

Standard Guide for the Determination of Iodine-129 In Uranium Oxide¹

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1. Scope

- 1.1 This method covers the determination of iodine-129 ($^{129}\mathrm{I}$) in uranium oxide by gamma-ray spectrometry. The method could also be applicable to the determination of $^{129}\mathrm{I}$ in aqueous matrices.
- 1.2 This standard does not purport to address all of the safety concerns, if any, associated with its use. It is the responsibility of the user of this standard to establish appropriate safety and health practices and to determine the applicability of regulatory limitations prior to use.

2. Referenced Documents

2.1 ASTM Standards:²

C1402 Guide for High-Resolution Gamma-Ray Spectrometry of Soil Samples

D1193 Specification for Reagent Water

D3648 Practices for the Measurement of Radioactivity

D3649 Practice for High-Resolution Gamma-Ray Spectrometry of Water

3. Summary of Practice

- 3.1 An aliquot of uranium oxide is dissolved in dilute nitric acid and the iodine is selectively extracted via liquid-liquid extraction. The iodine is further purified by selective precipitation and counted by gamma-ray spectrometry.
- 3.2 Gravimetric tracer recoveries using this method are typically between 75 and 90 %.
- 3.3 The minimum detectable activity (MDA) will vary with chemical yield, sample size, instrument background, counting time and counting efficiency. For a sample size of 100 mg U oxide, using a well shielded detector, a 1000 minute counting time, and 32 % detector efficiency at 30 keV, a MDA of ≤0.74 Bq/g (20 pCi/g) oxide was achieved.

4. Significance and Use

4.1 The determination of ¹²⁹I is not typically requested in nuclear fuel specifications however it is commonly requested for disposal of the spent fuel, or for disposal of excess uranium from national weapon complexes. This practice can provide results of sufficient quality for waste disposal repositories.

5. Interferences

- 5.1 Incomplete removal of uranium and its ²³⁴Th/^{234m}Pa daughters could lead to elevated Compton background in the low energy region of the gamma-ray spectrum, where the ¹²⁹I x-rays are counted.
- 5.2 Because the iodine yield monitor is added after the oxide dissolution, any loss of ¹²⁹I during the dissolution step will not be monitored and may lead to results that are biased low. To minimize any iodine loss, avoid prolonged heating of the sample and minimize the time the sample is in an acidic state.

6. Instrumentation

6.1 Extended-range or low-energy gamma ray spectrometry system. See C1402, D3648 or D3649 for a general description of gamma-ray spectrometry systems. The system used to measure the low-energy x-rays from ¹²⁹I should have a thin window to allow the efficient penetration and measurement of the low-energy x-rays.

7. Terms and Definitions

- 7.1 *ROI:* Region-of-Interest; the channels, or region, in the spectra in which the counts due to a specific radioisotope appear on a functioning, calibrated gamma-ray spectrometry system.
- 7.2 Reagent blank: reagent water processed the same as the samples; used in the determination of the minimum detectable activity.

8. Apparatus

- 8.1 Plastic bottles, 30 and 60-ml, or separatory funnels
- 8.2 Filter paper—25-mm diameter, 0.45µm pore size
- 8.3 Vacuum filter apparatus
- 8.4 pH paper with unit resolution

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² For referenced ASTM standards, visit the ASTM website, www.astm.org, or contact ASTM Customer Service at service@astm.org. For *Annual Book of ASTM Standards* volume information, refer to the standard's Document Summary page on the ASTM website.

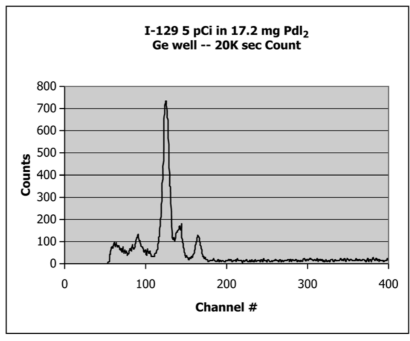


FIG. 1 Low-Energy Photon Spectrum of I-129 on a Ge Well Detector

9. Reagents and Materials

- 9.1 *Purity of Reagents*—Reagent grade chemicals shall be used in all tests. Unless otherwise indicated, it is intended that all reagents shall conform to the specifications of the Committee on Analytical Reagents of the American Chemical Society, where such specifications are available³.
- 9.2 *Purity of Water*—Unless otherwise indicated, references to water shall be understood to mean Type I water as defined in Specification D1193.
- 9.3 *1M Hydroxylamine-hydrochloride*—commercially available solution or dissolve 70 g of the powder in 500 mL of water, dilute to 1 litre final volume.
 - 9.4 Iodide carrier, 20 mg I per millilitre as KI.
 - 9.5 Nitric Acid, concentrated, ~16M
- 9.6 0.1M Nitric Acid—Add \sim 6 mL of concentrated HNO $_3$ to 950 mL of water, dilute with water to a final volume of 1 litre.
- 9.7 8M Nitric Acid—Add 500 mL of concentrated HNO₃ to 450 mL of water; dilute with water to a final volume of 1 litre.
 - 9.8 p-xylene.
- 9.9 *Palladium carrier*—~10 mg/mL, dilute a commercially prepared solution to the correct concentration
- 9.10 *Sodium bisulfite*, 0.1M—dissolve 10.4 g of powder in 500 mL of water, dilute to a final volume of 1 litre
- ³ Reagent Chemicals, American Chemical Society Specifications, American Chemical Society, Washington, DC. For suggestions on the testing of reagents not listed by the American Chemical Society, see Analar Standards for Laboratory Chemicals, BDH Ltd., Poole, Dorset, U.K., and the United States Pharmacopeia and National Formulary, U.S. Pharmacopeial Convention, Inc. (USPC), Rockville, MD.

- 9.11 *Sodium Carbonate*, 2M—dissolve 212 g of powder in 500 mL of water, dilute to 1 litre final volume.
- 9.12 Sodium Hydroxide, 4M—dilute a commercially prepared solution or dissolve 160 g of pellets in 700 mL of water, dilute to a final volume of 1 litre. This is a very exothermic reaction. The use of an ice bath can mitigate the magnitude of the exothermicity.
 - 9.13 Sodium Hypochlorite.

10. Calibration and Standardization

- 10.1 The gamma-ray spectrometry system should be calibrated for energy, resolution and efficiency according to the manufacturer instructions. The background counting rate for the instrument should be measured at a frequency determined by the user. See C1402, D3648 or D3649 for additional information. A typical spectrum for ¹²⁹I is shown in Fig. 1.
- 10.2 Confirm the concentration of the I⁻ carrier by adding 1.00 mL of the carrier solution to 15 mL of water. Add 1 mL of the 0.1M NaHSO₃, mix, heat gently and then add 2 mL of the Pd⁺² carrier. Collect the precipitate (PdI₂) on a tared 25-mm filter paper. Dry and reweigh the filter paper to confirm the expected precipitate weight. Repeat this confirmation several times to increase the precision of the determination.
- 10.3 Prepare an efficiency curve for the 30 keV x-rays comparing the relative efficiency versus weight of PdI_2 by precipitating equal quantities of ^{129}I with various weights of PdI_2 . A typical curve for a Ge well detector is shown in Fig. 2; note that this curve shows the net count rate versus weight of PdI_2 rather than calculated efficiency (the ^{129}I activity used to prepare this graph was 2.2 Bq (60 pCi)).

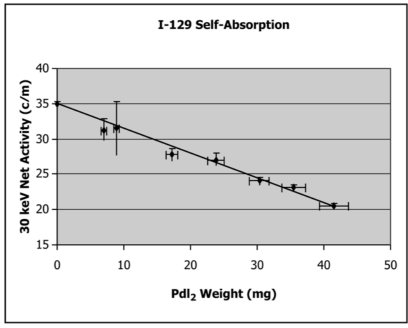


FIG. 2 Self-Adsorption of 30 keV X-ray versus Weight of Pdl₂ ·H₂O

11. Procedure

- 11.1 Weigh out no more than 100 mg of uranium oxide into a small beaker.
- 11.2 Dissolve the oxide in about 20 mL of 0.1M HNO₃. Heat gently, if required, to complete the dissolution.

Note 1—Avoid prolonged heating to minimize any iodine volatilization and possibly avoid heating altogether if only a small portion of the sample remains undissolved. Also, proceed directly to the next steps to minimize the time the sample is held under acidic conditions without the tracer present. Addition of the tracer prior to dissolution may not be appropriate since the sample iodine may not be in the same form and oxidation state as the tracer iodine.

- 11.3 Add 1 mL of the 4M NaOH. Swirl the solution to mix and check the pH. The solution should be strongly basic.
 - 11.4 Add 1 mL of the 2M NaCO₃. Swirl to mix the solution
 - 11.5 Add 1.00 mL of the 20 mg/mL I carrier. Swirl to mix.
- 11.6 Add 1 mL of the NaHClO $_3$ solution to the beaker to oxidize the iodine to periodate (IO $_4$ ⁻). Swirl to mix. Place the beaker on a hotplate and heat the solution to just below boiling. Remove from the hotplate and cool to room temperature.

CAUTION: The beaker and solution must be cool prior to the next step.

- 11.7 Carefully add 1 mL of the 8M HNO₃. Swirl the solution then check the pH. The solution should be strongly acidic.
- 11.8 Transfer the solution to a 60-mL plastic bottle or separatory funnel. Rinse the beaker a few times with small portions of water and add to the bottle.
 - 11.9 Add 10 mL of p-xylene to the bottle.
- 11.10 Add 3 mL of 1M NH_2OH -HCl to the bottle to reduce the periodate to iodine (I_2). Swirl to mix. The solution should be red-purple in color at this point.

- 11.11 Cap the bottle and shake for several minutes to extract the iodine into the organic layer. Let the solution stand and allow the organic layer to separate from the aqueous layer.
- 11.12 Remove the cap and draw off the top, organic layer with a disposable pipette. Transfer the organic layer to a 30-mL plastic bottle or clean separatory funnel.
- 11.13 Add 15 mL of water to the organic in the 30-mL bottle. Add 1 mL of the $0.1M \text{ NaHSO}_3$ to the bottle to reduce the iodine to iodide (Γ). Cap the bottle and shake for one minute until the organic layer is colorless. Let the solution stand and allow the organic layer to separate from the aqueous layer.
 - 11.14 Draw off the upper, organic layer and discard.
- 11.15 Transfer the aqueous layer to a 100-mL beaker and gently warm the solution on a hotplate.
 - 11.16 Add 2 mL of the Pd⁺² carrier solution to the beaker.
- 11.17 Allow the PdI_2 to precipitate and then filter the solution through a tared 25-mm filter paper.
- 11.18 Allow the filter paper to dry and then reweigh to determine the chemical yield of the separation.
- 11.19 Count the filter on an extended range or low-energy gamma-ray spectrometry system for the length of time required to meet the requested detection limit. Set the ROI for ¹²⁹ I to monitor the 29-34 keV Xe K x rays.

12. Calculation

12.1 CALCULATION OF CHEMICAL YIELD

 $Y = mg PdI_2$ recovered/mg PdI_2 expected based on calibration (10.2)

12.2 CALCULATION OF ACTIVITY

$$A_i = \left[(G_i - B_i) / (Y * E * A B_i * W) \right] \tag{1}$$

where

 A_i

= activity of ¹²⁹I in Bq per gram U oxide = gross counts per second in the ¹²⁹ I ROI G_I

background counts per second in the 129I ROI

= yield calculated above expressed as a fraction

 \boldsymbol{E} = detector efficiency for the 29-34 keV x-rays, ex-

pressed as a fraction, based on the weight of the PdI_2 = branching ratio for ^{129}I , expressed as a fraction

 AB_i

= weight of U oxide analyzed in grams

If the weight of uranium per gram of oxide is known the sample activity may be reported as Bq of 129I per gram of uranium by multiplying by the correct ratio.

12.3 CALCULATION OF MINIMUM DETECTABLE AC-**TIVITY**

 $MDA_i = (4.65*s_B + 2.71)/(E*Y*T*AB_i*W)$ (2)

where

 MDA_i = minimum detectable activity (Bq/g)

= standard deviation of the reagent blank counts in S_B

= sample counting time in seconds

13. Keywords

13.1 Gamma-ray spectrometry; liquid-liquid extraction;

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