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THORIUM, PLUTONIUM, AND URANIUM IN WATER

(WITH VACUUM BOX SYSTEM)

1. SCOPE

- 1.1. This is a method for the separation of thorium, plutonium and uranium in water. After completing this method, source preparation for measurement of thorium, plutonium and uranium by alpha spectrometry is performed by electrolytic deposition onto stainless steel planchets (Eichrom Method SPA02) or by rare earth fluoride micro precipitation onto polypropylene filters (Eichrom Method SPA01).
- 1.2. This method does not address all aspects of safety, quality control, calibration or instrument set-up. However, enough detail is given for a trained radiochemist to achieve accurate and precise results for the analysis of the analyte(s) from the appropriate matrix, when incorporating the appropriate agency or laboratory safety, quality and laboratory control standards.

2. SUMMARY OF METHOD

2.1. Thorium, plutonium and uranium are separated by Eichrom TEVA and UTEVA resins prior to measurement by alpha spectrometry. A calcium phosphate precipitation can be used to concentrate actinides from water samples. Tracers are used to monitor chemical recoveries and correct results to improve precision and accuracy.

3. SIGNIFICANCE OF USE

3.1. This is a rapid, reliable method for measurement of actinides in water samples that is more cost-effective and efficient than traditional ion exchange, solvent extraction and precipitation techniques.

4. INTERFERENCES

4.1. Nuclides with unresolvable alpha energies such as ²⁴¹Am and ²³⁸Pu, ²³⁷Np and ²³⁴U, or ²³²U and ²¹⁰Po must be chemically

Method No: ACW13VBS Revision: 1.4 Page 1 of 1 2 separated to enable measurement. This method separates these isotopes effectively.

- 4.2. The ²³²U tracer should be cleaned prior to use in this method (Eichrom Method TP01) to avoid false positive measurement of ²²⁸Th in the thorium fraction.
- 4.3. Neptunium will be in the Pu fraction if present. Since the energies of ²⁴²Pu and ²³⁷Np are overlapping, ²³⁶Pu tracer maybe used to achieve a cleaner separation of neptunium and plutonium peaks.
- 4.4. Very high levels of phosphate in the sample can lead to reduced recovery of actinides in calcium phosphate precipitation and column separations. Adjusting the amount of phosphate added to coprecipitate the actinides may be necessary in these cases.
- 4.5. The sample preparation procedure outlined in this method will adequately recover actinides from freshly collected, well preserved, homogenous water samples. Older, poorly preserved samples or samples with significant organic or solid matter may require more aggressive treatment to recover actinides which have precipitated or adsorbed to the walls of the storage container or solid matter. Rinsing the empty storage container with warm HNO₃, adjusting the HNO₃ concentration of the sample to 1M HNO₃ and boiling, and/or wet-ashing the calcium phosphate precipitate may be required for older, poorly preserved samples.
- 4.6. This method may also be applied to TEVA-TRU separation chemistry, as described in Eichrom Application Note AN-1413.

5. APPARATUS

- Analytical balance, 0.0001 g sensitivity
- Beakers, glass
- Cartridge reservoirs, 10 mL (Eichrom Part: AR-200-RV10) and 20mL (Eichrom Part: AR-200-RV20)
- Centrifuge tubes, 50mL and 250mL
- Centrifuge, with rotor and carriers for 50mL and 250mL tubes
- Fume hood
- Hotplate
- Inner support tube, Eichrom Part: AR-1000-TUBE-PE
- Stir rods, glass
- Vacuum Box System, Eichrom Part: AR-12-BOX or AR-24-BOX
- Vacuum box yellow outer tips, Eichrom Part: AR-1000-OT

- Vacuum Pump, dry pump, 115 V, 60 Hz Fisher Part: 01-092-25 or house vacuum
- Vortex mixer

6. REAGENTS

Note: Analytical grade or ACS grade reagents are recommended. Evaluation of key reagents, such as aluminum nitrate and ammonium hydrogen phosphate, for contribution to method background levels from naturally occurring radioactive materials is recommended.

	1-Octanol, C ₈ H ₁₇ OH
	Aluminum nitrate nonahydrate, AI(NO ₃) ₃ .9H ₂ O
	Ammonium hydrogen phosphate, (NH ₄) ₂ HPO ₄
	Ammonium hydroxide(57% NH ₄ OH or 28% NH ₃), concentrated NH ₄ OH
	Ammonium oxalate monohydrate, (NH ₄) ₂ C ₂ O ₄ ·H ₂ O
	Ammonium thiocyanate, NH₄SCN
	Appropriate tracers or standards (Th-229, U-232, Am-243, Pu-242 or Pu-236)
	Ascorbic acid powder, $C_6H_8O_6$
	Calcium nitrate, CaNO ₃
	Deionized water, All reagents are prepared with deionized water
	Ferric nitrate nonahydrate, Fe(NO ₃) ₃ ·9H ₂ O
	Hydrochloric acid (37%), concentrated HCl
	Hydrofluoric acid (49%), concentrated HF
	Hydrogen Peroxide (30%), concentrated H_2O_2
	Nitric acid (70%), concentrated HNO ₃
	Oxalic acid dihydrate, $H_2C_2O_4 \cdot 2H_2O$
,	Sodium nitrite, NaNO ₂
,	Sulfamic acid, H_3NSO_3
	<i>TEVA[®] resin,</i> 2mL prepacked cartridge <i>,</i> 50-100μm, Eichrom Part TE-R50-S
	Titanium (III) chloride, 10wt% TiCl₃ in 20-30wt% HCl
	<i>UTEVA[®] resin,</i> 2mL prepacked cartridge, 50-150μm, Eichrom Part UT-R50-S
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6.1. Ascorbic Acid (1M) - Dissolve 17.6g of ascorbic acid in 75mL of water. Dilute to 100 mL with water. **Prepare Fresh Weekly.**

- 6.2. Ammonium hydrogen phosphate (3.2M) Dissolve 104g of (NH₄)₂HPO₄ in 200mL of water. Heat gently to dissolve. Dilute to 250mL with water.
- 6.3. Calcium nitrate (1.25M) Dissolve 51g of Ca(NO₃)₂ in 100mL of water. Dilute to 250mL with water.
- 6.4. *Hydrochloric acid (1M)* Add 83mL of concentrated HCl to 900mL of water. Dilute to 1L with water.
- 6.5. Hydrochloric acid (5M) oxalic acid (0.05M) solution Dissolve
 6.3g oxalic acid dihydrate in 400mL water. Add 417mL
 concentrated HCI. Dilute to 1L with water.
- 6.6. *Hydrochloric acid (9M)* Add 750mL of concentrated HCI to 100mL of water. Dilute to 1L with water.
- 6.7. Nitric acid (0.05M) Hydrofluoric acid (0.05M) titanium chloride (0.02M) - Add 3.2mL of concentrated HNO₃, 1.8mL of concentrated HF and 30.8mL of 10% TiCl₃ in 800mL of water. Dilute to 1L with water
- 6.8. Nitric acid (3M)- Aluminum nitrate (1M) solution- Dissolve 375g of Al(NO₃)₃·9H₂O in 500mL of water, add 188mL of concentrated HNO_{3.} Dilute to 1L with water.
- 6.9. *Nitric acid solution (3M)* Add 188mL of concentrated HNO₃ to 700mL of water. Dilute to 1L with water.
- 6.10. Sodium nitrite (3M) solution- Dissolve 5.2g of NaNO₂ in 20mL of water. Dilute to 25mL with water. **Prepare fresh daily**.
- 6.11. Ferric Nitrate Solution (5 mg/mL Fe) in 0.1M HNO₃ To a 500mL volumetric flask, add 18g Fe(NO₃)₃·9H₂O, 400mL of water and 3.1mL of concentrated HNO₃. Swirl to dissolve. Dilute to 500mL with water.
- 6.12. Sulfamic acid (1.5M)- In a 500mL volumetric flask, add 73g of sulfamic acid to 400mL of water. Swirl to dissolve. Dilute to 500mL with water.
- 6.13. *Phenolphthalein solution-* dissolve 1g phenolphthalein in 10 mL 95% isopropyl alcohol. Dilute with 100mL of water.

7. PROCEDURE

7.1. Water Sample Preparation:

- 7.1.1. If required, filter the sample through a 0.45 micron filter.
- 7.1.2. If samples larger than 1L are analyzed, evaporate the sample to approximately 1L.
- 7.1.3. Aliquot 500 to 1000mL of the sample (or enough to meet required detection limit) into an appropriate size beaker.
- 7.1.4. Add 5mL of concentrated HNO_3 .

Note: If using self-cleaning ²³²U tracer (Eichrom Method TP01), vortex mix and centrifuge standard to ensure that ²²⁸Th and its daughters are effectively removed from ²³²U by the BaSO₄ precipitate.

- 7.1.5. Add appropriate tracers per lab protocol.
- 7.1.6. Calcium phosphate precipitation:
 - 7.1.6.1. Add 0.5mL of 1.25M $Ca(NO_3)_2$ to each sample.
 - 7.1.6.2. Place each beaker on a hot plate.
 - 7.1.6.3. Cover each beaker with a watch glass.
 - 7.1.6.4. Heat the samples at medium setting for 30-60 minutes.
 - 7.1.6.5. Remove the watch glass and turn the heat down.
 - 7.1.6.6. Add 2-3 drops of phenolphthalein indicator and 1mL of $3.2M (NH_4)_2HPO_4$ solution.
 - 7.1.6.7. While stirring, slowly add enough concentrated NH₄OH to reach the phenolphthalein end point and form a calcium phosphate precipitate. Heat the sample for another 20-30 minutes.
 - 7.1.6.8. Remove samples from the hot plate, cool to room temperature, and allow precipitate to settle until solution can be decanted (30 minutes to 2 hours) or centrifuge.
 - 7.1.6.9. Decant supernate and discard to waste.
 - 7.1.6.10. Transfer the precipitate to a centrifuge tube and centrifuge the precipitate for approximately 10 minutes at 2000 rpm.
 - 7.1.6.11. Decant supernate and discard to waste.



- 7.1.6.12. Wash the precipitate with an amount of water approximately twice the volume of the precipitate. Mix well on a vortex mixer. Centrifuge for 5-10 minutes. Discard the supernate.
- 7.1.6.13. If an ammonia odor persists repeat 7.1.6.12.
- 7.1.6.14. Dissolve precipitates with 5mL conc. HNO₃ and transfer to a 100mL glass beaker. Rinse centrifuge tube with 2-3mL conc. HNO₃ and transfer to the same 100mL beaker. Evaporate to dryness.
- 7.1.6.15. Dissolve each residue with 10mL of 3 M HNO_3 1.0 M $$\rm Al(NO_3)_3.$$

Note: An additional 5-10mL may be necessary if the precipitate volume is large.

Note: Make sure that all reagents and the load solution have cooled to room temperature. Warm solutions can cause reactions that will affect oxidation adjustments performed in the following steps.

Note: Pu must be present in Pu (IV) to be retained on TEVA resin. The following steps will ensure that Pu (III) and Pu (VI) are converted to Pu (IV).

7.1.6.16. Add 1mL of 1.5M sulfamic acid, 0.5mL of 5mg/mL Fe solution, and 1mL of 1M ascorbic acid to each sample. Swirl to mix and wait 5 minutes.

Note: If the additional 5-10mL was used to dissolve the sample in step 7.1.6.15, add a proportionately larger amount of sulfamic acid and ascorbic acid.

7.1.6.17. Add 1mL of 3M NaNO₂. Mix well and wait for 5 minutes.

Note: NaNO₂ is added to oxidize Pu(III) to Pu(IV).

- 7.1.7. Set up of TEVA and UTEVA cartridges in tandem on the vacuum box system
 - 7.1.7.1. Place the inner tube rack into the vacuum box with the centrifuge tubes in the rack. Fit the lid to the vacuum system box.
 - 7.1.7.2. Place the yellow outer tips into all 12 or 24 openings of the lid of the vacuum box. Fit an inner support tube into each yellow tip.

- 7.1.7.3. For each sample solution, fit a UTEVA cartridge on to the inner support tube. Attach TEVA cartridge to the top end of the UTEVA cartridge.
- 7.1.7.4. Attach syringe barrels (funnels/reservoirs) to the top end of the TEVA cartridge.

Note: The unused openings on the vacuum box should be sealed. Yellow caps can be used to plug unused white tips to achieve good seal during the separation. Alternatively, unused vacuum box holes can be sealed with scotch tape affixed to the vacuum box lid.

- 7.1.7.5. Connect the vacuum pump to the box. Turn the vacuum pump on and ensure proper fitting of the lid.
- 7.1.7.6. Add 5mL of 3M HNO₃ to each reservoir to precondition the TEVA and UTEVA cartridges. Adjust the vacuum pressure to achieve a flow-rate of 1.0 ml/minute.
- 7.1.8. Th and Pu separation from U using TEVA and UTEVA Resin
 - 7.1.8.1. Transfer each solution from step 7.1.6.17. into the appropriate reservoir. Allow solution to pass through both the cartridges at a flow rate of 1.0 mL/minute.
 - 7.1.8.2. Add 5mL of 3M HNO_3 to rinse to each sample beaker and transfer each solution into the appropriate reservoir. (the flow rate can be adjusted to 2mL/minute).
 - 7.1.8.3. Add 15mL of 3M HNO_3 into each reservoir. Allow liquid to pass through cartridges at 2mL/minute.
 - 7.1.8.4. Disengage vacuum. Separate TEVA cartridge from UTEVA cartridge. Place a new set of reservoirs on the UTEVA cartridges. Empty liquid from centrifuge tubes below each column (dispose as waste). Set TEVA cartridges aside, while Uranium is separated from UTEVA. Replace centrifuge tubes below each cartridge.
 - 7.1.8.5. Add 5mL of 3M HNO $_3$ into each UTEVA cartridge. Engage vacuum and allow liquid to pass through cartridges at 2mL/min.
 - 7.1.8.6. Add 5mL of 9M HCl into each UTEVA cartridge. Allow liquid to pass through cartridges at 2mL/min.
 - 7.1.8.7. Add 20mL of 5M HCI- 0.05M oxalic acid into each UTEVA cartridge. Allow liquid to pass through cartridges at 2mL/min.



Note: The rinses remove any residual neptunium, thorium and americium if present on UTEVA. Also removes any residual ferrous ion that might interfere with electrodeposition. ²¹⁰Po will have been removed by the TEVA cartridge.

- 7.1.8.8. Disengage the vacuum and place a clean, labeled tube below each cartridge. Replacing outer yellow vacuum box tips and inner tube supports at this time will help ensure clean uranium fractions in the following step.
- 7.1.8.9. Add 15mL of 1M HCl into each cartridge reservoir to strip the uranium from UTEVA. Engage vacuum and strip uranium at 1mL/min.
- 7.1.8.10. Set U samples aside for alpha source preparation.
- 7.1.9. Th, Pu Separation Using TEVA cartridge:
 - 7.1.9.1. Place TEVA cartridges along clean reservoirs on the appropriate openings on the vacuum box lid.
 - 7.1.9.2. Add 15mL of 3M HNO_3 to each cartridge reservoir. Allow solution to pass through cartridges at 2mL/min.
 - 7.1.9.3. Ensure that clean, labeled tubes are placed in the tube rack. Replacing outer yellow vacuum box tips and inner white vacuum box tips at this time will help ensure clean thorium fractions in the following step.
 - 7.1.9.4. Add 15mL of 9M HCl to each cartridge. Elute thorium at 1mL/min.
 - 7.1.9.5. Disengage vacuum and remove centrifuge tubes containing Th. Set Th samples aside for alpha source preparation.
 - 7.1.9.6. Ensure that clean, labeled tubes are placed in the tube rack under each cartridge. Replacing outer yellow vacuum box tips and inner support tubes at this time will help ensure clean plutonium fractions in the following step.
 - 7.1.9.7. Add 25mL of 0.05M HNO $_3$ /0.05M HF/0.02M TiCl $_3$ to each reservoir. Engage vacuum and elute plutonium at 1mL/min.



Note: Electrodeposition CANNOT be carried out on the Pu fraction because of $TiCl_3$ in the strip solution. If electrodeposition is desired then another suitable stripping agent may be used, such as 0.1M ammonium bioxalate.

7.1.9.8. Set Pu samples aside for alpha source preparation.

7.2. Prepare samples for the determination of actinides by alpha spectrometry using electrodeposition (Eichrom SPA02) or rare earth fluoride micro precipitation (Eichrom SPA01).

8. CALCULATIONS

Calculate the actinide activity as follows:

Calculate tracer yield:

$$\text{Yield} = \frac{\left(C_{s} - B_{s}\right)}{E_{s} \times A_{s}}$$

where:

C_s =measured actinide tracer, cpm

B_s =background, cpm

E_s =counting efficiency for tracer

A_s =tracer activity, dpm

Note: If any tracer may be present in the sample, a spiked and unspiked sample must be analyzed to determine chemical yield, where:

Yield = $\frac{\text{(spiked sample tracer cpm - unspiked sample tracer cpm)}}{\text{E} \times \text{actinide spike activity dpm}}$

Percent yield = Yield x 100

Calculate actinide isotope activity:

Sample dpm / g or dpm / L =
$$\frac{S - B}{E \times V \times Y}$$

where:

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- S = sample activity, cpm
- B = background, cpm
- E = counting efficiency = measured cpm/dpm of isotopic standard
- V = sample weight, g or volume, L
- Y = yield

Conversion of dpm/g to pCi/g: pCi/g = (dpm/g)/2.22

9. PERFORMANCE DATA

9.1. Chemical recoveries of tracers in various matrixes were measured on 1 liter samples.

	Spike*	DI Water	Tap Water
Th-229	97.2 ± 11%	$87.2\pm7\%$	102 ± 4%
Pu-242	99.7 ± 2%	90.8 ±6%	103 ± 5%
U-232	91.0±6%	80.3 ± 3%	87.2 ± 7%

*Load solution spiked with tracers. No precipitation carried out.

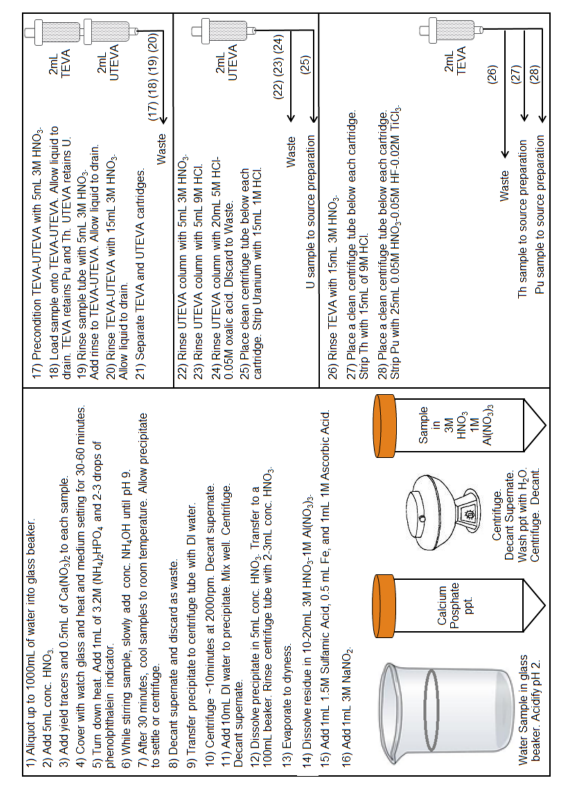
9.2. Decontamination factors were measured for the actinide elements shown below. The decontamination factors reported were calculated by dividing the activity spiked by the activity found in the fraction. Where the activity found was below the detection limit, the value is reported as ">."

	In Th	In Pu	In U fraction			
	fraction	Fraction				
U-232	>800	13,000	NA			
Pu-242	>1500	NA	20,000			
Am-241	>800	21,000	>2200			
Th-229	NA	>3600	>2000			
Np-237	>3000	NA	4600			

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10.REFERENCES

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