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Nuclear fuel technology — Chemical separation and purification of uranium and plutonium in nitric acid solutions for isotopic and isotopic dilution analysis by solvent extraction chromatography —

Part 2:

Samples containing plutonium and uranium in the nanogram range and below

Technologie du combustible nucléaire — Séparation et purification chimiques de l'uranium et du plutonium dans les solutions d'acide nitrique par extraction chromatographique par solvant pour les mesures isotopiques et les analyses par dilution isotopique https://standards.ite.h.a.ne.s.da.p.g.p.h

Partie 2: Échantillons ayant des teneurs en plutonium et en uranium de l'ordre du nanogramme et inférieures

[Revision of first edition (ISO 15366:1999)]

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Foreword

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ISO 15366 was prepared by Technical Committee ISO/TC 85, Nuclear Energy, Subcommittee SC 5, Nuclear Fuel Technology.

This first edition cancels and replaces the first edition of ISO 15366:1999.

Nuclear fuel technology — Chemical separation and purification of uranium and plutonium in nitric acid solutions for isotopic and isotopic dilution analysis by solvent extraction chromatography —

Part 2:

Samples containing plutonium and uranium in the nanogram range and below

SCOPE 1

This International Standard describes procedures to chemically separate and purify uranium and plutonium in dissolved solutions of irradiated light water reactor fuels and in samples of High Active Liquid Waste of spent fuel reprocessing plants, prior to their isotopic analysis by e.g. mass spectrometric method (ISO 8299) or alpha spectrometry (ISO 11483). This paper consists principally of two parts.

This part 2 describes a slightly different separation technique from part 1, based on the same chemistry, using smaller columns, different support material and special purification steps, applicable to samples containing plutonium and uranium amounts in the nanogram range and below. The detection limits were found to be 500 pg plutonium and 500 pg uranium. Annex describe the preparation of the columns and the column support materials.

Comparing with ISO CD 15366 part 1, as uranium and plutonium amounts are lowest, additional purification . ik Idardsite on an anion exchange resin is performed.

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PRINCIPLE OF THE METHOD Ambandation The chemical separation of small plutonium and uranium amounts (> 500 pg) is also based on a column extraction chromatography using tri-n-octylphosphine-oxide (TOPO) as extractant [1,2]. The necessary valency adjustment prior to the separation is done with Iron(II) sulphate and sodium nitrite. The extraction process is performed in disposable mini-columns loaded with a polyethylene or silica gel powder coated with the TOPO. Plutonium (IV) and uranium (VI) in 3 mol.¹⁻¹ nitric acid medium are selectively extracted into the TOPO while americium, the fission products and other interfering elements are not retained. Plutonium is eluted after reduction to the trivalent state with ascorbic acid [5]; uranium is eluted by an ammonium carbamate solution and finally purified from ammonium salts by an ion exchange separation [4] or can be fuming by 14 mol.l-¹nitric acid.

In order to ensure a favourable kinetics of chemical reactions the (gravity) column flow rates should not exceed 0,4 ml.min⁻¹.

Parallel measurement of blank and/or control sample is recommended to verify the analysis.

Blanks are run in parallel with the samples to verify the absence of significant external cross contamination or cross contamination between samples.

Control samples prepared from certified or analysed materials are also prepared and separated along with the sample to verify that suitable valency adjustment, isotopic equilibration and separation efficiency are achieved.

The whole process is carried out under clean conditions preferably in a laminar flow fume hood.

3 APPARATUS

3.1 Biological shielding, e.g. shielded glove box or fume cupboard, preferably a laminar flow fume cupboard.

3.2 Disposable polypropylene columns and frits, (see Figure A.1 in annex). Type: Micro-column. The packing and conditioning of the columns are described in the Annex. The chromatographic columns are to be disposed of in the radioactive waste after use.

3.3 Two hot plates.

3.4 Standard laboratory equipment (flasks and beakers, pipettes, glassware, stands and supports for columns, sample vials, fraction tubes, etc.). It is recommended to leach all equipment, which will be in contact with samples, eluates or eluents with nitric acid (4.1) and distilled water before use. After drying store it in plastic bags in a clean area.

3.5 PTFE-vials ¹⁾, dimensions: volume 15 ml, diameter 28 mm, height 37 mm. With screw caps and metal jackets.

These vials can be used at least 10 times, after a careful cleaning with hot nitric acid followed by a rinsing with demineralised or de-ionised water.

4 REAGENTS

Use only reagents of recognised suprapure grade or equivalent. All aqueous solutions shall be prepared with distilled water (resistivity 1 M Ω .m to 10 M Ω .m).

4.1 Concentrated nitric acid and nitric acid solutions, $c(HNO_3) \neq 6$ mol.¹ and 3 mol.¹.

4.2 Formic acid solution, c(HCOOH) = 1 mol

- 4.3 Ammonium carbamate [CAS No. 1111-78-0] solution, $c(NH_4CO_2NH_2) = 0.7 \text{ mol.}l^{-1}$.
- 4.4 Polyethylene powder²⁾ Hostalen® GUR X 1777 PE-UHMW Ultra-High-Molecular³⁾

grain size 25-95 μ m, pore size \leq 5 μ m (see Annex B, 2.5).

4.5 L(+)-Ascorbic acid solution $c(C_6H_8O_6) = 1 \times 10^{-5} \text{ mol.}\text{I}^{-1} \text{ or } c(C_6H_8O_6) = 1 \times 10^{-3} \text{ mol.}\text{I}^{-1}$ in formic acid solution (4.2)

4.6 Tri-n-octylphosphine-oxide, $c(TOPO) = 0.2 \text{ mol.I}^{-1}$ solution in cyclohexane.

Warning: this reagent is flammable and should always be handled in a well ventilated place and never in the vicinity of a naked flame.

4.7 Iron(II) sulphate solution, $c(FeSO_4) = 0,1 \text{ mol.}^{-1}$ or $c(FeSO_4) = 0,8 \text{ mol.}^{-1}$ in water. To be prepared freshly for each working session.

4.8 Sodium nitrite solution, $c(NaNO_2)=1 \text{ mol.}^{-1}$ or $c(NaNO_2)=5 \text{ mol.}^{-1}$ in water. To be prepared freshly for each working session.

¹⁾ Manufacturer can be : Savillex Corporation, 6133 Boker Road, Minnetonka, MN 55345-5910, USA, Art. no. 025R-SB.

²⁾ Silical gel may be used to. Refer to part 1.

³⁾ Manufacturer can be Hoechst, Werk Ruhrchemie

4.9 Aqua regia; mixture 3:1 (per volume) of concentrated hydrochloric acid and concentrated nitric acid.

4.10 Concentrated hydrochloric acid and hydrochloric acid solution $c(HCI) = 6 \text{ mol.}I^{-1}$

4.11 Anion exchange resin AG MP1, slurry in distilled water. Fill the mini-columns (10.2) with ca. 0,5 ml of the slurry. Condition the columns with two times 1 ml of distilled water and with three times 1 ml concentrated hydrochloric acid (11.10) immediately before use.

5 PROCEDURE ⁴⁾(Figure 5-1)

5.1 The sample should contain 500 pg to 1 μ g plutonium and 500 pg to 100 μ g uranium in a volume of 0,5 ml of 3 mol.I⁻¹ nitric acid solution (4.1). Whenever starting with dried samples apply the following dissolution procedure:

(a) Add 0,5 ml of 6 mol. Γ^1 nitric acid solution (4.1) to the dry samples and evaporate slowly on the hot plate, keeping the temperature slightly below the boiling point to avoid any splashing and bubbling until nitrate salts crystallise.

(b) Remove the sample vessels from the hot plate and redissolve the salts by adding 0,5 ml of 3 mol.¹ nitric acid solution (4.1), while still warm (40 \degree to 60 \degree). Shake the vessels for a few seconds.

5.2 Perform a redox valency cycle to ensure that all plutonium isotopes are in the tetravalent state before starting the separation, as follows:

(a) Add 50 µl of Iron(II) sulphate solution (47) to the sample

(b) Mix and wait for five minutes for a complete reduction of plutonium (VI) or plutonium (IV) to plutonium (III).

(c) Add 50 μ I sodium nitrite solution (4.8) to reoxidize plutonium to the tetravalent state and add further 100 μ I of 6 mol. Γ^1 nitric acid solution (4.1) to reach an acid concentration of 3 mol. Γ^1 . Mix again and wait for at least five minutes.

5.3 Transfer half of the pre-treated sample onto the column, wait approximately one minute, add the rest of the sample and let it flow through. This favours the retention of plutonium and uranium in the very upper layers of the column.

5.4 Rinse the sample vial with 0,5 ml of 3 mol. Γ^1 nitric acid solution (4.1) and transfer the solution to the column.

5.5 Wash out the fission products, including americium, from the column using 3 mol. Γ^1 nitric acid solution (4.1) in 4 successive aliquots of 1 ml.

5.6 Condition the column for the plutonium elution by adding twice $0,5 \text{ ml of } 1 \text{ mol.}^{-1}$ formic acid solution (4.2). Discard waste collected until now.

5.7 Elute the plutonium from the column with the ascorbic acid solution (4.5) with five successive aliquots of 1 ml. Place the vials containing the collected plutonium fractions on one of the hot plates.

5.8 Wash out the 'tail' of the plutonium with twice 1 ml of the ascorbic acid solution (4.5) in one aliquot and discard the plutonium 'tail' washings to the waste.

5.9 Condition the column for the elution of the uranium fraction by adding two aliquots each of 1 ml distilled or de-ionised water. Discard the water washings to the waste.

⁴⁾ This procedure is an example. If equivalent results could be expected, other conditions than these described in clause 5, can be applied for sample preparation.

5.10 Elute the uranium with 4 ml ammonium carbamate solution (4.3). Place the vials containing the collected uranium fractions on the other hot plate.

5.11 Let the plutonium and uranium fractions evaporate gently to dryness on the hot plates at 90 °C.

5.12 Remove the fractions from the hot plates, add 0,25 ml of concentrated nitric acid (4.1) and evaporate again to dryness. Repeat this step once. Proceed with the uranium fractions with clause. 5.17 if necessary.

5.13 Redissolve the plutonium fractions in 0,5 ml of 3 mol. Γ^1 nitric acid solution (4.1) and transfer the solutions to the PTFE-vials (3.5).

5.14 Rinse the vials of the plutonium fraction with 0,5 ml of 3 mol.l^{-1} nitric acid solution (4.1) and transfer the solutions to the corresponding PTFE-vials.

5.15 Place all PTFE-vials with the plutonium fractions on the hot plate, preferably in metal jackets for an even heating. Evaporate the solutions gently to dryness at 90 \degree avoiding any bubbling and splashing.

5.16 After cooling down to room temperature secure the cap of the PTFE-vials with the plutonium fractions and forward them for mass spectrometric measurement. The mass spectrometer operator will redissolve the residues in a minimum (2 μ l and less) of 3 mol. Γ^1 nitric acid solution immediately before loading the solutions onto the mass spectrometer. filaments. Performing this redissolution step by means of a microscope is recommended.

5.17 Redissolve all uranium fractions with 250 µl aqua regia (4.9) and evaporate to dryness.

5.18 Redissolve all uranium fractions with 250 µl concentrated hydrochloric acid (4.10) and evaporate to dryness.

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5.19 Redissolve all uranium fractions with 1 ml concentrated hydrochloric acid (4.10) and transfer slowly all solutions to the conditioned anion exchange resin columns (4.11).

5.20 Rinse all uranium fraction vials with 1 mic concentrated hydrochloric acid (4.10) and transfer slowly all solutions to the corresponding columns.

5.21 After the solutions (5.20) have completely passed the columns, wash each column with 0,5 ml of hydrochloride acid solution 6 mol. l^{-1} (4.1).

5.22 Elute the uranium into PTFE-vials (3.5) with four successive aliquots of 1 ml distilled water.

5.23 Place all PTFE-vials with the uranium fractions on the other hot plate, preferably in metal jackets for an even heating. Evaporate the solutions gently to dryness at 90 \degree avoiding any bubbling and splashing.

5.24 Remove the uranium fractions from the hot plates, add 0,25 ml of concentrated nitric acid (4.1) and evaporate again to dryness in the same way as in 5.23. Repeat this step once.

5.25After cooling down to room temperature secure the cap of the PTFE-vials with the uranium fractions and forward them for mass spectrometric measurement. The mass spectrometer operator will redissolve the residues in a minimum (2 μ I and less) of 3 mol. Γ^1 nitric acid solution immediately before loading the solutions onto the mass spectrometer. filaments. Performing this redissolution step by means of a microscope is recommended.



