Stability monitoring of selected spike isotopic reference materials for isotope dilution mass spectrometry

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Abstract
The provision of high quality spike isotopic reference materials is one of the objectives of the Joint Research Centre of the European Commission. They play an important part in measurements of nuclear materials for Nuclear Safeguards. Spike isotopic reference materials are prepared and certified according to the international standards. The assigned values for the isotope amount content and isotope ratios need to be verified at regular intervals. This is carried out by stability monitoring measurements. This paper will discuss the results of the stability assessment of some 242Pu and 243Am spike isotopic reference materials.

Keywords Plutonium · Americium · Isotopic reference materials · Spike · Stability monitoring · IDMS

Introduction
Confidence in comparability and reliability of measurement results in nuclear material analysis is established by production and dissemination of internationally accepted quality assurance tools such as certified reference materials (CRMs), reference measurements, and inter-laboratory comparisons (ILCs). They provide the basis for a strong verification system to safeguard nuclear activities in line with the international agreements [1, 2]. The fundamental role of CRMs in measurements is to establish traceability of a measured value to a primary unit of measurement, such as defined in the International System of Units (SI). Only measurement results that are traceable to a common reference, preferably the respective SI unit by unbroken chain of comparisons (i.e. traceability) can be regarded as truly comparable.

The Joint Research Centre of the European Commission in Geel (JRC-Geel) is a recognised producer of nuclear CRMs for Nuclear Safeguards and Nuclear Security [3]. They are part of a systematic programme to supply the international community with reference materials, in particular uranium and plutonium isotopic reference materials (IRMs) covering a wide range of concentrations and isotope ratios. IRMs are applied in mass spectrometry for calibration of instruments, in validation of analytical methods and for assessing the reproducibility and uncertainty of measurement results [4, 5]. One group of IRMs are the so-called “spike” reference materials applied in Isotope Dilution Mass Spectrometry (IDMS) for the determination of the amount content of nuclear materials [6–9]. Spikes are isotopically enriched materials with certified amount content and isotopic composition.

The preparation and certification of spike IRMs are demanding and challenging tasks that require state-of-the-art measurement procedures and equipment. For the last decade, the JRC-Geel has focused to align these procedures with the international standards such as the ISO 17034 and ISO Guide 35 [10–12]. A special emphasis was given to the assessment of the homogeneity and stability of spike reference materials and to include the corresponding uncertainty contributions in the final uncertainty of the reference material. This is important in view of providing fit-for-purpose and high quality IRMs with small uncertainties [13, 14]. IRMs are essential for laboratories striving to meet the international target values for measurement uncertainties in safeguarding nuclear materials (ITV2010) [15]. Various steps in the preparation and certification of IRMs are shown in Fig. 1.

As mentioned above, the stability of spike IRMs is an important aspect that needs to be addressed during the certification process. The term “stability” in this context refers...
to any changes that affect the assigned values of the reference material for the duration of the material’s shelf-time. The stability assessment is usually carried out in a dedicated experimental study over an extended period (e.g. 2 years) prior to the release of the reference material. Another way to establish stability behaviour is based on the experience with reference materials of similar characteristics (i.e. isotopic composition, concentration, and matrix). This data has been acquired at JRC-Geel over many years by participating in inter-laboratory comparisons (ILCs), between-lab verification campaigns and via in-house quality control. Some of the results have been published elsewhere [8, 16–18].

The behaviour of spike materials is difficult to predict reliably solely from stability studies over short periods. Evaporation over longer period can have an effect on the concentration.

For these reasons, spike IRMs, that have been produced more recently in compliance with the respective standards are subject to stability monitoring after release. Monitoring using the classical design is applied [12]. This involves the measurement of a number of units of a reference material at various points in time, for the duration of the material’s validity, as indicated on the reference material certificate.

Two $^{242}\text{Pu}$ materials, namely IRMM-049d and IRMM-049e were selected for this study. $^{242}\text{Pu}$ is commonly used as a spike in IDMS measurements for determination of the Pu content as this isotope is usually found only as a minor component in plutonium samples of the nuclear fuel cycle. Another selected material was the IRMM-0243 (referred to as STAM at CETAMA), which is a $^{243}\text{Am}$ spike IRM used for applications in nuclear waste characterisation, nuclear forensics and nuclear decay studies [9]. These IRM solutions are supplied in glass ampoules with a screw-cap. They were introduced at JRC-Geel to replace the flame sealing of ampoules, which is no longer applied due to safety reasons. The results of the stability studies of these three spike IRM will be presented in this paper.

**Experimental**

**Reference materials and test samples**

For IDMS measurements, various IRMs for isotopes $^{239}\text{Pu}$, $^{240}\text{Pu}$ and $^{244}\text{Pu}$ were available at JRC-Geel. The following
materials were chosen based on their availability, the certified isotope and their concentration.

- IRMM-1027o and IRMM-1027m LSD: certified mass of $^{239}$Pu, $^{235}$U and $^{238}$U per vial. About 2 mg plutonium, $(m(239\text{Pu})/m(\text{Pu}) = 0.978)$ and about 50 mg uranium $(m(235\text{U})/m(\text{U}) = 0.194)$ as dried nitrate embedded in cellulose acetate butyrate (CAB)
- IRMM-086: $^{239}$Pu solution, 3.7524 (22) µmol $^{239}$Pu g$^{-1}$, $m(239\text{Pu})/m(\text{Pu}) = 0.978$
- IRMM-083: $^{240}$Pu solution, 3.4064 (22) µmol $^{240}$Pu g$^{-1}$, $m(240\text{Pu})/m(\text{Pu}) = 0.990$
- IRMM-042a: $^{244}$Pu solution, 3.7507 (50) nmol $^{244}$Pu g$^{-1}$, $m(244\text{Pu})/m(\text{Pu}) = 0.980$
- $^{241}$Pu solution: 81.394 (53) nmol $^{241}$Pu g$^{-1}$, $m(241\text{Pu})/m(\text{Pu}) = 0.993$
- NBL 126 Pu: $^{239}$Pu solution, 7.0927(20) µmol $^{239}$Pu g$^{-1}$, $m(239\text{Pu})/m(\text{Pu}) = 0.980$, prepared gravimetrically from the $^{239}$Pu metal from the New Brunswick Laboratory Program Office (NBL PO) was used [17, 20].

In addition to the IRMs listed above, certified test items (with undisclosed values), EQRAIN and MOX-Pu4, were included in this study as part of the external quality control assessment. EQRAIN (Assessment of the Quality of Analysis Results in the Nuclear Industry) is the proficiency test for the analysis of $^{238}$U or $^{239}$Pu amount contents in nitrate solution organised at regular intervals by CEA/CETAMA [21, 22]. The JRC-Geel has participated in EQRAIN since 2008. MOX-Pu4 is a dried nitrate standard material prepared by the JAEA-PFDC (Japan Atomic Energy Agency, Plutonium Fuel and Development Centre).

An overview of IDMS measurements of IRMM-049d, IRMM-049e and IRMM-0243 materials using different IRMs and certified test items is shown in Fig. 2. Units analysed in the scope of the external quality assessment are highlighted in yellow.

### Chemical treatment and isotopic measurements

Blends (e.g. spike + sample) for IDMS measurements were prepared by metrological weighing using the substitution method [17]. In most cases, more than one blend was prepared from each unit of the material being tested for stability. Prior to separation, an isotopic equilibration (homogenisation) of the spike and sample isotopes and the valence state adjustment were achieved by performing a reduction–oxidation step. The separations of the plutonium and americium samples were carried out by anion-exchange (Bio-rad AG®1 × 4, 100–200 mesh, Cl$^-$ form) and extraction chromatography (UTEVA, DGA), respectively. Details about the chemical separation procedures are published elsewhere [16, 18].

Isotopic measurements were performed by total evaporation (TE) method on a multi-collector Triton TIMS (Thermo Fisher Scientific, Bremen, Germany) [23–27]. In this process...
method, the evaporation filament is heated up to maintain a steady intensity and isotopic ratios are measured until the entire sample is consumed (total evaporation). In this way, mass fractionation effects in the ion source are minimised. Isotopic standards (Pu IRMM-290b/A3, U IRMM-074/10) are measured to correct for these mass fractionation effects and to ensure the traceability to the International system of units (SI). A different approach was needed to correct for mass fractionation effects in Am measurements and is dealt with in "Results and Discussion" section.

The amount content was determined using the following IDMS equation [25]

\[
c_x = c_y \times \frac{m_y}{m_x} \times \frac{R_y - R_b}{R_y - R_x} \times \frac{\Sigma(R_i)_x}{\Sigma(R_i)_y} \quad (1)
\]

c_x is the element amount content of the "unknown" sample [mol g\(^{-1}\)], c_y is the element amount content of the spike [mol g\(^{-1}\)], m_x is the mass of the sample [g], m_y that of the spike [g], and R_x, R_y, and R_b are the isotope amount ratios of the sample (unspiked) [mol mol\(^{-1}\)] and the spike and the blend, respectively. \(\Sigma(R_i)_x\) and \(\Sigma(R_i)_y\) are the sums of all isotope amount ratios in the sample and spike, respectively.

Uncertainties associated with IDMS measurements were estimated according to the Guide to Expression of Uncertainty in Measurement (GUM) using the GUM Workbench©, a software developed by Metrodata GmbH (Germany) [13, 14]. All the uncertainties in this paper are expanded uncertainties with a coverage factor \(k=2\), corresponding to a level of confidence of about 95%.

### Results and Discussion

**IRMM-049d**

IRMM-049d is a highly enriched \(^{242}\)Pu spike material (94.6%) certified for \(^{242}\)Pu isotope amount content and Pu isotope amount ratios. 93 units of this material were produced in 2011. Each unit contains about 10 mL of 5 M nitric acid in screw-cap glass ampoules with the plutonium mass fraction of approximately 0.1 mg Pu g\(^{-1}\) solution [28, 29].

The results of the IDMS measurements of the \(^{242}\)Pu amount content in IRMM-049d are shown in Figs. 3 and 4. Each data point represents an independent measurement result of the selected unit of IRMM-049d (spiking/weighing, chemical separation and replicate measurements).

Uncertainties were estimated according to the Guide to Expression of Uncertainty in Measurement (GUM) using the GUM Workbench©, a software developed by Metrodata GmbH (Germany) [13, 14]. All the uncertainties in this paper are expanded uncertainties with a coverage factor \(k=2\), corresponding to a level of confidence of about 95%.

**Fig. 3** IDMS results of the \(^{242}\)Pu amount content (mol g\(^{-1}\)) measurements in the selected units of IRMM-049d using various spike materials: Eqrain (\(^{239}\)Pu)—green squares (filled green square), IRMM-1027m (\(^{239}\)Pu)—empty circles (open circle), \(^{241}\)Pu—brown circles (filled orange circle), JAEA MOX (\(^{239}\)Pu)—purple triangles (filled purple triangle) and IRMM-042a (\(^{244}\)Pu)—blue triangles (filled blue triangle). Error bars show the expanded uncertainty (\(k=2\)) of the measured result. The red line shows the certified value and the red dashed line the expanded uncertainty (\(k=2\)) of the certified value. (Color figure online)
also demonstrates that there was no significant change in the concentration of IRMM-049d over time.

IRMM-049e

As the IRMM-049d was approaching exhaustion, a new batch of this spike material was prepared in 2017 to ensure the continuous supply of the 242Pu spike material. The same 242Pu source solution was used as for its predecessor (IRMM-049d). 89 units of IRMM-049e material were made available with a similar concentration (i.e. ca. 1 mg Pu in 10 mL, 5 M HNO3) [18].

This batch of 242Pu spike IRM has been included in the stability monitoring after the material certification and release in 2017. The stability measurements were carried out for the amount content of plutonium and for the plutonium isotope amount ratios. This spike material was also included in various external EQRAIN campaigns (see Fig. 2). The results of the IDMS measurements of the 242Pu amount content in IRMM-049e are shown in Figs. 5 and 6. As for Figs. 3 and 4, each data point represents an independent measurement result.

It can be seen from Fig. 5 that the certified value for the 242Pu amount content in IRMM-049e was successfully verified using the selected spike materials. The average of all the measured results (22 units) was 3.5836 (46) × 10−7 mol g−1 with a relative difference of −0.022% from the certified value of 3.5844 (45) × 10−7 mol g−1. Figure 6 shows that there was no significant change in the concentration of IRMM-049e over time.

The evaluation of the results of stability monitoring is also done by calculating the compatibility score ($E_n$) using the equation below.

$$E_n = \frac{x_{\text{CRM}} - x_{\text{mon}}}{\sqrt{u^2_{\text{CRM}} + u^2_{\text{mon}}}}$$

(2)

This approach involves a comparison of each measured result ($x_{\text{mon}}$) with the certified value ($x_{\text{CRM}}$) and takes into account the standard uncertainties ($u_{\text{mon}}$, $u_{\text{CRM}}$) and an appropriate coverage factor. A coverage factor $k=2$ is chosen representing a level of confidence of approximately 95%. An absolute value of $E_n$ less than 2 means that there is no significant difference between the two values, $x_{\text{mon}}$ and $x_{\text{CRM}}$.

The compatibility scores in IRMM-049d and IRMM-049e are shown in Fig. 7. For all the results, a compatibility score (absolute value) smaller than 2 was obtained.

Two units of IRMM-049e were measured (#26 and #61) by TIMS for the determination of the Pu isotope amount ratios. The results of these measurements (October 22, 2020) are shown in Table 1.

All the results for the isotope ratios were in a good agreement with the certified value within measurement uncertainties, as well as highlighted by the compatibility scores (all |$E_n$| < 2).

![Fig. 4](image_url) 242Pu amount content (mol g−1) in the selected units of IRMM-049d as a function of time. Error bars show the expanded uncertainty ($k=2$) of the measured result. Red line shows the certified value and the red dashed line the expanded uncertainty ($k=2$) of the certified value. (Color figure online)
IRMM-0243 is an Am spike IRM, certified for the amount content of $^{243}\text{Am}$ and the $n(^{241}\text{Am})/n(^{243}\text{Am})$ isotope amount. An americium source solution with an isotopic composition of 88% $^{243}\text{Am}$ and 12% $^{241}\text{Am}$ was used for the preparation of this material. 587 units were produced in cooperation with CEA/CETAMA. The concentration of the americium is about 1.5 μg mL$^{-1}$ in 1 M nitric acid solution [9, 19]. This IRM was commercially released in 2017.

Two units of IRMM-0243 were selected (#197 and #313) for the measurement of the $^{243}\text{Am}$ amount content and the Am isotopic composition by IDMS and TIMS, respectively. A $^{241}\text{Am}$ spike, produced by ingrowth from a highly enriched $^{241}\text{Pu}$ material (99.3%) was used as a spike for IDMS measurements. Isotope ratio measurements were carried out on the Triton TIMS using the total evaporation (TE) method.
In the absence of a suitable Am isotopic standard at JRC-Geel, it was not possible to perform a correction for mass fractionation as is normally done for plutonium and uranium. Due to similarities in chemical behaviour and ionization energies, it can be assumed that Am behaves similarly to U or Pu during the total evaporation measurement and that the bias statements for U and Pu can be applied for Am [25]. For the \( n(238\text{Pu})/n(242\text{Pu}) \) ratio, the maximum expected bias of 0.033% \((k = 2, \text{for ratios spanning 2 mass units})\) was applied. In addition, an uncertainty of 0.020% \((k = 2, \text{for ratios spanning 2 mass units})\) derived from isotope measurements of the U and Pu quality control (QC) materials was added as a

| Isotope ratio | Unit no. | TIMS [mol mol\(^{-1}\)] | Certified value [mol mol\(^{-1}\)] | \( E_n \) |
|---------------|---------|--------------------------|-----------------------------------|--------|
| \( n(238\text{Pu})/n(242\text{Pu}) \) | 26      | 0.005066 (11)            | 0.0050666 (81)                    | − 0.09 |
| \( n(238\text{Pu})/n(242\text{Pu}) \) | 61      | 0.005068 (13)            |                                   | 0.18   |
| \( n(239\text{Pu})/n(242\text{Pu}) \) | 26      | 0.0022192 (37)           | 0.0022218 (26)                    | − 0.48 |
| \( n(239\text{Pu})/n(242\text{Pu}) \) | 61      | 0.0022213 (42)           |                                   | − 0.02 |
| \( n(240\text{Pu})/n(242\text{Pu}) \) | 26      | 0.046032 (89)            | 0.046033 (46)                     | − 0.02 |
| \( n(240\text{Pu})/n(242\text{Pu}) \) | 61      | 0.046035 (100)           |                                   | 0.13   |
| \( n(241\text{Pu})/n(242\text{Pu}) \) | 26      | 0.0021882 (61)           | 0.0021930 (23)                    | − 1.47 |
| \( n(241\text{Pu})/n(242\text{Pu}) \) | 61      | 0.0021878 (62)           |                                   | − 1.57 |
| \( n(244\text{Pu})/n(242\text{Pu}) \) | 26      | 0.00025724 (88)          | 0.00025766 (59)                   | − 0.79 |
| \( n(244\text{Pu})/n(242\text{Pu}) \) | 61      | 0.00025714 (110)         |                                   | − 0.90 |

The values in the table are decay corrected to the reference date of January 1, 2017. Expanded uncertainties \((k = 2)\) are shown in parentheses and apply to the last digits of the value

### Table 2 Results of the \(^{243}\text{Am} \) amount content in IRMM-0243

| Amount content | Unit No | IDMS [nmol g\(^{-1}\) sol] | Certified value [nmol g\(^{-1}\) sol] | \( E_n \) |
|----------------|---------|-----------------------------|-----------------------------------|--------|
| \( c^{(243}\text{Am}) \) | 197     | 5.698 (12)                  | 5.696 (11)                        | 0.25   |
| \( c^{(243}\text{Am}) \) | 313     | 5.698 (12)                  | 5.696 (11)                        | 0.29   |

The values are decay corrected to the reference date of January 1, 2017. Expanded uncertainties \((k = 2)\) are shown in parentheses and apply to the last digits of the value

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![Fig. 7 The compatibility score \((E_n)\) in IRMM-049d and IRMM-049e](image-url)
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| Isotope ratio | Unit no | TIMS [mol mol⁻¹] | Certified value [mol mol⁻¹] | $E_u$ |
|--------------|---------|-----------------|-----------------------------|------|
| $n(243\text{Am})/n(241\text{Am})$ | 197 | 0.136065 (60) | 0.136138 (54) | − 1.81 |
| $n(243\text{Am})/n(241\text{Am})$ | 313 | 0.136078 (60) | 0.136138 (54) | − 1.49 |
| $n(243\text{Am})/n(241\text{Am})$ | 3 | 0.136070 (60) | 0.136138 (54) | − 1.68 |

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