# **BS ISO 13167:2015**



BSI Standards Publication

**Water quality — Plutonium, americium, curium and neptunium — Test method using alpha spectrometry**



... making excellence a habit."

#### **National foreword**

This British Standard is the UK implementation of ISO 13167:2015.

The UK participation in its preparation was entrusted to Technical Committee EH/3/8, Radioactivity measurements methods.

A list of organizations represented on this committee can be obtained on request to its secretary.

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# INTERNATIONAL STANDARD

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# **Water quality — Plutonium, americium, curium and neptunium — Test method using alpha spectrometry**

*Qualité de l'eau — Plutonium, americium, curium et neptunium — Méthode d'essai par spectrométrie alpha*



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# <span id="page-5-0"></span>**Foreword**

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The committee responsible for this document is ISO/147, *Water quality*, Subcommittee SC 3, *Radioactivity measurements*.

# <span id="page-6-0"></span>**Introduction**

Radioactivity from several naturally-occurring and anthropogenic sources is present throughout the environment. Thus, water bodies (e.g. surface waters, ground waters, sea waters) can contain radionuclides of natural, human-made, or both origins.

- Natural radionuclides, including potassium 40, and those originating from the thorium and uranium decay series, in particular radium 226, radium 228, uranium 234, uranium 238, lead 210, can be found in water for natural reasons (e.g. desorption from the soil and wash-off by rain water) or can be released from technological processes involving naturally occurring radioactive materials (e.g. the mining and processing of mineral sands or phosphate fertilizer production and use).
- Human-made radionuclides, such as transuranium elements (americium, plutonium, neptunium, curium), tritium, carbon 14, strontium 90 and some gamma emitting radionuclides, can also be found in natural waters as a result of authorized routine releases into the environment in small quantities in the effluent discharged from nuclear fuel cycle facilities. They are also released into the environment following their used in unsealed form for medical and industrial applications. They are also found in the water as a result of past fallout contamination resulting from the explosion in the atmosphere of nuclear devices and accidents such as those that occurred in Chernobyl and Fukushima.

Drinking water may thus contain radionuclides at activity concentrations which could present a risk to human health. In order to assess the quality of drinking water (including mineral waters and spring waters) with respect to its radionuclide content and to provide guidance on reducing health risks by taking measures to decrease radionuclide activity concentrations, water resources (groundwater, river, lake, sea, etc.) and drinking water are monitored for their radioactivity content as recommended by the World Health Organization [WHO] and as may be required by some national authorities.

An international standard on a test method for isotopes of plutonium, americium, curium and neptunium activity concentrations in water samples is justified for test laboratories carrying out these measurements, which are sometimes required by national authorities, as laboratories may have to obtain a specific accreditation for radionuclide measurement in drinking water samples.

238Pu, 239Pu, 240Pu, 241Am, 242Cm, 243Cm, 244Cm, 237Np activity concentrations can vary according to any local effluent authorized discharges from nuclear plant and environmental characteristics. The guidance level for  $^{238}$ Pu,  $^{239}$ Pu,  $^{240}$ Pu,  $^{241}$ Am,  $^{242}$ Cm,  $^{243}$ Cm,  $^{244}$ Cm,  $^{237}$ Np in drinking water as recommended by WHO is 1 Bq/l for all of those isotopes, except for 242Cm (10 Bq/l) [WHO-Guidelines for drinking waters quality, 2011, Geneva].

NOTE The guidance level is the activity concentration (rounded to the nearest order of magnitude) with an intake of 2 l/day of drinking water for 1 year, that results in an effective dose of 0,1 mSv/year for members of the public, an effective dose that represents a very low level of risk that is not expected to give rise to any detectable adverse health effect.

This International Standard is one of a set of International Standards on test methods dealing with the measurement of the activity concentration of radionuclides in water samples.

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# <span id="page-8-0"></span>**Water quality — Plutonium, americium, curium and neptunium — Test method using alpha spectrometry**

# **1 Scope**

This International Standard specifies a test method for measuring actinides (238Pu, 239+240Pu, 241Am, 242Cm, 243+244Cm and 237Np) in water samples by alpha spectrometry following a chemical separation.

The method can be used for any type of environmental study or monitoring.

The volume of the test portion required depends on the assumed activity of the sample and the desired detection limit.

The detection limit of the test method is  $5 \times 10^{-3}$  to  $5 \times 10^{-4}$  Bq/l for a volume of the test portion of 0,1 l to 5 l with a counting time of two to ten days.

# **2 Normative references**

The following documents, in whole or in part, are normatively referenced in this document and are indispensable for its application. For dated references, only the edition cited applies. For undated references, the latest edition of the referenced document (including any amendments) applies.

ISO [5667-1](http://dx.doi.org/10.3403/00108511U), *Water quality — Sampling — Part 1: Guidance on the design of sampling programmes and sampling techniques*

ISO [5667-3](http://dx.doi.org/10.3403/00716647U), *Water quality — Sampling — Part 3: Preservation and handling of water samples*

ISO [11929,](http://dx.doi.org/10.3403/30166520U) *Determination of the characteristic limits (decision threshold, detection limit and limits of the confidence interval) for measurements of ionizing radiation — Fundamentals and application*

ISO [80000-10,](http://dx.doi.org/10.3403/30126401U) *Quantities and units — Part 10: Atomic and nuclear physics*

[ISO/IEC](http://dx.doi.org/10.3403/02033502U) 17025, *General requirements for the competence of testing and calibration laboratories*

# **3 Terms and definitions**

For the purposes of this document, the terms, definitions and symbols given in ISO [80000-10](http://dx.doi.org/10.3403/30126401U) and ISO [11929](http://dx.doi.org/10.3403/30166520U) apply.

# <span id="page-9-0"></span>**4 Symbols**

The following symbols apply.



# **5 Principle**

Actinide isotopes are deposited as a thin source for measurement by alpha spectrometry using a grid chamber or semi-conductor detector type equipment. The sources are usually prepared by electrodeposition or co-precipitation after chemical separation and purification of the actinide isotopes present in the test portion. $[1][2][3][4]$  $[1][2][3][4]$  $[1][2][3][4]$  $[1][2][3][4]$  $[1][2][3][4]$  $[1][2][3][4]$  $[1][2][3][4]$  $[1][2][3][4]$ 

Specific chemical separation and purification procedures are required in order to avoid interference from the presence of other  $\alpha$  emitters, and stable nuclides in the sample, in quantities that are often larger than the actinide isotopes of interest.

<span id="page-10-0"></span>These procedures allow the main sources of interference to be removed:

- the salt content of the water sample, especially hydrolysable elements, in order to prepare the thinnest deposited source*;*
- other  $\alpha$  emitting radionuclides, such as uranium and thorium isotopes, whose emissions may interfere with those of actinide isotopes of interest.

The total yield for each analysis (product of chemical separation yield and detection efficiency) is determined by adding a standard solution of tracer: 236Pu can be used for plutonium isotopes and 237Np, 242Pu can be used for plutonium isotopes only and 243Am can be used for americium and curium isotopes.

As a result, the procedure shall include a reduction/oxidation cycle to adjust the tracer and the analytes to the same oxidation state.

It is possible to quantify curium isotopes on the basis of 243Am tracer behaviour. This may lead to a potential bias that shall be quantified using a standard solution and/or participation in inter-laboratory comparison tests.

It is possible to quantify neptunium on the basis of  $^{236}$ Pu tracer behaviour. This may lead to a potential bias that shall be quantified using a standard solution and/or participation in inter-laboratory comparison.

NOTE 235Np, 236Np, 238Np and 239Np can be used as a yield tracers for 237Np (if available), and 245Cm as a yield tracer for other Cm isotopes but the test method of this International Standard does not cover these measurements.

# **6 Chemical reagents and equipment**

### **6.1 Chemical reagents**

The chemical reagents and equipment are described in **Annexes A and B** for chemical separation and in Annexes C and D for the preparation of the deposited source.

Except for the certified standard solutions, all the chemical reagents needed to carry out this procedure shall be analytical grade.

### **6.2 Equipment**

Usual laboratory apparatus and in particular the following equipment.

**6.2.1 Alpha-spectrometer**, of the grid chamber (with higher detection efficiency, but lower resolution) or semiconductor type (with lower detection efficiency, but higher resolution). Operation at constant temperature is recommended. Follow the manufacturer's instructions.

For semiconductor-type equipment, the measurements using alpha-spectrometry depend on the interaction of alpha-particles with an ion-implanted silicon detector. This interaction instantly changes the conductivity of the silicon, proportional to the energy of the incoming alpha-particle. If a well-resolved spectra is required, the detection system should be maintained at a pressure < 10 Pa. Resolution can be further enhanced through increasing distance between source and detector.

**6.2.2 Pipette**, suitable for the accurate transfer of (for example 100 μl) tracer solution with a total precision within  $\pm$  1 %.

**6.2.3 Balance**, capable of achieving ± 0,1 mg precision.

In addition, the equipment for chemical separation is listed in Annexes A and B and the equipment for the preparation of the deposited source is listed in Annexes C and D.

# <span id="page-11-0"></span>**7 Procedure**

# **7.1 Sampling**

Conditions of sampling shall follow ISO [5667-1.](http://dx.doi.org/10.3403/00108511U)

The sample should be filtered to remove solids and then acidified (pH < 2 with nitric acid) as soon as possible after sampling prior to analysis, as specified in ISO [5667-3.](http://dx.doi.org/10.3403/00716647U)

It is important that the laboratory receives a representative sample, unmodified during the transport or storage and in an undamaged container.

# **7.2 Concentration step and valence cycle**

The radioactive tracers are added during this initial treatment phase.

If required, a concentration step can be done by evaporation or co-precipitation. After evaporation, the residue is dissolved with acid mixture. The co-precipitation with the iron hydroxide, for example, [Fe(OH)3] at pH = 8 can be done by adding iron nitrate or chloride. After centrifugation or filtration, the precipitate is dissolved with acid mixture.

The procedure shall include a valence cycle, adjusting the tracer and the actinide isotopes to measure oxidation states, in order to achieve the identical chemical behaviour for all of them. For example, a primarily reduction step is carried out by adding  $NH<sub>2</sub>OH.HCl$ , NaHSO<sub>3</sub> or Na<sub>2</sub>S<sub>2</sub>O<sub>3</sub>, then an oxidation step is done with  $NaNO<sub>2</sub>$  or  $H<sub>2</sub>O<sub>2</sub>$ .

# **7.3 Chemical separation**

There are two commonly used techniques for the chemical separation of actinides: extraction on an ion exchange resin or specific extraction chromatographic resin. One method from each technique is presented in Annexes A and B: separation by anionic resin<sup>[[6\]](#page-28-5)</sup> or by extraction chromatographic resins. $[2],[8]$  $[2],[8]$ 

It is also possible to use a mix of Annexes A and B: use of an ion exchange resin followed by the use of a specific extraction chromatographic resin for americium phase purification only.

# **7.4 Preparation of the source to be measured**

# **7.4.1 General**

The source can be prepared by electrodeposition on a planchet, a stainless steel disk ([7.4.2](#page-11-1)), or by co-precipitation [\(7.4.3](#page-11-2)).

# <span id="page-11-1"></span>**7.4.2 Electrodeposition method**

Electrodeposition is carried out after the chemical separation of the actinides from interfering elements. It allows the electrochemical deposition of the actinides in an ultra-thin layer onto the planchet. The procedure described in **[Annex](#page-23-1) C** applies to the two chemical separation methods described in Annexes A and B.

NOTE Electrodeposition is not a selective method because some stable metal cations are likely to form insoluble hydroxides, which can be deposited at the same time as the actinides.

# <span id="page-11-2"></span>**7.4.3 Co-precipitation method**

Co-precipitation, using fluoride compounds, can be carried out after the chemical separation of the actinides from other interfering elements. It allows the precipitation of the radionuclide(s) in the form <span id="page-12-0"></span>of a thin layer of cerium or lanthanum fluoride on a filter. The procedure described in **[Annex](#page-26-1) D** can be applied to the two chemical separation methods described in **Annexes A and B**.

### **7.5 Background determination**

Measure the background using a blank sample prepared for the method chosen (for example: laboratory water). This blank sample should be prepared without tracer to be used for obtaining the background count rates.

### **7.6 Counting efficiency determination**

The counting efficiency is estimated by measuring the calibration source. It is needed to determine the chemical recovery.

### **7.7 Measurement**

The actinide activity concentration is calculated by counting the sample source for an appropriate counting time. The same instrumental parameters should be used for the sample, the background and the calibration source measurements.

The counting time required depends on the sample and background count rates and also the detection limit and decision threshold required.

It should be verified on the spectrum that no interferent is present and an adequate resolution is obtained.

# **8 Expression of results**

### **8.1 Calculation of the activity concentration**

The actinides activity concentration is calculated by integrating the number of counts in the corresponding peaks of the tracer and isotopes to be measured. The results of these integrations, divided by the counting time, are the gross count rates  $r_{\text{gT}}$  and  $r_{\text{g}}$  for the tracer and the actinide isotopes, respectively.

*r*<sub>gT</sub> and *r*<sub>g</sub> are corrected for the background contribution and, if needed, of the tailing contribution of the highest derived energy peaks that depends on the detector characteristics.

Background count rates are calculated from the alpha emission spectrum of a blank sample, and integrating the number of counts in the regions of interest (ROI) in which the peaks appear in the sample spectrum. The results of these integrations, divided by the counting time, are the background count rates, *r*0t and *r*0 for the tracer and the actinide isotopes, respectively.

The blank sample is obtained and measured, applying the procedure in use in the laboratory with tracer (for the test method qualification) or without tracer (for background quantification).

The activity concentration  $c_A$  of the actinides isotopes is calculated as given in Formula (1):

$$
c_A = (r_g - r_0) / (V \cdot R) = (r_g - r_0) \cdot w \quad \text{with} \quad w = \frac{1}{V \cdot R} \tag{1}
$$

The total measurement yield is determined from the activity *A* of the tracer added and from the net count rate in the corresponding peak as given in Formula (2):

$$
R = (r_{\rm gT} - r_{0\rm T})/A \cdot f \tag{2}
$$

With *f* being the correction factor for possible bias when calculating results for curium isotopes using 243Am tracer or 237Np results using 236Pu tracer. *f* is the ratio between the Pu/Np and/or Am/Cm chemical yield

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For plutonium isotopes and 241Am, the factor *f* is equal to 1.

The counting efficiency  $\varepsilon$  allows one to calculate the chemical yield  $R_c$ . This value is important for quality control.

### **8.2 Standard uncertainty**

According to the GUM (ISO/IEC Guide 98-3), the standard uncertainty of  $c_A$  is calculated by Formula (3):

$$
u(c_A) = \sqrt{w^2 \cdot (u^2 (r_g) + u^2 (r_0)) + c_A^2 \cdot u_{\text{rel}}^2 (w)} = \sqrt{w^2 \cdot (r_g / t_g + r_0 / t_0) + c_A^2 \cdot u_{\text{rel}}^2 (w)}
$$
(3)

where the uncertainty of the counting time is neglected.

The relative standard uncertainty of *w* is calculated by Formula (4):

$$
u_{\rm rel}^2(w) = u_{\rm rel}^2(R) + u_{\rm rel}^2(V) \tag{4}
$$

The relative standard uncertainty of *R* is calculated by Formula (5):

$$
u_{\rm rel}^2(R) = u_{\rm rel}^2 (r_{\rm gT} - r_{0\rm T}) + u_{\rm rel}^2(f) + u_{\rm rel}^2(A) = (r_{\rm gT} / t_{\rm g} + r_{0\rm T} / t_0) / (r_{\rm gT} - r_{0\rm T})^2 + u_{\rm rel}^2(f) + u_{\rm rel}^2(A)
$$
 (5)

 $u_{\text{rel}}(A)$  includes all the uncertainties related to the tracer activity: that is in the standard solution, preparation of the tracer solution and the addition of the tracer solution to sample.

For the calculation of the characteristic limits according to ISO [11929,](http://dx.doi.org/10.3403/30166520U)  $\tilde u(\tilde c_A)$  is needed, i.e. the standard uncertainty of *c*A as a function of its true value, calculated by Formula (6):

$$
\tilde{u}(\tilde{c}_A) = \sqrt{w^2 \cdot ((\tilde{c}_A / w + r_0) / t_g + r_0 / t_0) + \tilde{c}_A^2 \cdot u_{\text{rel}}^2(w)}
$$
(6)

### **8.3 Decision threshold**

In accordance with ISO [11929,](http://dx.doi.org/10.3403/30166520U) the decision threshold,  $c_A^*$ , is obtained from the above Formula (6) for  $\tilde{c}^{\,}_{\rm A}=0$  . This yields Formula (7):

$$
c_A^* = k_{1-\alpha} \cdot \tilde{u}(0) = k_{1-\alpha} \cdot w \cdot \sqrt{r_0 / t_g + r_0 / t_0}
$$
\n<sup>(7)</sup>

 $\alpha$  = 0,05 with  $k_{1-\alpha}$  = 1,65 is often chosen by default.

### **8.4 Detection limit**

In accordance with ISO [11929,](http://dx.doi.org/10.3403/30166520U) the detection limit,  $\,c_{\rm A}^{\,\,\#}$  , is calculated by Formula (8):

$$
c_A^{\#} = c_A^* + k_{1-\beta} \cdot \tilde{u}(c_A^{\#}) =
$$
  
\n
$$
c_A^* + k_{1-\beta} \cdot \sqrt{w^2 \cdot ((c_A^{\#}/w + r_0)/t_g + r_0/t_0) + c_A^{\#2} \cdot u_{rel}^2(w)}
$$
  
\n
$$
\beta = 0.05 \text{ with } k_{1-\beta} = 1.65 \text{ is often chosen by default.}
$$
\n(8)

The detection limit can be calculated by solving Formula (8) for  $c_A^{\#}$  or, more simply, by iteration with a starting approximation  $c_{\mathrm{A}}^{\phantom{\ast}\#} = 2 \cdot c$  $=2\cdot c_A^*$ .

<span id="page-14-0"></span>When taking  $\alpha = \beta$ , then  $k_{1-a} = k_{1-\beta} = k$  and the solution of Formula (8) is given by Formula (9):

$$
c_A^{\mu} = \frac{2 \cdot c_A^* + (k^2 \cdot w)/t_g}{1 - k^2 \cdot u_{\text{rel}}^2(w)}
$$
(9)

#### **8.5 Confidence limits**

In accordance with ISO [11929,](http://dx.doi.org/10.3403/30166520U) the lower,  $c_A^{\preceq}$ , and upper,  $c_A^{\preceq}$ , limits of the confidence interval are calculated using Formulae (10) and (11):

$$
c_A^{\mathcal{A}} = c_A - k_p \cdot u(c_A); p = \omega \cdot (1 - \gamma/2) \tag{10}
$$

$$
c_A^{\triangleright} = c_A + k_q \cdot u(c_A); q = 1 - \omega \cdot \gamma / 2 \tag{11}
$$

where

 $\omega = \Phi(y/u(y))$  with  $\Phi$  being the distribution function of the standardized normal distribution;

 $\omega = 1$  may be set if  $c_A \ge 4 \cdot u(c_A)$  and Formula (12) applies:

$$
c_A^{\alpha,\triangleright} = c_A \pm k_{1-\gamma/2} \cdot u(c_A) \tag{12}
$$

 $\gamma = 0.05$  with  $k_{1-\gamma/2} = 1.96$  is often chosen by default.

### **9 Quality assurance and quality control programme**

#### **9.1 General**

Quality control operations shall meet the requirements of [ISO/IEC](http://dx.doi.org/10.3403/02033502U) 17025.

#### <span id="page-14-1"></span>**9.2 Influence quantities**

Special precautions shall be taken to minimize the influence of quantities that can affect the measurement results:

- spike of tracer with an inappropriate activity (too small or too large) compared to the actinide activity to be quantified;
- poor chemical separation with interferences on the alpha spectrum;
- a small chemical yield (a value of at least 30 % is expected) which can be a consequence of important losses during the chemical separation steps or in the preparation of the source;

The chemical yield  $R_c$  can be calculated using Formula (13):

$$
R_{\rm c} = R/\epsilon \tag{13}
$$

presence in the test sample of a small amount of the actinide isotope that is used as the tracer, e.g. 242Pu. In this case, 236Pu can be used as tracer instead.

# <span id="page-15-1"></span><span id="page-15-0"></span>**9.3 Instrument verification and calibration**

Major instrument parameters (energy calibration, efficiency calibration, energy resolution, and background spectrum) shall be periodically checked within a quality assurance programme established by the laboratory and following the manufacturer's instructions.

Make sure the whole of the data acquisition chain is performing satisfactorily, using an alpha-emitting radioactive standard source (e.g. 239Pu).

# **9.4 Method verification**

The test method repeatability shall also be checked, e.g. by replicate measurements.

The acceptance limits of the test parameters contained in [9.2](#page-14-1) and [9.3](#page-15-1) shall be defined.

Periodical verifications of the accuracy of the method shall be run. These may be accomplished by:

- participating in inter-laboratory comparison tests;
- analysing reference materials.

### **9.5 Demonstration of analyst capability**

If an analyst has not used this procedure before, a precision and bias test shall be performed by running a duplicate measurement of a reference or spiked material. Acceptance limits shall be defined by the laboratory.

A similar test shall be performed by analysts routinely using this procedure with a periodicity defined by the laboratory. Acceptance limits shall be defined.

# **10 Test report**

The test report shall conform to [ISO/IEC](http://dx.doi.org/10.3403/02033502U) 17025 requirements and shall contain the following information:

- a) reference to this standard;
- b) identification of the sample;
- c) units in which the result is expressed;
- d) the test result,  $c_A \pm u(c_A)$  or  $c_A \pm U$ , with the associated *k* value.

Complementary information can be provided such as:

- probabilities *α*, *β* and (1 − *γ*);
- decision threshold and the detection limit;
- depending on the customer request there are different ways to present the result:
	- when the activity concentration  $c_A$  is compared with the decision threshold, in accordance with ISO [11929,](http://dx.doi.org/10.3403/30166520U) the result of the measurement should be expressed as  $\leq c_{\rm A}^*$  when the result is below the decision threshold;
	- when the activity concentration  $c_A$  is compared with the detection limit, the result of the measurement can be expressed as  $\leq c_A^\#$  when the result is below the detection limit;
- chemical yield value;
- any relevant information likely to affect the results.

If the detection limit exceeds the guideline value, it shall be documented that the method is not suitable for the measurement purpose.

# **Annex A**

# (informative)

# <span id="page-17-0"></span>**Chemical separation of actinides on anionic resin**

# **A.1 Principle**

The actinide isotopes are separated from the other radionuclides and from the matrix in an anionic complex form using an ion exchange resin in a column in a nitric acid medium and hydrochloric one.

# **A.2 Apparatus**

Usual laboratory apparatus and the following.

- **A.2.1 Hot plate.**
- **A.2.2 Glass column.**
- **A.3 Reagents**
- **A.3.1 Tracer solutions**.
- **A.3.2 Hydrochloric acid, concentrated**, (HCl) : 37 %.
- **A.3.3 Nitric acid, concentrated**, (HNO3) : 65 %.
- **A.3.4 Ammonia, concentrated**, (NH4OH) : 25 %.
- **A.3.5 Sodium nitrite** (NaNO<sub>2</sub>).
- **A.3.6 Anionic resin** 1 × 4 50/100 mesh or 1 × 8 100/200 mesh.
- **A.3.7 Nitric acid, solution**  $c(HNO_3)$ : 8 mol/l.
- **A.3.8 Hydrochloric acid, solution** *c*(HCl) : 2, 8 and 10 mol/l.
- **A.3.9 Hydroxylamine hydrochloride** NH2OH.HCl, 0,2 mol/l in hydrochloric acid (HCl), 2 mol/l.
- **A.3.10 Methanol**.
- **A.3.11 Solution A**: 10 % (vol) 10 mol/l HNO<sub>3</sub> and 90 % (vol) of methanol, freshly made.
- **A.3.12 Solution B:** 30 % (vol) 0,5 mol/l HNO<sub>3</sub> and 70 % (vol) of methanol, freshly made.

# **A.4 Procedure**

### **A.4.1 General**

Add a known activity of tracers to the water sample.

This procedure is carried out with two main steps: chemical separation and elution.

In all steps, if not specified, the flow rate should be of approximately 1 ml/min.

NOTE In some cases, the flow through resin columns is very poor because of sample composition. Working with vacuum is therefore very helpful to support the flow.

# **A.4.2 Separation of plutonium and neptunium**

Carry out the following steps.

- a) Fill the column with the resin (for example, a 150 ml volume column can be filled with about 100 ml of prepared resin).
- b) Prepare a 8 mol/l nitric acid solution.
- c) Convert the resin to the nitrate form by passing  $150$  ml of the 8 mol/l HNO<sub>3</sub>.
- d) Place a clean beaker under the column and load the actinide solution on the column.
- e) Wash the beaker and column with portions of 8 mol/l  $HNO<sub>3</sub>$  up to 150 ml with a flow of approximately 1 ml/min. This eluate E1 contains Am, Cm, Fe and U.
- f) Place a clean beaker under the column and wash the column with 80 ml of 8 mol/l HCl. This eluate E2 contains Th.

### <span id="page-18-1"></span>**A.4.3 Elution of plutonium and neptunium**

<span id="page-18-0"></span>**A.4.3.1** Carry out the following steps.

- a) Place a clean beaker under the column.
- b) Elute the plutonium and neptunium, E3, with 90 ml of 0,2 mol/ NH2OH.HCl in 2 mol/l HCl with a flow of approximately 0,5 ml/min.
- c) Evaporate the eluate E3 slowly until the volume is less than 25 ml and add carefully 2 ml of concentrated HNO<sub>3</sub>.

**A.4.3.2** When the presence of relative important amounts of other actinides is suspected, it is necessary to carry out a purification process that basically consists of the repetition of the following steps.

- a) Evaporate the eluate E3 to dryness.
- b) Add 50 ml of 8 mol/l  $HNO<sub>3</sub>$ .
- c) Add  $0.6$  g of NaNO<sub>2</sub> and heat to remove the nitrous fumes and leave to cool.
- d) Repeat the procedure from  $A.4.3.1$  to separate and elute the plutonium and neptunium.
- e) Prepare the source by electrodeposition or co-precipitation, as described in [Annex](#page-26-1) C and Annex D, respectively.

# **A.4.4 Separation of americium and curium**

Carry out the following steps.

- a) Evaporate the eluate E1 to dryness.
- b) Dissolve the residue with 50 ml of 10 mol/l HCl.
- c) Fill the column with 15 ml of anionic resin.
- d) Prepare 50 ml of a 10 mol/l HCl solution and wash the column to convert the resin to the chloride form. Discard the washings.
- e) Place a clean beaker under the column and pour the solution obtained in b) into the top of the column.
- f) Wash the beaker and column with portions of 10 mol/l HCl up to 50 ml. The eluate E4 is obtained from e) and f).
- g) Evaporate the eluate E4 to dryness.

# <span id="page-19-0"></span>**A.4.5 Purification of americium and curium**

Carry out the following steps.

- a) Dissolve the residue of eluate E4 with 10 ml of 10 mol/l HNO3.
- b) Stir and add 90 ml of methanol.
- c) Fill the column with 15 ml of anionic resin.
- d) Prepare 50 ml of a fresh solution A and add to the resin.
- e) Pour the solution into the top of the column.
- f) Wash the beaker and column with 50 ml of solution A.
- g) Place a clean beaker under the column and elute americium and curium with 100 ml of a fresh solution B as E5.
- h) Evaporate to dryness the eluate E5.
- i) Dissolve the residue with:
	- 1) 1 ml of concentrated  $HNO<sub>3</sub>$  in case of electrodeposition;
	- 2) 50 ml of 0,2 mol/l HCl in case of co-precipitation.
- j) Prepare the source by electrodeposition or co-precipitation, as described in  $\Delta$ nnex C and  $\Delta$ nnex D, respectively.

# **Annex B**

# (informative)

# <span id="page-20-0"></span>**Chemical separation of actinides by specific resins**

# **B.1 Principle**

This technique is based on the selective chromatographic extraction of plutonium isotopes, 241Am, curium isotopes and 237Np using a resin coated with a specific extractant CMPO/TBP. The chemical separation is fast and well suited for monitoring plutonium and americium activity in the environment.

# **B.2 Apparatus**

Usual laboratory apparatus and the following.

- **B.2.1 Scales**, to an accuracy of 0,1 mg.
- **B.2.2 Evaporator or hot plate**.
- **B.2.3 CMPO/TBP extractant-coated resin columns (2 ml volume in general)**.

# **B.3 Reagents**

- **B.3.1 Tracer solutions**.
- <span id="page-20-1"></span>**B.3.2** Nitric acid (HNO<sub>3</sub>): 3 mol/l in aluminium nitrate (Al(NO<sub>3</sub>)<sub>3</sub>): 0,5 mol/l.
- **B.3.3** Nitric acid (HNO<sub>3</sub>):  $2 \text{ mol/l}$  in sodium nitrite NaNO<sub>2</sub>:  $0.1 \text{ mol/l}$ .
- <span id="page-20-2"></span>**B.3.4** Nitric acid, concentrated,  $(HNO<sub>3</sub>)$ : 69 % minimum.
- **B.3.5 Different nitric acid (HNO<sub>3</sub>) solutions**: 3 mol/l, 2 mol/l, 0,5 mol/l, 1 mol/l.
- **B.3.6 Ascorbic acid**.
- **B.3.7 Hydrogen peroxide**  $(H_2O_2)$ : 30 %.
- **B.3.8 Hydrochloric acid, concentrated**, (HCl),: 37 %.
- **B.3.9 Hydrochloric acid solution** (HCl): 0,2 mo/l.
- **B.3.10** Ammonium hydrogen oxalate  $(NH_4HC_2O_4)$ : 0,1 mol/l.

# **B.4 Procedure**

### **B.4.1 General**

Add a known activity of tracers to the water sample acidified with nitric acid.

This procedure is carried out with two main steps: chromatographic extraction and elution of americium, curium, plutonium and neptunium. In all steps, if not specified, the flow rate should be of approximately 1 ml/min.

In some cases, a ferric hydroxide co-precipitation can be added before the separation steps to concentrate actinides.

NOTE In some case, the flow through resin columns is very poor because of sample composition. Working with vacuum is therefore very helpful to support the flow.

### **B.4.2 Extraction of americium and curium**

Carry out the following operations.

- a) Evaporate the solution to be analysed to dryness.
- b) Add 10 ml of 3 mol/l HNO<sub>3</sub> in 0,5 mol/l Al(NO<sub>3</sub>)<sub>3</sub> to dissolve the precipitate.
- c) Add 0,1 g of ascorbic acid to keep all Fe as Fe(II) (to ensure reduction of Pu and Np but also to be sure no Fe(III) could interfere with Pu(III) and Am(III) uptake), heat until total dissolution and leave to cool.
- d) Set up the CMPO/TBP column.
- e) Prepare the resin by passing 25 ml of 3 mol/l HNO<sub>3</sub> in 0,5 mol/l Al(NO<sub>3</sub>)<sub>3</sub>.
- f) Pour the solution into the top of the column.
- g) Wash the beaker with 5 ml of 3 mol/l HNO3, pour into the top of the column.
- h) Wash the column with 5 ml of 2 mol/l  $HNO<sub>3</sub>$ , 5 ml of 2 mol/l  $HNO<sub>3</sub>$  in 0,1 mol/l NaNO<sub>2</sub>. Discard the washings.
- i) Place a clean beaker under the column and elute americium and curium with 5 ml  $0.5$  mol/l HNO<sub>3</sub>, 3 ml 9 mol/l HCl and 20 ml of 4 mol/l HCl (E6).
- j) Evaporate the eluate E6 to dryness.
- k) Dissolve the residue with:
	- 1) 1 ml of concentrated  $HNO<sub>3</sub>$  in case of electrodeposition;
	- 2) 50 ml of 0,2 mol/l HCl in case of co-precipitation.
- l) Prepare the source by electrodeposition or co-precipitation, as described in [Annex](#page-23-1) C and [Annex](#page-26-1) D, respectively.

### **B.4.3 Elution of plutonium and neptunium**

Carry out the following operations.

- a) Place a clean beaker under the column.
- b) Elute plutonium and neptunium as E7 with 20 ml of 0,1 mol/l  $NH_4HC_2O_4$  in 1 mol/l HCl (U is not eluted).
- c) Evaporate the eluate E7 to dryness. The white precipitate is in the oxalate form.

Thorium is known to be very insoluble in water. If the presence of thorium is suspected in the sample, it is necessary, prior to Pu/Np elution, to add a step to wash the column with 20 ml of 4 mol/l HCl in 0,1 mol/l HF to elute thorium as it interferes with plutonium in *α*-spectrometry.

### **B.4.4 Oxalate decomposition**

Carry out the following operations.

- a) Add 1 ml of 69 % HNO<sub>3</sub>, and five drops of 30 % H<sub>2</sub>O<sub>2</sub>, evaporate to dryness.
- b) Add 1 ml of 69 % HNO<sub>3</sub>, and 1 ml of 37 % HCl, evaporate to dryness.
- c) Add 1 ml of 69 % HNO<sub>3</sub>, evaporate to dryness.
- d) Repeat the step c) until the precipitate disappears.
- e) Dissolve the residue in 4 ml of 1 mol/l HNO<sub>3</sub> (Solution E8).
- f) Evaporate the solution E8 to dryness.
- g) Dissolve the residue with:
	- 1) 1 ml of concentrated  $HNO<sub>3</sub>$  in case of electrodeposition;
	- 2) 50 ml of 0,2 mol/l HCl in case of co-precipitation.
- h) Prepare the source by electrodeposition or co-precipitation, as described in [Annex](#page-26-1) C and Annex D, respectively.

# <span id="page-23-1"></span>**Annex C**

# (informative)

# <span id="page-23-0"></span>**Preparation of the source by electrodeposition**

# **C.1 Principle**

Using a DC generator to apply different voltages between two electrodes leads to the reduction of the metal cations dissolved in the electrolyte. The reduction that takes place at the cathode leads to the formation of a deposit of actinides in hydroxide form.

# **C.2 Apparatus**

- **C.2.1 Electrodeposition apparatus**, generally composed of the following.
- **C.2.1.1 Glass or polyethylene electrodeposition cell**.
- <span id="page-23-3"></span>**C.2.1.2 Platinum wire** (anode).
- <span id="page-23-2"></span>**C.2.1.3 Stainless steel disk** (cathode) with a diameter adapted to the electrodeposition cell.
- **C.2.1.4 DC power supply**.
- **C.2.2 Hot plate**.
- **C.2.3 Petri dish**.

# **C.3 Reagents**

- **C.3.1 Nitric acid, concentrated**  $(HNO<sub>3</sub>)$ : 65 %.
- **C.3.2 Sodium sulfate** (Na2SO4): 0,3 mol/l.
- **C.3.3 Sulfuric acid, concentrated**  $(H_2SO_4)$ : 95 % to 97 %.
- **C.3.4 Thymol blue**  $(C_{27}H_{30}O_{5}S)$ : 0,04 %.
- **C.3.5** Ammonia, concentrated  $(NH_4OH)$ : 25 %.
- **C.3.6 Sulfuric acid**  $(H_2SO_4)$ : 1 %.
- **C.3.7 Ammonia** (NH4OH): 1 %.

# **C.4 Procedure**

# **C.4.1 Assembly of the electrodeposition cell**

This procedure refers to the electrodeposition apparatus designed for deposits on small diameter stainless steel disks (see [Figure](#page-25-0) C.1). The electrodeposition procedure for this type of apparatus is as follows.

- a) Place the clean and degreased stainless steel disk  $(C.2.1.3)$  $(C.2.1.3)$  in the cap assembly.
- b) Assemble the cell with the cap.
- c) Fill the cell with water to check for leaks.
- d) Empty the cell.
- e) Fix the platinum anode  $(C.2.1.2)$  $(C.2.1.2)$  vertically to its support. The distance between the anode and the disk (cathode) is approximately 3 mm.

### **C.4.2 Analyte deposition**

Using the preparation obtained from the elution and purification phase (see  $A.4.3$  and  $A.4.5$ ) or from the extraction and oxalate decomposition step (see  $\underline{B.3.2}$  and  $\underline{B.3.4}$  $\underline{B.3.4}$  $\underline{B.3.4}$ ), carry out the following operations.

- a) Evaporate to near dryness; care should be taken to avoid evaporating the sample to complete dryness.
- b) Add 1 ml of concentrated HNO<sub>3</sub>, and evaporate near dryness. Repeat this step three times.
- c) Add 1 ml of 0,3 mol/l Na<sub>2</sub>SO<sub>4</sub> and evaporate to dryness.
- d) Add 0.5 ml of 95 % to 97 % H<sub>2</sub>SO<sub>4</sub>.
- e) Add 10 ml of distilled water and 3 drops of 0,04 % thymol blue and stir.
- f) Adjust the pH between 2,1 and 2,3 by adding concentrated ammonia solution (colour changes from red to orange, a pH-meter can also be used).
- g) Introduce this solution in the electrodeposition cell.
- h) Rinse the beaker with a few millilitres of  $1\%$  H<sub>2</sub>SO<sub>4</sub>, whose pH is previously adjusted to 2,3; add to the electrodeposition cell.
- i) Set up the anode and carry out the electrodeposition at a constant current (about 1,0 A) for approximately 1 h. Low temperatures improve the efficiency of electrodeposition.
- j) One minute before switching off the generator, add 1 ml of 25 % NH4OH.
- k) Wait 1 min, remove the anode before switching off the generator and quickly empty the cell.
- l) Dismantle the apparatus, rinse the disk in water, then in 1 % NH4OH, and dry it using flow of hot air or on a hot plate.
- m) Identify the disk and place it in a plastic Petri dish.
- n) The source is ready to be measured by  $\alpha$  spectrometry.

Cooling the cell during electrodeposition could avoid an increase of temperature that can lead to excessive bubbling.



#### **Key**

- 1 anode
- 2 scintillation vial<br>3 disk
- disk
- 4 cap assembly
- 5 base

# <span id="page-25-0"></span>**Figure C.1 — Diagram of an electrodeposition cell**

# <span id="page-26-1"></span>**Annex D**

# (informative)

# <span id="page-26-0"></span>**Preparation of the source by co-precipitation**

# **D.1 Principle**

The actinides isotopes are recovered by precipitation using lanthanide fluoride.

# **D.2 Apparatus**

The equipment used for the co-precipitation method generally includes the following:

- **D.2.1 Cellulose ester membrane filter** (pore diameter: 0,22 μm).
- **D.2.2 Test dish or stainless steel disk**, with a diameter adapted to the spectrometry apparatus.
- **D.2.3 Filtration system**.
- **D.3 Reagents**
- **D.3.1 Hydrochloric acid** (HCl): 0,2 mol/l.
- **D.3.2 Ammonia, concentrated** (NH4OH): 25 %.
- **D.3.3 Lanthanum oxide** (La<sub>2</sub>O<sub>3</sub>): 99 % or nitrate of cerium (III) hexahydrated (Ce NO<sub>3</sub>)O<sub>3</sub>.6H<sub>2</sub>O).

**D.3.4** Hydrofluoric acid (HF): 40 % (a salt of fluoride of ammonium (NH<sub>4</sub>F) can be used instead).

**D.3.5 Solution C**: 29 mg of lanthanum oxide  $(La_2O_3)$  dissolved in 100 ml of hydrochloric acid (HCl) 0,2 mol/l or 155 mg of nitrate of cerium (III) hexahydrated dissolved in 100 ml of water.

# **D.4 Procedure**

Using the preparation obtained from the purification phase (see  $A.4.3$ ) or from the oxalate decomposition step (see  $\underline{B.3.4}$  $\underline{B.3.4}$  $\underline{B.3.4}$ ), carry out the following operations.

- a) Adjust the pH to 1,9 by adding NH4OH.
- b) Add, for a deposit of 20 cm2, the equivalent of 250 μg of lanthanum (i.e. 1 ml of solution C of lanthanum) or of cerium (for example 0,5 ml of solution C of cerium).
- c) Add 1 ml of HF or 1 ml of NH4F solution.
- d) Stir, then leave to settle for 10 min.
- e) Recover the precipitate by light vacuum filtering on the membrane filter. Ensure complete transfer of the precipitate by successive washings with water.

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- f) Leave the precipitate to dry in open air, then attach the membrane to the test dish or stainless steel dish.
- g) The source is ready to be measured by *α* spectrometry.

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