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Assessing rare earth elements in Indian coal ash using Westcott formalism in NAA and leaching studies

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This study employs Westcott formalism coupled with the internal monostandard approach in k_0 -NAA to assess Rare Earth Elements (REEs) in Indian coal ash samples. This method mitigates neutron flux perturbations during irradiation and enables *in situ* detector efficiency calibration for quantitative analysis. Deviations in thermal capture cross-sections for non-1/v nuclides in REEs were addressed using the Westcott $g(T_n)$ -factor obtained from the ENDF/B-VIII.0 data. REEs content in Indian coal ash was found to be 350–500 mg kg⁻¹, aligning with global standards. Leaching feasibility for medium-lived REEs with hydrochloric acid was demonstrated, supporting effective recovery. These findings suggest Indian coal ash as a potential secondary resource for REEs amid global supply challenges.

1 Introduction

Rare Earth Elements (REEs) are incredibly versatile due to their unique chemical, magnetic, and luminescent properties, making them indispensable for the development of environmentally friendly and sustainable technologies. These elements are crucial for a wide range of applications, including defence systems, renewable energy technologies, rechargeable batteries, lighting products, catalytic converters, and electric vehicles. Nevertheless, the uncertainty surrounding the supply of rare earths poses a growing obstacle to the progression and maturation of cutting-edge and futuristic technologies. The interplay between disruptive potential and the essential requirement of REEs for defence, renewable energy sources and other applications like rechargeable batteries, lighting products, catalytic converters, electric vehicles *etc.* has elevated their critical importance and generated a substantial demand for REEs.^{1–3} The global REE market has been characterized by significant supply chain challenges. Historically, the USA led REE mining from 1965 to 1984, but China has since become the dominant producer, controlling 70% of global REE mining as of 2022. The 40% reduction in REE exports by China in 2010 led to a significant price increase and highlighted the vulnerability of global supply chains.^{4–8}

The absence of immediate substitutes of REEs and heavy dependence on imports hindered the global advancement of emerging technologies. As a result of this, multiple REEs such as yttrium (Y), neodymium (Nd), europium (Eu), terbium (Tb),

dysprosium (Dy), and erbium (Er) have been identified as critical by the European Commission, National Environmental Research Council, and the U.S. Department of Energy due to their high potential in clean energy applications and significant supply risk. This has spurred global research initiatives aimed at indigenous production and sustainable recovery of these elements.^{9,10} Global Mine production of REEs saw a rise from 280 000 tons of REO equivalent in 2021 to an estimated 300 000 tons in 2022. China's Ministry of Industry and Information Technology raised 2022 quotas for rare-earth mining and separation to 210 000 tons and 202 000 tons of REO equivalents, respectively. Even though China shows dominance in the rare-

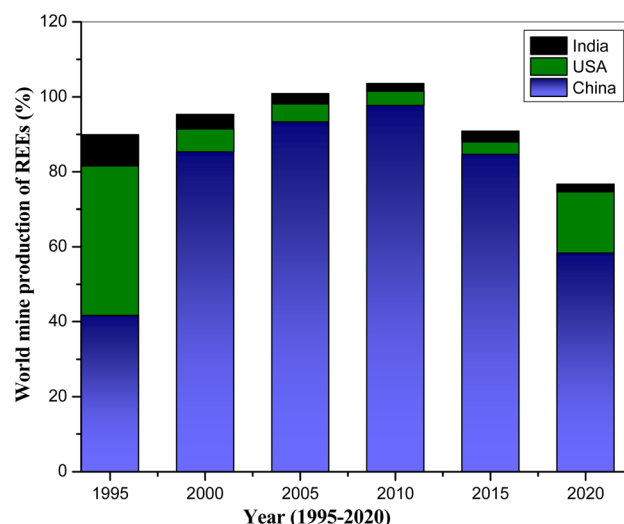


Fig. 1 REEs Mining from 1995 to 2020 (USGS).

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earth but its global REEs mining decreased from 90% in 2010 to the 70% in 2022 (Fig. 1).¹¹

Global rare earth oxide (REO) reserves stand at an estimated 130 million tons, distributed globally as shown in Fig. 2. India stands at the fifth position, with the leading by China. If the annual demand growth rate remains at approximately 10%, global REO reserves could be depleted by the middle of the 21st century.¹¹

The increasing demand for REEs has sparked a global search for new primary sources and alternative reserves. From the research, it was found that various REEs reserves like mud, end-use products, coal and its combustion residues are potential secondary sources for the REEs production.¹²⁻¹⁸ In India, thermal power generation reliant on coal represents the predominant source of power capacity expansion, accounting for over 69% of the overall electricity production. Indian coal, with its high ash content of 30–60%, presents a significant opportunity for REE recovery, especially given the contrast with imported coals (2–20% ash content). Various studies have demonstrated that REEs in coal ash are often uniformly dispersed within a glassy amorphous aluminosilicate matrix, necessitating strong acidic or alkaline solutions for their extraction. Recent research has focused on optimizing leaching conditions to maximize REE recovery from coal ash.¹⁴⁻²¹ Shanshan Cao *et al.*²¹ studied the about the various influencing factors towards the leaching of most abundant REEs (La, Ce and Nd) from coal fly ash in the HCl medium. Similar study has been performed for the leaching behaviour of medium-lived REEs from coal ash in the same HCl medium using their corresponding radioactive tracers produced *via* (n,γ) capture reaction.²⁰⁻²⁴ Radioactive tracer methodology for leaching study offers several advantages like sensitivity, selectivity, minimal spectral interference, dilution free *etc.* over conventional analysis techniques like ICPMS (Inductively Coupled Plasma Mass Spectrometry) and ICPOES (Inductively Coupled Plasma Optical Emission Spectroscopy). Extraction of REEs from the coal combustion residue may enhance its applicability and effective utilization of surplus amount of coal ash.¹⁸ The profiling of REEs in coal ash samples becomes mandatory towards their extraction with economical and efficient methods. The present

research study focuses on the application of non-destructive k_0 -Neutron Activation Analysis (NAA) to ascertain the concentration of REEs in coal ash samples sourced from different thermal power plant in India due to its several advantages over ICPOES, ICPMS. In NAA, most of the nuclides exhibit a thermal neutron capture cross-section that follows the $1/\nu$ law, where the cross-section is inversely proportional to the neutron velocity. However, a few nuclides deviate from this $1/\nu$ behaviour, referred to as non- $1/\nu$ nuclides. Despite the advantages of considering $1/\nu$ and non- $1/\nu$ nuclides, there is a paucity of articles applying the Westcott formalism in k_0 -NAA. The present study employs the Westcott formalism in k_0 -NAA due to its ability to account for deviation from the $1/\nu$ law in certain REEs, such as ^{151}Eu and ^{176}Lu . This approach, coupled with the internal monostandard method, minimizes neutron flux perturbations during irradiation and self-attenuation of gamma radiation during counting, ensuring accurate and reliable quantification of REEs in coal ash. Towards elemental characterization, recently determined parameters including the neutron temperature, modified spectral index $\left(r(\alpha)\sqrt{\frac{T_n}{T_0}}\right)$, and epi-thermal flux shape factor (α) at the Pneumatic Fast Transfer System (PFTS) location of the KAMINI reactor were employed.²⁵⁻³³

This study aims to quantify the concentration of REEs in coal ash samples sourced from various thermal power plants in India using non-destructive k_0 -NAA with Westcott formalism. Additionally, the feasibility of leaching medium-lived REEs using hydrochloric acid is demonstrated, supporting the potential recovery of these critical elements from coal ash.

2 Experimental

2.1 Materials and methods

Four coal ash samples, sourced from thermal power plants in India were obtained for REEs analysis using NAA technique. The quantification of activated samples was executed using high-resolution gamma spectrometry by employing a 30% p-type co-axial High Purity Germanium (HPGe) detector acquired from M/s Canberra Eurisys. The detector offers energy resolution of 1.85 keV at the prominent 1332 keV gamma energy line of ^{60}Co , complemented by an associated 8k multichannel analyser (MCA) system equipped with Aptec spectra software. To ensure precision in energy calibration, the detector was calibrated using standard radioactive sources, namely ^{241}Am and ^{152}Eu , procured from M/s Amersham, Inc.

2.2 Sample irradiation and radioactive assay

Four coal ash samples and NIST-SRM 1633b coal ash, each weighing approximately 50 mg were prepared in duplicate using clean polythene sheets. The first set of four sample were subjected to short-time irradiation (30 min) and the other set for long-time irradiation (6 h) at the Pneumatic Fast Transfer System (PFTS) position of the KAMINI reactor, operating at 20 kW reactor power. However, two sample of different coal ash, along with the NIST SRM, and a gold (Au) standard were taken

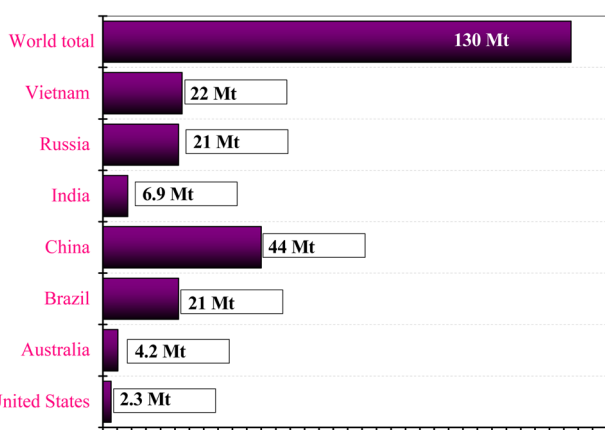


Fig. 2 Rare Earth reserves across the world.



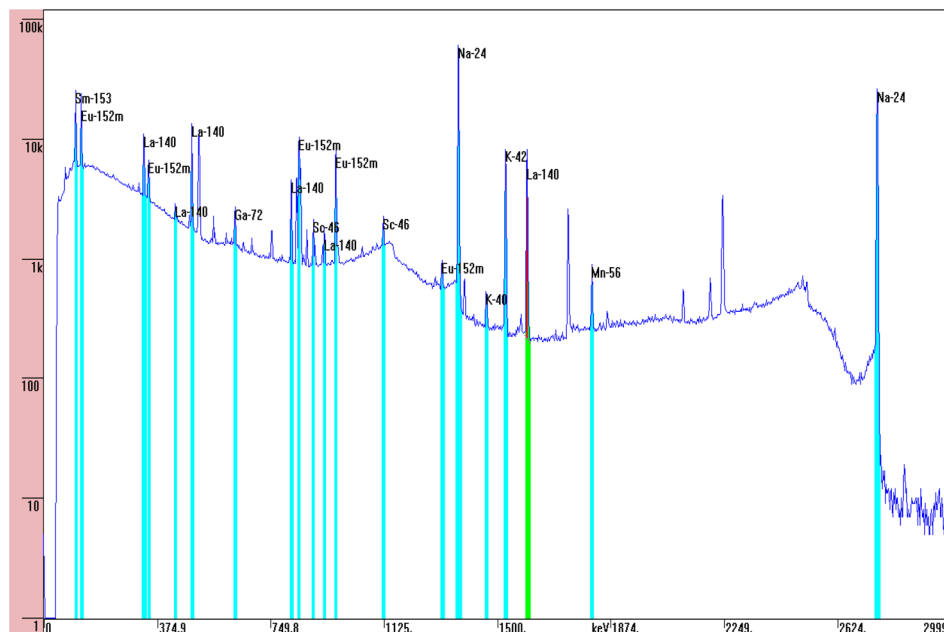


Fig. 3 Gamma spectrum of neutron irradiated coal ash sample.

in each irradiation. Short irradiated samples were counted for 15 min to 1 h after a 30 min cooling time; whereas the long irradiated samples were counted for 1 h to 14 h, after 1 day cooling time at a distance of 30 cm from 30% p-type co-axial High Purity Germanium (HPGe) detector. The recently characterized neutron spectrum parameters at PFTS $r(\alpha)\sqrt{\frac{T_n}{T_0}}$ and α value were 0.037 ± 0.001 and -0.0494 ± 0.0071 respectively. The average value of $g_{Lu}(T_n)$ was found to be 1.8939 ± 0.0130 using $1/\nu$ nuclides (^{94}Zr , ^{58}Fe and ^{197}Au). The Maxwellian neutron temperature corresponding to the $g_{Lu}(T_n)$ was found to be $47.2\text{ }^\circ\text{C}$.²⁵ These characterized parameters were used in the quantification of REEs. Fig. 3 illustrates a representative gamma spectrum obtained from the HPGe detector for the 6 h neutron irradiated coal ash sample, which was counted for the 5000 s after 1 day cooling time. The gamma spectrum of the neutron-irradiated coal ash sample indicates the presence of REEs like Sm, Eu, La etc. along with other matrix elements, identified by characteristic gamma energies of their activated products. The activation product of sodium as matrix element i.e.²⁴Na emits 1368.6 keV and 2754 keV gamma rays, which causes high Compton background. To mitigate this, the same sample was again counted after the decay of short lived ²⁴Na ($t_{1/2} = 14.96\text{ h}$).

2.3 Leaching of REEs using radioactive tracers

In addition to the above, present study also demonstrates the feasibility of leaching for medium-lived REEs from coal ash in HCl medium using their corresponding radioactive tracers produced via NAA. The NIST 1633b Standard Reference Material (SRM) bituminous coal fly ash, weighing approximately 100 mg, was subjected to neutron irradiation with a flux of $1 \times 10^{11}\text{ n cm}^{-2}\text{ s}^{-1}$ at the PFTS location of the KAMINI reactor, operated

at 20 kW for a duration of 6 h. Subsequently, the irradiated sample was analysed using HPGe detector for estimation of activation products of various rare earths. The irradiated samples were subjected to leaching in a 100 mL of 8 M HCl solution at a temperature of $90\text{ }^\circ\text{C}$, with constant agitation at 200 revolutions per minute (RPM), for a period of 7 hours.²¹ At different time intervals during the leaching process, samples were collected from the leachate in 5 mL gamma vials and subsequently analysed using HPGe detector. The initial two samples were collected at 30 minutes intervals due to high activity and a higher leaching rate. As the leaching rate decreased, the time interval was increased to 1 hour to ensure sufficient activity in the diluted samples.

2.4 Elemental quantification using Westcott formalism in k_0 -NAA

The concentration of the element i i.e. C_i (mg kg^{-1}) using Westcott formalism in k_0 -NAA can be obtained using the following equation²⁵

$$C_i(\text{mg kg}^{-1}) = \frac{\left(\frac{\text{cps}}{SDCW} \right)_i \cdot \frac{1}{k_{0,i}} \cdot \frac{\left(G_{th}g(T_n) + G_e r(\alpha) \sqrt{\frac{T_n}{T_0}} s_0(\alpha) \right)_{\text{Au}} \cdot \varepsilon_{\text{Au}}}{\left(G_{th}g(T_n) + G_e r(\alpha) \sqrt{\frac{T_n}{T_0}} s_0(\alpha) \right)_i \cdot \varepsilon_i} \times 10^6}{\left(\frac{\text{cps}}{SDCW} \right)_{\text{Au}}} \quad (1.0)$$

where cps = counts per second, S is the saturation factor $S = (1 - \exp(-\lambda t_{\text{irr}}))$, D is the decay correction factor $D = \exp(-\lambda t_{\text{d}})$, C is the counting correction factor $C = (1 - \exp(-\lambda t_{\text{c}})) / \lambda t_{\text{c}}$, $k_{0,\text{Au}}$ is the literature reported $k_{0,\text{Au}}$ -factors and ε_{γ} is the detector efficiency, G_{th} and G_e are the thermal and epi-thermal self-shielding



correction factors respectively, $g(T_n)$ is the Westcott g -factor at a neutron temperature T_n , $s_0(\alpha)$ is the modified reduced resonance integral and $r(\alpha)\sqrt{\frac{T_n}{T_0}}$ is modified spectral index.²⁵⁻³¹ The ratio of the modified reduced resonance integral (for a $1/E^{l+\alpha}$ epithermal spectrum) to the 2200 ms^{-1} cross-section is given by

$$s_0(\alpha) = s_0 \left(\overline{E_r} \right)^{-\alpha} (1 \text{ eV})^\alpha \quad (1.1)$$

where s_0 is the corresponding quantity for an ideal $1/E$ epithermal neutron flux distribution and E_r is the effective resonance energy. For " l/v " (n,γ) reactions, s_0 can be calculated as:

$$s_0 = \frac{2}{\sqrt{\pi}} (Q_0 - 0.429) \quad (1.2)$$

The ratio of mass (m) of an element (x) to mass of the internal mono standard element (y) in the coal fly ash was calculated by the following equation in the internal monostandard using Westcott formalism in k_0 -NAA. For obtaining the absolute concentration, the internal mono standard concentration was determined using Energy Dispersive X-ray Fluorescence (EDXRF) as well as relative NAA with minimal sample.

$$\frac{m_x}{m_y} = \frac{\left(\frac{\text{cps}}{\text{SDC}} \right)_x \cdot k_{0,y}}{\left(\frac{\text{cps}}{\text{SDC}} \right)_y \cdot k_{0,x}} \cdot \frac{\left(G_{\text{th}}g(T_n) + G_e r(\alpha) \sqrt{\frac{T_n}{T_0}} s_0(\alpha) \right)_y \cdot \varepsilon_y}{\left(G_{\text{th}}g(T_n) + G_e r(\alpha) \sqrt{\frac{T_n}{T_0}} s_0(\alpha) \right)_x \cdot \varepsilon_x} \quad (1.3)$$

The following Table 1 nuclear data was used for the estimation REEs using above discussed standardization methods.

3 Results and discussion

The REEs in the NIST SRM 1633b CFA were determined using k_0 -NAA as well as internal monostandard approach using Westcott formalism and presented in the Table 2. The uncertainty associated with the result is calculated using error propagation method from the replicate experiments. Iron (Fe), which possesses favourable nuclear properties like appropriate half-life ($T_{1/2}$ of ^{59}Fe 44.5 day), high gamma intensity (1099 keV, 56.5%; 1291 keV, 43.2%), and is abundant within the NIST SRM, was selected as the internal standard for this analytical procedure. This choice enabled for the normalization and accurate quantification of REEs within the coal fly ash sample. Calibration of the high-purity germanium (HPGe) detector for k_0 -NAA was accomplished using a standard point geometry source containing ^{152}Eu . k_0 -NAA requires known activity of ^{152}Eu with identical geometry to achieve the full energy peak efficiency calibration. The uncertainties associated in the efficiency calibration using ^{152}Eu were found to be less than 2%. The calibration process covers the energy range of 121.8 to 1408 keV, as illustrated in Fig. 4. The gamma energies *i.e.* less than the 121.8 keV are not available in the case of ^{152}Eu for the detector efficiency calibration in lower energy region. Nuclides emitting gamma radiation with energies lower than 121.8 keV, such as ^{153}Sm (69.7 keV, 103.2 keV), ^{165}Dy (94.7 keV), ^{166}Ho (80.6 keV), and ^{170}Tm (84.3 keV), posed a challenge for the quantification within the framework of k_0 -NAA.

These challenges were resolved in the case of internal monostandard approach *i.e.* the in-site relative detector efficiency calibration has been carried out in the range of 69.7 to 2754 keV using the gamma energy of activation products itself *i.e.* ^{24}Na (1368 and 2754 keV), ^{46}Sc (889.3, 1120.5 keV), ^{140}La

Table 1 Relevant nuclear data^{27,31} for the estimation of REEs using k_0 -NAA

Target isotope	Formed isotope	$T_{1/2}$	E_γ (keV)	$k_{0,\text{Au}}$	Q_0	$(\overline{E_r})$ (eV)	s_0	$s_0(\alpha)$
^{139}La	^{140}La	1.678 day	1596.2	1.34×10^{-1}	1.24	76	0.92	1.13
^{140}Ce	^{141}Ce	32.51 day	145.4	3.66×10^{-3}	0.83	7200	0.45	0.70
^{141}Pr	^{142}Pr	19.12 h	1575.6	6.12×10^{-3}	1.51	296	1.22	1.62
^{146}Nd	^{147}Nd	10.98 day	91.1	1.02×10^{-3}	2.00	874	1.77	2.48
^{146}Nd	^{147}Nd	10.98 day	531	4.56×10^{-4}	2.00	874	1.77	2.48
^{152}Sm	^{153}Sm	46.50 h	69.7	3.52×10^{-2}	14.4	8.53	15.77	17.53
^{152}Sm	^{153}Sm	46.50 h	103.2	2.31×10^{-1}	14.4	8.53	15.77	17.53
$^{151}\text{Eu}^a$	^{152m}Eu	9.312 h	121.8	1.48	—	0.448	1.20	1.15
$^{151}\text{Eu}^a$	^{152m}Eu	9.312 h	841.6	3.02	—	0.448	1.20	1.15
^{152}Gd	^{153}Gd	240.4 day	97.4	5.86×10^{-3}	0.77	16.7	0.38	0.44
^{158}Gd	^{159}Gd	18.56 h	363.5	8.49×10^{-4}	29.9	48.2	33.26	40.28
^{159}Tb	^{160}Tb	72.3 day	298.6	8.25×10^{-2}	17.9	18.1	19.72	22.75
^{159}Tb	^{160}Tb	72.3 day	879.4	9.42×10^{-2}	17.9	18.1	19.72	22.75
^{164}Dy	^{165}Dy	2.334 h	94.7	3.57×10^{-1}	0.19	224	-0.27	-0.35
^{165}Ho	^{166}Ho	26.83 h	80.6	4.94×10^{-2}	10.9	12.3	11.82	13.38
^{170}Er	^{171}Er	7.516 h	308.3	1.04×10^{-2}	4.42	129	4.50	5.73
^{169}Tm	^{170}Tm	128.6 day	84.3	3.26×10^{-2}	13.7	4.8	14.98	16.19
^{174}Yb	^{175}Yb	4.185 day	282.5	1.46×10^{-2}	0.46	602	0.03	0.05
^{174}Yb	^{175}Yb	4.185 day	396.3	3.12×10^{-2}	0.46	602	0.03	0.05
$^{176}\text{Lu}^a$	^{177}Lu	6.73 day	112.9	4.15×10^{-2}	—	0.158	1.67	1.52
$^{176}\text{Lu}^a$	^{177}Lu	6.73 day	208.4	7.14×10^{-2}	—	0.158	1.67	1.52

^a s_0 Values for non- $1/v$ nuclides were taken from ref. 31.



Table 2 REEs estimation in NIST SRM 1633b using NAA. Accuracy and precision of k_0 -NAA and IM-NAA methods were found to be nearly identical. Analysis of certain elements like Sm, Dy, Ho, and Tm were feasible only by IM-NAA

Element	Conc. (mg kg^{-1}) $\pm u_{\text{lab}}$		Conc. (mg kg^{-1}) $\pm u_{\text{cert}}$	Deviation (%)		E_n number	
	k_0 -NAA	IM-NAA		Certificate value	k_0 -NAA	IM-NAA	k_0 -NAA
La	92.0 ± 3.2	91.0 ± 2.1	94		2.1	3.2	—
Ce	185 ± 5.8	196 ± 9.6	190		2.6	-3.2	—
Pr	21.8 ± 1.1	20.0 ± 1.0	21		-3.8	4.8	—
Nd	86.5 ± 4.1	86.0 ± 3.1	85		-1.8	-1.2	—
Sm	—	21.0 ± 0.9	20			-5.0	—
Eu	4.20 ± 0.20	4.00 ± 0.15	4.1		-2.4	2.4	—
Gd	13.6 ± 0.6	12.6 ± 0.6	13		-4.6	3.1	—
Tb	2.47 ± 0.11	2.64 ± 0.09	2.6		5.0	-1.5	—
Dy	—	16.3 ± 0.9	17			4.1	—
Ho	—	3.36 ± 0.10	3.5			4.0	—
Er	8.30 ± 0.30	8.95 ± 0.40	8.7^a		4.6	-2.9	—
Tm	—	2.00 ± 0.08	2.1			4.8	—
Yb	7.27 ± 0.20	7.69 ± 0.30	7.6		4.3	-1.2	—
Lu	1.15 ± 0.05	1.16 ± 0.06	1.2		4.2	3.3	—
As	134.3 ± 2.9	134.7 ± 2.4	136.2 ± 2.6		1.4	1.1	-0.48
Cr	197.2 ± 3.8	196.3 ± 4.2	198.2 ± 4.7		0.5	1.0	-0.16
Th	26.1 ± 1.2	25.1 ± 1.2	25.7 ± 1.3		-1.6	2.3	0.23
U	8.90 ± 0.30	8.71 ± 0.20	8.79 ± 0.36		-1.3	0.9	0.23
K (%)	1.97 ± 0.04	1.96 ± 0.05	1.95 ± 0.03		-1.0	-0.5	0.40
Na (%)	0.202 ± 0.005	0.203 ± 0.004	0.201 ± 0.003		-0.5	-1.0	0.17
							0.40

^a GeoRem database (INAA).

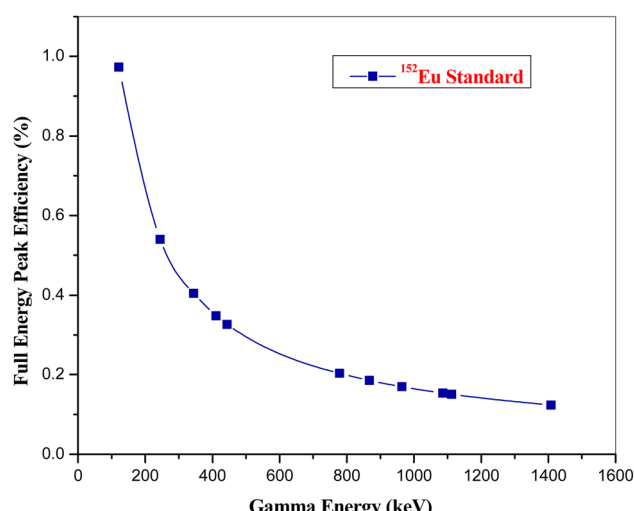


Fig. 4 Full energy peak efficiency calibration using ^{152}Eu standard.

(328.8, 487.0, 815.8 and 1596.5 keV), ^{147}Nd (91.1 and 531.0 keV), ^{152}Eu (121.9, 963 and 1408 keV), ^{153}Sm (69.7 and 103 keV), ^{175}Yb (282.5 and 396.5 keV) and ^{177}Lu (112.9 and 208.4 keV). The known activity of the activation products is not required for the *in situ* relative detector efficiency calibration, which makes the process geometrically independent. The nuclide emits minimum a pair of gamma energy was used for the efficiency calibration in the energy range of interest. The efficiency ratios for a nuclide were calculated using the ratios of peak area and intensity. The fitted ratios were obtained corresponding to the same energy as the measured ratios using the following equation.

$$\left(\frac{\varepsilon_1}{\varepsilon_2} \right) = \exp \left[A_1 [\ln(E_1) - \ln(E_2)] + A_2 [(\ln E_1)^2 - (\ln E_2)^2] \right. \\ \left. + \dots + A_n [(\ln E_1)^n - (\ln E_2)^n] \right] \quad (1.4)$$

The coefficients $A_1, A_2, A_3, \dots, A_n$ were determined using the least square fitting method. The full energy peak efficiency calibration obtained by using the following equation.

$$f(x_i) = \ln(\varepsilon) = \sum_{i=0}^n A_i [\ln(E)]^i \quad (1.5)$$

where ε is the efficiency and E is the Gamma ray energy. The deviation between the calculated and fitted efficiency ratios was found to be less than 2% in the high energy region, while in the lower energy region, it was found to be in the range of 5–10%. A typical plot of *in situ* relative efficiency calibration has been shown in Fig. 5. The relative efficiency pattern was found to be similar to the intrinsic efficiency calibration in the low energy region with the peaking at the region of 110–120 keV. This relative efficiency calibration data was used in the internal monostandard approach in k_0 -NAA.

In the context of REEs estimation, it is observed that the majority of elements exhibit a thermal capture cross-section behaviour proportional to $1/v$, where ' v ' represents the neutron velocity. However, two specific nuclear reactions, namely $^{176}\text{Lu}(n,\gamma)^{177}\text{Lu}$ and $^{151}\text{Eu}(n,\gamma)^{152m,152}\text{Eu}$, deviate from the $1/v$ behaviour, as depicted in Fig. 6. This deviation from the expected $1/v$ behaviour due to the resonance involved into the thermal energy region is quantified using a parameter known as the Westcott factor ($g(T_n)$). The Westcott factor serves as



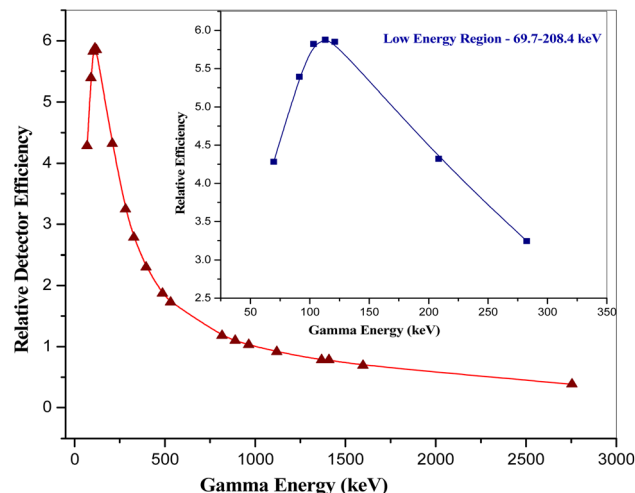
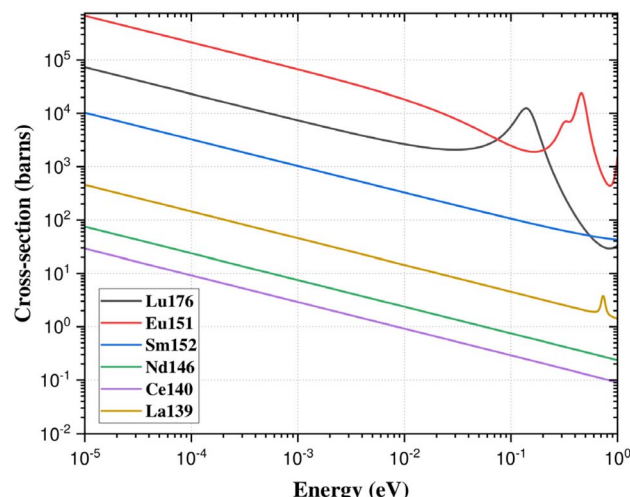
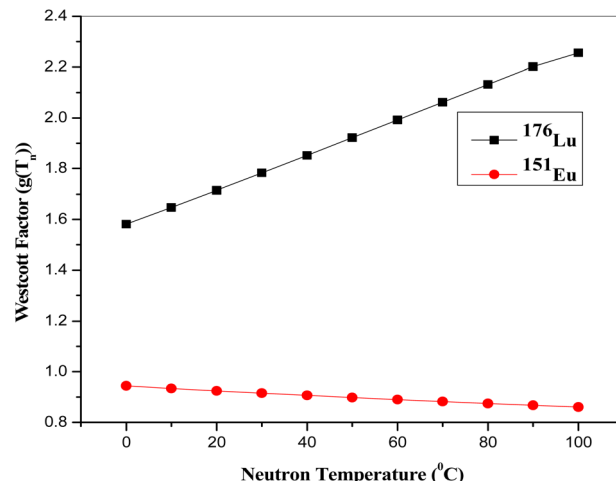
Fig. 5 *In situ* relative detector efficiency calibration.

Fig. 6 Variation of capture cross-section with neutron energy of REEs in thermal region.

a quantitative measure of the departure from the $1/\nu$ behaviour and is known to linearly correlate with the neutron temperature. For the assessment and characterization of non- $1/\nu$ behaviours, detailed calculations for the above nuclear reactions were performed using cross-section data of ENDF/B-VIII.0 data (USA, 2018) nuclear data library in the temperature range of 0 to 100 °C (Fig. 7). The $g(T_n)$ values were particularly calculated from the linear equation, using the recently measured neutron temperature in the PFTS irradiation channel of the KAMINI reactor.²⁵ For nuclides that adhere to the $1/\nu$ relationship, $g(T_n)$ is equal to 1.

The spectral interferences were found in the low energy region due to emission of same gamma energy by other radioactive nuclide formed in the sample matrix. The gamma line 103.2 keV of ^{153}Sm is interfered by the ^{153}Gd (240.4d, $E_\gamma = 103.2$ keV) as well as $^{232}\text{Th}(\text{n},\gamma) \rightarrow ^{233}\text{Th} \rightarrow ^{233}\text{Pa}$ ($t_{1/2} = 26.97$ day and $E_\gamma = 103.9$ keV) and $^{238}\text{U}(\text{n},\gamma) \rightarrow ^{239}\text{U} \rightarrow ^{239}\text{Np}$ ($t_{1/2} = 2.357$ day and $E_\gamma = 103.7$ keV). Other similar cases are the following γ -line

Fig. 7 Westcott factor for the non- $1/\nu$ nuclides using ENDF/B-VIII.0 data (USA, 2018).

interferences of ^{239}Np on the 80.6 keV of ^{166}Ho and ^{153}Gd on the 69 keV of ^{153}Sm . To mitigate these interferences, samples were analysed at different time intervals. Correction factors for the interfering gamma emissions were applied, utilizing the measured activity corresponding to the non-interfering gamma ray. The magnitude of the interference depends upon the half-life and gamma intensity, in the case of ^{153}Sm (103.2 keV) the impact of ^{239}Np was found to be more pronounced than that of ^{233}Pa and ^{153}Gd .

The presence of uranium (U) and thorium (Th) in the coal ash samples could become a possible source of the error for the estimation of lower REEs like ^{140}La , ^{141}Ce , ^{143}Ce and ^{147}Nd due to high fission yield. The average concentration of U and Th in the coal ash was found to be 10 and 30 mg kg⁻¹ respectively. However, due to the low concentration, small sample size and low flux at the PFTS location, their impact on determining the concentration of lower REEs was deemed to be negligible. The quality assurance of k_0 -NAA and IM-NAA with the Westcott formalism has been tested by calculating the % deviation and E_n number with that of the certificate values. The following equation was used to calculate the E_n number.³⁴

$$E_n = \frac{X_{\text{lab}} - X_{\text{cert}}}{\sqrt{(u_{\text{lab}})^2 + (u_{\text{cert}})^2}} \quad (1.6)$$

where, X_{lab} is the measured elemental concentration in the present work, X_{cert} is the certified elemental concentration and u_{lab} is the uncertainty in the measured elemental concentration expanded at 95% confidence interval and u_{cert} represents uncertainty of the certificate values in the same confidence interval. The uncertainty in the measured elemental concentration at 95% confidence interval were found to be within $\pm 5\%$ for most of the elements. Due to the high abundance or greater sensitivity, the deviation from the certificate concentration for La, Ce, Nd, and Eu was found to be minimal. In contrast, elements that emit low-energy gamma rays, such as Sm, Dy, Ho, and Tm, exhibit slightly higher deviations, which also may be

Table 3 REEs in the Indian coal ash samples

REEs	Coal ash samples conc. (mg kg^{-1}) $\pm u_{\text{lab}}$)					Worldwide conc. (mg kg^{-1})
	Fly ash	Pond ash	Clinker ash	Bottom ash		
La	120 \pm 4	96 \pm 3	60 \pm 3	108 \pm 3	15.5–134.4	
Ce	208 \pm 4	176 \pm 3	141 \pm 4	212 \pm 4	30.7–266.4	
Pr	29.0 \pm 0.9	22.0 \pm 0.6	12.5 \pm 0.2	19.5 \pm 0.4	3.3–29.8	
Nd	68.2 \pm 1.2	46.0 \pm 1.2	63.0 \pm 1.0	84.0 \pm 1.4	12.7–114.7	
Sm	20.3 \pm 0.4	15.3 \pm 0.3	16.6 \pm 0.2	16.9 \pm 0.3	2.8–22.9	
Eu	4.30 \pm 0.12	2.47 \pm 0.10	0.80 \pm 0.02	3.80 \pm 0.20	0.56–6.3	
Gd	9.9 \pm 0.3	5.1 \pm 0.3	13.1 \pm 0.4	10.6 \pm 0.2	2.85–30.7	
Tb	2.50 \pm 0.10	1.80 \pm 0.06	1.90 \pm 0.09	2.40 \pm 0.06	0.45–4.7	
Dy	14.0 \pm 0.3	16.0 \pm 0.5	13.5 \pm 0.4	19.0 \pm 0.4	2.61–25.1	
Ho	2.50 \pm 0.10	1.80 \pm 0.08	2.50 \pm 0.20	2.30 \pm 0.15	0.59–4.5	
Er	5.10 \pm 0.20	6.50 \pm 0.25	6.70 \pm 0.15	4.5 \pm 0.10	1.79–14	
Tm	2.50 \pm 0.10	1.70 \pm 0.09	1.30 \pm 0.09	0.90 \pm 0.03	0.27–1.5	
Yb	8.90 \pm 0.30	5.40 \pm 0.20	7.00 \pm 0.15	7.2 \pm 0.25	1.8–11	
Lu	1.40 \pm 0.05	1.10 \pm 0.03	0.77 \pm 0.01	0.94 \pm 0.03	0.3–1.6	

due to the higher uncertainty associated in the gamma self-attenuation correction. The % deviations for the elements determined in 1633b CFA with respect to certificate values were also found to be within $\pm 5\%$, whereas the $|E_n|$ values for the certified concentration were less than one. The findings of this study are consistent with the assigned values, falling within the expected range of uncertainty margins and validates the methodology.

3.1 Rare earth in Indian coal ash sample

Rare earths are a group of 17 elements comprising of 15 “Lanthanides”, lanthanum (La) to lutetium (Lu) and two others, yttrium (Y) and scandium (Sc). The present study focused only on lanthanides, among them promethium (*Pm*) does not occur in the Earth's crust. Similar methodology as discussed above was adopted for the assay of REEs in fly ash samples and the results so obtained were presented in Table 3. The REEs concentration varies from 0.7 (Lu) to 210 (Ce) mg kg^{-1} . The

concentration of different REEs in the different ash samples are plotted in Fig. 8 and found identical patterns of REEs in all the samples. Total REEs concentration in the Indian coal ash varies in the range of 350–500 mg kg^{-1} .

Lin *et al.*, 2017 (ref. 34) suggested that the estimated cut-off grade for beneficial recovery of REEs in coal could be 115–130 mg kg^{-1} and 677–762 mg kg^{-1} on the ash basis. Coal combustion byproducts typically contain 200–500 mg kg^{-1} of REEs, with a global average of 445 mg kg^{-1} in coal fly ash. Based on this data, REEs in Indian fly ash is close to the beneficial recovery and falls in the world range.³⁵ Among the REEs, the cerium (Ce) was found to be most abundant in all samples; its concentration varies from 41–44%, while lutetium (Lu) was found to be the least in the range of 0.19–0.28%. The abundance of individual REE with respect to the total REEs in the coal ash samples is shown in Fig. 9. The light REEs (LREE) from La–Sm found dominance (86–90%) over the medium REEs (MREE) Eu–Dy (6.4–8.6%) and heavy REEs (HREE) Ho–Lu (3.2–5.4%).

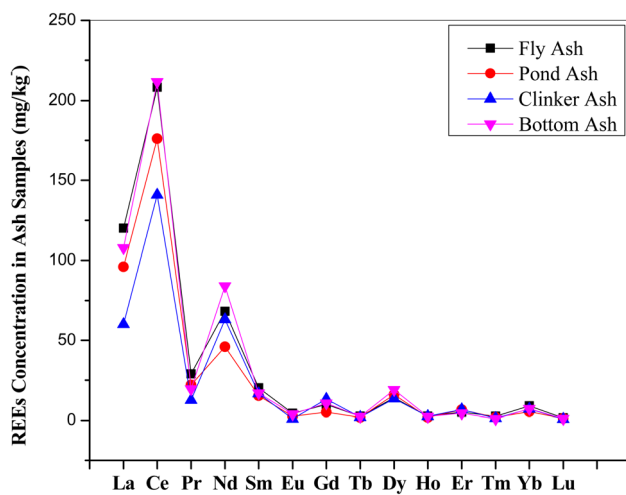


Fig. 8 REEs concentration in the coal ash samples.

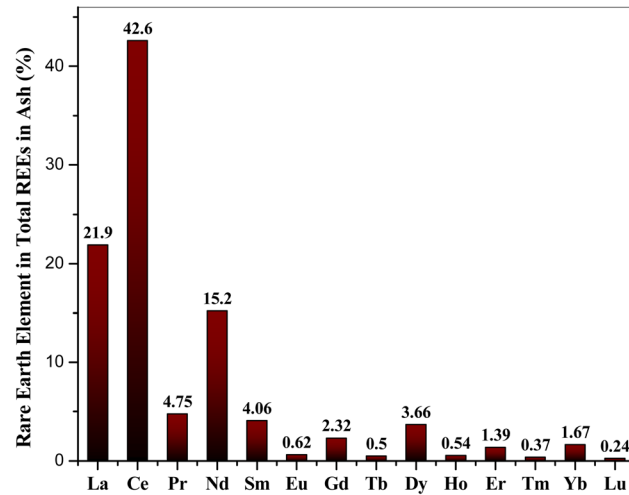


Fig. 9 Contribution of individual REE for total REEs in the ash samples.



3.2 Grouping of the coal ash samples by hierarchical clustering

Hierarchical clustering, a powerful statistical technique, facilitates the systematic grouping of akin data points within a hierarchical framework. Within the Python code framework, this technique is realized through the utilization of a linkage matrix. A pivotal component of hierarchical clustering is the utilization of a distance metric, typically Euclidean distance, to quantify dissimilarity between data points. In the code, the Euclidean distances between every pair of data points are computed. In this specific implementation, the 'single' linkage method is employed. This linkage method computes the distance between clusters by assessing the minimum pairwise distance among the constituent data points within and between clusters. This approach encourages the emergence of elongated clusters, unveiling latent patterns within the data. The linkage matrix, a key outcome of the hierarchical clustering procedure, encodes the intricate hierarchy of formed clusters. At each iteration, this matrix delineates the clusters scheduled for amalgamation, detailing their inter-cluster distance and the count of data points encompassed within the resulting amalgamated cluster. The dendrogram, a graphical manifestation of the hierarchical clustering process, plays an integral role in conveying the clustering outcomes. It graphically represents the hierarchical relationships between ash samples based on their total content of REEs concentrations (Fig. 10). This dendrogram provides invaluable insights into the degree of similarity among ash samples, thereby enabling data-driven decision-making informed by the latent structure inherent in the dataset.

In the specific dendrogram under consideration, a noteworthy observation is the complete isolation of the clinker ash sample from all other samples, signifying its distinctiveness. This distinction suggests that the distribution of REEs within the clinker ash sample significantly deviates from the distribution observed in the remaining samples. Within this dendrogram, an evident similarity emerges between the bottom ash and fly ash samples, as they are seen to unite at an early stage in the diagram. Progressing upward, the subsequent union links the clade containing the bottom and fly ash samples with the clade housing the pond ash sample. This structural arrangement conveys that all

samples within this cluster exhibit greater similarity to each other compared to any samples connecting at higher dendrogram levels. Additionally, a notable inference arises from the positioning of the pond ash sample, which is found to be closer in similarity to the bottom and fly ash samples than to the clinker ash sample. A similar pattern has been observed with respect to the calculated total standard deviation also.

3.3 Leaching of REEs from coal ash using radioactive tracers

Leaching of REEs from coal ash has been carried out using radioactive isotopes formed by irradiating the sample into KAMINI reactor. Radiation detection methods are highly sensitive, rapid and assay could be carried out on direct solid samples unlike conventional analytical techniques. Thus, the irradiated coal ash samples were counted directly using HPGe detector to obtain the initial reference activities, which are normalized to the end of irradiation (EOI). The initial activity (A_{EOI}) of the medium-lived REEs was determined, employing the efficiency of a point geometry source with ^{152}Eu and by incorporating all the decay corrections.

$$A_{EOI} = \frac{\text{cps}}{I_{\gamma} \times \varepsilon \times D \times C} \quad (1.7)$$

where I_{γ} = gamma ray intensity and ε = detector efficiency, similar to the initial activity, the activity was calculated for each collected leachate sample, accounting for radioactive decay and volume corrections. The percentage leaching of each REE has been calculated using the following equation.

$$\text{Leaching percentage} = \frac{[A_{EOI}]_{\text{leachate}}}{[A_{EOI}]_{\text{Initial}}} \times 100 \quad (1.8)$$

In the course of our investigation, it was observed that among the diverse medium-lived REEs, specifically La with 55–60% and Eu with 65–70%, were found to be present in the leached fraction of the sample in the detection range as shown in Fig. 11. Conversely, all other REEs were detected at levels lower than the

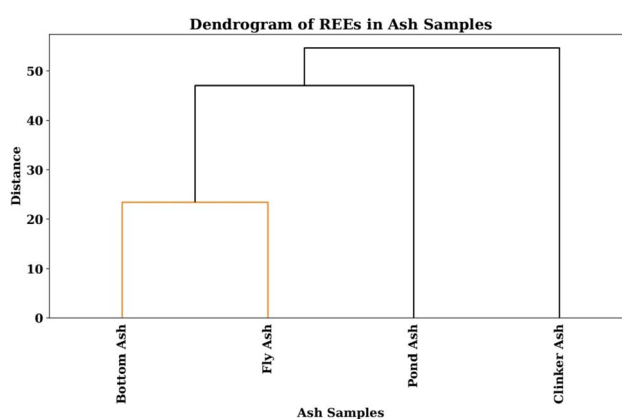


Fig. 10 Dendrogram of REEs in ash samples.

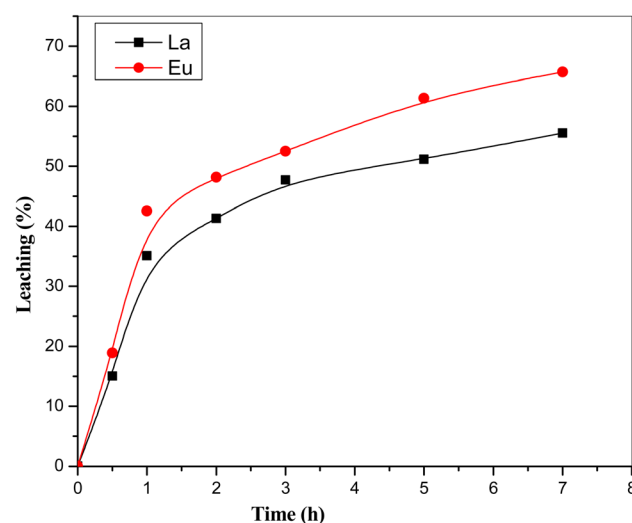


Fig. 11 Leaching of La and Eu from coal ash sample using radioactive tracer method.



Table 4 Normalized activity of REEs before and after leaching

Element	Activity (Bq)		
	Before leaching	After leaching	% Leaching
La	1.443	0.589	59.2
Ce	0.007	0.004	51.3
Pr	0.799	0.356	55.4
Nd	0.012	0.004	64.0
Sm	1.383	0.578	58.2
Eu	33.26	12.46	62.6
Yb	0.185	0.073	60.6
Lu	0.036	0.019	48.4

detection limit of our analytical methodology due to lower counting time. Following the leaching process, the remaining sample was subjected to a thorough drying procedure and subsequently packed in a consistent geometric configuration and subjected to analyse by the same detector. The analysis of the remaining activity associated with the medium-lived REEs was performed and the results have been presented in Table 4.

It is worth noting that the data from Table 4 reveal a substantial leaching efficiency for the medium-lived REEs in the presence of 8 M HCl. Specifically, it was observed that the leaching process facilitated the removal of approximately 50–70% of the medium-lived REEs isotopes from the sample with the uncertainty $\pm 5\%$, thereby indicating the efficacy of 8 M HCl as a leaching agent in this context. Similar study has been carried out with 4 M HCl also and found the almost same leaching pattern. These leaching parameters like leaching time and concentration of HCl were fixed as per the Shanshan Cao *et al.*²¹ The results on leaching behaviour using the radioactive tracer technique were in good agreement with the reported literature by Shanshan Cao *et al.*²¹ However, the study²² was limited to report the leaching behaviour of only three REEs *i.e.* La, Ce and Nd, whereas the present study could explore eight REEs including Pr, Sm, Eu, Yb and Lu. Thus, the present study established the strengths of radioactive tracer method, which is highly sensitive, rapid and independent of sample physical state (solid/liquid). These findings shed light on the behaviour of these REEs in the leaching process and contribute valuable insights to our understanding of their chemical interactions and separations.

4 Conclusions

The study involved the objective of quantifying the content of REEs in coal samples using NAA coupled with the Westcott formalism. Coal fly ash samples from different thermal power plants in India were utilized, and high-resolution gamma spectrometry was employed to quantify the REE content of the samples. By coupling the internal standard and *in situ* relative detector efficiency strategies, the analytical precision and accuracy in the assessment of REE concentrations in complex matrices are substantially enhanced, facilitating a more comprehensive and reliable characterization of these critical

elements. Westcott factor play a crucial role in estimating the elemental concentrations of REEs, providing insights into the neutron capture characteristics of these nuclides. This scientific approach allows for a more accurate understanding of neutron interactions with REEs and their subsequent quantification. This scientific investigation highlights the challenges posed by spectral interferences in gamma-ray spectroscopy and provides insights into the correction methods employed to ensure accurate quantification. The study found that the individual REEs concentration varied from 0.7 mg kg^{-1} to 210 mg kg^{-1} . The LREE found dominance over the MREE and heavy HREE. The results of this study could aid in the development of eco-friendly and sustainable technological advancements by providing a better understanding of the REE content in coal ash samples. This study provides valuable insights into the leaching behaviour using radioactive tracer of medium-lived REEs from coal ash, contributing to our understanding of REE recovery and utilization from this unconventional source.

Data availability

The data that support the findings of this study are available at <https://www.nndc.bnl.gov/endf> and in the ref. 27–31.

Conflicts of interest

All co-authors have gone through and agreed with the contents of the manuscript and there is no conflict of interest to declare.

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