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Determination of isotopic content and concentration of uranium and plutonium in nitric acid solution — Mass spectrometric

iTeh Methodard PREVIEW

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Détermination de la teneur isotopique et chimique en uranium et plutonium <u>Id/une_solution</u> d'acide nitrique — Méthode par spectrométrie https://standards/dehmassepg/standards/sist/67faf975-c150-477b-834ac5c5e456c45a/iso-8299-1993



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https://standards.iteh.ai/catalog/standards/sist/67faf975-c150-477b-834a-Annex A forms an integral part of this International Standard so-8299-1993

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International Organization for Standardization

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Determination of isotopic content and concentration of uranium and plutonium in nitric acid solution — Mass spectrometric method

1 Scope

This International Standard specifies a method to determine the isotopic content and the concentration of uranium and plutonium in input solutions of irradiated fuels in light water reactors (boiling water or pressurized water) and in final products at spent fuel reprocessing plants. The method is applicable to other fuels, but the chemical separation and spike solution shall, if necessary, be adapted to suit each type of fuel. separated isotopes (e.g. ²⁴⁴Pu, ²³⁹Pu, ²³⁵U and ²³⁶U) can be used when available and appropriate.

The isotopic composition of the spikes and the isotopic abundance of the ²³⁸U and ²³⁹Pu isotopes in the sample shall be accurately known. If the ²⁴²Pu and ²³⁶U isotopes are used as spikes, it is also necessary to measure the isotopic abundance of these isotopes in the sample.

Tards. A chemical separation is necessary to eliminate interfering elements (e.g. ²⁴¹Am, fission products) following a reduction-oxidation cycle to ensure isotopic ISO 8299:199 exchange between spikes and sample.

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2 Principle of the method

The method is based on isotope ratio measurements by thermal ionization mass spectrometry.

The isotopic composition of uranium and plutonium is determined through isotopic ratio measurements.

 $^{238}\mathrm{Pu}$ is determined by means of alpha spectrometry if interferences from $^{238}\mathrm{U}$ cannot be eliminated.

The sample is diluted in order to economize on the amount of spike needed and to minimize the level of biological protection which is necessary.

A highly accurate gravimetric dilution is necessary when elemental concentrations are measured.

Because plutonium tends to hydrolyse at low acidities, a solution of nitric acid (HNO_3) with a molarity greater than 1 mol/l should be used for the dilution of the sample.

When elemental concentrations are measured, accurately weighed quantities of spike isotopes are added in quantities comparable to the quantities of ²³⁸U and ²³⁹Pu isotopes in the diluted sample. Although ²³³U and ²⁴²Pu isotopes are normally used as spikes, other

c5c5c456c45a/iso-82Thel isotopic composition of fresh product material is normally measured without purification.

Plutonium aged more than 1 week should be purified to remove ²⁴¹Am.

The method includes the following steps:

- a) dilution by weight;
- b) spike addition by weight;
- c) isotope exchange chemistry;
- d) ion exchange purification/separation;
- e) preparation of filaments;
- f) mass spectrometric measurements and calculation of isotopic and elemental concentrations.

Care shall be taken to avoid cross contamination. For this purpose, it is recommended that disposable laboratory ware (e.g. vessels and columns) be used. Throughout the method it is necessary to follow good analytical practices regarding cleanliness, accuracy of measurement, avoiding evaporation errors, etc.

3 Reagents

3.1 Isotopic reference materials, which cover the isotopic range of interest and are certified to 0.1 % or better for the major isotope ratios, such as CBNM-047a, NBL 128, CEA MIRF, NBL 137 (ex NBS Pu947) for plutonium and CBNM-072, CBNM-199, NBL 117, NBL U500 to NBL U930 for uranium.

3.2 Natural uranium metal, of purity certified to 0,05 % or better, such as NBL 112 (ex NBS U960), EC-101 or CEA-MU-2.

3.3 Plutonium metal, of purity certified to 0,05 % or better, such as NBL 126, EC-201, CEA-MP-2 or NBS 949, with a ²³⁹Pu isotopic abundance of 90 % or more, known also to + 0,05 % or better.

3.4 ²³³U spike solution, of certified isotopic and chemical composition, such as NBL 111A, CBNM-040-1 or NBS U995.1)

3.5 Pu spike, of certified isotopic and chemical composition, such as CBNM-041-1 (²⁴²Pu solutions) or NBL 131 (ex NBS Pu996) (²⁴⁴Pu nitrate).¹⁾

CBNM-046-2 (may be used in place of 3.4, 3.5 or mixtures thereof).¹⁾

ISO 8299:1 Resolution: greater than 400 at 1 % of the peak **3.7 Concentrated nitric acid** (HNO₃), 14 mol/catalog/standards/heightalThiscresolution should be measured on the c5c5e456c45a/iso-825U-and 238U masses. ution.

3.8 Nitric acid, 7 mol/l solution.

3.9 Nitric acid, 4 mol/l solution.

3.10 Nitric acid, 3 mol/l solution.

Nitric acid, 1 mol/l solution. 3.11

3.12 Nitric acid, 0,2 mol/l solution, uranium-free.

3.13 Ferrous sulfate septahydrate (FeSO₄.7H₂O), 42,5 g/l solution in H_2SO_4 (0,05 mol/l), freshly prepared.

3.14 Sodium nitrite (NaNO₂), 250 g/l solution, freshly prepared.

3.15 Dowex AG 1 × 4 resin, 100/200 mesh (conditioned in 7 mol/l nitric acid solution).

In a 1 000 ml beaker, wash 250 ml of resin Dowex 1×4 , successively with:

- 500 ml water: twice,

- 500 ml of 0,2 mol/l nitric acid solution (3,12); twice.
- 500 ml of 4 mol/l nitric acid solution (3.9): twice,
- 500 ml of 7 mol/l nitric acid solution (3.8): five times, until a sample of the supernating solution no longer yields a chloride precipitate after additon of silver nitrate.
- Use the resin within 30 days. Beyond this time, wash the unused resin with water and discard.

A blank determination should be performed to show that the quantity of the uranium or plutonium present in the reagents or picked up during the separation procedure is less than 0,2 % of the uranium or plutonium expected to be present in a sample determination.

Apparatus and materials

4.1 Mass spectrometric equipment

3.6 Mixed ²³³U/²⁴²Pu **spike solution**, of certified thermal ionization type, designed for precise isotopic and chemical composition, such as a measurement of isotopic composition having at least the following features:

Abundance sensitivity: less than 10^{-5} for 237 U.

Vacuum: pressures less than 10^{-4} Pa in the source and less than 10^{-5} Pa in the analyser. For instance, a modern apparatus can reach 10^{-5} Pa in the source and about 10^{-6} Pa to 10^{-7} Pa in the analyser.

Ionization: thermal ionization source fitted with two or three filament assemblies with disposable filament inserts. The ionization filament (50 µm thick and 0,75 mm wide) is made of rhenium, highly purified especially with respect to the alkali elements (see clause 11) and uranium. The sample filament(s) is (are) made either of rhenium like the ionization filament or tungsten (25 µm thick). Tantalum sample filaments may also be used when plutonium is analysed.

Detection: single or multidetector assembly preferably consisting of a Faraday cup detector or detectors, a secondary electron multiplier or Daly detector for the measurement of minor isotopes and one or several suitable voltage-to-frequency

¹⁾ If certified spikes 3.4, 3.5 or 3.6 are not available, spike solutions may be prepared and standardized as described in annex A.

converter(s) and a high-precision digital measuring system for ion current integration.

4.1.2 Filament preheating and degassing device, for cleaning the mounted unloaded filaments, including if possible a small attraction plate to collect alkali ions.

4.1.3 Filament preparation device, for precise loading of the samples onto cleaned filaments and reproducible drying of the samples without crosscontamination.

4.1.4 Optical pyrometer, desirable to determine the temperature of the ionizing filament. Alternatively, the intensity of the $^{187}\mathrm{Re}^+$ ions can be carefully calibrated as a function of the temperature of the ionizing filament; the $^{187}\mathrm{Re}^+$ ion intensity can then be used to measure and control the temperature of this filament.

4.2 Instrument calibration

Use internationally accepted isotopic reference materials certified to 0,1 % or better for the ratio of the major isotopes.

Calibrate the analyses of uranium against a certified **RD PREVIEW** 1:1 mixture of ²³⁵U and ²³⁸U (e.g. NBL U500) or of **4.6 Analytical balances**, for the gravimetric of ²³³U and ²³⁸U (e.g. NBL 117, CBNM-199) and **4.6 Analytical balances**, for the gravimetric of ²³⁹U and ²⁴⁰Pu (e.g. NBL 137, ex NBS Pu947) or of ²³⁹Pu and ²⁴⁰Pu (e.g. NBL 137, ex NBS Pu947) or of ²³⁹Pu and ²⁴⁴Pu (e.g. NBL 128) or of ²³⁹Pu and ²⁴⁴Pu (e.g. NBL 128) or of ²³⁹Pu and ²⁴⁴Pu (e.g. NBL 128) or of ²³⁹Pu and ²⁴⁴Pu (e.g. 299:19**4.7 Disposable plastic pipettes and containers**.

CBNM-047a). https://standards.iteh.ai/catalog/standards/sist/67faf975-c150-477b-834a-

Treat, prepare and measure n samples of the certified reference material ($n \ge 10$) in the same way that unknown samples are analysed.

Calculate the arithmetic mean, $\overline{R}_{i/i'}$ of the *n* measurements of the ratio of the ion currents produced by isotopes i and j. Compare this mean to the certified value, R_{ill}(s), of the isotope ratio and calculate a discrimination factor, B, equal to

$$B = [(\overline{R}/R_s)_{i|j} - 1]/(M_j - M_i) \qquad \dots (1)$$

where M_i and M_i are the relative atomic masses (weights) of isotopes j and i.

The number of calibration measurements, n, is selected to ensure that the standard error on B is equal to or less than 0,000 1.

The calibration is repeated at least once a year or whenever a significant drift is detected, in accordance with 4.3, or suspected, for example after changing a major component of the instrument or the method of preparation of the samples.

4.3 Linearity of the method and stability of the calibration

Verify that the mass spectrometer and the method

of analysis give linear response and accurate results over the range of isotopes and isotope ratios to be analysed.

For this purpose, regularly treat, prepare and measure samples of certified isotopic reference materials of different isotopic composition in the same way as the unknown samples currently analysed.

Calculate the results of the analyses of the reference materials as described in clause 8.

Statistically significant differences in time or in the range of isotope ratios between the measured and the certified isotope ratios indicate a drift of the calibration or a deviation from linearity.

In such cases corrective actions should be undertaken.

4.4 Shielded cell, equipped with manipulators or tongs for carrying out the chemical preparations on highly radioactive solutions remotely.

4.5 Glove box, for handling diluted solutions for operations of spiking and subsequent chemical treatment.

4.6 Analytical balances, for the gravimetric di-

4.9 Flasks and beakers.

4.10 Disposable ion exchange column

The procedure specified in this International Standard is intended for use with a column made of a 16 mm diameter polyethylene tube, tapered at one end and filled with a 10 mm \pm 1 mm high bed of Dowex 1 x 4 resin of 100/200 mesh (3.15). A column of different dimensions may be used, but the volumes of eluents must be adapted.

5 Chemical preparation

5.1 Spike preparation

Accurately prepare either

- two separate standardized spike solutions, one containing 150 μg/g to 200 μg/g of ²³³U, the other 1 μg/g to 2 μg/g of ²⁴²Pu, or
- a single standardized spike solution of mixed spike containing 150 μ g/g to 200 μ g/g of ²³³U and 1 μ g/g to 2 μ g/g of ²⁴²Pu.

The volume to be prepared depends on the availability of the isotopes. An example of preparation and standardization procedure is described in annex A.

5.2 Dilution of the input solution sample

A typical input solution contains 250 g/l to 300 g/l of uranium and 2 g/l to 3 g/l of plutonium when the burn-up is about 30 000 MWd/t. This highly radioactive solution is handled in a hot cell where it is diluted by weight 200- to 2 000-fold with 1 mol/l nitric acid (3.11).

Carry out at least two replicate dilutions on each batch of input solution.

As an example, a typical dilution procedure may be carried out as follows.

Transfer into a tared 100 ml flask, 2 ml of the input solution A, weighed to the nearest 0,1 mg (m_1) and containing 500 mg to 600 mg of uranium and 4 mg to 6 mg of plutonium.

Dilute to 100 ml with 1 mol/l nitric acid, determine the mass, m_2 , of the diluted solution B to the nearest 10 mg and mix well.

Into another tared 100 ml flask, transfer 5 ml of solution B, weighed to the nearest 0,1 mg (m'_1) .

Dilute to 100 ml with 1 mol/l nitric acid, determine, to <u>ISO 82774056</u> solution D onto the ion exchange column, the nearest 10 mg, the mass, <u>m'2'</u>, of the diluted solution C which is obtained, and mix well. <u>c5c5e456c45a the 7 mol/l nitric acid solution</u>.

The dilution factor, F_{i} is equal to

$$F = \frac{m_2 m'_2}{m_1 m'_1}$$
 (2)

Transfer a sample of about 6 ml of solution C to a suitable facility.

5.3 Preparation for content assay

The following operations are performed at least once after each replicate dilution of the input solution in accordance with 5.2.

5.3.1 Spiking

The quantity of 233 U (and 242 Pu) added shall be comparable to the quantity of 238 U (and 239 Pu) present in solution C.

Transfer into a tared 60 ml plastic container 2 ml of the spike solution, weighed to the nearest 0,1 mg $(m_{\rm S})$.

Add 2 ml of solution C, weighed to the nearest 0,1 mg, $(m_{\rm C})$, and mix thoroughly without splashing.

Ensure that all liquid is included in the mixing by wetting the interior container walls.

5.3.2 Isotopic exchange

If there is a possibility of Pu polymer being present in the sample or in the spike, it is advisable to add a drop of 0,1 mol/l hydrofluoric acid and reflux the spiked sample and then to complex excess fluoride with Al⁺⁺⁺ before proceeding (for example, with aluminium nitrate).

A redox valency cycle is performed to ensure that all plutonium isotopes are in the tetravalent state before the ion exchange separation is done.

Add 1 ml of the ferrous sulfate solution (3.13) into the 60 ml container.

Mix and wait 5 min for a complete reduction of Pu(VI) to Pu(III) or Pu(IV).

Add 1 ml of sodium nitrite solution (3.14) to reoxidize all plutonium to the tetravalent state.

Stir for 5 min.

5.3.3 Ion exchange separation

Add 8 ml of concentrated nitric acid (3.7) into the **60 ml container** and mix to obtain 14 ml of solution D in 7 mol/l nitric acid (3.8).

Wash the column three times with 15 ml of 4 mol/l nitric acid solution (3.9). Discard the eluate which contains americium and fission products.

Wash again with 15 ml of 4 mol/l nitric acid solution. Collect the eluate D' in a glass beaker; it will be used for the uranium assay.

Wash with 15 ml of 4 mol/l nitric acid solution and discard the eluate.

Then wash with 10 ml of 0,2 mol/l nitric acid solution (3.12), collect the eluate D" in a glass beaker; it will be used for the plutonium assay.

Evaporate the eluates D' and D" to dryness.

Redissolve in 50 μ l of concentrated nitric acid and fume to dryness to eliminate organic residues. Repeat once or twice.

Redissolve finally with a few drops of 0,2 mol/l nitric acid solution. The solutions obtained are ready to be loaded on the filaments used for mass spectrometry.

Operations 5.2 and 5.3 are presented in the form of a diagram in figure 1.



Figure 1 — U and Pu content assay

5.4 Preparation for isotopic assay

Carry out the operations described in 5.2.

Transfer 4 ml of solution C into a 60 ml plastic container.

Perform a redox cycle as described in 5.3.2.

Add 8 ml of concentrated nitric acid and mix.

Transfer the solution E obtained onto the ion exchange column.

Wash three times with 15 ml of 4 mol/l nitric acid solution and discard the eluate.

Wash again with 10 ml of 4 mol/l nitric acid solution and collect the eluate E' in a glass beaker for the isotopic assay of uranium. Wash four more times with 15 ml of 4 mol/l nitric acid solution to eliminate uranium and to avoid the interference of 238 U on the isotopic assay of plutonium. Discard the eluate. The repeated washings are necessary to achieve a decontamination factor of about 10⁴ for uranium.

Now wash with 10 ml of 0,2 mol/l nitric acid and collect the eluate E" in a glass beaker for the isotopic assay of plutonium.

Evaporate the eluates E' and E'' to dryness.

Redissolve in a few drops of 0,2 mol/l nitric acid. The solutions obtained are ready to be loaded on the filaments used for mass spectrometry.

Operations 5.2 and 5.4 are presented in the form of a diagram in figure 2.

6 Preparation of filament

6.1 Purification of filaments

Purify the filaments by degassing in a vacuum chamber for 10 min to 30 min at approximately 2 000 °C.

6.2 Loading procedure

Load 5 μ l to 10 μ l of the uranium and plutonium solutions obtained in the ion exchange separation which is equivalent to 0,5 μ g to 2 μ g uranium and 0,05 μ g to 0,5 μ g of plutonium (the amount depends on the type of mass spectrometer used) onto a suitably prepared sample filament.

Try to deposit the sample in the centre of the filament in the form of a precise dot. Heat the filament progressively, for example, by means of an electric current, to 90 °C to 100 °C in order to evaporate the solvent smoothly.

Then bring the filament to a dark red glow for 10 s to destroy the nitrate and to fix the oxides on the filament.

Avoid temperatures higher than 600 °C.

The temperature of the filament during the final stages of sample mounting is a critical factor and shall be carefully controlled to prevent significant variations in the fractionation between runs.

NOTE 1 It is important to reproduce closely all the parameters, which control the quality of the deposit, from one analysis to another: acidity, element concentrations (\pm 30 %), chemical purity especially with regard to the alkali elements, amount (\pm 30 %) and position of deposit and heating procedure.



Figure 2 — U and Pu isotopic assay

7 Isotopic mass spectrometry measurement

7.1 Position the filament assemblies in the mass spectrometer source (most modern instruments allow the simultaneous installation of several assemblies in the source), close the source and pump down to obtain a source pressure of less than 10^{-5} Pa $(10^{-7}$ torr).

7.2 Perform the measurement according to the instruction manual of the mass spectrometer.

7.3 Degas the sample by heating the sample filaments at a preselected temperature. Turn off heating of all filaments after 15 min and pump without heating for about 30 min for accountability analysis and about 15 min for process control measurements. Degassing is considered essential for measurements with high-sensitivity electron detectors where low-level interference from organic and inorganic molecules is a major problem. Degassing is also essential with conventional Faraday cup detectors when there are small amounts of organic impurities in the sample.

NOTE 2 Sample degassing may be considered optional **8.1** C for U and Pu isotopic analysis, but is strongly recommended site of the s

7.4 Resume heating of the ionization filament to 99:1993, 4, 5, 6, 8, 8', 9, 0, obtain the signal of the main isotope of the filament ads/sistopes ²³³U, ²³⁴U, ²⁴⁴F (e.g. the ¹⁸⁷Re peak) and focus to obtain maximum/iso-82³⁴Pu, ²⁴⁴F and ²⁴⁴F intensity.

7.5 Adjust the temperature of the ionization filament to 2 080 °C \pm 50 °C. Control the temperature using an optical pyrometer or by measuring the ¹⁸⁷Re⁺ ion current. If, at this stage, the ion current of the main isotope of the ionization filament (e.g. the ¹⁸⁷Re ion current) is unstable, interrupt the analysis here.

7.6 Otherwise, increase the sample filament current stepwise to yield a preset U^+ or Pu^+ signal.

Focus for maximum signal intensity.

7.7 Wait 20 min to 30 min to obtain a steady emission and readjust the sample filament heating currents to maintain the U^+ or Pu^+ signal intensity.

Focus again for maximum intensity.

7.8 Adjust the sum of the ion currents of the isotopes to be analysed to a preset value equal to approximately 80 % of the range of the current amplifier system (e.g. about 10^{-12} A).

Start the measurement after the ion currents begin to decay in a slow and smooth fashion.

Use exactly the same time and collection schedules for the analyses of samples and for calibration measurements.

7.9 In single collector operation, the "peakjumping" technique of ratio measurements is normally preferred to the "mass-scanning" technique. The baseline is therefore determined at least once per cycle of isotope ratio measurements.

7.10 The peak top observation time is governed by many factors including response of the measuring circuit, emission stability, rate of signal decay, settling time of the magnet switching circuit and desired precision and accuracy of the measurement. In general, there is a minimum delay of 2 s for circuits to stabilize after peak switching occurs when the isotope ratios range between 0,02 and 50. For accurate measurement of less abundant isotopes, longer delay times can be required to minimize the effects of system response and should be determined for each measuring circuit/mass spectrometer.

8 Calculation of the results

8.1 Calculation of isotope ratios

Average raw ion current ratios are obtained by interpolation and using appropriate scale factors. Use 3, 4, 5, 6, 8, 8', 9, 0, 1, 2 and 4' to designate the isotopes ²³³U, ²³⁴U, ²³⁸U, ²³⁸Pu, ²³⁹Pu, ²⁴⁰Pu, ²⁴¹Pu, ²⁴⁴Pu respectively and \overline{R}_{ijj} to designate the average raw isotope current ratio for isotopes *i* and *j*.

The isotope ratios $\overline{R'}_{i|j'}$ corrected for mass discrimination, are calculated as follows:

$$\overline{R'}_{i|j} = \overline{R}_{i|j} \left[1 + (M_j - M_i)B \right]^{-1} \qquad \dots (3)$$

where B is the mass discrimination factor.

8.2 Calculation of the isotopic abundance

The isotopic abundance A_i for the isotope i in a sample, is calculated as follows:

$$A_{i} = \frac{R'_{i|j}}{\sum_{k} \overline{R'}_{k|j}} \qquad \dots (4)$$

where all isotope ratios are referred to the same base isotope *j*.

For example, in a uranium sample which is normally free from $^{232}\text{U},~^{233}\text{U}$ and ^{237}U isotopes, the isotopic abundance of $^{2'35}\text{U}$ is given by

$$A_5 = \frac{\overline{R'}_{5/8}}{\overline{R'}_{4/8} + \overline{R'}_{5/8} + \overline{R'}_{6/8} + \overline{R'}_{8/8}} \qquad \dots (5)$$

For a plutonium sample normally free from ²³⁶Pu and ²⁴⁴Pu isotopes, the isotopic abundance of ²³⁹Pu is equal to

$$A_{9} = \frac{\overline{R'}_{9/9}}{\overline{R'}_{8'/9} + \overline{R'}_{9/9} + \overline{R'}_{0/9} + \overline{R'}_{1/9} + \overline{R'}_{2/9}} \qquad \dots (6)$$

The atomic percentage, A_{t_i} , for isotope *i* is equal to

$$At_i = A_i \times 100^{\circ} \qquad \dots (7)$$

NOTES

3 In case of an insufficient separation of U from Pu in samples containing both elements, the isotopic ratio $R'_{8/9}$ should be determined by means of alpha spectrometry, be-cause of possible interferences from 238 U in mass spec-trometry. The alpha activity ratio of the isotopes 238 Pu and (239 Pu + 240 Pu) of the unspiked sample solution C should be measured within the week following the purification (see clause 5), especially if the abundance of ²⁴¹Pu is high, since ²⁴¹Am interferes and will grow in again from the beta decay of ²⁴¹Pu.

4 The isotopic ratio $\overline{R'}_{8/9}$ is calculated preferably as follows, using the ratio of the activities of the isotope ²³⁸Pu and the isotopes (²³⁹Pu and ²⁴⁰Pu) measured by alpha spectrometry.

$$M_{8'}(^{238}Pu) = 238,049 6$$

$$M_{9}(^{239}Pu) = 239,052 2$$

$$M_{0}(^{240}Pu) = 240,053 8$$

$$M_{1}(^{241}Pu) = 241,056 8$$

$$M_{2}(^{242}Pu) = 242,058 7$$

$$M_{4'}(^{244}Pu) = 244,064 2$$

8.4 Calculation of content

The contents of uranium, C_A^U , and plutonium, C_A^{Pu} , in grams per kilogram of uranium and plutonium, respectively, are given by the following formulae:

Is a b), especially if the abundance of ²¹⁷Pu is high, since
m interferes and will grow in again from the beta decay
The isotopic ratio
$$\overline{R'}_{8/9}$$
 is calculated preferably as follows,
ing the ratio of the activities of the isotope ²³⁹Pu and the
opes (²³⁹Pu and ²⁴⁰Pu) measured by alpha spectrometry.
 $R'_{8'/9} = R(\alpha)T_8\left(\frac{1}{T_9} + \frac{R'_{0/9}}{T_0}\right)$ **iTeh STA** (8) **DARP** $P_{Pu} = C_S^{Pu} \times \frac{G_{2S}}{G_{2S}} \times \frac{M_9}{M_2} \times \frac{m_S}{m_C} \times$
(10)

are the isotopic mass percentages of the isotope ²⁴²Pu in the spike solution S, and

in

 $M_{\rm e}(^{236}\rm U) = 236,045~6$

 $M_8(^{238}\cup) = 238,050.8$

of the isotope 239Pu in the diluted sample solution C and original sample solution A, respectively;

are the atomic masses of ²³³U, ²³⁸U, ²³⁹Pu M_3 , M_8 , M_9 and M_2 and ²⁴²Pu, as given in 8.3;

are the masses of the $m_{\rm C}$ and $m_{\rm S}$ diluted sample solution C and of the spike solution S, respectively, used to prepare the spiked sample solution M (mixture);

 $(\overline{R'}_{i/j})_{M}, (\overline{R'}_{i/j})_{S} \text{ and } (\overline{R'}_{i/j})_{C}$ are the isotope ratios corrected for mass discrimination as described in 8.1 in the spiked sample solution M (mixture), in the spike solution S

9.2 Isotopic analysis

The coefficients of variation of the repeatability to be expected under the above conditions for the isotopic analyses are listed in table 1.

Accuracy of the method 10

10.1 Element assay

A plant laboratory analysed 21 solutions of plutonium nitrate by the isotope dilution method described in this International Standard. The mean of two independent analyses was compared with the mean result of two independent potentiometric titrations, performed at the same laboratory.

The average relative difference between the results of the two methods was equal to 0,03 %. The standard deviation of the relative differences was equal to 0,27 %.

Table 1 — Expected coefficients of variation of
repeatability and reproducibility of isotopic
analyses by thermal ionization mass spectrometry

solution C or original sample solution A, re- spectively;) PREV Isotope iteh.ai)	Abundance %	Repeatability ¹⁾ (% relative)	Repro- ducibility ²⁾ (% relative)
<i>F</i> is the dilution factor _{299:19} on a weight basis.	235U 93 115(67faf975-c150	0,2)-477 <mark>8-8</mark> 34a-	0,7 0,5	0,9 0,9
c5c5e456c45a/iso-88.5 Isotope decay correction	299-1993 ²³⁸ U	3 97 to 99	0,3 0,01	0,4 0,015
The report of the analyses of plutonium containing	²³⁸ Pu	0,3	2	3
solutions or mixed samples of uranium and plutonium shall include the date of the mass spectrometric	²³⁹ Pu	1,5 50 to 80	0,7 0,1	0,15
measurements in order to apply decay correction if necessary, e.g. for safeguards verification or umpire	²⁴⁰ Pu ²⁴¹ Pu	10 to 30 3 to 15	0,2 0,3	0,3 0,4
purposes. The decay of ²⁴¹ Pu and other Pu isotopes requires the correction of changes in the element	²⁴² Pu ²³⁹ Pu + ²⁴¹ Pu	1 to 5 65 to 83	0,3 0,1	0,4 0,15

requires the correction of changes in the element concentration and isotopic content of plutonium as well as uranium, because the uranium daughter growth should not be neglected. The analyses of reference materials also require appropriate decay corrections.

9 **Repeatability of the measurements**

9.1 Element assay

The coefficient of variation of the repeatability of duplicate determinations of uranium and plutonium contents in light water reactor spent fuel solutions observed at an industrial plant over a typical reprocessing campaign is about 0,2 %.

10.2 Isotopic analysis

ditions possible (as defined in ISO 3724).

The coefficients of variation of the reproducibility expected for the isotopic analyses are listed in table 1.

1) Coefficient of variation of replicate analyses performed at

2) Coefficient of variation of replicate analyses performed at

different laboratories or times, under the most different con-

the same time at a single laboratory (as defined in ISO 3724).

Interferences 11

lons with mass 233, 234, 235, 236 and 238 cause interference in the analysis of uranium if they have not been removed, or if they have been introduced as