC-103), Column Tip Closures (AC-130), and 2-piece 25 mL Extension Funnels (AC-120) are also available.

The first step in slurry packing columns is to wet the resin by mixing with an appropriate aqueous phase. For most Eichrom resins, a solution of 0.05M HNO<sub>3</sub> is ideal for column packing. Add a volume of resin sufficient to pack your columns to 3-5x that volume of 0.05M HNO<sub>3</sub>. Mix the resin by vortexing, swirling or gently tumbling. Avoid vigorous shaking, which can lead to air bubbles that can degrade column flow and separation efficiency. If a portion of the resin floats on the surface of the 0.05M HNO<sub>3</sub>, centrifuge the sample. Repeat mixing and centrifuging as necessary to achieve a well wetted resin with minimal amount of floating material.

Some more hydrophobic resins will not wet well in 0.05M HNO<sub>3</sub>. Table 1 lists some difficult to wet resins and alternative matrices to facilitate wetting. The resins in Table 1 can be wetted by replacing the 0.05M HNO<sub>3</sub> with the alternative slurry matrix and following the steps above. Once DGA, LN2 and LN3 have been wetted with the alternative slurry matrix, centrifuge and decant the aqueous phase and replace with 0.05M HNO<sub>3</sub> for storage and column packing. Higher concentrations of acid may enable faster wetting, but may be more dense than the resin. Once wetted, the high acid concentrations can be remove or diluted to allow resin to sink.

Wetted

resin.

slurry

ideal for

packing

columns.

#### Table 1. Slurry Matrices for Difficult to Wet Resins

| Resin                                                                 | Slurry Matrix          |  |  |  |  |
|-----------------------------------------------------------------------|------------------------|--|--|--|--|
| Prefilter                                                             | 0.05M HNO <sub>3</sub> |  |  |  |  |
| Ni Resin                                                              | 0.15M Ammonium Citrate |  |  |  |  |
| DGA, Normal                                                           | 2-4M HNO <sub>3</sub>  |  |  |  |  |
| DGA, Branched                                                         | 2-4M HNO <sub>3</sub>  |  |  |  |  |
| Cu Resin*                                                             | 0.05 - 2M HCI          |  |  |  |  |
| LN2                                                                   | 1M HNO <sub>3</sub>    |  |  |  |  |
| LN3                                                                   | 2M HNO <sub>3</sub>    |  |  |  |  |
| *Cu Resin will float on the surface of the solution even when wetted. |                        |  |  |  |  |

Floating or poorly wetted resin, difficult to slurry pack into columns. The packing method is written assuming a 0.05M HNO<sub>3</sub> slurry matrix. For Ni Resin and Cu Resin, replace 0.05M HNO<sub>3</sub> with the appropriate alternative. Pre-packed Eichrom 2 mL columns contain 1.6 mL of resin. This method was written to replicate this fill volume.

As the resin is wetting in the appropriate matrix, add empty 2 mL columns to a column rack or other support. Add a small volume of 0.05M HNO<sub>3</sub> to soak the bottom frit and remove air bubbles. Soak until no can be seen escaping from the frits.



Add enough top frits for each column to a centrifuge tube with a small volume of 0.05M HNO<sub>3</sub>. Soak the frits to remove air bubbles. Swirl or vortex to mix, but avoid vigorous shaking.



Decant the 0.05M HNO<sub>3</sub> from the empty columns. Mix the slurry of resin and 0.05M HNO<sub>3</sub> to suspend the resin. Add the resin slurry to each column until the reservoir above the column is ~half full. Allow the resin to settle (~1 hr).



Full 2 mL columns should have a bed height of  $4.1 \pm 0.2$  cm. Add additional slurry to meet this height or remove excess resin using a plastic transfer pipet. Leave enough 0.05M HNO<sub>3</sub> above the packed bed to fill the column and a portion of the reservoir.

Place a pre-soaked frit into each column. Using a glass or plastic stir rod, push the frit to the top of the packed bed. Decant the 0.05M HNO<sub>3</sub> above the top frit and rinse away any residual resin from above the top frit using 0.05M HNO<sub>3</sub>.





If storing the columns for future use, fill the reservoir above the top frit ~half full with  $0.05M \text{ HNO}_3$  and place a top cap on each column. If using the column immediately, snap off the bottom tip, allow any excess  $0.05M \text{ HNO}_3$  to drain, and begin the column preconditioning step.

**Dry Packing Columns** Some difficult to wet resins can also be dry-packed into columns:

1) Place 2 mL columns with bottom frits in column rack.

- 2) Weigh  $0.65 \pm 0.05$  g of dry resin into each column.
- 3) Tap to settle the resin.

4) Place a top frit on each column and push the frit to the top of the resin bed.

5) Rinse away any excess resin above the top frit.

6) Add preconditioning solution to the column reservoir. Over pressure or vacuum may be required to initiate column flow.

7) Allow solution to drain through column.

**Cichrom** 1955 University Lane Lisle, IL 60532, USA +1 (630)963-0320 www.eichrom.com

## Rapid Determination of 89/90Sr in Steel Samples

#### AN-1801-10

Strontium is separated and measured from 1-2 gram steel samples. Samples are digested Summary of Method with concentrated nitric, hydrochloric, and hydrofluoric acids. The digestate is evaporated to dryness, the residue dissolved in HNO<sub>3</sub>/H<sub>3</sub>BO<sub>3</sub>, and a calcium fluoride precipitate is used to concentrate the strontium and remove matrix. An optional NaOH fusion may also be performed, post sample digestion, to dissolve concrete or stone included in the sample. Strontium is separated from matrix impurities and potentially interfering radionuclides in the sample using stacked 2 mL and 1 mL cartridges of Eichrom Sr Resin. Radiostrontium is measured on a low background gas flow proportional counter. Average chemical recovery of strontium, determined by gravimetric yield of stable strontium carrier, was 90-94%. Measured values of 90Sr agreed to within 3% of reference values for 60 minute count times. The minimum detectable activity for <sup>90</sup>Sr in 2 g samples with 60 minute count times was 0.56 Bg/g. A single operator can

Add 15 g NaOH.

Dilute to 100 mL.

Add 2 mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> and

prepare batches of 12 samples for the measurement of <sup>90</sup>Sr in less than 8 hours.

#### Reagents

Sr Resin, 2 mL Cartridges (Eichrom SR-R50-S) Sr Resin, 1 mL Cartridges (Eichrom SR1ML-R50-S) Nitric Acid (70%) Hydrochloric Acid (37%) Hydrofluoric Acid (49%) or Ammonium Bifluoride **Deionized Water** 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> Optional NaOH Fusion. Strontium Carrier (10 mg/mL) 2M AI(NO<sub>3</sub>)<sub>3</sub> Fuse at 600C for 20 min. Sr-90 standard Oxalic acid Dissolve in DI water. Boric acid Transfer to 250 mL c-tube. 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>\* Sodium Hydroxide\*

#### Equipment

8.5 mL 3.2 M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>. Vacuum Pump Centrifuge Centrifuge. Decant Muffle Furnace\* supernate. Dissolve ppt. in Hot Plate 80 mL 1.5M HCI. Analytical Balance Teflon Beakers (Zr Crucibles\*) 50 mL and 250 mL Centrifuge Tubes Cupped Stainless Steel Planchets (~5 mL volume) Gas Flow Proportional Counter Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20 mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT)

#### **Figure 1. Sample Preparation**

Add 1-2 g steel sample to Teflon beaker\*. \*If using optional fusion, omit HF/H<sub>3</sub>BO<sub>3</sub> and use Zr crucible.

> Add 6 mg Sr Carrier, 5 mL 70% HNO<sub>3</sub>, 20 mL 37% HCl, and 5 mL 49% HF. Digest on Hotplate to dryness.

Add 1 mL 70% HNO3. 10 mL 37% HCl, and 1 mL 49% HF. Digest on Hotplate to dryness.

Add 5 mL 3M HNO<sub>3</sub>-0.25M H<sub>3</sub>BO<sub>3</sub> and 5 mL 37% HCl. Digest on Hotplate to dryness.

Add 25 mL 1M HCl. Warm to dissolve. Transfer to 250 mL centrifuge tube. Repeat 2 additional times.

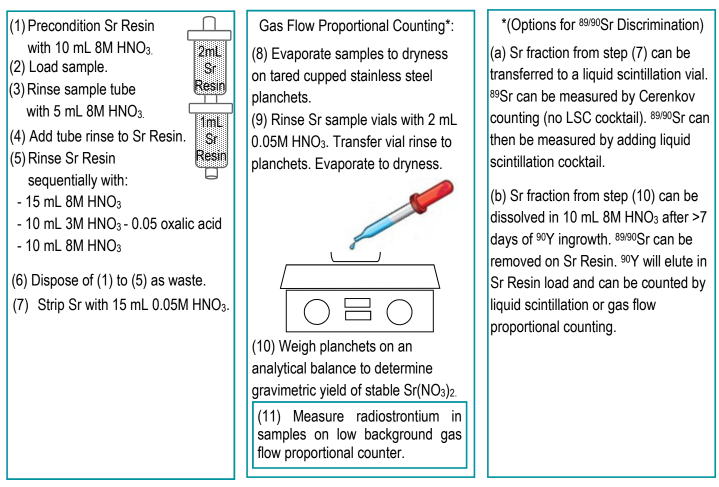
Add 2 mL 1.25M Ca(NO3)<sub>2</sub>. Dilute to 170 mL with 0.01M HCl.

Add 25 mL 49% HF. Mix Well. Allow to sit 20 min. Centrifuge. Discard supernate.

Dissolve precipitate in 5 mL 3M HNO<sub>3</sub>-0.25M H<sub>3</sub>BO<sub>3</sub>, 5 mL 8M HNO<sub>3</sub>, 5 mL 2M Al(NO<sub>3</sub>)<sub>3</sub>. Mix. Warm if necessary. Centrifuge. Check for solids.

Continue to Sr Separation.

### Figure 2. Strontium Separation and Measurement



\*Additional discussion of <sup>89/90</sup>Sr separation and measurement options can be found in Eichrom Application Note AN-1624-10.

| <br>Details   | Sample<br>replicates | Reference<br>(mBq/sample) | Measured<br>(mBq/sample) | Average<br>% Diff. | Sr Carrier<br>% Yield |
|---------------|----------------------|---------------------------|--------------------------|--------------------|-----------------------|
| 90Sr          | 10                   | 1.415                     | 1.41 <u>+</u> 0.04       | 2.6                | 90.1 <u>+</u> 2.4     |
| <br>89Sr+90Sr | 8                    | 3.816                     | 3.97 <u>+</u> 0.09       | 4.1                | 94.1 <u>+</u> 2.8     |

#### Method Performance for 2 g Steel Samples

### References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchison, Robin. C. Utsey, Ralf Sudowe, Daniel R. McAlister, "Rapid method to determine <sup>89/90</sup>Sr in steel samples," *J. Radioanal. Nucl. Chem.*, *314(1)*, *439-450* (2017).

## Rapid Determination of Pu in Steel Samples

Figure 1. Sample Preparation

Add 1-2 g steel sample to Teflon beaker\*.

#### AN-1802-10

**Summary of Method** Plutonium is separated and measured from 1-2 gram steel samples. Samples are digested with concentrated nitric, hydrochloric, and hydrofluoric acids. The digestate is evaporated to dryness, the residue dissolved in HNO<sub>3</sub>/H<sub>3</sub>BO<sub>3</sub>, and a CaF<sub>2</sub>/LaF<sub>3</sub> precipitate is used to concentrate the Pu and remove matrix. An optional NaOH fusion may also be performed, post sample digestion, to dissolve concrete or stone included in the sample and deal more rigorously with refractory Pu. Plutonium is separated from matrix impurities and potentially interfering radionuclides in the sample using 2 mL cartridges of Eichrom TEVA Resin. Plutonium is measured by alpha spectrometry following rare earth fluoride microprecipitation onto Eichrom Resolve filters. The chemical recovery of Pu, determined by <sup>242</sup>Pu tracer, was 90–99%. Measured values of Pu typically agreed to within 7-8% of reference values for 16 hour count times. The minimum detectable activity for Pu in 2 g samples with 16 hour count times was 0.25 mBq/g.

A single operator can prepare batches of 12 samples for the measurement of Pu in less than 8 hours.

#### Reagents

Muffle Furnace\*

Analytical Balance

Teflon Beakers (Zr Crucibles\*)

Alpha Spectrometry System

50 mL and 250 mL Centrifuge Tubes

Yellow Outer Tips (Eichrom AR-1000-OT)

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX)

Cartridge Reservoir, 20 mL (Eichrom AR-200-RV20)

Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE)

Resolve Filters in Funnel (Eichrom RF-DF25-25PP01)

Hot Plate

\*If using optional fusion, omit HF/H<sub>3</sub>BO<sub>3</sub> and use Zr crucible. TEVA Resin, 2 mL Cartridges (Eichrom TE-R50-S) Nitric Acid (70%) Add <sup>242</sup>Pu tracer, 5 mL 70% HNO<sub>3</sub>, Hydrochloric Acid (37%) Hydrofluoric Acid (49%) or Ammonium Bifluoride 20 mL 37% HCl, and 5 mL 49% HF. Lanthanum Carrier (10 mg/mL) Digest on Hotplate to dryness. Cerium Carrier (10 mg/mL) Add 1 mL 70% HNO3. Deionized Water 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> Optional NaOH Fusion. <sup>242</sup>Pu Tracer 10 mL 37% HCl, and 1 mL 49% HF. 2M AI(NO<sub>3</sub>)<sub>3</sub> Add 15-20 g NaOH. Boric acid Digest on Hotplate to dryness. NaNO<sub>2</sub> Fuse at 600C for 20 min. Ascorbic Acid 30% H<sub>2</sub>O<sub>2</sub> 10-20% (w:w) TiCl<sub>3</sub> in HCl Add 5 mL 3M HNO<sub>3</sub>-0.25M H<sub>3</sub>BO<sub>3</sub> Dissolve in DI water. 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>\* and 5 mL 37% HCl. Transfer to 250 mL c-tube. Sodium Hydroxide\* Digest on Hotplate to dryness. Dilute to 100 mL. Add 2 mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub>, 5 mg La, and Add 25 mL 1M HCl. Warm to dissolve. Equipment 8.5 mL 3.2 M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>. Transfer to 250 mL centrifuge tube. Vacuum Pump Repeat 2 additional times. Centrifuge

Centrifuge. Decant – supernate. Dissolve ppt. in 80 mL 1.5M HCI.

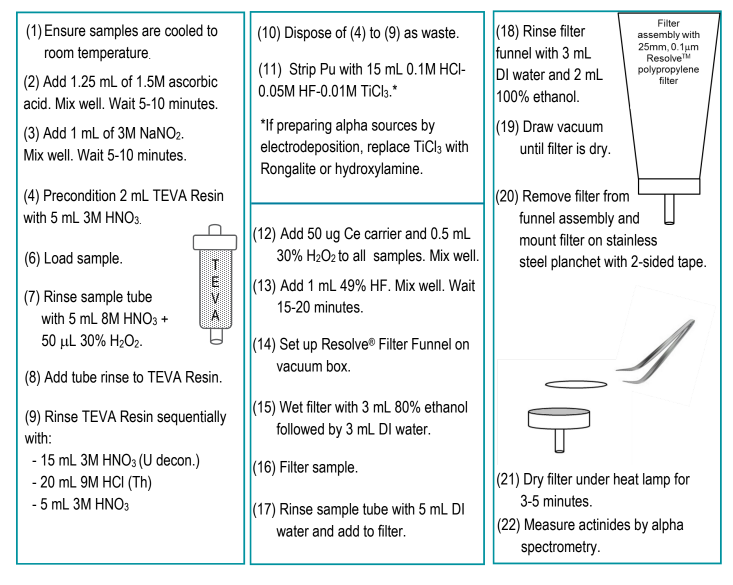
### Dilute to 170 mL with 0.01M HCI. Add 2 mL 1.25M Ca(NO3)<sub>2</sub>,5 mg La, and 3 mL 20% TiCl<sub>3</sub>.

Add 25 mL 49% HF. Mix Well. Allow to sit 20 min. Centrifuge. Discard supernate.

Dissolve precipitate in 7 mL 3M HNO<sub>3</sub>-0.25M H<sub>3</sub>BO<sub>3</sub>, 6 mL 8M HNO<sub>3</sub>, 7 mL 2M Al(NO<sub>3</sub>)<sub>3</sub>. Mix. Warm if necessary. Centrifuge. Check for solids.

Continue to Pu Separation.

### Figure 2. Load Solution Preparation and Plutonium Separation



#### Method Performance for Pu in Steel Samples

| Details                         | Sample<br>replicates | Reference<br>(mBq/sample) | Measured<br>(mBq/sample) | Average<br>% Diff. | 242Pu tracer<br>% Yield |
|---------------------------------|----------------------|---------------------------|--------------------------|--------------------|-------------------------|
| 238Pu in 2 g Steel              | 5                    | 37.0                      | 37.7 <u>+</u> 1.6        | 4.2                | 89.3 <u>+</u> 2.3       |
| 239Pu in 2 g Steel              | 5                    | 24.5                      | 24.4 <u>+</u> 1.6        | 6.6                | 96.5 <u>+</u> 3.4       |
| 239Pu (refractory) in 2 g Steel | 5                    | 24.5                      | 23.4 <u>+</u> 0.9        | 3.8                | 98.9 <u>+</u> 6.6       |
| 239 Pu in 5 g Steel             | 4                    | 37.0                      | 38.3 <u>+</u> 1.0        | 2.6                | 92 <u>+</u> 14          |

### References

1) Sherrod L. Maxwell, Brian K. Culligan, Jay B. Hutchison, Robin. C. Utsey, Ralf Sudowe, Daniel R. McAlister, "Rapid method to determine plutonium isotopes in steel samples," *J. Radioanal. Nucl. Chem.*, 314(2), 1103-1111 (2017).

## Rapid Determination of <sup>226</sup>Ra in Steel Samples

#### AN-1803-10

**Summary of Method** <sup>226</sup>Ra is isolated from 1 gram steel samples. Samples are digested with concentrated nitric, hydrochloric, and hydrofluoric acids. The digestate is evaporated to dryness, the residue dissolved in HNO<sub>3</sub>/H<sub>3</sub>BO<sub>3</sub> and calcium fluoride precipitate is used to concentrate the radium and remove matrix. Radium is separated from matrix impurities and potentially interfering radionuclides in the sample using cation exchange and DGA Resin. Radium is measured by alpha spectrometry following barium sulfate microprecipitation onto Eichrom Resolve Filters. The chemical recovery, determined by <sup>133</sup>Ba tracer, was 89–95%. Measured values of <sup>226</sup>Ra agreed to within 5% of reference values for 16 hour count times. The minimum detectable activity for <sup>226</sup>Ra in 1 g samples with 16 hour count times was 0.5 mBq/g. A single operator can prepare batches of 12 samples for the measurement of <sup>226</sup>Ra in less than 8 hours.

#### Reagents

Cation Exchange Resin (Eichrom C8-B500-F-H) DGA Resin, Normal 2 mL Cartridges (Eichrom DN-R50-S) Ammonium Hydroxide (Listed as 28% NH<sub>3</sub> or 56% NH<sub>4</sub>OH) Nitric Acid (70%) Deionized Water <sup>133</sup>Ba Tracer 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> Barium Carrier (1 mg/mL) Isopropyl Alcohol Ammonium Sulfate Denatured Ethanol Hydrochloric Acid (37%) Hydrogen Peroxide (30%)

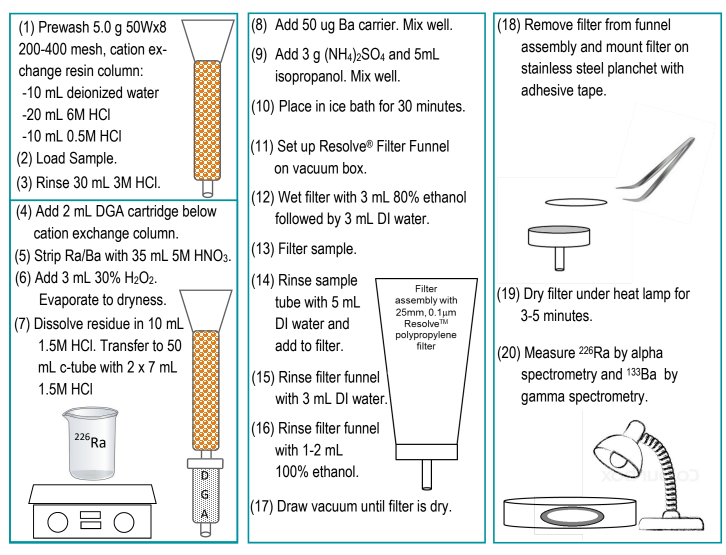
#### Equipment

Plastic Chromatography Column (Eichrom AC-50E-5M) Column Extension Funnel (Eichrom AC-20X-20M) Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20 mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filter in Disposable Funnel (Eichrom RF-DF-25-25PP01) 50 mL Centrifuge Tubes Centrifuge Hotplate 150 mL Glass beakers Vacuum Pump Heat Lamp Stainless Steel Planchets with adhesive tape Alpha Spectrometry System Gamma Spectrometry System (<sup>133</sup>Ba tracer)

## Figure 1. Sample Preparation Add 1 g steel sample to Teflon beaker. Add <sup>133</sup>Ba tracer, 5 mL 70% HNO<sub>3</sub>, 20 mL 37% HCl, and 5 mL 49% HF. Digest on Hotplate to dryness. Add 5 mL 70% HNO<sub>3</sub>, 10 mL 37% HCl, and 1 mL 49% HF. Digest on Hotplate to dryness. Add 5 mL 3M HNO<sub>3</sub>-0.25M H<sub>3</sub>BO<sub>3</sub> and 5 mL 37% HCl. Digest on Hotplate to dryness. Add 10 mL 0.25M HCI. Warm to dissolve. Transfer to 50 mL centrifuge tube. Repeat 2 additional times. Add 2 mL 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> and 6 mL 49% HF. Mix Well. Allow to sit 20 min. Centrifuge. Discard supernate. Dissolve precipitate in 10 mL 1M HCI -0.25M H<sub>3</sub>BO<sub>3</sub> and 10 mL 1M HCl. Mix. Warm if necessary. Centrifuge. Check for solids.

Continue to Ra Separation.

# Figure 2. Column Purification and Alpha Source Preparation



#### Method Performance for 1 gram Steel Samples

| Sample     | Reference    | Measured          | Average | 133Ba tracer      |
|------------|--------------|-------------------|---------|-------------------|
| replicates | (mBq/sample) | (mBq/sample)      | % Diff. | % Yield           |
| 5          | 36.8         | 36.5 <u>+</u> 0.8 | 1.9     | 95.4 <u>+</u> 5.9 |
| 5          | 73.7         | 74.9 <u>+</u> 3.1 | 3.7     | 88.8 <u>+</u> 1.8 |
| 5          | 184          | 183 <u>+</u> 5    | 1.9     | 90 <u>+</u> 13    |

#### References

1) Sherrod L. Maxwell, Brian K. Culligan, Robin C. Utsey and Daniel R. McAlister, "Rapid Method to Determine <sup>226</sup>Ra in Steel Samples," *J. Radioanal. Nucl. Chem.*, 314(2), 1417-1423 (2017).

# Rapid Determination of Pu/Np and Am/Cm in Granite

#### AN-1804-10

**Summary of Method** Pu/Np and Am/Cm are separated and measured from 1 gram samples of granite. Samples are finely ground and fused in a zirconium crucible for 15 minutes at 600°C with 15 grams of NaOH. The fusion cake is dissolved in water, and actinides are concentrated and separated from the matrix using a calcium phosphate precipitate enhanced with iron. A secondary precipitation with calcium fluoride removes additional matrix (including silicates) and decreases the volume of precipitate. The calcium fluoride precipitate is dissolved with nitric acid-boric acid-aluminum nitrate to form the load solution. Analytes are separated from remaining matrix and potentially interfering radionuclides using stacked 2 mL TEVA and DGA Resin cartridges. Actinides are measured by alpha spectrometry after CeF<sub>3</sub> microprecipitation onto Resolve<sup>(R)</sup> Filters. An additional separation of Am/Cm from rare earth elements using TEVA resin and ammonium thiocyanate may be required for samples with significant rare earth content. The rugged sample preparation technique enables high tracer recovery and excellent analytical results, even when refractory materials are present.

#### Reagents

TEVA Resin, 2 mL Cartridges (Eichrom TE-R50-S) DGA Resin, 2 mL Cartridges (Eichrom DN-R50-S) Lanthanum and Cerium Carriers (10 mg/mL) Iron Carrier (50 mg/mL Fe, as ferric nitrate) <sup>242</sup>Pu (or <sup>236</sup>Pu if Np is measured) tracer <sup>243</sup>Am tracer 10-20% TiCl<sub>3</sub> Ammonium Thyiocyanate HF(49%) 30% H<sub>2</sub>O<sub>2</sub> Nitric Acid (70%) Hydrochloric Acid (37%) **Deionized Water** 1.25M Ca(NO<sub>3</sub>)<sub>2</sub> 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>  $2M AI(NO_3)_3$ Boric Acid Sodium Hydroxide Ascorbic Acid NaNO<sub>2</sub>

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20 mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PP01) 50 mL and 250 mL Centrifuge Tubes 250 mL Zirconium crucibles with zirconium lids Alpha Spectrometry System Stainless Steel planchets with two sided tape Centrifuge Muffle Furnace Hot Plate/Heat Lamp Analytical Balance Vacuum Pump

### Figure 1. Sample Preparation

1 g finely ground sample in zirconium crucible Add <sup>242</sup>Pu or <sup>236</sup>Pu and <sup>243</sup>Am tracers.

Heat samples to dryness on hot plate.

Add 15 g of NaOH. Cover crucibles with zirconium lids. Fuse at 600°C for 15-20 minutes.

Carefully remove samples from furnace and cool in fume hood. Add 25-50 mL of water and heat on hot plate to dissolve fusion cake.

Transfer to a 250 mL centrifuge tube. Rinse crucible with water. Dilute to 180 mL with water.

Cool to room temperature. Add 125 mg Fe, 4 mg La, and 50 mg Ca. Mix. Add 5 mL 3.2M (NH<sub>4</sub>)<sub>2</sub>HPO<sub>4</sub>. Mix. Add 4 mL 20% TiCl<sub>3</sub>. Mix. Centrifuge 10 min. Decant supernate.

Dissolve precipitate in 80 mL 1.5M HCI. Dilute to 170 mL. Add 2 mL 20% TiCl<sub>3</sub>, 25 mg Ca, and 20 mL 49% HF. Mix. Cool in ice bath 10 min. Centrifuge 10min. Decant supernate.

Dissolve precipitate in 7 mL 3M HNO<sub>3</sub>-0.25M H<sub>3</sub>BO<sub>3</sub>, 6 mL 7M HNO<sub>3</sub>, and 7 mL 2M Al(NO<sub>3</sub>)<sub>3</sub>. Adjust valence with 1 mg Fe, 1.25 mL 1M ascorbic acid. Mix. Wait 5-10 min. Add 1 mL 3.5M NaNO<sub>2</sub> and 1.5 mL 70% HNO<sub>3</sub>.

# Figure 2. Actinide Separation on TEVA-DGA and Source Preparation

| (1) Precondition stacked 2 mL                                       | Optional Am/Cm rare earth separation.                          | (22) Filter sample. Filter          |
|---------------------------------------------------------------------|----------------------------------------------------------------|-------------------------------------|
| TEVA and DGA cartridges                                             | (10) Add 2 mL 70% HNO <sub>3</sub> + 50 uL 10%                 | assembly with 25mm, 0.1μm           |
| with 10 mL 3M HNO <sub>3</sub> .                                    | H <sub>2</sub> SO <sub>4</sub> to Am/Cm. Evaporate to dryness. | (23) RINSE Sample Resolve™          |
| (2) Load sample solution.                                           | (11) Ash to dryness with 3 mL 70%                              | tube with 5 mL DI                   |
| (3) Rinse sample tube                                               | HNO <sub>3</sub> + 2 mL 30% H <sub>2</sub> O <sub>2</sub> .    | water. Add to filter.               |
| with 5 mL 6M HNO <sub>3</sub> . Add                                 | (12) Dissolve Am/Cm in 5 mL                                    | (24) Rinse funnel                   |
| tube rinse to cartridges.*                                          | 4M NH₄SCN-0.1M Formic acid.                                    | with 3 mL DI water                  |
| (4) Rinse cartridges with                                           | (13) Precondition 2 mL TEVA with 5 mL                          | and 2 mL 100                        |
| 10 mL 3M HNO <sub>3</sub> .                                         | 4M NH₄SCN-0.1M Formic acid.                                    | ethanol.                            |
| (5) Separate TEVA and DGA                                           | (14) Load Am/Cm on TEVA.                                       | (25) Draw vacuum                    |
| cartridges.                                                         | (15) Rinse Am/Cm beaker with 5 mL                              | until filter is dry.                |
| (6) Rinse TEVA cartridge with:                                      | 4M NH <sub>4</sub> SCN-0.1M Formic acid.                       | (26) Remove filter from funnel.     |
| -15 mL 3M HNO <sub>3</sub> (U decon.)                               | Add to TEVA.                                                   | Mount filter on stainless steel     |
| -20 mL 9M HCI (Th)                                                  | (16) Rinse TEVA w/ 10 mL 1.5M                                  | planchet with 2-sided tape.         |
| -5 mL 3M HNO <sub>3</sub>                                           | NH <sub>4</sub> SCN-0.1M Formic acid.                          | /                                   |
| (7) Strip Pu (and Np) from TEVA                                     | (17) Strip Am/Cm from TEVA with                                |                                     |
| cartridge with 20 mL 0.1M HCl-<br>0.05MHF-0.01M TiCl <sub>3</sub> . | 20 mL 1M HCI.                                                  |                                     |
|                                                                     | (18) Add 0.5 mL 30% H <sub>2</sub> O <sub>2</sub> to Pu, and   |                                     |
| (8) Rinse DGA cartridge with:                                       | 0.2  mL 30% H <sub>2</sub> O <sub>2</sub> to Am/Cm samples.    |                                     |
| -10 mL 3M HCl                                                       | (19) Add 50ug Ce to Pu and Am/Cm                               |                                     |
| -3 mL 1M HNO <sub>3</sub>                                           | samples. Mix well. Add 1 mL 49% HF.                            |                                     |
| -20 mL 0.1M HNO <sub>3</sub> (U decon.)                             | Mix well. Wait 15-20 minutes.                                  |                                     |
| -10 mL 0.05M HNO <sub>3</sub>                                       | (20) Set up Resolve <sup>®</sup> Filter Funnel on              | (27) Dry filter under heat lamp for |
| -20 mL 3M HNO <sub>3</sub> -0.25M HF (Th)                           | vacuum box.                                                    | 3-5 minutes.                        |
| -5 mL 4M HCl                                                        | (21) Wet filter with 3 mL 80% ethanol                          | (28) Measure actinides by alpha     |
| (9) Strip Am and Cm from DGA with                                   | followed by 3 mL DI water.                                     | spectrometry.                       |
| 10 mL 0.25M HCI.                                                    |                                                                |                                     |

\*Adding 50uL 30% H<sub>2</sub>O<sub>2</sub> to the tube rinse can improve Uranium recoveries and decontamination in Pu(Np) fractions.

#### Method Performance for 1 gram Granite Samples

| Analyte           | Sample replicates | Reference<br>(mBq/g) | Measured<br>(mBq/g) | Average<br>% Diff. | Tracer<br>% Yield |
|-------------------|-------------------|----------------------|---------------------|--------------------|-------------------|
| <sup>239</sup> Pu | 8                 | 29.4                 | 29.2 <u>+</u> 1.4   | 4.3                | 92.1 <u>+</u> 5.5 |
| <sup>239</sup> Pu | 6                 | 21.2                 | 20.1 <u>+</u> 1.2   | 5.4                | 97.2 <u>+</u> 5.8 |
| <sup>238</sup> Pu | 6                 | 25.2                 | 25.0 <u>+</u> 2.2   | 6.9                | 97.2 <u>+</u> 5.8 |
| <sup>237</sup> Np | 6                 | 37.0                 | 37.1 <u>+</u> 1.7   | 3.5                | 97.2 <u>+</u> 5.8 |
| <sup>241</sup> Am | 4                 | 37.0                 | 37.7 <u>+</u> 3.3   | 7.0                | 90.7 <u>+</u> 5.1 |
| <sup>244</sup> Cm | 4                 | 33.1                 | 34.4 <u>+</u> 2.0   | 5.2                | 90.7 <u>+</u> 5.1 |

### References

1) Maxwell, S.L. Culligan, B. Hutchinson, J.B. Sudowe, R. McAlister, D.R. "Rapid Method to Determine Pu, Np, Am/Cm in Granite Samples," *J. Radioanal. Nucl. Chem.* 140, 102-108 (2018).

# Alpha Spectrometry Source Preparation: Rare Earth Fluoride Microprecipitation

#### AN-1805-11

eichrom

**Summary of Method** Rare earth fluoride microprecipitation is an alternative to electrodeposition for alpha spectrometry source preparation, which provides adequate alpha peak resolution for most analytical applications, while greatly reducing the time for sample preparation. Alpha spectrometry sources can often be prepared directly from the eluate used to recover the actinide fraction from the chromatographic column used to separate the actinides from the sample matrix and potentially interfering nuclides, eliminating the numerous evaporation and digestion steps normally required for electrodeposition, reducing the alpha spectrometry source preparation time from 3-8 hours to 30-60 minutes, and reducing the emission of corrosive acid fumes through the laboratory fume hood vents.

Lanthanum, cerium or neodymium carrier and hydrofluoric acid are normally used to produce the rare earth fluoride precipitate. Ammonium bifluoride may be used instead of HF. Additionally, for laboratories which are restricted from the use of fluoride, Ce(OH)<sub>4</sub> precipitation (AN-1807) may be a suitable alternative.

Rare earth fluoride precipitates will nearly quantitatively carry trivalent and tetravalent actinides, while rejecting pentavalent and hexavalent actinides. Therefore, the addition of  $TiCI_3$  is required to reduce U(VI) to U(IV) to prepare uranium samples. Samples of the other actinides may be further purified from U during the rare earth fluoride precipitation by the addition of  $H_2O_2$ , which will ensure U(VI) that will not be carried on rare earth fluorides.

Eichrom's Resolve Filters (RF-DF25-25PE01) are manufactured specifically for alpha spectrometry source preparation. The manufacture and quality control procedures ensure a uniform surface for the collection of the rare earth fluoride precipitate, reducing self attenuation of the alpha emissions, which can degrade peak resolution. Other filter membranes may not be suitable for alpha source preparation or may require the addition of substrate to achieve adequate resolution.

Sources prepared by rare earth precipitation and mounted to stainless steel planchets with doubled-side tape or glue typically sit closer to the detector in alpha spectrometry systems than electrodeposition sources. The difference in distance from the source to the detector can lead to a 5-10% higher efficiency for the measurement of microprecipitation sources. Since most laboratories calibrate their alpha spectrometry systems with electrodeposited sources, the efficiency difference must be considered when determining the absolute recovery of the chemical yield tracers.

Rare earth fluoride microprecipitation onto Eichrom Resolve Filters produces alpha spectra which are suitable for most analytical applications. However, electrodeposition may be required for some applications, such as the preparation of calibration sources and the measurement of nuclides with difficult to resolve alpha peaks.

#### Reagents

Lanthanum, Cerium or Neodymium Carrier (10 mg/mL) HF(49%) 30% H<sub>2</sub>O<sub>2</sub> Deionized Water Denatured Ethanol 10-20% TiCl<sub>3</sub> (for Uranium fractions)

#### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Yellow Outer Tips (Eichrom AR-1000-OT) Resolve Filters in Funnel (Eichrom RF-DF25-25PE01) 50 mL Centrifuge Tubes Alpha Spectrometry System Heat Lamp Vacuum Pump Stainless Steel planchets with two sided tape (A.F. Murphy part no. F-2-C)

#### 4 Deve Earth Eluarida Alpha Spectrometry Source Drevention

| Figure I. Rate Earth i                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      | -iuonue Alpha Spectrometr                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               | y Source Preparation                                                                                                                                                                                                                                                                                                                                                                                                                             |
|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Uranium Samples*(1) Obtain a purified sample of U in a50 mL centrifuge tube using anappropriate separation method.Samples are typically in 10-20 mL of1M HCl or 0.1M ammoniumbioxalate. Other matrices andvolumes should be tested prior toapplication. Concentrations of >0.1MHNO3 may interfere with uraniumreduction by TiCl3 and lead to poorrecoveries in the rare earth fluorideprecipitation.(2) Add 0.25 mL of 20% TiCl3 and100 μg of La, Ce or Nd carrier. Mix.(3) Add 1 mL 49% HF. Mix well. Wait | Pu/Np Samples*(1) Obtain a purified sample of Pu/Np in a 50 mL centrifuge tube using<br>an appropriate separation method.Samples are typically in 15-20 mL of<br>dilute HCI-HF with a reducing agent.(2) Add 50 µg of La, Ce or Nd carrier<br>and 0.5 mL 30% H2O2. Mix.(3) Add 1 mL 49% HF. Mix well. Wait<br>15-20 minutes. Proceed to step (4).Am/Cm, An(III), and Ln(III) Samples*(1) Obtain a purified sample of Am/<br>Cm, An(III) or Ln(III) in a 50 mL<br>centrifuge tube using an appropriate<br>separation method. Samples are | <ul> <li>(5) Wet filter with<br/>3 mL 80% ethanol<br/>followed by 3 mL<br/>DI water.</li> <li>(6) Filter sample.</li> <li>(7) Rinse sample<br/>tube with 5 mL DI<br/>water. Add to filter.</li> <li>(8) Rinse funnel<br/>with 3 mL DI water<br/>and 2 mL ethanol.</li> <li>(9) Draw vacuum until filter is dry.</li> <li>(10) Remove filter from funnel. Mount<br/>filter on stainless steel planchet with<br/>2-sided tape or glue.*</li> </ul> |
| <ul> <li>15-20 minutes. Proceed to step (4).</li> <li>Thorium Samples*</li> <li>(1) Obtain a purified sample of Th in<br/>a 50 mL centrifuge tube using an<br/>appropriate separation method.</li> <li>Samples are typically in 10-15 mL of<br/>6-9M HCI.**</li> <li>(2) Add 50 μg of La, Ce or Nd carrier.</li> <li>Dillute to 40 mL with DI H<sub>2</sub>O. Mix.</li> <li>(3) Add 3 mL 49% HF. Mix well. Wait<br/>15-20 minutes. Proceed to step (4).</li> </ul>                                          | <ul> <li>typically in 15-20 mL of 0.1-4M HCl.<br/>Samples with high native rare earth<br/>content will require removal of rare<br/>earths using TEVA-SCN (AN-1806).</li> <li>(2) Dilute to reduce HCl concentration<br/>to &lt; 2M, add 50 μg of La, Ce or Nd<br/>carrier and mix.**</li> <li>(3) Add 1 mL 49% HF. Mix well. Wait<br/>15-20 minutes. Proceed to step (4).</li> <li>(4) Set up Resolve<sup>®</sup> Filter Funnel on<br/>vacuum box.</li> </ul>                                                                           | <ul> <li>(11) Dry filter under heat lamp for 5-10 minutes.</li> <li>(12) Measure actinides by alpha spectrometry. (samples can be dissolved off the filter using 1M HNO<sub>3</sub>-0.25M boric acid if additional purification is needed)</li> </ul>                                                                                                                                                                                            |

\*Some users prefer to dry the filters before mounting. With the polyethylene Resolve Filters®, this can lead to curling, making the filters more difficult to mount. Mounting the filters prior to drying is recommended.

\*\*Preparing filters from >2M HCI can lead to poor resolution. Dilution or neutralization with NH<sub>4</sub>OH can improve performance. Typical Performance of CeF<sub>3</sub> Microprecipitation onto Eichrom Resolve Filters

| Nuclide              | μg Ce | Matrix                                | Yield | <b>Resolution (FWHM)</b> |
|----------------------|-------|---------------------------------------|-------|--------------------------|
| <sup>230</sup> Th    | 50    | 30 mL 4.5M HCl                        | >95%  | 20-30 keV                |
| <sup>238/234</sup> U | 100   | 20 mL 1M HCl                          | >95%  | 30-40 keV                |
| <sup>239</sup> Pu    | 50    | 20 mL 0.1M HCl-0.05MHF-0.01MTiCl $_3$ | >95%  | 30-40 keV                |
| <sup>241</sup> Am    | 50    | 15 mL 4M HCl                          | >95%  | 22-28 keV                |

### References

1) Claude W. Sill, "Precipitation of Actinides as Fluorides or Hydroxides for High-Resolution Alpha Spectrometry," Nuclear and Chemical Waste Management, 7, 201-215 (1987).

2) ASTM C1163-14, Standard Practice for Mounting Actinides for Alpha Spectrometry Using Neodymium Fluoride

# Actinide/Rare Earth Separation (TEVA-SCN)

#### AN-1806-10

**Summary of Method** Am/Cm or other trivalent actinide(s) are separated from trivalent rare earth cations prior to preparation of rare earth fluoride microprecipitation sources for alpha spectrometry. Some samples (soil, rock, building materials, etc.) may have a high native content of rare earth metal ions, which cannot be adequately separated from the trivalent actinides during the normal analytical scale chromatographic separations used to purify these elements. The mass of the native rare earths can degrade the alpha spectra of the nuclides through mass self-attenuation. For these samples, an additional separation of the actinides from the rare earths using TEVA Resin in the thiocyante (SCN) form will improve the resolution of the alpha spectra.

After purification of the Am/Cm (or other trivalent actinides) on TRU or DGA Resin, the actinide fraction is digested with HNO<sub>3</sub>-H<sub>2</sub>SO<sub>4</sub>, evaporated to dryness, and dissolved in 4M NH<sub>4</sub>SCN-0.1M formic acid. The sample is then loaded onto a 2 mL cartridge of TEVA resin, which retains the actinides, while the rare earth elements are not retained. The Am/Cm (or other trivalent actinide) are then recovered in 1M HCl and prepared for alpha spectrometry by rare earth fluoride microprecipitation (AN-1805).

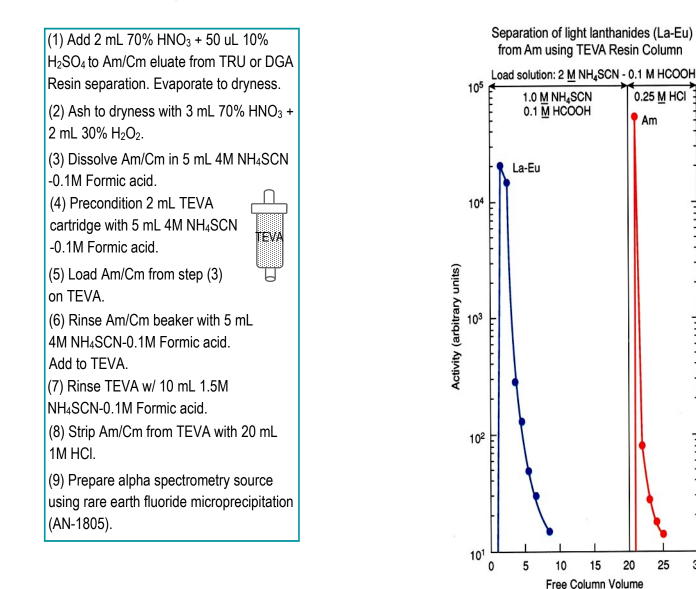
#### Reagents

TEVA Resin, 2 mL Cartridges (Eichrom TE-R50-S) Ammonium Thiocyanate (NH<sub>4</sub>SCN) Nitric Acid (70%) Hydrochloric Acid (37%) Sulfuric Acid (98%) Deionized Water Formic Acid

#### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20 mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50 mL Centrifuge Tubes Hot Plate Vacuum Pump

### Figure 1. Actinide/Rare Earth Separation on TEVA Resin



#### References

1) SEPERATION OF AMERICIUM FROM RARE EARTHS, Eichrom Method SPA-03.

30

# Alpha Spectrometry Source Preparation: Cerium Hydroxide Microprecipitation

#### AN-1807-11

**Summary of Method** Cerium hydroxide microprecipitation is an alternative to rare earth fluoride microprecipitation and electrodeposition for alpha spectrometry source preparation, providing adequate alpha peak resolution for most analytical applications, while greatly reducing the time for sample preparation relative to electrodeposition. Alpha spectrometry sources can often be prepared directly from the eluate used to recover the actinide fraction from the chromatographic column used to separate the actinides from the sample matrix and potentially interfering nuclides, eliminating the numerous evaporation and digestion steps normally required for electrodeposition, reducing the alpha spectrometry source preparation time from 3-8 hours to 30-60 minutes, and eliminating the emission of corrosive acid fumes through the laboratory fume hood vents.

Cerium hydroxide is an alternative to rare earth fluoride microprecipitation for labs looking to avoid the use of HF. Cerium hydroxide precipitates will nearly quantitatively carry actinides in all oxidation states from mineral acid solutions, but <u>will not work from bioxalate or other complexing agents</u>. Additional U decontamination of Th, Np/Pu and Am/ Cm samples achieved by the rare earth fluoride precipitation (AN-1805) will not occur using the cerium hydroxide precipitate has a yellow color, providing visual confirmation of the collection of the precipitate on the Resolve Filter and easy identification of the surface of the filter containing the precipitate.

Cerium carrier, hydrogen peroxide and a pH indicator are added to each sample fraction from the appropriate separation method. After mixing to distribute the carrier, ammonium hydroxide is added to adjust the pH. The optimal pH and the appropriate pH indicator will depend on the actinide metal ion being collected. U and Th show the highest recovery from pH 5-7, utilizing the bromocresol purple pH indicator. However, U and Th recoveries do not decrease dramatically if the pH is increased to 8-10. Am and Pu/Np recoveries are most consistent utilizing thymol blue, with a color change from pH 8-10. The higher pH range is important to ensure high recoveries of Am. Since Pu and Np are often measured together, with a single <sup>236</sup>Pu yield tracer, it is important that their recoveries are very similar. The pH of 8-10 is important to ensure similar recoveries of Pu and Np. At lower or higher pH, Np recovery can diverge significantly from Pu. [2]

Eichrom's Resolve Filters (RF-DF25-25PE01) are manufactured specifically for alpha spectrometry source preparation. The manufacture and quality control procedures ensure a uniform surface for the collection of the rare earth fluoride precipitate, reducing self attenuation of the alpha emissions, which can degrade peak resolution. Other filter membranes may not be suitable for alpha source preparation or may require the addition of substrate to achieve adequate resolution.

Sources prepared by microprecipitation precipitation and mounted to stainless steel planchets with double-sided tape or glue typically sit closer to the detector in alpha spectrometry systems than electrodeposition sources. The difference in distance from the source to the detector can lead to a 5-10% higher efficiency for the measurement of microprecipitation sources. Since most laboratories calibrate their alpha spectrometry systems with electrodeposited sources, the efficiency difference must be considered when determining the absolute recovery of the chemical yield tracers.

#### Reagents

Cerium Carrier (10 mg/mL) Deionized Water 30% Hydrogen Peroxide (H<sub>2</sub>O<sub>2</sub>) Denatured Ethanol Ammonium Hydroxide (NH<sub>4</sub>OH) Bromocresol Purple or Thymol Blue

#### Equipment

| Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX)                              |                        |  |  |  |
|--------------------------------------------------------------------------|------------------------|--|--|--|
| Yellow Outer Tips (Eichrom AR-1000-OT)                                   |                        |  |  |  |
| Resolve Filters in Funnel (Eichrom RF-DF25-25PE01)                       |                        |  |  |  |
| Stainless Steel planchets with two sided tape (A.F Murphy part no F-2-C) |                        |  |  |  |
| Alpha Spectrometry System                                                | 50 mL Centrifuge Tubes |  |  |  |
| Heat Lamp                                                                | Vacuum Pump            |  |  |  |

| Figure 1. Cerium Hydroxide Alpha Spectrometry Source Preparation*                                                                                                                                                                                                                                                                                                                                                            |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              |                                                                                                                                                                                 |  |  |  |
|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--|--|--|
| Uranium Samples<br>(1) Obtain a purified sample of U in a<br>50 mL centrifuge tube using an<br>appropriate separation method.<br>Samples are typically in 10-20 mL of<br>1M HCI.                                                                                                                                                                                                                                             | <ul> <li>(2) Add 25-50 μg of Ce carrier,</li> <li>0.2 mL of 30% H<sub>2</sub>O<sub>2</sub> and thymol blue.</li> <li>Mix.</li> <li>(3) Adjust to pH 8-10 (light blue color). Mix well. Proceed to step (4).</li> </ul>                                                                                                                                                                                                                                                                                                       | <ul> <li>(8) Rinse funnel<br/>with 3 mL DI water<br/>and 2 mL ethanol.</li> <li>(9) Draw vacuum<br/>until filter is dry.</li> </ul>                                             |  |  |  |
| <ul> <li>(2) Add 25-50 μg of Ce carrier,</li> <li>0.2 mL of 30% H<sub>2</sub>O<sub>2</sub> and bromocresol purple. Mix.</li> <li>(3) Adjust to pH 5-7 (blue/purple color). Mix well. Proceed to step (4).</li> </ul>                                                                                                                                                                                                         | Am/Cm, An(III), and Ln(III) Samples<br>(1) Obtain a purified sample of Am/<br>Cm, An(III) or Ln(III) in a 50 mL<br>centrifuge tube using an appropriate<br>separation method. Samples are                                                                                                                                                                                                                                                                                                                                    | (10) Remove filter<br>from funnel. Mount<br>filter on stainless<br>steel planchet with<br>2-sided tape.**                                                                       |  |  |  |
| Thorium Samples(1) Obtain a purified sample of Th ina 50 mL centrifuge tube using anappropriate separation method.Samples are typically in 20 mL of6-9M HCI.(2) Add 25-50 µg of Ce carrier,0.2 mL of 30% H <sub>2</sub> O <sub>2</sub> and bromocresolpurple. Mix.(3) Adjust to pH 5-7 (blue/purplecolor). Mix well. Proceed to step (4).Pu/Np Samples(1) Obtain a purified sample of Pu/Np in a 50 mL centrifuge tube using | <ul> <li>typically in 15-20 mL of 0.1-4M HCI.</li> <li>Samples with high native rare earth content will require removal of rare earths using TEVA-SCN (AN-1806).</li> <li>(2) Add 25-50 μg of Ce carrier,</li> <li>0.2 mL of 30% H<sub>2</sub>O<sub>2</sub> and thymol blue. Mix.</li> <li>(3) Adjust to pH 8-10 (light blue color). Mix well. Proceed to step (4).</li> <li>(4) Set up Resolve<sup>®</sup> Filter Funnel on vacuum box.</li> <li>(5) Wet filter with 3 mL 80% ethanol followed by 3 mL DI water.</li> </ul> | (11) Dry filter under heat lamp for<br>3-5 minutes.<br>(12) Measure actinides by alpha<br>spectrometry.<br>*Results in Table 1 are for typical<br>sample matrix from separation |  |  |  |
| Np in a 50 mL centrifuge tube using<br>an appropriate separation method.<br>Samples are typically in 15-20 mL of<br>dilute HCI-HF with a reducing agent.                                                                                                                                                                                                                                                                     | <ul><li>(6) Filter sample.</li><li>(7) Rinse sample tube with 5 mL DI water. Add to filter.</li></ul>                                                                                                                                                                                                                                                                                                                                                                                                                        | method. Performance with other<br>matrices or volumes should be<br>verified prior to implementation.                                                                            |  |  |  |

\*Some users prefer to dry the filters before mounting. With the polyethylene Resolve Filters®, this can lead to curling, making the filters more difficult to mount. Mounting the filters prior to drying is recommended.

Typical Performance of Ce(OH)<sub>4</sub> Microprecipitation onto Eichrom Resolve Filters

| Nuclide              | рН   | µg Ce | Matrix                                   | Yield  | Resolution (FWHM) |
|----------------------|------|-------|------------------------------------------|--------|-------------------|
| <sup>230</sup> Th    | 5-7  | 25    | 20 mL 9M HCl                             | 95-99% | 25-35 keV         |
| <sup>238/234</sup> U | 5-7  | 25    | 20 mL 1M HCl                             | 93-97% | 25-35 keV         |
| <sup>237</sup> Np    | 8-10 | 25    | 20 mL 0.15M HCI-0.05M KF-0.04M Rongalite | 89-93% | 25-35 keV         |
| <sup>239</sup> Pu    | 8-10 | 25    | 20 mL 0.15M HCI-0.05M KF-0.04M Rongalite | 86-92% | 25-35 keV         |
| <sup>241</sup> Am    | 8-10 | 25    | 15 mL 4M HCl                             | 91-95% | 25-35 keV         |

#### References

1) Claude W. Sill, "Precipitation of Actinides as Fluorides or Hydroxides for High-Resolution Alpha Spectrometry," Nuclear and Chemical Waste Management, 7, 201-215 (1987).

2) Hiromu Kurosaki, Rebbeca J. Mueller, Susan B. Lambert, Govind R. Rao, "Alternate method of source preparation for alpha spectrometry: no electrodeposition, no hydrofluoric acid," *J. Radioanal. Nucl. Chem.*, 311, 323-329 (2017).

# **Zirconium Separation** on ZR Resin

#### AN-1808-10

Summary of Method ZR resin contains a hydroxamate extractant which exhibits a high selectivity for Zr(IV), Ti(IV) and Nb(V) over Y(III), Sc(III) and Fe(III). From 0.01-10M HCI, Zr, Ti and Nb are strongly retained by the ZR resin, while Y and Sc are poorly retained. Fe(III) is strongly retained from 0.01-1M HCI and can be eluted from the ZR resin with 2-3M HCI. Zr can be recovered from the ZR resin with 0.1M oxalic acid, while Ti and Nb elution requires >0.25M oxalic acid.

emerging PET nuclides from their target materials, such as Zr(IV) from Y(III) and Ti(IV)

The unique selectivity of ZR resin makes it a useful material for the separation of from Sc(III). The target materials can be dissolved in high concentrations of hydrochloric acid and the dissolved target loaded onto ZR resin. Zr(IV) or Ti(IV) is retained, while the bulk target mass, Y(III) or Sc(III) passes through the ZR resin. Rinsing the ZR with 2-10M HCl completes removal of the target material and any Fe(III) present in the sample. The ZR resin can then be rinsed with more dilute HCl to reduce the residual acidity, Zr(IV) can be stripped using 0.1M oxalic acid, and Ti(IV) can be stripped with 0.25M oxalic acid. Further purification of the Zr(IV) or Ti(IV) can be achieved by loading the Zr(IV) or Ti(IV) onto strong base anion exchange resin from dilute oxalic acid-HCl.

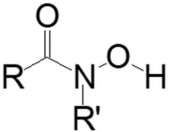
The easily hydrolyzed Zr(IV), Ti(IV) and Nb(V) should be stored in solution containing trace HF or oxalic acid to prevent loss of material to vials or formation of colloidal aggregates.

#### Reagents

ZR Resin 2 mL Cartridges (Eichrom ZR-R10-S) 1 mL Cartridges (Eichrom ZR1-R10-S) 0.3 mL Cartridges (Eichrom ZR0.3-R10-S) Bulk Resin (Eichrom ZR-B25-S) Hydrochloric Acid (37%) **Oxalic Acid Deionized Water** Hydrofluoric Acid (49%) - Optional

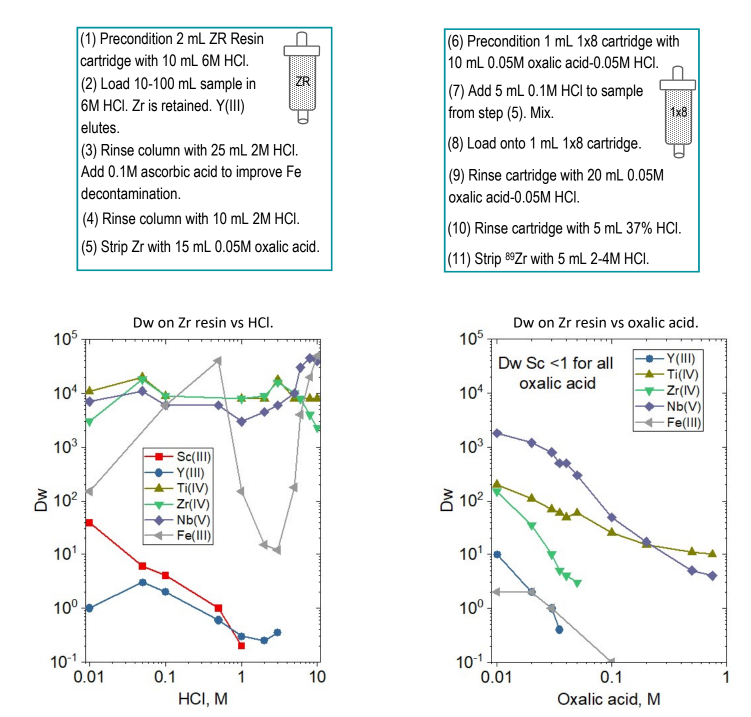
#### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20 mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50 mL Centrifuge Tubes Vacuum Pump



Hydroxamate Extractant

### Zirconium Separation on ZR Resin and Anion Exchange



### References

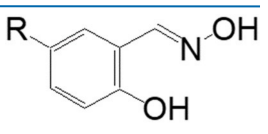
1) Dirks, et al., "On the Development and Characterization of a hydroxamate based extraction chromatographic resin," 61st Radiobioassy and Radiochemical Measurements Conference, October 25-30, 2015, Iowa City, Iowa.

2) Triskem INFOS, No 15, February 2016. http://www.triskem-international.com/scripts/files/59d1f4fc31f796.50370140/ tki\_15\_en\_web.pdf

# Copper Separation on CU Resin

#### AN-1809-10

**Summary of Method** CU Resin contains a benzaldoxime extractant adsorbed on an inert polymeric support. CU resin can be used to separate copper from other transition metals, such as zinc or nickel target material used in the production of Cu-64 and Cu-67. CU resin will selectively retain Cu from pH 2-5 HCl, HNO<sub>3</sub> or H<sub>2</sub>SO<sub>4</sub>, while Ni(II), Zn(II), Cd(II), Co(II), Fe(II), Fe(III), and Cr(III) are rejected. Cu can then be recovered from the Cu resin using 1-8 M HCl. Additional purification of Cu can be achieved by



#### **Benzaldoxime extractant**

stripping the Cu resin with 8M HCl through a strong base anion exchange resin (AG1x8). The Cu will be retained on the AG1x8 and can then be recovered in dilute HCl.

The CU is very hydrophobic and can be difficult to wet in dilute acid. Soaking the CU resin in 2M HCl improves the wetting. However, the wetted resin will still float on top of the liquid, making it difficult to slurry pack the CU resin. It is therefore recommended that the CU resin be used in prepacked cartridges or dry packed columns. Wet the columns or cartridges with 5-10 bed volumes of 2M HCl and then precondition the CU resin with dilute acid prior to loading the Cu sample. To initiate flow on the dry packed column or cartridge, a vacuum box, peristaltic pump or luer syringe will be required.

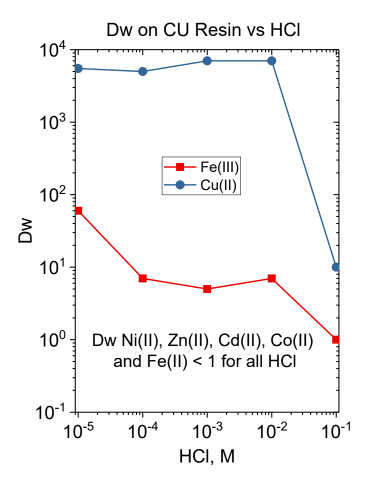
#### Reagents

#### CU Resin

2 mL Cartridges (Eichrom CU-R10-S) 1 mL Cartridges (Eichrom CU1-R10-S) 25 g bulk resin, 100-150 μm (Eichrom CU-B25-A) 25 g bulk resin, 50-100 μm (Eichrom CU-B25-S) Anion Exchange Resin (Eichrom A8-B500-F-CL) Hydrochloric Acid (37%) Ammonium Hydroxide (56%) Deionized Water

### Equipment

Vacuum Box (Eichrom AR-24-BOX or AR-12-BOX) Cartridge Reservoir, 20 mL (Eichrom AR-200-RV20) Inner Support Tubes-PE (Eichrom AR-1000-TUBE-PE) Yellow Outer Tips (Eichrom AR-1000-OT) 50 mL Centrifuge Tubes Vacuum Pump



# Figure 1. Cu Separation

(1) Dissolve Cu sample in HCl. Evaporate to dryness. Dissolve in 0.001M HCl. Adjust to pH 2-3 as necessary.\*

(2) Wet 1 mL CU resin cartridge with 10 mL 2M HCI.



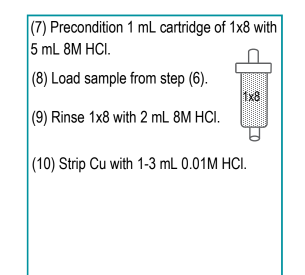
(3) Precondition CU resin with 10 mL 0.01M HCI.

(4) Load sample.

(5) Rinse CU resin with 10 mL 0.01M HCI.

(6) Strip Cu with 2-3 mL 8M HCl.

\*Sulfate may also be used and may provide buffering capacity, simplifying the pH adjustment.



### References

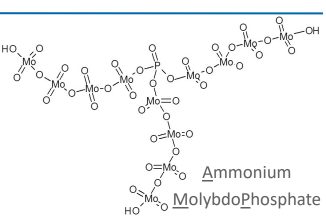
1) C. Dirks, B. Scholten, S. Happel, A. Zulauf, A. Bombard, H. Jungclas, "Characterization of a Cu selective resin and its application to the production of 64Cu," *J. Radioanal, Nucl. Chem.*, 286, 671-674 (2010).

2) Triskem INFOS, No 6, July 2011. http://www.triskem-international.com/scripts/files/59d1f4fc2c2091.54193347/ tki6\_en\_binderonline\_1.pdf

# Cs Separation on AMP-PAN and KNiFC-PAN Resins

#### AN-1810-10

**AMP-PAN** contains an inorganic ion exchange material (ammonium molybdophosphate, AMP) dispersed in an inert Ho polymeric support (polyacrylonitrile, PAN). The AMP has been shown to exhibit high selectivity for Cs from a wide range of solutions, including high acid concentrations and high salt concentrations (sea water). The AMP is imbedded into the PAN to improve the flow characteristics of packed columns. The material exhibits fast kinetics and high radiation stability, with no change in uptake observed for radiation doses of up to 1000 kGy [1]. Recovery of Cs from the AMP-PAN resin requires elution with 10 bed volumes of 5M NH<sub>4</sub>Cl or NH4NO<sub>3</sub>.



AMP-PAN has been used to remove Cs-137 from radioactive acidic waste streams containing high levels of sodium and potassium [2]. Actual waste and waste simulants were loaded onto 1.5 mL columns at 0.7 mL/min. In the first cycle, 0.15% breakthrough of Cs-137 was measured after 1500 mL of feed (99.85% Cs-137 removal). After regenerating the column by eluting Cs-137 with 50 mL of 5M NH<sub>4</sub>NO<sub>3</sub>, 0.53% breakthrough of Cs-137 was measured after 1250 mL of feed (99.47% removal of Cs-137). Average recovery of Cs-137 in the 5M NH<sub>4</sub>NO<sub>3</sub> regeneration cycles was 87%.

AMP-PAN has also been used to recover Cs-137 from sea water samples [3]. 5 mL columns of AMP-PAN were used to process 20 L samples of sea water which had been acidified to pH 1-2. Stable Cs, measured by ICP-MS, was used to trace the chemical recovery during the column separation. Flow rates of 35 mL/min were used. Recovery of Cs was  $93.5 \pm 5.0\%$ .

#### Reagents

#### AMP-PAN Resin 5 mL Cartridges (HC5-R10-M) 2 mL Columns (HC-C50-M) 5 mL Columns (HC5-C20-M) 8 mL Columns (HC8-C20-M) 10 mL Columns (HC10-C20-M) Nitric Acid (70%) Deionized Water Ammonium chloride or Ammonium Nitrate

### Equipment

Vacuum Box (AR-24-BOX or AR-12-BOX)\* Cartridge Reservoir, 20mL (AR-200-RV20)\* Inner Support Tubes-PE (AR-1000-TUBE-PE)\* Yellow Outer Tips (AR-1000-OT)\* 50 mL Centrifuge Tubes Vacuum Pump\* \*Or appropriately sized gravity flow column and accessories (see reverse).

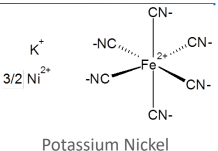
#### References

1) F. Sebesta, V. Stefula, "Composite ion exchanger with ammonium molybdophosphate and its properties" *J. Radioanal, Nucl. Chem.*, 140(1), 15-21 (1990).

2) Brewer, et al. "AMP-PAN column tests for the removal of Cs-137 from actual and simulated INEEL high-activity wastes," *Czechoslov J Phys*, 49(S1), 959-964 (1999).

3) Pike, et al. "Extaction of cesium in seawater off Japan using AMP-PAN resin and quantification via gamma spectroscopy and inductively coupled plasma mass spectrometry," *J Radioanal Nucl Chem*, 296(1), 369-374 (2012).
4) Triskem INFOS, No 10, July 2013. http://www.triskem-international.com/scripts/files/59d1f4fc2ec7b3.42683976/ tki10\_binder\_en\_web.pdf

**KNIFC-PAN** contains an inorganic ion exchange material (potassium nickel ferrocyanate, KNIFC) dispersed in an inert polymeric support (polyacrylonitrile, PAN). The KNIFC has been shown to exhibit high selectivity for Cs from a wide range of solutions, including sea water and other environmental waters. The KNIFC is imbedded into the PAN to improve the flow characteristics of packed columns.



FerroCyanate (KNiFC)

KNiFC-PAN has been used to remove cesium from sea water samples [5]. 100 L samples of sea water were processed through 25 mL columns of KNiFC-PAN at flow rates of up to 300 mL/min. Stable Cs was added as a yield tracer (measured by ICP-MS). Yields for cesium were 92.9 <u>+</u> 1.1% for 100 L samples of sea water acidified to pH 1. For 100 L samples of sea water (unacidified), cesium yields were 90.2 <u>+</u> 2.7%.

#### Reagents

KNiFC-PAN Bulk Resin (NC-B50-M) Nitric Acid (70%) Hydrochloric Acid (37%) Deionized Water

### Equipment

Empty Columns 2 mL snap tip (AC-141-AL) 2 mL cap tip (AC-100-MT-PP) 5 mL (AC-50E-5M) 20 mL (AC-20E-20M) Column Reservoir For 2 mL columns (AC-120-TK) 250 mL For 5 and 20 mL columns (AC-20X-20M) Column Rack 15 hole for 2 mL columns (AC-103) 12 hole for 5 and 20 mL columns (AC-20M-RACK) 50 mL Centrifuge Tubes

#### **Comparison of AMP-PAN and KNiFC-PAN Resins**

| Paramater             | AMP-PAN                                                  | KNIFC-PAN            |
|-----------------------|----------------------------------------------------------|----------------------|
| Cs Capacity           | 64 mg / g dry resin                                      | 256 mg / g dry resin |
| Density               | 0.27 g/mL                                                | 0.20 g/mL            |
| Recommended sample pH | 1 - 2                                                    | 1 - 7                |
| Sample Types          | Waste, Environmental                                     | Environmental        |
| Regeneration          | 5M NH <sub>4</sub> Cl or NH <sub>4</sub> NO <sub>3</sub> | None                 |

### References

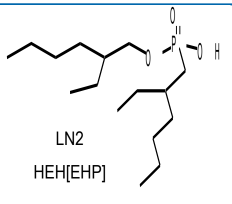
5) Kamenik, et al. "Fast concentration of dissolved forms of cesium radioisotopes from large sea water samples," *J. Radioanal. Nucl. Chem.*, 292(2), 841-846 (2012)

# Ce Separation from Rare Earth Nitrate Solutions

#### AN-1811-10

**Summary of Method** Cerium is oxidized from Ce(III) to Ce(IV) using sodium bromate and then selectively extracted from rare earth nitrate solutions using a column of LN2 resin. LN2 is an extraction chromatographic resin containing 2-ethyl-1-hexyl(2-ethyl-1-hexyl)phosphonic acid (HEH[EHP].

Ce can be oxidized to Ce(IV) from solutions of nitric acid and rare earth nitrate using NaBrO<sub>3</sub>, while the other rare earth metal ions remain in the trivalent oxidation state. The oxidation of Ce(III) to Ce(IV) and the retention on LN2 increases with the concentration of nitrate. The oxidation will not work from chloride solutions. Berkelium (Bk) can also be oxidized to Bk(IV) and separated from other trivalent actinides and rare earths using very similar conditions.



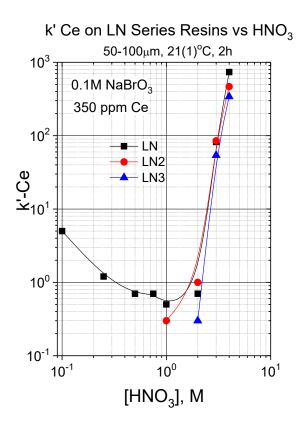
Once oxidize, the Ce(IV) or Bk(IV) are retained on the LN2 resin from 2-3M HNO<sub>3</sub>/Rare Earth Nitrate solutions, while trivalent metal ions are not retained. After rinsing with HNO<sub>3</sub> to remove any residual trivalent metal ions, the Ce or Bk can be recovered from the LN2 by elution with 0.25-0.50M HCl or HNO<sub>3</sub> + reducing agent ( $H_2O_2$ , hydroxylamine or ascorbic acid). Removal of Ce from 500 mL 2M HNO<sub>3</sub> + 0.75 M Y/Yb(NO<sub>3</sub>)<sub>3</sub> was >99.9% using a 10 mL column of LN2 resin[1].

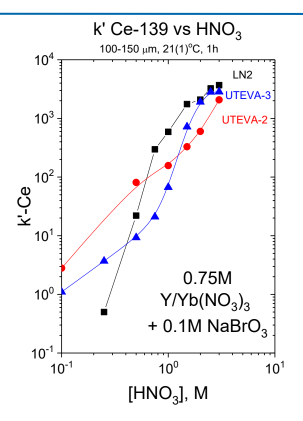
#### Reagents

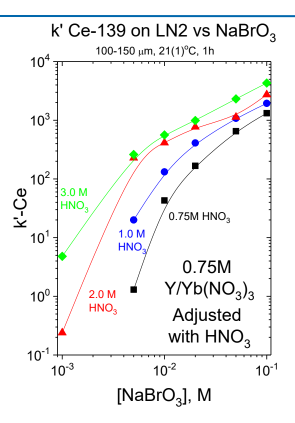
LN2 Bulk Resin (L2-BO1-S) Nitric Acid (70%) Hydrochloric Acid (37%) Sodium Bromate (NaBrO<sub>3</sub>) H2O2 (30%), Hydroxylamine ·HCl or Ascorbic Acid Deionized Water

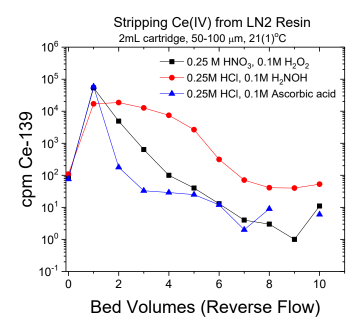
### Equipment

Empty Columns 2 mL snap tip (AC-141-AL) 2 mL cap tip (AC-100-MT-PP) 5 mL (AC-50E-5M) 20 mL (AC-20E-20M) Column Reservoir For 2 mL columns (AC-120-TK) 250 mL For 5 and 20 mL columns (AC-20X-20M) Column Rack 15 hole for 2 mL columns (AC-103) 12 hole for 5 and 20 mL columns (AC-20M-RACK) 50 mL Centrifuge Tubes

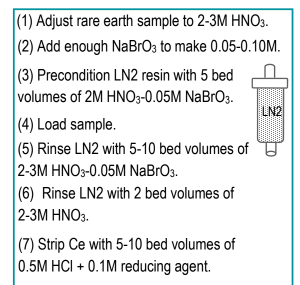








### **Ce Separation**



#### References

1) D.R. McAlister and E.P. Horwitz, unpublished data (2013).

# Fe Separation from Rare Earth Chlorides

#### AN-1812-10

**Summary of Method** Fe(III) is removed from rare earth chloride solutions by extraction of [FeCl<sub>4</sub>] on TEVA resin. The anionic ferric chloride complex is strongly retained by the TEVA Resin, while the rare earth chlorides are rejected. Hydrogen peroxide is added to the sample to ensure Fe(III), as Fe(II) is not extracted. The TEVA column can be regenerated by eluting Fe with five bed volumes of 0.1M HNO<sub>3</sub>. 99.7% removal of Fe from 500 mL 0.75M YCl<sub>3</sub>-1M HCl was achieved on a 10 mL column of TEVA resin (3 mL/min flowrate) [1].

### Reagents

TEVA Bulk Resin (TE-B25-S)Nitric Acid (70%)Hydrogen Peroxide (30% H2O2)Hydrochloric Acid (37%)Deionized Water

# Fe(III) Separation on WBEC Resin

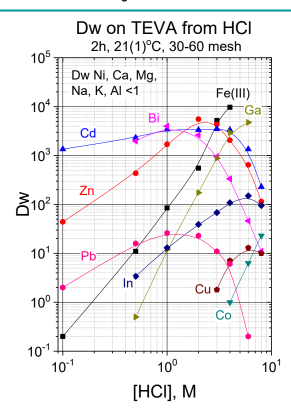
(1) Precondition WBEC column with 3 bed volumes of 2.5M HCl.

(2) Load Sample of rare earth chloride in >0.1M HCI (>2.25M total chloride ).



(4) Rinse column with 5 bed volumes of 6M HCl.

(5) Strip Fe in reverse direction with 5 bed volumes of  $0.10M \text{ HNO}_3$  to regenerate.

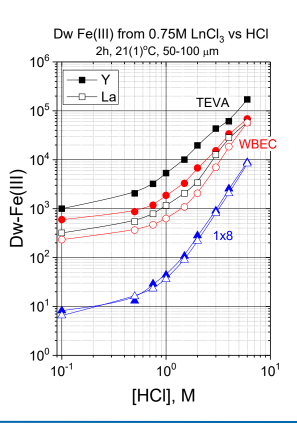


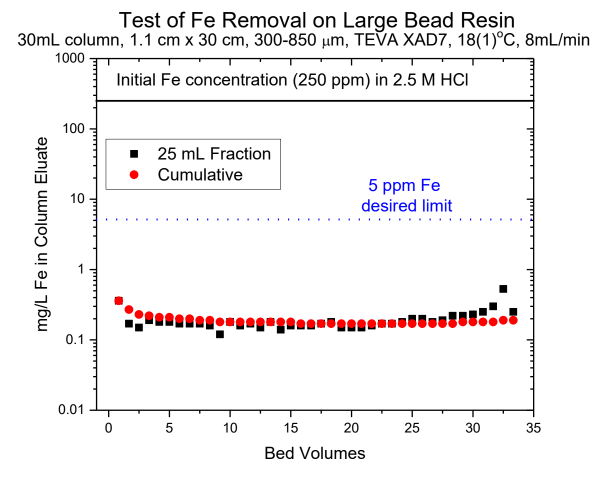
# Equipment

Empty Columns 2 mL snap tip (AC-141-AL) 2 mL cap tip (AC-100-MT-PP) 5 mL (AC-50E-5M) 20 mL (AC-20E-20M) Column Reservoir For 2 mL columns (AC-120-TK) 250 mL For 5 and 20 mL columns (AC-20X-20M) Column Rack 15 hole for 2 mL columns (AC-103)

12 hole for 5 and 20 mL columns (AC-20M-RACK)

50 mL Centrifuge Tubes





#### References

1) D.R. McAlister and E.P. Horwitz, unpublished data (2013).

eichrom\* 1955 University Lane Lisle, IL 60532, USA +1 (630)963-0320 www.eichrom.com

# **Tc Separation on WBEC Resin**

#### AN-1813-10

**Summary of Method** Pertechnetate, Tc(VII), is removed from dilute acid solution with WBEC Resin. The WBEC resin contains a tertiary amine exractant (Alamine 336) on an inert polymeric support. The Alamine 336 acts as an anion exchanger when protonated in dilute acidic media. However, from basic media, the Alamine 336 is deprotanated and no longer acts as an anion exchanger. This behavior allows anions, such as pertechnetate to be efficiently stripped from the WBEC resin using 1M NH<sub>4</sub>OH, whereas a quaternary amine, such as Aliquat 336 (TEVA) will continue to act as an anion exchanger from basic media and requires 8-10 M HNO<sub>3</sub> to strip pertechnetate.

#### Reagents

WBEC Bulk Resin (WB-B25-S) Nitric Acid (70%) Hydrogen Peroxide (30% H<sub>2</sub>O<sub>2</sub>) Ammonium Hydroxide (56%)

# Tc Separation on WBEC Resin

(1) Add 1-2 mL 30%  $H_2O_2$  per 100 mL of sample to

ensure Tc(VII). Adjust to 0.01M HNO<sub>3</sub>. Mix well.

(2) Precondition WBEC column with

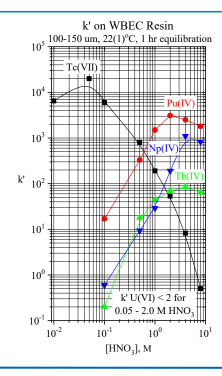
3 bed volumes of 0.01M HNO<sub>3</sub>.



(3) Load Sample.

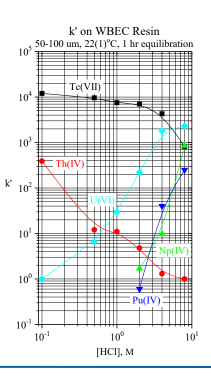
(4) Rinse column with 10 bed volumes of 0.01M HNO<sub>3</sub>.

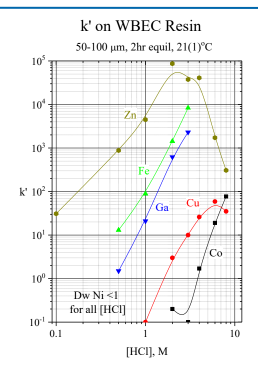
(5) Strip Tc with 5 bed volumes of 1M NH<sub>4</sub>OH.

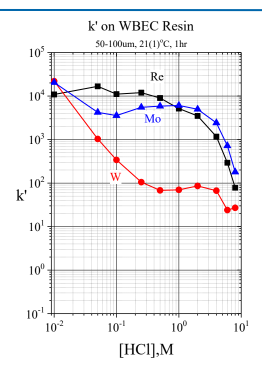


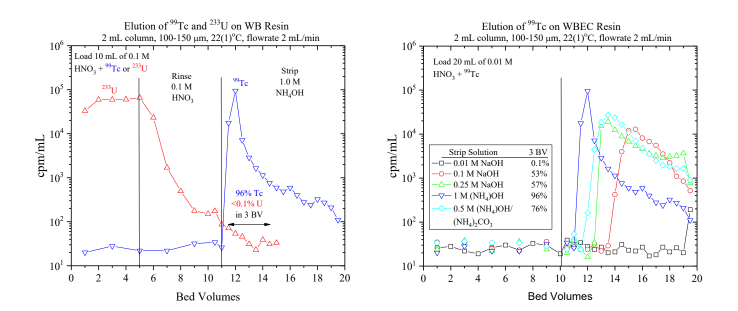
### Equipment

Empty Columns 2 mL snap tip (AC-141-AL) 2 mL cap tip (AC-100-MT-PP) 5 mL (AC-50E-5M) 20 mL (AC-20E-20M) Column Reservoir For 2 mL columns (AC-120-TK) 250 mL For 5 and 20 mL columns (AC-20X-20M) Column Rack 15 hole for 2 mL columns (AC-103) 12 hole for 5 and 20 mL columns (AC-20M-RACK) 50 mL Centrifuge Tubes









#### References

G.D. Jarvinen, K.M. Long, G.S. Goff, W.H. Runde, E.J. Mausolf, K.R. Czerwinski, F. Poineau, D.R. McAlister, E.P. Horwitz, "Separation of Pertechnetate from Uranium in a Simulated UREX Processing Solution Using Anion Exchange Extraction Chromatography," Solv. Extr. Ion Exch., 31, 416-429, (2013).

eichrom

1955 University Lane Lisle, IL 60532, USA +1 (630)963-0320 www.e