



BSI Standards Publication

Nuclear energy — Evaluation of homogeneity of Gd distribution within gadolinium fuel blends and determination of Gd_2O_3 content in gadolinium fuel pellets by measurements of uranium and gadolinium elements

National foreword

This British Standard is the UK implementation of ISO 16424:2012.

The UK participation in its preparation was entrusted to Technical Committee NCE/9, Nuclear fuel cycle technology.

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

This publication does not purport to include all the necessary provisions of a contract. Users are responsible for its correct application.

© The British Standards Institution 2012. Published by BSI Standards Limited 2012

ISBN 978 0 580 73017 7

ICS 27.120.30

Compliance with a British Standard cannot confer immunity from legal obligations.

This British Standard was published under the authority of the Standards Policy and Strategy Committee on 31 December 2012.

Amendments issued since publication

Date	Text affected
------	---------------

**Nuclear energy — Evaluation of
homogeneity of Gd distribution
within gadolinium fuel blends and
determination of Gd_2O_3 content
in gadolinium fuel pellets by
measurements of uranium and
gadolinium elements**

*Énergie nucléaire — Évaluation de l'homogénéité de la distribution
du Gd dans les mélanges de combustibles au gadolinium et
détermination de la teneur en Gd_2O_3 dans les pastilles combustibles
au gadolinium par mesurage des éléments uranium et gadolinium*





COPYRIGHT PROTECTED DOCUMENT

© ISO 2012

All rights reserved. Unless otherwise specified, no part of this publication may be reproduced or utilized in any form or by any means, electronic or mechanical, including photocopying and microfilm, without permission in writing from either ISO at the address below or ISO's member body in the country of the requester.

ISO copyright office
Case postale 56 • CH-1211 Geneva 20
Tel. + 41 22 749 01 11
Fax + 41 22 749 09 47
E-mail copyright@iso.org
Web www.iso.org

Published in Switzerland

Contents

Page

Foreword	iv
1 Scope	1
2 Normative references	1
3 Principle	1
4 Apparatus	2
4.1 High dispersion direct reading ICP-AES.....	2
4.2 Analytical balance.....	2
4.3 Small stainless steel spoon.....	2
5 Reagents	2
6 Reference solutions	2
6.1 Gadolinium element reference solutions.....	2
6.2 Uranium element reference solutions.....	3
7 Sample preparation	3
7.1 Sample preparation for evaluation of Gd homogeneity in gadolinium fuel blend.....	3
7.2 Sample preparation for determining the Gd ₂ O ₃ content of Gd fuel pellets.....	3
8 Calibration and analysis of the samples	3
8.1 Calibration of Gd peak line intensity.....	3
8.2 Calibration of U peak line intensity.....	3
8.3 Evaluation of Gd homogeneity in gadolinium fuel blend.....	3
8.4 Determination of Gd ₂ O ₃ content in gadolinium fuel pellet.....	4
9 Precision	6
10 Test report	6
Annex A (informative) Calibration and Gd and U measurement uncertainties	7
Annex B (informative) Derivation of O/M and O/U ratio formulas	10
Annex C (informative) Evaluation of Gd₂O₃ measurement precision	12
Bibliography	15

Foreword

ISO (the International Organization for Standardization) is a worldwide federation of national standards bodies (ISO member bodies). The work of preparing International Standards is normally carried out through ISO technical committees. Each member body interested in a subject for which a technical committee has been established has the right to be represented on that committee. International organizations, governmental and non-governmental, in liaison with ISO, also take part in the work. ISO collaborates closely with the International Electrotechnical Commission (IEC) on all matters of electrotechnical standardization.

International Standards are drafted in accordance with the rules given in the ISO/IEC Directives, Part 2.

The main task of technical committees is to prepare International Standards. Draft International Standards adopted by the technical committees are circulated to the member bodies for voting. Publication as an International Standard requires approval by at least 75 % of the member bodies casting a vote.

Attention is drawn to the possibility that some of the elements of this document may be the subject of patent rights. ISO shall not be held responsible for identifying any or all such patent rights.

ISO 16424 was prepared by Technical Committee ISO/TC 85, *Nuclear Energy, Nuclear Technologies, and Radiological Protection*, Subcommittee SC 5, *Nuclear Fuel Cycle*.

Nuclear energy — Evaluation of homogeneity of Gd distribution within gadolinium fuel blends and determination of Gd₂O₃ content in gadolinium fuel pellets by measurements of uranium and gadolinium elements

1 Scope

This International Standard is applicable to the evaluation of the homogeneity of Gd distribution within gadolinium fuel blends, and the determination of the Gd₂O₃ content in sintered fuel pellets of Gd₂O₃+UO₂ from 1 % to 10 %, by measurements of gadolinium (Gd) and uranium (U) elements using ICP-AES.

After performing measurements of Gd and U elements using ICP-AES, if statistical methodology is additionally applied, homogeneity of Gd distribution within a Gd fuel pellet lot can also be evaluated. However, this International Standard covers the statistical methodology only on a limited basis.

NOTE 1 ISO 16796 also provides a method for Gd₂O₃ content determination by atomic emission spectrometry using an inductively coupled plasma source (ICP-AES). The methodology of ISO 16796 is different from the one of this International Standard.

NOTE 2 In this International Standard, gadolinium fuel blend represents a mixture of uranium dioxide (UO₂) powder and gadolinium oxide (Gd₂O₃) powder. The physically blended and homogenized powder may additionally contain in it rather large quantities of uranium oxide (U₃O₈) powder particles and/or the M₃O₈ powder particles obtained by oxidation of Gd pellets. In this International Standard, the symbol “M” in the chemical formula “M₃O₈” and in the terminology “O/M ratio” represents metallic elements U and Gd.

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 amendments) applies.

ISO 3696:1987, *Water for analytical laboratory use — Specification and test methods*

3 Principle

If the Gd and U element contents and the oxygen to metal atomic ratio (commonly referred to as O/M ratio) in a gadolinium fuel pellet are measured or determined, the Gd₂O₃ content of that pellet can be determined by calculation based on the stoichiometry of the pellet. The stoichiometric compositions for Gd and U will depend upon pellet manufacturing specification. If the specification requires that the Gd₂O₃ content in the pellet be 6 % as mass fraction, after manufacturing, the ratio of total Gd mass to total U mass in that pellet will be close to 0,063.

The Gd and U element content values measured from a powder blend can make it possible to evaluate whether Gd distribution in the powder is sufficiently homogeneous. Moreover, the two values make it possible to estimate accurately the actual Gd₂O₃ content of the pellet after sintering. The estimated Gd₂O₃ content can be used to anticipate whether the Gd pellets to be produced will meet Gd₂O₃ content specifications or not.

Impurity interferences have not been observed for the usual samples of the nuclear grade material. Very small quantities of impurity elements which might be contained in the samples do not affect the principle of this International Standard.

NOTE Even in the case of M_3O_8 powder or mixture of UO_2 powder and Gd_2O_3 powder, if the Gd and U element contents and the O/M ratio are measured or determined, the Gd_2O_3 content (or Gd content) of that powder can be determined by calculation based on the stoichiometry of the powder. Using the Gd_2O_3 content (or Gd content) thus obtained and the O/M ratio, the uranium concentration factor of the powder can also be calculated so as to obtain accurate uranium accounting data.

4 Apparatus

4.1 High dispersion direct reading ICP-AES

The measured values should be indicated down to at least two decimal places when element concentrations in the range from 1 mg/l to 100 mg/l are measured. For determination of Gd and U element concentrations, 354,580 nm and 398,579 nm peaks in atomic emission spectrum are used, respectively. However, instead of the two peaks, other peaks are also available.

4.2 Analytical balance

The sensitivity of the balance is $\pm 0,1$ mg.

4.3 Small stainless steel spoon

The small spoon is able to accommodate powder in the range from 1 mg to 100 mg.

4.4 Hot plate.

4.5 Glass beakers or polytetrafluoroethylene (PTFE) beakers.

4.6 Micropipettes.

4.7 Volumetric flasks.

4.8 Percussion mortar.

5 Reagents

5.1 **Nitric acid**, aqueous solution at a volume fraction of 50 %.

5.2 **Water**, distilled water or water complying with grade 3 of ISO 3696:1987.

6 Reference solutions

6.1 Gadolinium element reference solutions

Three Gd reference solutions whose Gd concentrations are 5 mg/l, 10 mg/l and 15 mg/l, respectively, are used for calibration. These Gd reference solutions¹⁾ are the ones that have been manufactured by diluting a reference material with the water specified in 5.2.

1) A certified gadolinium reference material from NIST (National Institute of Standards and Technology, USA) is an example of a suitable product available commercially. This information is given for the convenience of users of this document and does not constitute an endorsement of this product by ISO.

6.2 Uranium element reference solutions

Three U reference solutions whose U concentrations are 50 mg/l, 70 mg/l and 100 mg/l, respectively, are used for calibration. These U reference solutions²⁾ are the ones that have been manufactured by diluting a reference material with the water specified in 5.2.

7 Sample preparation

7.1 Sample preparation for evaluation of Gd homogeneity in gadolinium fuel blend

Using a small spoon, take randomly at least five different powder samples from the blended powder, then dissolve each of the samples in the nitric acid solution of 5.1. Number each beaker individually. The U concentrations of the sample solutions prepared should be somewhere in the range from 50 mg/l to 100 mg/l.

To evaluate the homogeneity of the Gd distribution within a Gd fuel pellet lot, take at least five different pellet samples from the lot and then crush each of the pellets into small pellet fragments as described in 7.2.

NOTE 1 If about 2 mg of the powder sample is dissolved in 20 ml of the nitric acid solution, the required uranium concentration range will be obtained. Alternatively, if about 100 mg of sample is dissolved and then diluted by proper pipetting and addition of distilled water, the concentration range can also be obtained.

7.2 Sample preparation for determining the Gd₂O₃ content of Gd fuel pellets

After crushing a gadolinium fuel pellet using a percussion mortar, randomly take a small sample from the pellet fragments, and dissolve it in the nitric acid solution. The U concentration should be somewhere in the range from 50 mg/l to 100 mg/l.

8 Calibration and analysis of the samples

8.1 Calibration of Gd peak line intensity

The three Gd reference solutions of 6.1 are used to calibrate the intensity of the Gd peak line. The Gd peak line intensity is repeatedly measured to establish an intensity-versus-concentration curve (regression curve) needed for determination of Gd concentrations. See A.1.

8.2 Calibration of U peak line intensity

The three U reference solutions of 6.2 are used to calibrate the intensity of the U peak line. The U peak line intensity is repeatedly measured to establish an intensity-versus-concentration curve (regression curve) needed for determination of U concentrations. See A.2.

8.3 Evaluation of Gd homogeneity in gadolinium fuel blend

8.3.1 Estimated Gd₂O₃ content

Measure the concentrations of Gd and U elements contained in the sample solutions (at least five different solutions) that were prepared according to 7.1. Each sample solution is repeatedly measured six times, and the six measurement values of Gd and U each are averaged. Each sample solution will be given a Gd

2) A certified uranium reference material from NIST (National Institute of Standards and Technology, USA) is an example of a suitable product available commercially. This information is given for the convenience of users of this document and does not constitute an endorsement of this product by ISO.

average and a U average. After all of the Gd concentration average and U concentration average sets have been obtained, the estimated Gd₂O₃ content for each sample is calculated by the following formula:

$$w_{\text{Gd}_2\text{O}_3, \text{est}} = \frac{\left(\frac{1}{2}\right)(100)\left[\frac{(15,999\ 4 \times 3)}{157,25} + 2\right](\bar{\rho}_{\text{Gd}})}{(\bar{\rho}_{\text{Gd}} + \bar{\rho}_{\text{U}}) + (2)(15,999\ 4) \left(\frac{\bar{\rho}_{\text{Gd}}}{157,25} + \frac{\bar{\rho}_{\text{U}}}{238,03}\right)}$$

$$= \frac{(0,5)(100)(2,305\ 2\ \bar{\rho}_{\text{Gd}})}{(\bar{\rho}_{\text{Gd}} + \bar{\rho}_{\text{U}}) + (2)(0,1017\ \bar{\rho}_{\text{Gd}} + 0,067\ 2\ \bar{\rho}_{\text{U}})}$$

where

$w_{\text{Gd}_2\text{O}_3, \text{est}}$ is the estimated Gd₂O₃ content, mass fraction (%) as given by Gd₂O₃/(Gd,U)O₂;

157,25 is the atomic mass of gadolinium (Gd);

238,03 is the atomic mass of uranium (U);

15,9994 is the atomic mass of oxygen (O);

$\bar{\rho}_{\text{Gd}}$ is the Gd concentration average in mg/l, indicated down to two decimal places;

$\bar{\rho}_{\text{U}}$ is the U concentration average in mg/l, indicated down to two decimal places.

If five different sample solutions have been prepared and measured, five estimated values of Gd₂O₃ content will be obtained by this procedure.

NOTE The estimated Gd₂O₃ content represents the most possible Gd₂O₃ content of the expected Gd pellet to be manufactured from the powder blend. As a general rule, after the powder blend has been sintered, the O/M ratio of manufactured Gd pellet will be 2 or very close to 2. Therefore, the actual Gd₂O₃ content of the pellet will be very close to this estimated Gd₂O₃ content.

8.3.2 Evaluation of homogeneity of Gd distribution

Evaluate the homogeneity of Gd distribution within the blended powder by applying a statistical methodology to the estimated contents of Gd₂O₃ obtained in Clause 8.3.1. The average and the standard deviation calculated from at least five estimated values of Gd₂O₃ content will be used for the evaluation.

EXAMPLE If the homogeneity specification of the blended powder requires that one standard deviation of estimated Gd₂O₃ content values of five blended powder samples randomly taken shall be within 1 % relative from the mean of the five values, and if the mean and the standard deviation of the five estimated Gd₂O₃ contents are 6,03 % and 0,04 %, respectively, it can be said that the blended powder meets the homogeneity specification, because all of the related inequalities, i.e. $6,03 - (6,03 \times 0,01) < 6,03 - 0,04$ and $6,03 + 0,04 < 6,03 + (6,03 \times 0,01)$, are correct.

NOTE The statistical methodology and limitations will depend on the quality assurance policy and goals of the organization concerned. To find detailed information on the applicable statistical methods other than the above evaluation example, see ISO 3951-4.

8.4 Determination of Gd₂O₃ content in gadolinium fuel pellet

8.4.1 Preliminary Gd₂O₃ content

Repeatedly measure six times Gd and U concentrations in the sample solution prepared according to Clause 7.2, and calculate the concentration average of Gd and U each. The preliminary Gd₂O₃ content in the pellet can be determined by the following formula:

$$w_{\text{Gd}_2\text{O}_3, \text{pre}} = \frac{(100)(157,25 \times 2 + 15,999\ 4 \times 3)(Z)}{2(1 - Z)(238,03 + 15,999\ 4 \times 2) + (157,25 \times 2 + 15,999\ 4 \times 3)(Z)}$$

$$Z = \frac{\left(\frac{\bar{\rho}_{\text{Gd}}}{157,25}\right)}{\left(\frac{\bar{\rho}_{\text{Gd}}}{157,25} + \frac{\bar{\rho}_{\text{U}}}{238,03}\right)}$$

where

$w_{\text{Gd}2\text{O}3,\text{pre}}$ is the preliminary Gd_2O_3 content, mass fraction in %;

$\bar{\rho}_{\text{Gd}}$ is the Gd concentration average in mg/l, indicated down to two decimal places;

$\bar{\rho}_{\text{U}}$ is the U concentration average in mg/l, indicated down to two decimal places.

NOTE The preliminary Gd_2O_3 content represents the interim value before obtaining the final Gd_2O_3 content value of the Gd pellet. From the aspect of the stoichiometry of the Gd pellet, this preliminary Gd_2O_3 content value is not complete theoretically.

8.4.2 O/M ratio of the Gd pellet

The O/M ratio, B , of the pellet is determined by the following formula:

$$B = 2 - \frac{(m_2 - m_1)}{(m_2)(0,0592) + (m_1)(w_{\text{Gd}2\text{O}3,\text{pre}})(0,0264)(0,01)}$$

where

B is the O/M ratio, and will be indicated down to at least two decimal places;

m_1 is the sample mass before equilibration in grams, indicated down to four decimal places;

m_2 is the sample mass after equilibration in grams, indicated down to four decimal places;

$w_{\text{Gd}2\text{O}3,\text{pre}}$ is the preliminary Gd_2O_3 content (mass fraction of Gd_2O_3 in %) obtained in 8.4.1.

NOTE 1 The above formula is used to determine the O/M ratio by an atmospheric equilibration method. For application of this method, a Gd pellet goes through an oxidation-reduction process at $900\text{ °C} \pm 20\text{ °C}$ for 4 h in a mixed gas of argon (or nitrogen) at a volume fraction of 96 % and hydrogen at a volume fraction of 4 %; the mixed gas also contains a small volume fraction of water vapor. The Gd pellet masses before and after equilibration by oxidation-reduction and Gd_2O_3 content value are used to determine the O/M ratio. See ASTM C1430-07.^[4]

NOTE 2 The derivation of the formulas for O/M and O/U ratios of Gd pellet can be seen in Annex B.

8.4.3 Final Gd_2O_3 content

The Gd_2O_3 content in the sampled pellet is finally determined by the following formula:

$$\begin{aligned} w_{\text{Gd}2\text{O}3,\text{fin}} &= \frac{\left(\frac{1}{2}\right)(100)\left[\frac{(15,999\ 4 \times 3)}{157,25} + 2\right](\bar{\rho}_{\text{Gd}})}{(\bar{\rho}_{\text{Gd}} + \bar{\rho}_{\text{U}}) + (B)(15,999\ 4) \left(\frac{\bar{\rho}_{\text{Gd}}}{157,25} + \frac{\bar{\rho}_{\text{U}}}{238,03}\right)} \\ &= \frac{(0,5)(100)(2,305\ 2\ \bar{\rho}_{\text{Gd}})}{(\bar{\rho}_{\text{Gd}} + \bar{\rho}_{\text{U}}) + (B)(0,101\ 7\ \bar{\rho}_{\text{Gd}} + 0,067\ 2\ \bar{\rho}_{\text{U}})} \end{aligned}$$

where

- $w_{\text{Gd}_2\text{O}_3, \text{fin}}$ is the final Gd_2O_3 content in %, i.e. mass fraction of Gd_2O_3 (%) as given by $\text{Gd}_2\text{O}_3/(\text{Gd,U})\text{O}_2$;
- $\bar{\rho}_{\text{Gd}}$ is the Gd concentration average that was obtained in 8.4.1;
- $\bar{\rho}_{\text{U}}$ is the U concentration average that was obtained in 8.4.1;
- B is the O/M ratio that was obtained in 8.4.2.

Although nominal O/M ratio 2 is substituted into the above formula in order to determine the final Gd_2O_3 content, instead of the O/M ratio B that was directly measured in 8.4.2, the results will almost be the same. Therefore, the use of nominal O/M ratio may be permitted. In this case, the formula will become the same as the one that was introduced to obtain the estimated Gd_2O_3 content in 8.3.1.

9 Precision

The possible uncertainty of Gd_2O_3 content determination using the method of this International Standard, excluding systematic uncertainty such as calibration uncertainty, is about 0,2 % relative (0,012 % absolute) at nominal Gd_2O_3 content of 6 %.

To find more information regarding precision of Gd_2O_3 content determination using the method of this International Standard, see Annex C.

10 Test report

The test report shall include at least the following information:

- a) identification of sample, i.e. lot number of powder or pellet, etc.;
- b) the reference of the method used;
- c) the measurement results and their units;
- d) any unusual features noted during the measurements;
- e) any operations not included in this International Standard.

Annex A (informative)

Calibration and Gd and U measurement uncertainties

A.1 Calibration of Gd line intensity and residual mean square

Table A.1 — Gd line intensity-versus-concentration of Gd reference solution

Measurements	Measured intensity 5 mg/l solution	Measured intensity 10 mg/l solution	Measured intensity 15 mg/l solution
1st measurement	572 478,511	1 127 995,396	1 682 373,714
2nd measurement	568 482,882	1 118 771,581	1 692 754,967
3rd measurement	567 587,661	1 124 687,706	1 678 890,058
4th measurement	573 010,558	1 119 946,173	1 674 248,737
5th measurement	567 155,019	1 114 052,914	1 685 414,131
6th measurement	568 116,061	1 131 790,625	1 688 138,089
Mean Value	569 471,782	1 122 874,065	1 683 636,616
Standard Deviation	2 580,558	6 515,181	6 615,467

Using the least square method, the regression curve equation of Gd line intensity-versus-concentration can be obtained as follows:

$$\rho_{Gd} = a + bI_{Gd} = -0,1 + 8,975 \times 10^{-6}I_{Gd}$$

where

I_{Gd} is the measured Gd line intensity in generic units;

ρ_{Gd} is the Gd concentration of the solution in mg/l;

a and b are the coefficients of the regression curve ($a = -0,100$ and $b = 8,975 \times 10^{-6}$).

The residual mean square and regression standard deviation are calculated as follows:

$$\begin{aligned} (\sigma_{Gd})^2 &= \{\sum(\rho_{Gd\ i} - a - bI_{Gd\ i})^2\}/(n - 2), \text{ (} n \text{ is 3, and } \sum \text{ is the summation from } i = 1 \text{ to 3.)} \\ &= 0,000\ 728 \text{ (residual mean square)} \end{aligned}$$

$$\therefore \sigma_{Gd} = 0,027 \text{ (mg/l) (regression standard deviation)}$$

A.2 Calibration of U line intensity and residual mean square

Table A.2 — U line intensity-versus-concentration of U reference solution

Measurements	Measured intensity 50 mg/l solution	Measured intensity 70 mg/l solution	Measured intensity 100 mg/l solution
1st measurement	53 409,807	73 762,528	107 245,053
2nd measurement	53 509,371	73 878,279	107 085,192
3rd measurement	53 300,306	73 633,468	108 205,616
4th measurement	52 518,142	73 598,608	107 540,834
5th measurement	53 102,651	74 037,009	107 383,643
6th measurement	53 208,716	74 061,585	107 607,147
Mean Value	53 174,832	73 828,580	107 511,248
Standard Deviation	352,271	197,747	390,098

Using the least square method, the regression curve equation of U line intensity-versus-concentration can be obtained as follows:

$$\rho_U = a + bI_U = 1,627 + 9,173 \times 10^{-4}I_U$$

where

I_U is the measured U line intensity in generic units;

ρ_U is the U concentration of the solution in mg/l;

a and b are the coefficients of the regression curve ($a = 1,627$ and $b = 9,173 \times 10^{-4}$).

The residual mean square and regression standard deviation are calculated as follows:

$$(\sigma_U)^2 = \{\sum(\rho_{U_i} - a - bI_{U_i})^2\}/(n - 2), \text{ (} n \text{ is 3, and } \sum \text{ is the summation from } i = 1 \text{ to 3.)}$$

$$= 0,647\ 036 \text{ (residual mean square)}$$

$$\therefore \sigma_U = 0,804 \text{ (mg/l) (regression standard deviation)}$$

NOTE 1 If only one of the three reference solutions is used for calibration of the Gd line intensity, the regression standard deviation σ_{Gd} (0,027 mg/l) can simply be regarded as calibration uncertainty of the Gd line intensity. Similarly, the regression standard deviation σ_U (0,804 mg/l) can also be regarded as calibration uncertainty of the U line intensity.

NOTE 2 The calibration uncertainties in this International Standard are mainly due to the equipment's random errors and pipetting errors that arise while manufacturing the reference solutions from the certified reference materials.

A.3 Measurement uncertainties of Gd and U concentrations

After the correlation coefficient between a and b , and the uncertainties of regression coefficients a and b were calculated and the rules for error propagation were applied to the equations $\rho_{Gd} = -0,1 + 8,975 \times 10^{-6} I_{Gd}$ and $\rho_U = 1,627 + 9,173 \times 10^{-4} I_U$ to obtain the combined uncertainties of Gd and U concentration measurements. The combined uncertainties thus obtained can be seen in [Tables A.3](#) and [A.4](#).

Table A.3 — Measurement uncertainty of Gd concentration

Intended Gd concentration of reference solution mg/l	Calibration uncertainty mg/l	Random uncertainty mg/l	Combined uncertainty mg/l
5	0,024 3	0,009 5	0,026 1
10	0,014 6	0,023 9	0,028 0
15	0,023 7	0,024 3	0,033 9

Table A.4 — Measurement uncertainty of U concentration

Intended U concentration of reference solution mg/l	Calibration uncertainty mg/l	Random uncertainty mg/l	Combined uncertainty mg/l
50	0,696 0	0,131 9	0,708 4
70	0,473 2	0,074 1	0,478 9
100	0,765 5	0,146 1	0,779 3

NOTE When calculating the above measurement uncertainties, the uncertainties of the certified reference materials that were used to manufacture the reference solutions were disregarded, because they were small.

Annex B (informative)

Derivation of O/M and O/U ratio formulas

B.1 Derivation of O/M ratio formula

Based on the stoichiometry of sintered Gd fuel pellet, the following two simultaneous Formulae (B.1) and (B.2) can be established. From the equations, the formula for the O/M ratio, B , is derived as follows:

$$m_1 = (m_U + m_{Gd}) + \{(m_U/238,03) + (m_{Gd}/157,25)\} (B) (15,999\ 4) \quad (B.1)$$

$$m_2 = (m_U + m_{Gd}) + \{(m_U/238,03) + (m_{Gd}/157,25)\} (2) (15,999\ 4) \quad (B.2)$$

where

m_1 is the mass of the Gd pellet sample before equilibration, in grams;

m_2 is the mass of the Gd pellet sample after equilibration, in grams;

m_U is the quantity of uranium contained in the sample, in grams;

m_{Gd} is the quantity of gadolinium contained in the sample, in grams;

238,03 is the atomic mass of uranium;

157,25 is the atomic mass of gadolinium;

15,999 4 is the atomic mass of oxygen;

B is the O/M ratio of Gd pellet before equilibration

$$m_2 - m_1 = \{(m_U/238,03) + (m_{Gd}/157,25)\} (2) (15,999\ 4)$$

$$- \{(m_U/238,03) + (m_{Gd}/157,25)\} (B) (15,999\ 4)$$

$$\therefore B = 2 - (m_2 - m_1) / [\{(m_U/238,03) + (m_{Gd}/157,25)\} (15,999\ 4)]$$

$$= 2 - (m_2 - m_1) / \{0,067\ 2 (m_U) + 0,101\ 7 (m_{Gd})\}$$

$$m_{Gd} = \{(314,5)/(362,498\ 2)\} (w_{Gd2O3}) (0,01) (m_1) = 0,008\ 676 (w_{Gd2O3}) (m_1)$$

w_{Gd2O3} = the Gd₂O₃ mass fraction (%) as given by Gd₂O₃/(Gd,U)O₂ before equilibration

$$m_2 = \{1 + (2 \times 15,999\ 4)/(238,03)\} (m_U) + \{1 + (2 \times 15,999\ 4)/(157,25)\} (m_{Gd})$$

$$\therefore m_U = 0,881\ 5 m_2 - 0,920\ 42 (0,01) (w_{Gd2O3}) (m_1)$$

$$\therefore B = 2 - (m_2 - m_1) / \{(0,059\ 2)(m_2) + (0,026\ 4) (w_{Gd2O3}) (m_1)\} \times 0,01$$

B.2 Derivation of O/U ratio formula

If the O/U ratio is defined as the ratio of the number of oxygen atoms to the number of uranium atoms contained in the Gd fuel pellet, the formula for the O/U ratio, D , is derived as follows:

$$\begin{aligned}
 N_{\text{oxygen}} &= \{m_1 - (m_U + m_{\text{Gd}})\} / (15,999\ 4) \\
 &= m_1 \{0,062\ 5 + (0,003\ 301) (0,01) (w_{\text{Gd}203})\} - m_2 (0,055\ 10) \\
 N_{\text{uranium}} &= \{(0,881\ 50) (m_2) - (0,920\ 42) (0,01) (w_{\text{Gd}203}) (m_1)\} / (238,03) \\
 \therefore D &= [\{(0,062\ 5) + (0,003\ 301) (0,01) (w_{\text{Gd}203})\} (m_1) - (0,055\ 10) (m_2)] / \{(0,003\ 703) (m_2) - (0,003\ 867) \times (0,01) (w_{\text{Gd}203}) (m_1)\}
 \end{aligned}$$

where

- N_{oxygen} is the number of oxygen atoms contained in Gd fuel pellet;
- N_{uranium} is the number of uranium atoms contained in Gd fuel pellet;
- D is O/U ratio

Annex C (informative)

Evaluation of Gd₂O₃ measurement precision

C.1 Experiment for Gd₂O₃ measurement precision evaluation

A gadolinium oxide pellet having 6 % (mass fraction) of nominal Gd₂O₃ content was used in an experiment to evaluate the precision of Gd₂O₃ content determined by measurements of the Gd and U elements. A small quantity of sample was taken from that crushed pellet and the sample was dissolved in a nitric acid solution at a volume fraction of 50 %. Then, the Gd concentration and the U concentration of the solution were measured six times by ICP-AES, respectively. The data obtained from the experiment can be seen in [Table C.1](#).

Table C.1 — Measurement data obtained from experiment for precision evaluation

Measurements	Gd concentration ρ_{Gd} , mg/l	U concentration ρ_U , mg/l
1st measurement	5,32	85,74
2nd measurement	5,38	85,46
3rd measurement	5,33	85,49
4th measurement	5,36	85,52
5th measurement	5,36	85,55
6th measurement	5,37	85,41
Mean Value	5,353	85,528
Standard Deviation	0,023	0,14

From the two averaged values, i.e. the averaged Gd concentration value 5,353 mg/l and the averaged U concentration value 85,528 mg/l, the Gd₂O₃ content of the sample pellet was determined at 5,963 % (mass fraction). When determining the Gd₂O₃, nominal value 2 was used as O/M ratio B. From the above data, it can be said that because the average of six measurement values has been used, the standard deviation of the Gd concentration mean is $0,023/(6)^{1/2} = 0,009\ 4$ (mg/l). By similar procedure, the standard deviation of the U concentration mean is $0,114/(6)^{1/2} = 0,046$ (mg/l).

C.2 Calculation of combined uncertainty using error propagation rules

It was assumed that measurements of the Gd and U concentrations and determination of the O/M ratio are independent from one another. It was also assumed that the uncertainty of the O/M ratio determination is 0,01 for convenience of calculation. Strictly speaking, from the equation used to obtain the O/M ratio in 8.4.2, the O/M ratio determination is dependent upon the measurements of Gd and U concentrations. However, because the extent of uncertainty of O/M ratio determination does not exceed 0,01 at any rate empirically, such assumptions will be conservative when evaluating the extent

of combined uncertainty. This will make the calculation of combined uncertainty much easier. Under these conditions, the combined uncertainty was calculated as follows:

$$u_c^2 = (\partial C/\partial \bar{\rho}_{Gd})^2 u_{Gd}^2 + (\partial C/\partial \bar{\rho}_U)^2 u_u^2 + (\partial C/\partial B)^2 u_{O/M}^2 \quad (C.1)$$

where C represents the equation to obtain Final Gd₂O₃ content ($w_{Gd2O3,fin}$) in 8.4.3.

$$\partial C/\partial \bar{\rho}_{Gd} = \{[115,26 \bar{\rho}_{Gd} + \bar{\rho}_U + B(0,1017 \bar{\rho}_{Gd} + 0,0672 \bar{\rho}_U)] + 115,26 \bar{\rho}_{Gd} (1 + 0,1017 B)\} /$$

$$\{\bar{\rho}_{Gd} + \bar{\rho}_U + B(0,1017 \bar{\rho}_{Gd} + 0,0672 \bar{\rho}_U)\}^2$$

$$\partial C/\partial \bar{\rho}_U = (115,26 \bar{\rho}_{Gd})(1 + 0,1017 B) / \{\bar{\rho}_{Gd} + \bar{\rho}_U + B(0,1017 \bar{\rho}_{Gd} + 0,0672 \bar{\rho}_U)\}^2$$

$$\partial C/\partial B = (115,26 \bar{\rho}_{Gd})(0,1017 \bar{\rho}_{Gd} + 0,0672 \bar{\rho}_U) / \{\bar{\rho}_{Gd} + \bar{\rho}_U + B(0,1017 \bar{\rho}_{Gd} + 0,0672 \bar{\rho}_U)\}^2$$

u_c is the combined uncertainty;

u_{Gd} is 0,009 4 (the measurement uncertainty of Gd concentration mean obtained in C.1);

u_u is 0,046 (measurement uncertainty of U concentration mean obtained in C.1);

$u_{O/M}$ is 0,01 (determination uncertainty of O/M ratio assumed in C.1).

$$\bar{\rho}_{Gd} = 5,353; \bar{\rho}_U = 85,528; B = 2$$

(These values are originated from C.1).

$$u_c^2 = (1,183)^2(0,0094)^2 + (0,065)^2(0,046)^2 + (0,363)^2(0,01)^2 \quad (C.2)$$

$$= 0,000124 + 0,00000894 + 0,0000132$$

$$= 0,000146$$

∴ Combined uncertainty, u_c , is 0,012 % absolute as one standard deviation

In conclusion, the measurement precision of Gd₂O₃ content, excluding systematic uncertainty such as calibration uncertainty, is about 0,012 % absolute (0,2 % relative) at nominal Gd₂O₃ content of 6 %.

NOTE Based on the Formulae (C.1) and (C.2), the most dominant factor contributing to the precision of the Gd₂O₃ content determination by this International Standard is the measurement uncertainty of the Gd concentration in the nitric acid solution.

C.3 Measurement data for sample pellets of nominal Gd₂O₃ contents 4 % and 8 %

Table C.2 — Measurement data for a sample pellet of nominal Gd₂O₃ content 4 %

Measurements	Gd concentration ρ_{Gd} , mg/l	U concentration ρ_U , mg/l
1st measurement	3,83	94,09
2nd measurement	3,82	93,78
3rd measurement	3,84	93,70

Table C.2 (continued)

Measurements	Gd concentration ρ_{Gd} , mg/l	U concentration ρ_{U} , mg/l
4th measurement	3,81	92,00
5th measurement	3,83	93,67
6th measurement	3,80	93,47
Mean Value	3,822	93,452
Standard Deviation	0,015	0,739

From the data in [Table C.2](#), the Gd₂O₃ content of the sample pellet was determined at 3,983 % (mass fraction). Nominal value, 2, was used as O/M ratio *B*.

Table C.3 — Measurement data for a sample pellet of nominal Gd₂O₃ content 8 %

Measurements	Gd concentration ρ_{Gd} , mg/l	U concentration ρ_{U} , mg/l
1st measurement	7,35	83,91
2nd measurement	7,38	84,20
3rd measurement	7,33	83,54
4th measurement	7,34	83,85
5th measurement	7,28	83,11
6th measurement	7,23	83,21
Mean Value	7,318	83,637
Standard Deviation	0,054	0,426

From the data in [Table C.3](#), the Gd₂O₃ content of the sample pellet was determined at 8,135 % (mass fraction). Nominal value, 2, was used as O/M ratio *B*.

Bibliography

- [1] ISO 16796:2004, *Nuclear energy – Determination of Gd₂O₃ content in gadolinium fuel blends and gadolinium fuel pellets by atomic emission spectrometry using an inductively coupled plasma source ICP-AES*
- [2] ISO 3696:1987, *Water for analytical laboratory use — Specification and test methods*
- [3] ISO 3951-4:2011, *Sampling procedures for inspection by variables — Part 4: Procedures for assessment of declared quality levels*
- [4] ASTM C1430-07, *Standard Test Method for Determination of Uranium, Oxygen to Uranium (O/U), and Oxygen to Metal (O/M) in Sintered Uranium Dioxide and Gadolinia-Uranium Dioxide Pellets by Atmospheric Equilibration*

British Standards Institution (BSI)

BSI is the national body responsible for preparing British Standards and other standards-related publications, information and services.

BSI is incorporated by Royal Charter. British Standards and other standardization products are published by BSI Standards Limited.

About us

We bring together business, industry, government, consumers, innovators and others to shape their combined experience and expertise into standards-based solutions.

The knowledge embodied in our standards has been carefully assembled in a dependable format and refined through our open consultation process. Organizations of all sizes and across all sectors choose standards to help them achieve their goals.

Information on standards

We can provide you with the knowledge that your organization needs to succeed. Find out more about British Standards by visiting our website at bsigroup.com/standards or contacting our Customer Services team or Knowledge Centre.

Buying standards

You can buy and download PDF versions of BSI publications, including British and adopted European and international standards, through our website at bsigroup.com/shop, where hard copies can also be purchased.

If you need international and foreign standards from other Standards Development Organizations, hard copies can be ordered from our Customer Services team.

Subscriptions

Our range of subscription services are designed to make using standards easier for you. For further information on our subscription products go to bsigroup.com/subscriptions.

With **British Standards Online (BSOL)** you'll have instant access to over 55,000 British and adopted European and international standards from your desktop. It's available 24/7 and is refreshed daily so you'll always be up to date.

You can keep in touch with standards developments and receive substantial discounts on the purchase price of standards, both in single copy and subscription format, by becoming a **BSI Subscribing Member**.

PLUS is an updating service exclusive to BSI Subscribing Members. You will automatically receive the latest hard copy of your standards when they're revised or replaced.

To find out more about becoming a BSI Subscribing Member and the benefits of membership, please visit bsigroup.com/shop.

With a **Multi-User Network Licence (MUNL)** you are able to host standards publications on your intranet. Licences can cover as few or as many users as you wish. With updates supplied as soon as they're available, you can be sure your documentation is current. For further information, email bsmusales@bsigroup.com.

BSI Group Headquarters

389 Chiswick High Road London W4 4AL UK

Revisions

Our British Standards and other publications are updated by amendment or revision.

We continually improve the quality of our products and services to benefit your business. If you find an inaccuracy or ambiguity within a British Standard or other BSI publication please inform the Knowledge Centre.

Copyright

All the data, software and documentation set out in all British Standards and other BSI publications are the property of and copyrighted by BSI, or some person or entity that owns copyright in the information used (such as the international standardization bodies) and has formally licensed such information to BSI for commercial publication and use. Except as permitted under the Copyright, Designs and Patents Act 1988 no extract may be reproduced, stored in a retrieval system or transmitted in any form or by any means – electronic, photocopying, recording or otherwise – without prior written permission from BSI. Details and advice can be obtained from the Copyright & Licensing Department.

Useful Contacts:

Customer Services

Tel: +44 845 086 9001

Email (orders): orders@bsigroup.com

Email (enquiries): cservices@bsigroup.com

Subscriptions

Tel: +44 845 086 9001

Email: subscriptions@bsigroup.com

Knowledge Centre

Tel: +44 20 8996 7004

Email: knowledgecentre@bsigroup.com

Copyright & Licensing

Tel: +44 20 8996 7070

Email: copyright@bsigroup.com



...making excellence a habit.™