RSC Advances



PAPER

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2024, 14, 31422

Inductively coupled plasma optical emission spectroscopic determination of trace rare earth elements in highly refractory gadolinium zirconate $(Gd_2Zr_2O_7)^{\dagger}$

Abhijit Saha, 60 * Khushboo Kumari, Sadhan Bijoy Deb* and Manoj Kumar Saxena

Gadolinium zirconate ($Gd_2Zr_2O_7$) belongs to the category of burnable absorber (BA) material in nuclear reactors. The high neutron-absorption cross sections of Gd isotopes (^{155}Gd and ^{157}Gd) implement negative reactivity in the reactor core to control the excess reactivity of the fuel at the beginning of the fission cycle. The presence of other rare earth elements, which can come from the starting material and/ or may be taken up during the synthesis steps, will affect the negative reactivity calculation. Thus the chemical quality assurance of $Gd_2Zr_2O_7$ concerning the other trace rare earth impurities is indispensable. Here in this work, we decided to quantify thirteen trace rare earth elements in $Gd_2Zr_2O_7$ by inductively coupled plasma optical emission spectroscopy (ICP-OES). One of the matrix elements viz., zirconium was separated via precipitation using D_iL -mandelic acid. The trace rare earths were determined in the presence of gadolinium matrix in solution. It was found that all thirteen rare earth elements can be quantified in the range of 0.25 to 2.5 mg L^{-1} in the presence of 516 mg L^{-1} of Gd with a relative standard deviation of 1-3%. This corresponds to the determination of a minimum of 0.25 mg analyte per g of $Gd_2Zr_2O_7$. The method detection limits of all thirteen rare earth elements range between 0.01 and 0.075 mg g^{-1} . The proposed methodology was validated by analyzing synthetic standards and real samples with spike addition.

Received 11th July 2024 Accepted 27th September 2024

DOI: 10.1039/d4ra05017g

rsc.li/rsc-advances

1. Introduction

In the scenario of depleting fossil fuel resources, nuclear reactors are one of the potential candidates for fulfilling the huge worldwide energy demand. The nuclear fuel meat which is the target component in fission reaction offers excess reactivity at the beginning of the reactor cycle. This excess reactivity is required to increase the fuel burn-up and to recompense for the loss of reactivity due to fuel depletion and fission product buildup.1-3 In nuclear reactors, this excess reactivity is being controlled by burnable absorbers (BAs) which are materials containing non-fissile nuclei with large neutron-absorption cross-sections (σ_a). Typically, a BA should burn out at the same rate as fuel to keep the net reactivity constant with time.1 When the BA burns out too quickly, a positive reactivity swing in the fuel cycle will hamper the reactor operation in the later stages. Similarly, if the BA burns out too slowly, then a negative reactivity of the remaining BA will hinder the end of the fuel's operational cycle. Other than

their control over excess reactivity, varieties of BAs are reported to offer benefits like tritium production and burning of long-lived radionuclides from spent nuclear fuel.4-6 There are two modes of using BA material in the reactor namely, homogeneous and heterogeneous modes.7-11 In homogeneous mode, the BA material is either homogeneously dispersed in the fuel or used as a coating. On the other hand, in heterogeneous mode, the BA material is dispersed in an inert matrix and the discrete rods are placed in the fuel assembly. The homogeneous mode has the advantage of even power distribution over the heterogeneous one.2 But the former mode has direct affect on the sinterability and thermal conductivity of the fuel.2 Hence the heterogeneous mode is more widely explored in the nuclear industry as it does not affect the fuel properties. Also in this mode, the BA material property can be tuned according to the reactor's requirement.1-3

^a, Radioanalytical Chemistry Division, Bhabha Atomic Research Centre, Mumbai, India 400085. E-mail: abhi.gallary@gmail.com; sbdeb@barc.gov.in

 $[\]dagger$ Electronic supplementary information (ESI) available. See DOI: $\label{eq:def:https://doi.org/10.1039/d4ra05017g} \text{ }$

Paper RSC Advances

Gd₂Zr₂O₇ is one such BA material that was prepared by various scientists and tested for its performance. 16-18 This pyrochlore is highly refractory and has low swelling properties, excellent irradiation, and chemical stability. 12-15 In India, this material has been proposed to be the BA rod material for compact hightemperature reactor (CHTR, a light water reactor). The chemical quality assurance (CQA) of any BA material includes the checking of burnable poison concentration and material properties before being used in the reactor. Thus the characterization of trace critical impurities (elements other than Gd which have high σ_a values) which can affect the negative reactivity calculation of the BA material is one of the important aspects of CQA. These impurities can come directly from the precursor viz., Gd₂O₃ and ZrO₂, or from the production steps. The precursor materials can ingress impurities with high σ_a values like rare earth elements and hafnium. Hence we focused our research on quantifying thirteen trace rare earth impurities viz., La, Ce, Pr, Nd, Sm, Eu, Tb, Dy, Ho, Er, Tm, Yb, and Lu in Gd₂Zr₂O₇.

The determination of thirteen trace rare earth impurities in Gd₂Zr₂O₇ presented challenges in the selection of suitable digestion procedure, difficulties in the separation of trace rare earth impurities from matrix elements, and matrix interference on analyte signals. The method commenced with the microwave-assisted dissolution of this highly refractory material. The dissolution procedure was optimized concerning the various acid combinations and proportions. Once the material was dissolved, zirconium(IV) was selectively precipitated using D,L-mandelic acid. This step also eliminated hafnium from the solution and hence we did not carry out hafnium estimation. The separation of thirteen trace rare earth elements from the Gd matrix is a highly difficult task due to their similar chemical behavior. Hence, we decide to analyze all thirteen trace rare earth elements in the presence of gadolinium. The estimation thus needed an analytical technique with high sample throughput, high sensitivity, and low detection limit. A technique like inductively coupled plasma mass spectrometry (ICP-MS)19-22 qualifies for all the above-mentioned requirements. However, the high matrix concentration can clog its cones and affect the ion beam in the ion optics region which may lead to unreliable results. In the case of X-ray fluorescence (XRF) spectroscopy,23,24 the low fluorescence yield of lanthanide L lines will be further reduced in presence of a high Gd matrix. On the other hand, ICP optical emission spectroscopy (ICP-OES)25-29 can handle higher matrix concentration compared to both ICP-MS and XRF techniques. Hence the digested sample solution after zirconium precipitation was directly analyzed by ICP-OES. The ICP-OES procedure was optimized for the determination of minimum analyte concentration in the presence of maximum matrix concentration. The matrix interferences on all thirteen analyte spectra were examined and corrected for the required ones. The method was established by analyzing synthetic standards of Gd and Zr matrix containing thirteen trace rare earth elements. The proposed analytical methodology was validated by analyzing real samples via spike addition and recovery studies.

2. Experimental

2.1. Instrumentation

A high resolution ICP-OES of model no: Ultima 2 from HORIBA Scientific (Jobin Yvon Technology), France was employed for the quantification of analytes. The spectrometer has a spectral range of 160–800 nm. The complete instrumental details are tabulated in Table 1. An Anton Par Multiwave 5000 microwave digestion system with $8\times$ rotor was used for dissolution of ${\rm Gd_2Zr_2O_7}$. A Mettler Toledo microbalance, model: XP26, was used for weighing the solids.

2.2. Reagents and solutions

Suprapur® sulfuric acid (96% assay), hydrochloric acid (30% assay), nitric acid (65% assay) and hydrofluoric acid (40% assay) were purchased from Merck. Ultrapure water (18.2 M Ω cm) from Milli-Q system of Millipore Corporation was used throughout the experiments. 1000 mg L $^{-1}$ ICP standard solutions (Alfa Aesar) of thirteen rare earth elements was used to prepare the synthetic standard solutions and for spike addition. High purity Gd $_2$ O $_3$ (99.999%, Sigma-Aldrich) and 10 000 mg L $^{-1}$ ICP standard solution of Zr (Aldrich Chemical Company, Inc.) were utilized for preparing synthetic standards. D,L-Mandelic acid (99%) was procured from Sigma-Aldrich. All the solutions were prepared in Nalgene® PP volumetric flasks.

2.3. Preparation of synthetic standard and sample solutions

About 29.5, 59, and 119 mg of Gd₂O₃ were weighed in the microbalance and taken in different beakers. In these beakers, calculated volumes of 1000 mg L⁻¹ rare earth standards and 10 000 mg L⁻¹ Zr standard were added to prepare synthetic standards according to Table T1 (see ESI†). The mixtures were dissolved by adding 2 mL of conc. HNO3 under heating at 80 °C for 5 min. The clean solution was then evaporated near to dryness and re-dissolved by conc. HCl. This step was repeated for 2-3 times to eliminate the nitric acid. The solution was then made up to 20 mL by 6.0 mol L^{-1} HCl solution. To this solution, 50 mL of 16% (w/v) mandelic acid was added at room temperature and the total volume was made 100 mL by Milli-Q water. The temperature was then slowly raised to 85 °C, and held there for 30 min. The white precipitate was then filtered and washed by using 2% (v/v) HCl and 5% (w/v) mandelic acid solution. The raffinate was then evaporated to dryness and then heated in a muffle furnace at 850 °C for 4 h. The residue was then completely dissolved by 5 mL of conc. HNO₃ under heating at 80 °C for 5 min. The solution was then made up to 100 mL in a volumetric flask with Milli-Q water for ICP-OES analysis.

About 100 mg of $Gd_2Zr_2O_7$ was taken in NXF100 Teflon vessel along with 3 mL of conc. H_2SO_4 , 5 mL of conc. HCl, 1 mL of conc. HF and 1 mL of Milli-Q water. In case of spike addition appropriate volumes of 1000 mg L $^{-1}$ individual rare earth ICP-standard solutions were added along with the sample and reagents into the NXF100 Teflon vessel. All eight vessels along with their attachments were kept in the microwave at 700 W for 30 min and then at 1400 W for 60 min. The complete dissolution parameters have been discussed in Table 2 in the

Table 1 ICP-OES specifications and operating parameters

Spectrometer	
Optical design	1 m Czerny Turner
Grating	Holographic, dual back-to-back gratings
Groove density	4320 g mm ⁻¹ and 2400 g mm ⁻¹
Resolution	1st and 2nd order with optic
	resolution <0.005 nm for 160-
	320 nm and <0.01 nm for 320–800 nm
Thermal regulation	Controlled to 32.5 °C
Thermal regulation	Controlled to 32.5
Plasma parameters	
Radio frequency	40.68 MHz
Applied power	1000 W
Reflected power	<1 W
Ar gas flow rates	
Plasma gas	13 L min ⁻¹
Nebulizer gas	$1.5~\mathrm{L~min^{-1}}$
Sheath gas	$0.2~\mathrm{L~min^{-1}}$
Acquisition parameters	
Nebulizer	Concentric
Spray chamber	Cyclonic
Sample rate	$0.9~\mathrm{L~min}^{-1}$
Detection	Dual PMT detection with high
	dynamic detection system (HDD
Gain	3
Entrance slit	20 μm
Exit slit	15 μm
Measurement	Peak top, background
Integration time	3 s
No. of replicates	5

results and discussion section. The digested sample was then taken out in a beaker and heated at 275 °C in a fume hood for 1 h. The fumes came out during this heating procedure were trapped in a NaOH solution. This method removes all the HF and HCl from the solution leaving all the elements in the $\rm H_2SO_4$ medium. To this 20 mL of 6.0 mol $\rm L^{-1}$ HCl solution was added and then zirconium was precipitated using mandelic acid. The rest of the procedure followed was as discussed above. The proposed analytical methodology is presented as a scheme in Fig. 1.

3. Results and discussion

3.1. Optimization of microwave digestion procedure

The highly refractory Gd₂Zr₂O₇ needs an extreme environment to get completely dissolved. One such method is the fusion method, but this has the possibility of impurity ingression from a large excess of fusion mixture and loss of sample due to sputtering. On the other hand, a microwave digestion system using Suprapur acids provides a clean and closed environment with high temperature and high pressure for the digestion of materials. Hence, we chose the microwave digestion procedure for digesting our sample. A combination of conc. H₂SO₄, conc. HCl and conc. HF was used to digest the sample. Sulphuric acid was chosen because of its high boiling point which enabled us to reach either a high temperature of 240 °C (maximum pressure 45 bar) or high pressure of 60 bar (maximum temperature 180 °C) inside the Teflon vessels. Both hydrochloric and hydrofluoric acids are highly corrosive and hence they break the metal-oxide bonds to dissolve the material. The volumes of these acids were optimized by following these points: (i) minimum volume of H2SO4 was used so that maximum volumes of corrosive acids can be used; (ii) maximum volume of HCl was utilized because in this medium zirconium was precipitated; (iii) minimum volume of HF was added so that its removal became easy. Since the presence of fluoride ion in the solution could hamper zirconium precipitation, its complete removal was absolutely necessary. The low boiling point of HF (b.p. is near RT of pure HF) compared to H_2SO_4 (b.p. ~ 337 °C) allowed the partial removal of the former acid at 275 °C.30 The heating was continued for prolonged time (1 h) to ensure the complete removal of fluoride ions as HF. Using all this information we optimized the proportion of acids as shown in Table 2. The power input was kept constant in all the trials.

3.2. Selection of emission lines

The rare earth elements are rich in atomic and ionic emission lines. However, the ionic lines have the highest theoretical intensities compared to the atomic ones. Hence, we selected the ionic lines for all thirteen rare earth elements. The lines were selected based on the criteria that they should have the maximum intensity irrespective of interference from Zr emission lines but have negligible interference from Gd emission lines. The selected lines are tabulated in Table T2 (ESI†) along with the interference information. Each selected line also has minimum interference of

Table 2 Optimization of the microwave assisted digestion parameters of Gd₂Zr₂O₇

				Reagents				
Sample amount (mg)	Power	Ramp (min)	Hold time	Conc. H ₂ SO ₄	Conc. HCl	Conc. HF	Milli-Q water	Remarks
~100	First at 700 W and	15	30 min at 700 W and	1	9	0	0	Not dissolved
	then at 1400 W		60 min at 1400 W	1	8	0.5	0.5	Not dissolved
				1	7	1	1	Not dissolved
				2	6	1	1	Partially dissolved
				3	5	1	1	Completely dissolved
				4	4	1	1	Completely dissolved

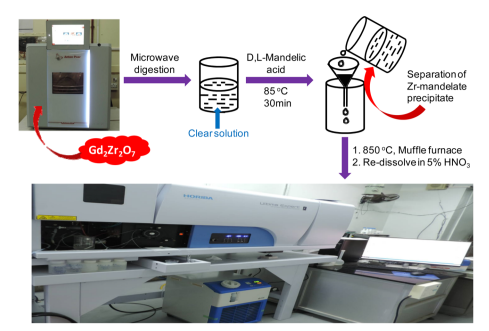


Fig. 1 Schematic representation of the proposed analytical methodology for trace rare earth determination in Gd₂Zr₂O₇.

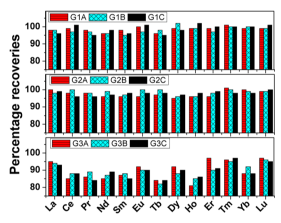


Fig. 2 Pictorial representation of percentage recoveries of thirteen rare earth elements in nine synthetic standards as prepared in Table T1 (ESI \dagger). The three sets G1, G2, and G3 contain 258, 516, and 1032 mg L $^{-1}$ gadolinium as matrix during ICP-OES determination. The sub-sets under these three sets were named as A, B, and C which contain thirteen trace rare earth elements according to Table T1 (ESI \dagger).

the remaining lanthanide emission lines. This has been observed by plotting the calibration spectra of all thirteen rare earth elements in the range 0.1–5.0 mg $\rm L^{-1}$ in the presence of each other, as shown in Fig. S1 (ESI†). All thirteen rare earths had a regression coefficient of \geq 0.99.

3.3. Optimization of analyte and matrix concentrations

To overcome the hurdle of separating thirteen trace rare earth elements from gadolinium we thought of quantifying the analytes in the presence of the matrix element. However, one of the matrix elements *i.e.*, zirconium was separated by using D,L-mandelic acid in the digested sample solution following the same procedure as Kumins *et al.*³¹ The effect of zirconium(IV)

separation on the quantification of analytes was first examined by carrying out recovery studies on synthetic standards in Table T1 (see ESI†). The matrix concentrations in all the synthetic standards were maintained by calculating the mass fraction of every element and the dilution factor. The calculations are given in ESI.† The gadolinium and zirconium concentrations in all three synthetic standard sets viz., G1, G2, and G3 were 258 & 150, 516 & 300, and 1032 & 600 mg L^{-1} respectively. The analyte concentrations were varied in the range of 0.25 to 2.5 mg L^{-1} in all three sets. Once Zr(w) was precipitated and filtered out, the raffinate was analyzed for the concentration of thirteen rare earths in the presence of gadolinium. The percentage recoveries of all thirteen rare earth elements in the nine synthetic standards are pictorially represented in Fig. 2. The recovered amounts of the individual elements in all nine synthetic standards are represented in Table T3 (ESI†). In the case of both sets G1 and G2, recoveries of ≥95% were obtained for all the thirteen rare earth elements. The raffinate was also treated with oxalic acid to precipitate Gd(III). Both the precipitate viz., Zr(IV)mandelate and Gd(III)-oxalate were converted to their respective oxides in a Muffle furnace at 850 °C. The WDXRF spectra of both ZrO₂ and Gd₂O₃ are shown in Fig. S2 and S3 (ESI†) respectively. The ZrO₂ spectrum is free from any rare earth elements whereas the Gd₂O₃ spectrum shows the presence of all other rare earth elements of interest. Hence both the recovery and XRF studies confirmed that the precipitation of Zr(IV) has no adverse effect on the recovery of analytes.

The emission spectra of all thirteen elements at $0.25~{\rm mg~L}^{-1}$ level in the absence and presence of gadolinium matrix are given in Fig. S4 and S5 (ESI†). From these figures, it can be seen that elements like Ce, Pr, Nd, Sm, Tb, Ho, and Yb need background correction. The background count for each element in the presence of a matrix was determined by analyzing pure

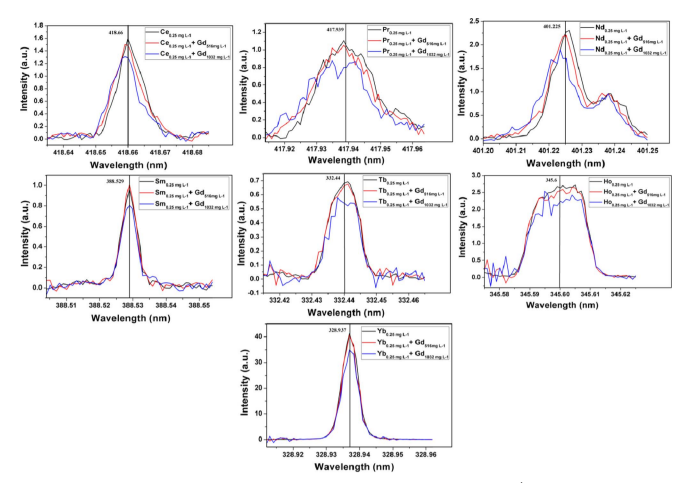


Fig. 3 Background intensity corrected emission spectra of Ce, Pr, Nd, Sm, Tb, Ho and Yb at 0.25 mg mL^{-1} level in absence and presence of gadolinium matrix.

synthetic solutions of 516 and 1032 mg L^{-1} of gadolinium. The gadolinium emission lines responsible for these blank counts are mentioned in Table T4 (ESI†). In the case of elements like Ce, Pr, Nd, and Sm the nearby single emission line of gadolinium increases the blank intensity. However, the emission lines of Tb and Ho are affected by two or more gadolinium emission lines. Only the emission intensities of Tm and Yb have direct interference from gadolinium emission lines (see Table T2 in ESI†). However, the emission spectra of Tm in the presence or absence of gadolinium matrix were found to be the same at 95% confidence interval, as shown in Fig. S4 (ESI†). The heavy rare earth elements have nearly filled or filled 4f orbitals which results in their highly intense emission lines. The intensity of Tm 313.125 nm line is so high that it remains unaltered by Gd 313.081 line at 0.25 mg L^{-1} level in the presence of 516 and 1032 mg L⁻¹ of gadolinium. Hence Tm needs no background correction due to the presence of gadolinium matrix. The background corrected spectra of Ce, Pr, Nd, Sm, Tb, Ho, and Yb in the presence of 516 and 1032 mg L⁻¹ of gadolinium are represented in Fig. 3. In the case of all thirteen rare earth elements the minimum analyte concentration of 0.25 mg L^{-1} was recovered up to \geq 95% in presence of 516 mg L⁻¹ of gadolinium. The higher matrix concentration of 1032 mg L^{-1} resulted in recoveries in between 80-97% for all thirteen

analytes. Since the background intensities of Nd, Pr, Tb, and Ho in the presence of 1032 mg $\rm L^{-1}$ of Gd is nearly equal to or more than the matrix-free analyte intensities at 0.25 mg $\rm L^{-1}$ level, their recoveries were severely affected. Other elements like Ce, Sm, and Yb also resulted in poor recoveries at 1032 mg $\rm L^{-1}$ Gd concentration level due to high background intensities. Thus the optimization procedure revealed that a minimum of 0.25 mg $\rm L^{-1}$ of all thirteen rare earth elements can be precisely determined in the presence of 516 mg $\rm L^{-1}$ of Gd. This corresponded to the determination of 0.25 mg analyte per g of $\rm Gd_2Zr_2O_7$ by the proposed methodology.

3.4. Challenges of the procedure

The precise ICP-OES determination of trace rare earth elements in such a refractory matrix needs an efficient and clean dissolution. This difficulty was resolved by microwave assisted digestion of $Gd_2Zr_2O_7$ using Suprapur acids. The separation of the matrix elements $\emph{viz.}$, Gd and Zr would nullify the spectroscopic and non-spectroscopic matrix interferences on the impurity analysis. Zr(rv) was separated by mandelate precipitation whereas Gd(m) separation was not performed due to its chemical similarities with trace rare earth impurities. This can be attributed as the limitation of the proposed methodology. The ICP-OES determination of trace rare earth impurities in

Table 3 IDL and MDL values

Elements	$IDL \left(\times 10^{-3}\right) \left(mg \ L^{-1}\right)$	$MDL \left(\times 10^{-3}\right) \left(mg \ g^{-1}\right)$
La	1.1	40
Ce	3.2	65
Pr	3.8	75
Nd	2.1	56
Sm	2.5	60
Eu	0.5	25
Tb	2.0	52
Dy	1.0	35
Но	0.8	30
Er	1.5	47
Tm	0.4	20
Yb	0.2	12
Lu	0.2	10

presence of gadolinium matrix was done by optimizing the matrix concentration, suitable choice of analyte emission lines, and background correction.

3.5. Detection limit of the analysis

The detection limit was determined for both the 5% (v/v) HNO_3 solution and sample blank known as instrument detection limit (IDL) and method detection limit (MDL) respectively. A 516 mg L^{-1} Gd solution was used as the sample blank. The 3σ

values of both the blank solutions and the sensitivities (s) of the calibration curves in Fig. S1† were used to calculate the detection limits ($3\sigma/s$). The values of both IDL and MDL are mentioned in Table 3.

3.6. Analysis of real sample

Two Gd₂Zr₂O₇ samples were digested in duplicate using a microwave digestion procedure. In the case of each sample, one of the two sets contained only sample and the other set was spiked with the analytes of interest before digestion. All digested samples were subjected to the zirconium precipitation and then subsequent sample treatment. The analysis results are presented in Table 4. The recoveries of all thirteen analytes were found to be ≥95% in both samples. The percentage relative standard deviation (RSD) on the determined values was found to vary between 1 and 3%. These results validate the proposed methodology for its further use in routine analysis. The individual trace rare earth concentrations result in cumulative concentrations of 4.18 and 8.13 mg g⁻¹ in Gd₂Zr₂O₇-1 and Gd₂Zr₂O₇-2 samples respectively. Such information is very much helpful in ascertaining the purity of the compound. The individual concentrations of the trace rare earths having high σ_a values viz., Sm, Eu, Tb, Dy, Ho, Er, Yb etc. will be helpful in calculating the total negative reactivity imposed by the two compounds.

Table 4 Analysis of real samples $(N = 5)^a$

Elements	$Gd_2Zr_2O_7$ -1		$\mathrm{Gd_{2}Zr_{2}O_{7}}\text{-}2$		
	Added (mg g^{-1})	Found (mg g^{-1})	Added (mg g^{-1})	Found (mg g^{-1})	
La	0	BDL	0	0.277 ± 0.006	
	0.25	0.248 ± 0.008 (99)	0.25	$0.517 \pm 0.010 (96)$	
Ce	0	0.824 ± 0.009	0	1.214 ± 0.016	
	0.25	1.068 ± 0.012 (98)	0.25	1.456 ± 0.015 (97)	
Pr	0	BDL	0	0.312 ± 0.008	
	0.25	0.242 ± 0.007 (97)	0.25	0.559 ± 0.009 (99)	
Nd	0	0.343 ± 0.012	0	0.547 ± 0.009	
	0.25	0.588 ± 0.008 (98)	0.25	0.786 ± 0.013 (96)	
Sm	0	0.408 ± 0.015	0	0.573 ± 0.009	
	0.25	0.648 ± 0.010 (96)	0.25	0.814 ± 0.014 (96)	
Eu	0	0.377 ± 0.011	0	0.622 ± 0.015	
	0.25	0.615 ± 0.015 (95)	0.25	$0.873 \pm 0.017 (100)$	
Tb	0	0.766 ± 0.016	0	1.445 ± 0.022	
	0.25	1.011 ± 0.015 (98)	0.25	$1.687 \pm 0.020 (97)$	
Dy	0	0.654 ± 0.011	0	0.844 ± 0.018	
	0.25	0.897 ± 0.018 (97)	0.25	$1.088 \pm 0.020 \ (98)$	
Но	0	0.542 ± 0.010	0	0.776 ± 0.016	
	0.25	0.780 ± 0.016 (95)	0.25	1.021 ± 0.018 (98)	
Er	0	0.288 ± 0.008	0	0.463 ± 0.009	
	0.25	0.536 ± 0.013 (99)	0.25	0.701 ± 0.011 (95)	
Tm	0	BDL	0	0.357 ± 0.008	
	0.25	0.246 ± 0.007 (98)	0.25	$0.609 \pm 0.009 (101)$	
Yb	0	BDL	0	0.313 ± 0.007	
	0.25	0.252 ± 0.010 (101)	0.25	$0.562 \pm 0.009 (100)$	
Lu	0	BDL	0	0.386 ± 0.009	
	0.25	$0.251 \pm 0.008 (100)$	0.25	0.632 ± 0.011 (98)	

^a The percentage recoveries are given in parenthesis. BDL = below detection limit.

4. Conclusion

The determination of thirteen trace rare earth impurities in Gd₂Zr₂O₇ has been reported for the first time in literature. Since the measurement was carried out by separating only zirconium, the presence of gadolinium matrix was optimized to obtain accurate results. The use of ICP-OES was found to provide precise analyte concentrations even in the presence of high matrix levels. The measurement is necessary to calculate the exact negative reactivity of the BA material before putting it into the reactor. This developed methodology can be applied to characterize any $A_2Zr_2O_7$ type of compounds (where A = rare earth element) which have applications in the fields of solid oxide fuel cells, nuclear waste storage, thermal barrier coating, and nuclear reactor BA or control rod materials. The microwaveassisted dissolution of this refractory BA material is also reported first time in the literature. The developed digestion procedure may also help in carrying out other studies on this BA material. The methodology is free from generating any waste. To the best of the authors' knowledge, this is the first report on analyzing trace rare earth impurities in Gd₂Zr₂O₇ by ICP-OES.

Data availability

The authors confirm that the data supporting the findings of this study are available within the article and ESI.†

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors are thankful to Dr P. K. Mohapatra, Associate Director, RC & IG, BARC for his constant encouragement throughout the work.

References

- 1 J. A. Evans, M. D. DeHart, K. D. Weaver and D. D. Keiser Jr, *Nucl. Eng. Des.*, 2022, **391**, 111726.
- 2 H. S. Kim, C. Y. Joung, B. H. Lee, S. H. Kim and D. S. Sohn, *J. Nucl. Mater.*, 2008, 372, 340–349.
- 3 M. T. Malachevsky, D. R. Salvador, S. Leiva and C. A. D'Ovidio, *J. Ceram.*, 2015, **2015**, 298690.
- 4 D. Lanning, D. Baldwin, M. Cunningham and S. Marschman, *Post irradiation examination report for the tritium-producing burnable absorber rods irradiated in the Watts Bar Nuclear Plant*, ed. U. S. Nrc, Pacific Northwest National Laboratory, 2002, p. 72.
- 5 E. P. Naessens, K. S. Allen and B. E. Moretti, *Nucl. Sci. Eng.*, 2006, **152**(3), 306–313.

- 6 I. Shaaban and M. Albarhoum, Appl. Radiat. Isot., 2017, 125, 188–195.
- 7 T. Cardinaels, J. Hertog, B. Vos, L. de Tollenaere, C. Delafoy and M. Verwerft, *J. Nucl. Mater.*, 2012, 424(1-3), 289-300.
- 8 T. Jevremovic, *Nuclear Principles in Engineering*, Springer, 2nd edn, 2009.
- 9 H. van Dam, Nucl. Energy, 2000, 27, 63-69.
- 10 J. Porta, S. Baldi, J. P. Chauvin and Ph. Fougeras, *Prog. Nucl. Energy*, 2001, 38, 355–358.
- 11 M. Asou and J. Porta, *Nucl. Eng. Des.*, 1997, **168**(1-3), 261-270.
- 12 V. D. Risovany, E. E. Varlashova and D. N. Suslov, *J. Nucl. Mater.*, 2000, 281, 84–89.
- 13 G. Panneerselvam, R. Venkata Krishnan, M. P. Antony, K. Nagarajan, T. Vasudevan and P. R. Vasudeva Rao, *J. Nucl. Mater.*, 2004, 327, 220–225.
- 14 H. S. Kim, C. Y. Joung, S. H. Kim, B. H. Lee, Y. W. Lee, D. S. Sohn and S. H. Lee, *J. Korean Ceram. Soc.*, 2002, 39(11), 1108–1112.
- 15 A. Saha, S. B. Deb, B. K. Nagar and M. K. Saxena, Spectrochim. Acta Part B At. Spectrosc., 2014, 94–95, 14–21.
- 16 W. J. Weber, J. W. Wald and Hj. Matzke, J. Nucl. Mater., 1986, 138, 196–209.
- 17 W. L. Gong, W. Lutze and R. C. Ewing, *J. Nucl. Mater.*, 2000, 277, 239–249.
- 18 A. J. Feighery, J. T. S. Irvine and C. Zheng, *J. Solid State Chem.*, 2001, **160**, 302–306.
- 19 A. Saha, S. B. Deb and M. K. Saxena, *J. Anal. At. Spectrom.*, 2016, 31, 1480–1489.
- 20 J. W. Olesik and S. Jiao, *J. Anal. At. Spectrom.*, 2017, **32**, 951–966.
- 21 C. Agatemor and D. Beauchemin, *Anal. Chim. Acta*, 2001, **706**(1), 66–83.
- 22 A. Saha, S. B. Deb, B. K. Nagar and M. K. Saxena, *At. Spectrosc.*, 2013, 34(4), 125–132.
- 23 K. Sanyal, A. Saha, A. Sarkar, S. B. Deb, R. V. Pai and M. K. Saxena, *ACS Omega*, 2023, **8**(44), 41402–41410.
- 24 C. Bowers, J. Chem. Educ., 2019, 96(11), 2597-2599.
- 25 B. Ticová, K. Novotný and V. Kanický, *Chem. Pap.*, 2019, 73, 2913–2921.
- 26 J. C. Fariñas, I. Rucandio, M. S. Pamarer-Alfonso, M. E. Villanueva-Tagle and M. T. Larrea, *Talanta*, 2016, 154, 53–62.
- 27 F. Ardini, F. Soggia, F. Rugi, R. Udisti and M. Grotti, *Anal. Chim. Acta*, 2010, **678**(1), 18–25.
- 28 J. A. Gásquez, E. DeLima, R. A. Olsina, L. D. Martinez and M. de la Guardia, *Talanta*, 2005, **67**(4), 824–828.
- 29 L. Bendakovská, A. Krejčová, T. Černohorský and J. Zelenková, *Chem. Pap.*, 2016, **70**(9), 1155–1165.
- 30 N. N. Greenwood and A. Earnshaw, *Chemistry of elements*, Pergamon Press, Oxford, 1984, p. 921.
- 31 C. A. Kumins, Anal. Chem., 1947, 19(6), 376-377.