RSC Advances



PAPER

View Article Online
View Journal | View Issue



Cite this: RSC Adv., 2025, 15, 4546

Ratiometric discrimination of Th⁴⁺ ions by a fluorogenic quinoline appended phenanthridine sensor and its applications†

Selin Manoj Kumar,^a Sathishkumar Munusamy,^b Saravanan Enbanathan, ^b Sivaranjani Ranganadin^a and Sathiyanarayanan Kulathu Iyer ^b *

A new phenanthridine appended quinoline-based chemoreceptor 5-(5-(quinolin-8-yl)thiophen-2-yl)tetrahydrodibenzo[a,i]phenanthridine (PHQBA) was successfully synthesized and characterized by 1 H, 13 C, and HRMS spectral analyses. The promising chelation-induced process in PHQBA was accelerated by Th $^{4+}$ ions, which impart robust ratiometric green fluorescence at 515 nm. The host–guest complex formed in a 1:1 binding stoichiometry between Th $^{4+}$ ions and PHQBA was demonstrated by Jobs plot experiments. The Benesi–Hildebrand (BH) plot was employed to compute the binding constant for the complexation of PHQBA + Th $^{4+}$, which was determined to be 3.77 \times 10 5 M $^{-1}$. To comprehend the detection mechanism of PHQBA, DFT, and TDDFT, eased computational studies were conducted and well supported by the experimental results. Moreover, the limit of detection (LOD) of PHQBA was determined as 223 nM, which defines the remarkable optical sensitivity of the sensor PHQBA. Further, portable paper strip detection and real-time determination of Th $^{4+}$ ions in real water samples ensure the practical applications of PHQBA.

Received 17th December 2024 Accepted 3rd February 2025

DOI: 10.1039/d4ra08840a

rsc.li/rsc-advances

Introduction

Thorium is a potentially important radioactive metal in the earth's crust that is extensively used in modern industrial platforms. It has plenty of applications in the formulation of lantern mantles, lenses, carbon lamps, ceramics, and welding appliances.² Since thorium exists in a +4-oxidation state, most nuclear fuel recycling is accomplished in a medium with high acidity.3,4 Undesirably, thorium ions create serious contamination in waterbodies such as seas, lakes, and rivers which immensely affects living creatures. Also, the toxicity of thorium ions can cause renal injury and dermatitis.5 According to the reports of the World Health Organisation (WHO), the permissible limit of thorium is 0.6 μg per kg per day.6 Therefore, there is a need to develop a precise method for the quantification of thorium ions. Many spectroanalytical techniques are available, but it can be challenging to identify target analytes due to their complex operating procedures. Examples of these techniques include electrothermal atomic absorption spectroscopy (ETASS),7 inductively coupled plasma emission spectroscopy (ICP-AES),8 accelerator mass spectrometry (AMS),9 laser ablation

microprobe mass analyzer (LAAMA),10 atomic absorption spectroscopy (AAS)11 and neutron activation analysis (NAA).12 While comparing these approaches, fluorescent sensing is an emerging alternative to conventional methods 13-18 for the detection of analytes due to their selectivity, sensitivity, rapid response, and ease of access. Detection of heavy metal ions follows various photochemical mechanisms including Forster resonance energy transfer (FRET),19 intramolecular charge transfer (ICT),20 exited state intramolecular charge transfer (ESIPT),21 and twisted intramolecular charge transfer (TICT).22 Among all these photochemical processes, chelation enhanced fluorescence process predominantly produces a well-defined spectral response that accelerates sensitive ratiometric detection of analytes. 23-30 Due to the well-functionalized design 31,32 of phenanthridine-based chemosensing systems, it has gained significant attention, which plays a vital role in spectroscopic properties. The advantages of these sensors include bright emission,33 large Stokes shift,34 and ease of emission wavelength modulation35,36 upon analyte interaction. These features are necessary for the rational construction of ratiometric sensors. While comparing turn-on and turn-off fluorescent probes, ratiometric chemosensors could provide precise calibration by decoding dual emission ratio signals, which could improve the sensitivity with great accuracy during the determination of target analytes.37-41 Hence, designing a ratiometric fluorescent sensor to identify Th4+ ions with greater sensitivity and selectivity is crucial. In our group, phenanthridine-derived chemosensors have recently been employed as effective tools for

^aDepartment of Chemistry, School of Advanced Sciences and Vellore Institute of Technology, Vellore-632014, India. E-mail: sathiya_kuna@hotmail.com

^bDepartment of Chemistry, University of Missouri-Columbia, Columbia, MO, USA. E-mail: pra3sat@gmail.com

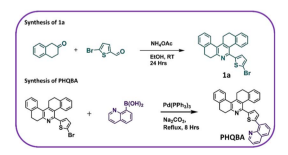
[†] Electronic supplementary information (ESI) available. See DOI https://doi.org/10.1039/d4ra08840a

the selective detection of thorium ions. These sensors exhibit a turn-on fluorescence response⁴²⁻⁴⁴ upon interaction with thorium ions, enabling low detection limits. Ratiometric fluorescent probes are particularly advantageous as they provide signal ratios, enhancing the dynamic range and offering built-in correction for environmental effects. The observed emission color change is valuable for ratiometric fluorescence detection. To improve the optical performance of the sensor, we have rationally incorporated quinoline conjugation into the phenanthridine system. In our endeavor to develop a novel phenanthridine-derived ratiometric thorium ion sensor, PHQBA has been constructed. In the detection process, the chelation of heavy metal (Th4+) to the electron-rich PHQBA ligand, results in the formation of a new PHQBA + Th⁴⁺ complex. As a result, ratiometric fluorescence has been accomplished. Fascinatingly, the sensor PHQBA exclusively detects Th⁴⁺ ions even in the presence of other cations, thereby ensuring anti-interference properties. The detection mechanism of PHQBA is well strengthened by DFT, HRMS, and NMR titration experiments. Further, the Jobs plot and Benesi-Hildebrand (B-H) plot depict the binding stoichiometry and association constant of Th⁴⁺ ions. On account of their greater emission properties, portable fluorescent paper strips were exploited for the selective and sensitive discrimination of Th⁴⁺ ions. In addition, Th4+ ions were successfully detected in environmental water samples with excellent recovery.

Results and discussion

Design and synthesis of PHQBA

In general, phenanthridine-based fluorescent sensors possess well-defined spectral responses, which is highly advantageous for analyte detection owing to their ability to hinder the selfabsorption and inner filtering effects. 45-47 The designing of a thiophene-appended quinoline moiety to the phenanthridine system facilitates the chelation-enhanced fluorescence (CHEF) process with Th⁴⁺ ions. As shown in Scheme 1, the design of the chemoreceptor PHQBA includes two steps. In step one, compound 1a was successfully synthesized by the reaction of 5bromo-2-thiophenecarboxaldehyde and 2-tetralone. Finally, quinoline moiety was introduced to the phenanthridine system by a Suzuki coupling reaction to extend the conjugation of the phenanthridine system (PHQBA) for achieving ratiometric fluorescence detection (Scheme 1). The compounds were characterized by ¹H, ¹³C NMR, and HRMS analyses (Fig. S1-S5†).



Scheme 1 Synthesis of sensor PHQBA

Absorption studies of PHQBA

Optical selectivity is considered to be a crucial tool for the evaluation of a chemosensor. In this regard, the influence of absorption on the detection properties of PHQBA was initially inspected in the presence of various cation solutions such as Ca²⁺, Ba²⁺, Al³⁺, Cd²⁺, Cu²⁺, Th⁴⁺, Co²⁺, Fe²⁺, Fe³⁺, Hg²⁺, Ni²⁺, Cr³⁺, Pb²⁺, Pd²⁺ Zn²⁺ La³⁺, and Yb³⁺ in ACN/H₂O (8:2 v/v). As illustrated in Fig. 1a, PHQBA displayed an explicit absorption band at 278 and 380 nm, which could be accredited to π – π * and $n-\pi^*$ electronic transition respectively. Competitive cations have no significant impact on the absorption of PHQBA. However, the introduction of the Th⁴⁺ ion to the **PHQBA** solution resulted in the appearance of a new hypsochromic absorption band at 312 nm. The resulting absorption band with a hypsochromic shift⁴⁸ demonstrated the active chelation of Th⁴⁺ ions with the sensor **PHQBA**. Further scrutiny of **PHQBA**'s detection properties with Th⁴⁺ ions was carried out through the emission titration experiments (Fig. 1b). The gradual addition of thorium ions (0-11 equiv.) to the PHQBA solution resulted in the decreased absorption intensity at 380 nm and the subsequent enhancement aroused at 312 nm. The shifts in the absorption bands to produce isosbestic points at 290 and 342 nm revealed the formation of a new complex PHQBA + Th⁴⁺ These findings prove that the sensor PHQBA has selective and sensitive detection capability with Th4+ ions. To investigate the effect of pH on the detection properties of PHQBA, absorption experiments were performed in the presence and absence of Th⁴⁺ ions. As shown in Fig. S12,† the absorption intensity of PHQBA is nearly stable over a broad pH range of 6-12 and shows an enhanced intensity at pH 2-5, likely due to the protonation of the PHQBA. Upon the addition of Th⁴⁺ ions, the absorption intensity of the PHQBA + Th⁴⁺ complex persists stable within the pH range of 6-9 and began to decrease beyond pH 9, due to the hydrolysis of thorium ions. These results indicate that the sensor is capable of responding to Th⁴⁺ ions across a pH range of 2-12.

Emission studies of PHQBA

Inspired by the absorption experiments, the detection ability of sensor PHQBA was examined in ACN/H2O (8:2, v/v) by treating with competing metal ion solutions including Ca²⁺, Ba²⁺, Al³⁺, Cd²⁺, Cu²⁺, Th⁴⁺, Co²⁺, Fe²⁺, Fe³⁺, Hg²⁺, Ni²⁺, Cr³⁺, Pb²⁺, Pd²⁺ Zn²⁺, La³⁺, and Yb³⁺. As illustrated in Fig. 2a, the free sensor

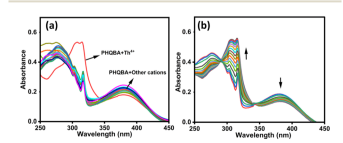


Fig. 1 (a) Absorption spectra of PHQBA (2 \times 10⁻⁵ M) in the presence of competing cations (1 \times 10⁻⁴ M); (b) absorption spectra of PHQBA in the presence of varying concentrations of thorium ions (0-11 equiv.).

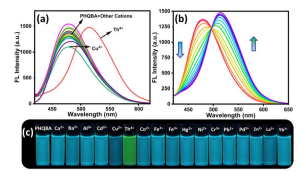


Fig. 2 (a) Fluorescence spectra of PHQBA (2×10^{-5} M) in the presence of various cations (1×10^{-4} M); (b) fluorescence spectra of PHQBA with the gradual addition of Th⁴⁺ (0–11 equiv.); (c) emission color changes of PHQBA in the presence of series of cations ($\lambda_{ex}=389$ nm).

PHQBA showed a maximum emission intensity of 478 nm. After the addition of Th4+ ions, the emission band at 478 nm redshifted to 515 nm, which could be attributed to the synergic binding of Th⁴⁺ ion with the sensor PHQBA. Notably, other cations did not cause any significant spectral alterations as they led to no discernible binding interactions with the sensor PHQBA. To explore the sensitive detection properties of PHQBA with Th⁴⁺ ions, emission titration experiments were performed in ACN/H₂O (8:2, v/v) medium. Thereby, the gradual addition of Th⁴⁺ ions (0-11 equiv.) to the **PHQBA** solution resulted in diminished emission intensity at 478 nm, and the successive emission enhancement was attained at 515 nm (Fig. 2b). The emergence of a clear iso-emissive point at 496 nm indicated the formation of a new complex PHQBA + Th⁴⁺. As a result, the emission color of PHQBA transformed from blue to green (Fig. 2c). The calculated quantum yield values of **PHOBA** and its complex PHQBA + Th4+ were found to be 0.12 and 0.14 respectively. The slight enhancement in the quantum yield corresponds to the redshifted ratiometric fluorescence of PHQBA on chelating with Th⁴⁺ ions. These findings provided further evidence that sensor PHQBA had strong fluorescence detection abilities towards Th⁴⁺ ions. To clearly distinguish the emission of PHQBA and its complex PHQBA + Th⁴⁺, the chromaticity plot (CIE) was plotted⁴⁹ using the emission spectra. As demonstrated in Fig. S8,† the color coordinates of the free probe (PHQBA) were found to be in a blue region with the coordinates of X = 0.145, Y = 0.258, further, the introduction of Th⁴⁺ the color coordinates of PHQBA were shifted towards the green region with the coordinates of X = 0.215, Y = 0.499. These results denote that PHQBA acts as a versatile sensor for the ratiometric detection of Th4+ ions. The limit of detection is an essential parameter for evaluating the performance of the chemosensor. The limit of detection was determined by the standard formula $3\sigma/\text{slope}^{50}$ where σ is the standard deviation of blank measurements of **PHQBA** (n = 10). The corresponding slope value was obtained from the concentration-dependent linear plot (Fig. S6†). The limit of detection was calculated to be 223 nM, which is comparatively lower than the previously reported thorium sensors (Table S1†). Thus, sensor PHQBA can

be used as a versatile sensitive tool for the quantitative ratiometric determination of Th⁴⁺ ions with outstanding sensitivity. Interestingly, the present work relies on evaluating the ratio of fluorescence intensities at two distinct wavelengths (515 and 478 nm). This self-calibration eliminates variations caused by external factors like light source intensity, detector sensitivity, and sample concentration, making the results more robust and reliable.

Interference, reaction kinetics, and pH effect of PHQBA on detecting Th⁴⁺

To explore the anti-interference capabilities of **PHOBA** (2 \times 10⁻⁵), interference studies were fluorometrically carried out based on the emission ratio of F_{515}/F_{478} nm. The effect of interference was investigated in the presence of various competing cations (1×10^{-4}) such as Ca^{2+} , Ba^{2+} , Al^{3+} , Cd^{2+} , Cu²⁺, Th⁴⁺, Co²⁺, Fe²⁺, Fe³⁺, Hg²⁺, Ni²⁺, Cr³⁺, Pb²⁺, Pd²⁺, Zn²⁺, La³⁺, and Yb³⁺. No significant interference was observed. Further, the addition of Th⁴⁺ ion to the **PHQBA** containing other analyte solutions exhibited enhanced emission intensity in the emission ratio of F_{515}/F_{478} nm, which designates that the sensor PHQBA's sensing capability was sufficient to distinguish Th⁴⁺ ions from other competing cations and supports the potential identification of Th4+ ions coexistence with other cations (Fig. 3a). Further, to explore the influence of pH on the detection properties of PHQBA, pH experiments were conducted in the range of 2-12. According to Fig. S11,† the emission of PHQBA is consistent across a broad pH range of 6-12 and is observed to be enhanced in an acidic medium spanning from 2-5. The protonation of the N atoms in quinoline and pyridine within the PHQBA system may cause increased emission intensity in the acidic medium. On the other hand, thorium(IV) nitrate has the potential to undergo hydrolysis and breakdown when subjected to higher alkaline conditions (11-12). When Th4+ ions were added to the PHQBA solution, the emission intensity began to increase in the pH range of 6-10 and decreased at pH 11-12. Therefore, the ideal pH range for Th⁴⁺ ion detection ranges between 6 and 10. The pH-responsive characteristic of the PHQBA sensor makes it perfect for detecting Th4+ under optimal physiological conditions. Rapid detection of analytes is one of the remarkable characteristics of an ideal chemosensor.51 For this, time-dependent emission studies were explored. According to Fig. 3b, the free sensor

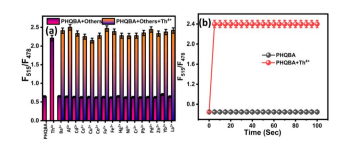


Fig. 3 (a) Interference of PHQBA (2 \times 10⁻⁵ M) in the presence of other cations (1 \times 10⁻⁴ M) and coexisting with Th⁴⁺; (b) time-dependent emission response of PHQBA with Th⁴⁺ ($\lambda_{\rm ex}=389$ nm).

Paper

PHQBA showed feeble emission in the ratio of F_{515}/F_{478} nm. Further, on the addition of Th⁴⁺ ion to the **PHQBA** solution, the emission intensity significantly elevated within five seconds. Hence, the sensor **PHQBA** was employed as an inevitable tool for the rapid detection of Th⁴⁺ ions.

Inspection of the detection mechanism

The sensor PHQBA encompasses a phenanthridine fluorophore, which is extensively conjugated with thiophene and quinoline moiety. Thus, it produces a blue emissive band. Since the Th⁴⁺ is electron deficient, it will have a binding interaction with electron-rich heteroatoms [N, N, S] in the PHQBA system. By the effect of synergic chelation, sensor PHQBA generates distinctive hypsochromic absorption and ratiometric green emission signals with Th4+ ions. Moreover, the binding stoichiometry of PHOBA with Th4+ ions was successively assessed by jobs plot experiments. As depicted in Fig. 4a, the maximum fluorescence intensity was attained when the molar fraction reached 0.5, revealing that the most likely binding mode was a 1:1 stoichiometric ratio for the PHQBA + Th⁴⁺ complex. To authenticate the stoichiometric binding of the sensor PHOBA with Th⁴⁺ ions, the Benesi-Hildebrand plot was linearly fitted between $[1/(F_0 - F)]$ and $1/[Th^{4+}]$. The association constant of the formed complex was calculated using the formula: $K_a =$ intercept/slope.52 The Ka value for PHQBA was determined as $3.77 \times 10^5 \text{ M}^{-1}$. Such a significant value denotes the robust binding of sensor PHQBA with Th⁴⁺ ions (Fig. 4b).⁵³ To further understand the coordination mode of PHQBA with Th4+, a series of ¹H NMR titration experiments were conducted in DMSO- d_6 . As shown in Fig. 4c, the base spectrum represents the free sensor. Upon the sequential addition of Th⁴⁺ ions (0-11 equivalents), significant changes were observed in the pyridyl proton of PHQBA (a) at 9.59 ppm, including a marked decrease in intensity. Furthermore, the incremental addition of Th⁴⁺ ions induced slight upfield shifts at 8.97 ppm (b), 7.06 ppm (d), and 6.97 ppm (e). A minor upfield shift was also noted for the c proton, which shifted from 8.29 ppm to 8.28 ppm. Notably, the aliphatic protons of PHQBA (g, h, and f) were significantly impacted, with their intensities progressively decreasing as the

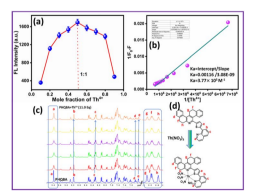


Fig. 4 (a) Jobs plot of PHQBA + Th^{4+} ; (b) Benesi–Hildebrand plot of PHQBA + Th^{4+} ; (c) series of ^{1}H NMR titration spectra of PHQBA with Th^{4+} ions (0–11 equiv.).

Th⁴⁺ concentration increased. These observations collectively indicate the effective chelation of Th^{4+} ions by **PHQBA**. Additionally, the mass of the complex was confirmed by high-resolution mass spectroscopy. According to the mass spectral data, the observed mass of the complex **PHQBA** + $\mathrm{Th}(\mathrm{NO_3})_4$ was 973.1636, which matched its calculated mass (973.1633) (Fig. S7†). To confirm nitrate binding, we have recorded FTIR spectra. As depicted in Fig. S13,† the observed stretching frequency at 1025 cm⁻¹ confirms the presence of $\mathrm{NO_3}^-$ in the **PHQBA** + Th^{4+} system.⁵⁴

DFT calculations

To gather clear insights into the theoretical mechanistic aspects of Th⁴⁺ binding, DFT calculations were performed by employing B3LYP/6-31G*(d,p) levels in Gaussian 16 software. 55-58 For the thorium atom, the SDD basis set was employed⁵⁹ as an effective core potential. The optimized electronic structures of PHQBA and its complex PHQBA + Th4+ are depicted in Fig. S10.† In the PHQBA + Th⁴⁺ complex, the receptor PHQBA utilizes N atoms of pyridine, quinoline, and sulfur of thiophene to enable efficient chelation with Th4+. The oscillatory strength, Z-matrix, calculated energy gap, and frontier molecular orbitals of PHQBA and its complex PHQBA + Th⁴⁺ are tabulated in Tables S2-S4.† The frontier molecular orbital images of PHOBA and its complex PHQBA + Th⁴⁺ are illustrated in Fig. 5. In the HOMO (highest occupied molecular orbitals) of PHQBA, the electron density predominately resided throughout the system, which denotes the higher electron density of the PHQBA ligand. In LUMO (lowest unoccupied molecular orbitals) the electron density was located at the phenanthridine skeleton, implying the adequate charge transfer effect in the PHQBA system. The calculated energy gap between the HOMO and LUMO orbitals of PHQBA was 3.3753 eV. Moreover, in the HOMO of the PHQBA + Th⁴⁺ complex, the phenanthridine thiophene moiety had the enriched electron density, whereas, in the LUMO orbitals, the electron density was predominately found in the quinoline moiety. Further, the calculated energy gap between the HOMO and LUMO of the PHQBA + Th4+ complex was found to be 3.723 eV, suggesting that the chelation of the Th⁴⁺ ion effectively enhances the energy gap of the Th⁴⁺ complex. The introduction of Th⁴⁺ ions into the detection medium has the potential to alter the local vicinity of PHQBA molecules. Such alterations can exert a significant influence on the electronic structure and

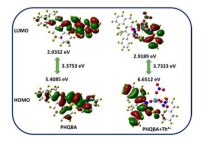


Fig. 5 The frontier molecular orbitals of PHQBA and PHQBA + ${\rm Th}^{4+}$ complex.

optical characteristics of the material. Perturbations in the environmental conditions can lead to shifts in the energy levels associated with electronic transitions, thereby inducing a red shift in emission. Concurrently, the overall band gap of the PHQBA + Th⁴⁺ complex may increase as a consequence of the perturbed electronic structure induced by the surrounding medium. Based on this ratiometric fluorescence sensing mechanism, the frontier molecular orbitals are mainly distributed at quinoline moiety in the Th⁴⁺ complex, while the excited electron of the nitrogen unit is back to the ground state resulting in ratiometric fluorescence. These findings denote that sensor PHQBA could synergically coordinate with the Th⁴⁺ atom to form a 1:1 stoichiometric complex with the sensor PHQBA.

Portable test strips aided detection

Portable test strip facilitated detection is a facile and intellectual approach for the visual discrimination of target analytes. ⁶² In our endeavor to assess the practical value of **PHQBA**, portable test strip experiments have been conducted in our laboratory. The selectivity of **PHQBA** (2×10^{-5}) in the presence of various ion solutions (1×10^{-4}) such as Ca^{2+} , Ba^{2+} , Al^{3+} , Cd^{2+} , Cu^{2+} , Co^{2+} , Fe^{2+} , Fe^{3+} , Hg^{2+} , Ni^{2+} , Cr^{3+} , Pb^{2+} , Pd^{2+} , Zn^{2+} , La^{3+} , and Yb^{3+} were tested appropriately. When seen under the illumination of UV-365 nm, **PHQBA** showed a remarkable selectivity to Th^{4+} ions over other cations. Only after the exposure of Th^{4+} ions, the blue emission of **PHQBA** altered to green, as shown in Fig. 6. However, other cation solutions did not show appreciable emission changes in the **PHQBA**-impregnated strips. Hence, it can be used as a portable sensor for the ratiometric discrimination of Th^{4+} ions.

Reproducible nature of PHQBA

Reversibility and reusability are characteristic features of a noble chemosensor. In this regard, the reproducible properties of **PHQBA** were explored in ACN/ H_2O (8:2, v/v). Sensor **PHQBA** demonstrated an intensive emission band at 478 nm, which produced a characteristic blue fluorescence. Then, the addition of Th⁴⁺ ions to the **PHQBA** solution resulted in ratiometric green fluorescence. Since ethylenediaminetetraacetic acid (EDTA) is a potential chelating agent, ⁶³ Th⁴⁺ ion-facilitated

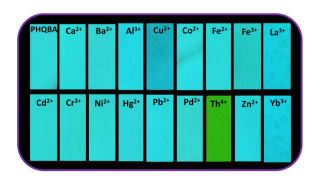


Fig. 6 PHQBA (2 \times $10^{-5})$ impregnated test strips for Th $^{4+}$ ion (1 \times $10^{-4})$ detection.

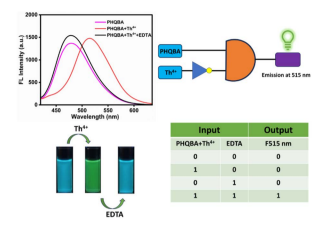


Fig. 7 Reversibility studies of **PHQBA** in the presence of Th⁴⁺ with EDTA and their logic gates representation.

ratiometric fluorescence was substantially reversed by the subsequent additions of EDTA. As depicted in Fig. 7, the emission changes were incredibly imitated in fluorescence spectra, in which the introduction of EDTA solution in the **PHQBA** + Th⁴⁺ complex, resulted in the regeneration of the probe's emission at 478 nm. These outcomes imply that the Th⁴⁺ ion has been significantly eradicated from the sensor and that it finely achieves the reversibility process induced by the efficient interaction of EDTA with the **PHQBA** + Th⁴⁺ complex. These reproducible and reversible processes (blue to green fluorescence) were repeated for 4 cycles (Fig. S9†) with successive alternative additions of Th⁴⁺ and EDTA. The outcomes revealed that the sensor **PHQBA** can be reused and also sensitively employed for the detection of Th⁴⁺ ions.

Real-time applications

The versatile sensing properties of **PHQBA** inspired us to explore its practical efficacy in various environmental water samples.⁶⁴ The known concentration of Th⁴⁺ ion solution was spiked with realistic water samples (drinking water, lake water, and tap water collected from Vellore). Then the experiment was repeated three times with each sample. The average recoveries and relative standard deviations (R. S. D.) of the corresponding spiked samples are tabulated in Table 1. As represented in Table 1, the average recoveries of various water samples ranged from 90.1 to 98.5%, and the corresponding relative standard deviation values were found to be 0.97 to 1.19%.

Table 1 Tabulation of real water sample analysis^a

Sample	Added (μM)	Found (µM)	Recovery (%)	RSD (%)
Lake water	0.5	0.46	92.2	1.01
	1.0	0.90	90.1	0.97
Tap water	0.5	0.47	94.3	0.99
	1.0	0.96	96.5	1.19
Drinking water	0.5	0.49	98.5	1.09
	1.0	0.97	97.6	1.11

^a Samples were collected in and around Vellore.

Based on these analytical findings, sensor **PHQBA** was effectively applied to determine Th⁴⁺ ion in real water samples with reliability and accuracy. Hence, the sensor **PHQBA** emerged as a valuable candidate for the real-time analysis of Th⁴⁺ ions in real water samples.

Conclusion

Paper

A new phenanthridine-derived quinoline-based fluorescent sensor was successfully developed for the ratiometric discrimination of Th⁴⁺ ions. On interaction with Th⁴⁺ ions, the emission of PHQBA ratiometrically changed from blue to green with the facilitation of chelation enhanced fluorescence process. The versatile sensing properties of PHQBA endorse hypsochromic absorption, and the redshifted emission bands during the interaction with Th4+ were thoroughly investigated by absorption and emission experiments. The binding interaction of Th⁴⁺ ions with PHQBA was explored by 1H NMR, HRMS, and DFT analysis. The portable test strips confirmed that the sensor PHQBA can selectively detect Th4+ ions without the aid of any sophisticated instrumental facility. Also, the sensor PHQBA was analyzed for the real-time detection of Th4+ ions in real water samples. The versatile features of PHQBA such as reversibility/ reusability, selectivity, and practical utility will accelerate the advancement of fluorescent sensors in adaptable analytical applications. Developing ultrasensitive chemosensors for the detection of radioactive ions is a critical research area with extensive implications in environmental monitoring, nuclear safety, and non-proliferation initiatives. Systematic monitoring of thorium ions is particularly essential to ensure compliance with nuclear regulations and to safeguard public health and safety. Advanced tools capable of detecting ultra-low concentrations of thorium in soil, water, and air are crucial for identifying contamination and assessing its environmental impact. In our laboratory, ongoing research focuses on innovative structural modifications to develop highly sensitive detection methods capable of identifying radioactive elements at extremely low concentrations. These advanced analytical techniques aim to significantly enhance the accuracy, sensitivity, and efficiency of analyte detection, even in complex and challenging environments, thereby paving the way for more effective and reliable monitoring solutions in the future.

Data availability

The data supporting this article have been included as part of the ESI.†

Conflicts of interest

The authors declare no financial interests.

Acknowledgements

Selin Manoj Kumar sincerely thanks Vellore Institute of Technology for providing a Teaching Cum Research Assistant (Project No.: VIT/HR/2022/19062). The DST-FIST NMR facility at

VIT University is duly acknowledged. Finally, the authors thank Dr R. Srinivasan, SSL-VIT, for language editing.

References

- 1 M. Cuney, Non-Renewable Resource Issues: Geoscientific and Societal Challenges, 2012, pp. 91–129.
- 2 M. Corn, *Handbook of Hazardous Materials*, Academic Press, 2012.
- 3 C. S. K. Raju and M. Subramanian, *J. Hazard. Mater.*, 2007, **145**, 315–322.
- 4 S. Yang, N. Tan, X. Yan, F. Chen, W. Long and Y. Lin, *Mar. Pollut. Bull.*, 2013, **74**, 213–219.
- 5 S. Yang, N. Tan, X. Yan, F. Chen, W. Long and Y. Lin, *Mar. Pollut. Bull.*, 2013, 74, 213–219.
- 6 D. Brugge, J. L. deLemos and B. Oldmixon, *Rev. Environ. Health*, 2005, 20, 177–194.
- 7 E. Zambrzycka-Szelewa and B. Godlewska-Zylkiewicz, *Spectrochim. Acta, Part B*, 2024, **213**, 106859.
- 8 M. Ioannidou, G. Zachariadis, A. Anthemidis and J. Stratis, *Talanta*, 2005, **65**, 92–97.
- 9 A. M. Clark, A. D. Nelson, T. L. Bailey, D. Blankstein, C. Boomershine, G. M. Brown, P. C. Burns, S. Carmichael, L. K. Callahan and J. Koros, *Nucl. Instrum. Methods Phys. Res.*, B, 2024, 548, 165253.
- 10 X. Qiu, Z. Hu, T. He, T. Luo, W. Zhang, M. Li, K. Zong, Z. Wang and Y. Liu, J. Anal. At. Spectrom., 2024, 39, 545–557.
- 11 J. Sardans, F. Montes and J. Peñuelas, *Spectrochim. Acta, Part B*, 2010, **65**, 97–112.
- 12 M. Ghosh, M. Sarma, S. Dagupta, J. Datta and K. Swain, *Anal. Chem. Lett.*, 2024, **14**, 382–394.
- 13 K. P. Carter, A. M. Young and A. E. Palmer, *Chem. Rev.*, 2014, **114**, 4564–4601.
- 14 A. P. De Silva, H. N. Gunaratne, T. Gunnlaugsson, A. J. Huxley, C. P. McCoy, J. T. Rademacher and T. E. Rice, *Chem. Rev.*, 1997, 97, 1515–1566.
- 15 M. Dutta and D. Das, *TrAC, Trends Anal. Chem.*, 2012, 32, 113–132.
- 16 T. Rasheed, M. Bilal, F. Nabeel, H. M. Iqbal, C. Li and Y. Zhou, *Sci. Total Environ.*, 2018, **615**, 476–485.
- 17 D. Jothi, S. Munusamy, S. Enbanathan and S. K. Iyer, *RSC Adv.*, 2022, **12**, 8570–8577.
- 18 S. M. Kumar, D. Jothi, S. Munusamy, S. Enbanathan and S. K. Iyer, *J. Photochem. Photobiol. Chem.*, 2023, 434, 114269.
- 19 Z. Aydin, J. Turk. Chem. Soc., Sect. A, 2020, 7, 277-286.
- 20 X. M. Wang, H. Yan, Y. Chen and H. B. Bao, *Adv. Mater. Res.*, 2011, 239, 1105–1108.
- 21 D. Mohanasundaram, R. Bhaskar, G. G. V. Kumar, J. Rajesh and G. Rajagopal, *Microchem. J.*, 2021, **164**, 106030.
- 22 H.-I. Un, C.-B. Huang, C. Huang, T. Jia, X.-L. Zhao, C.-H. Wang, L. Xu and H.-B. Yang, *Org. Chem. Front.*, 2014, 1, 1083–1090.
- 23 L. Gao, C. Deng, J. Xiong, P. Zhu, Q. Chen and K. Tan, *Microchem. J.*, 2019, **150**, 104096.
- 24 Z. Zhang, J. Feng, P. Huang, S. Li and F.-Y. Wu, Sens. Actuators, B, 2019, 298, 126891.

- 25 G. Lu, Z. Jia, M. Yu, M. Zhang and C. Xu, *Molecules*, 2023, 28, 7818.
- 26 M. Royzen, Z. Dai and J. W. Canary, *J. Am. Chem. Soc.*, 2005, 127, 1612–1613.
- 27 H. Wang, H. Xing, W. Liu, Y. Hao, L. Zhang, Z. Yang, Q. Hu, S. Shuang, C. Dong and X. Gong, *Sens. Actuators*, B, 2022, 352, 130991.
- 28 N. Tohora, S. Ahamed, M. Mahato, J. Chourasia, S. Ali and S. K. Das, *J. Photochem. Photobiol. Chem.*, 2024, 457, 115921.
- 29 Y. Li, L. Li, X. Pu, G. Ma, E. Wang, J. Kong, Z. Liu and Y. Liu, *Bioorg. Med. Chem. Lett.*, 2012, 22, 4014–4017.
- 30 M. Gao, P. Xie, L. Wang, X. Miao and F. Guo, Res. Chem. Intermed., 2015, 41, 9673-9685.
- 31 Y. Chen, F. Li and Z. Bo, *Macromolecules*, 2010, **43**, 1349–1355
- 32 S. Yashmin, S. Mondal, R. Das, P. Banerjee and A. T. Khan, *Org. Biomol. Chem.*, 2022, **20**, 7302–7315.
- 33 D. Wu, B. Fang, M. Zhang, W. Du, J. Zhang, X. Tian, Q. Zhang, H. Zhou, J. Wu and Y. Tian, *Dyes Pigm.*, 2018, 159, 142–150.
- 34 S. Karthik, J. Ajantha, S. Easwaramoorthi and T. Gandhi, *New J. Chem.*, 2020, 44, 9530–9539.
- 35 R. M. Gadirov, L. G. Samsonova, K. M. Degtyarenko, A. E. Kurtsevich, I. K. Yakushchenko and T. N. Kopylova, *J. Fluoresc.*, 2021, 31, 1333–1342.
- 36 S. Swaminathan, S. Munusamy, D. Jothi and S. K. Iyer, *ChemistrySelect*, 2021, **6**, 858–864.
- 37 L. Liu, L. Ga and J. Ai, *Biosens. Bioelectron.*, 2022, 213, 114456.
- 38 S. M. Kumar, S. Munusamy, S. Manickam, D. Jothi, S. Enbanathan and S. K. Iyer, J. Mol. Liq., 2023, 381, 121828.
- 39 X. Pei, Y. Pan, L. Zhang and Y. Lv, *Appl. Spectrosc. Rev.*, 2021, **56**, 324–345.
- 40 S. Manoj Kumar and S. Kulathu Iyer, *J. Org. Chem.*, 2024, **89**, 5392–5400.
- 41 S. Enbanathan, S. Munusamy, D. Jothi, S. Manojkumar, S. Manickam and S. K. Iyer, *RSC Adv.*, 2022, **12**, 27839–27845.
- 42 S. Seenan and K. I. Sathiyanarayanan, *Inorg. Chem. Commun.*, 2021, 132, 108825.
- 43 S. Sawminathan and S. K. Iyer, *Spectrochim. Acta, Part A*, 2022, **265**, 120403.
- 44 D. Jothi, S. Manickam, S. Sawminathan, S. Munusamy, S. A. Kumar and S. K. Iyer, *Dyes Pigm.*, 2022, **197**, 109826.

- 45 W. Li, Y. Fu, T. Liu, H. Li and M. Huang, *Spectrochim. Acta, Part A*, 2023, **288**, 122147.
- 46 N. N. Zhadin and R. R. Alfano, *J. Biomed. Opt.*, 1998, 3, 171–186.
- 47 S. Srinivas and R. Mutharasan, *Biotechnol. Bioeng.*, 1987, 30, 769–774.
- 48 A. Elabd, RSC Adv., 2016, 6, 45525-45532.
- 49 R. Yao, Z. Li, G. Liu, C. Fan and S. Pu, *Talanta*, 2021, 234, 122612.
- 50 S. M. Kumar, D. Jothi, S. Munusamy, S. Enbanathan and S. K. Iyer, *J. Photochem. Photobiol. Chem.*, 2023, 434, 114269.
- 51 N. Duan, S. Yang, H. Tian and B. Sun, *Food Chem.*, 2021, 358, 129839.
- 52 S. M. Kumar, S. Munusamy, S. Manickam, D. Jothi, S. Enbanathan and S. K. Iyer, *J. Mol. Liq.*, 2023, 381, 121828.
- 53 R. Wang and Z. Yu, *Acta Phys.-Chim. Sin.*, 2007, 23, 1353–1359.
- 54 A. Knyazev, G. Fukin, R. Rumyantcev, M. Komshina, I. Savushkin and A. Paranyuk, *Polyhedron*, 2016, 117, 600– 603.
- 55 B. Pavankumar, P. Ranjan, R. R. Panicker, P. C. Jha, C. Brahmananda Rao, R. Desikan and A. Sivaramakrishna, *ChemistrySelect*, 2024, **9**, e202304509.
- 56 J. E. Del Bene, W. B. Person and K. Szczepaniak, *J. Phys. Chem.*, 1995, **99**, 10705–10707.
- 57 H. Kruse, L. Goerigk and S. Grimme, J. Org. Chem., 2012, 77, 10824–10834.
- 58 M. P. Andersson and P. Uvdal, *J. Phys. Chem.*, 2005, **109**, 2937–2941.
- 59 R. Sk, S. A. Kumar, K. Vijayakrishna, A. Sivaramakrishna, C. V. S. B. Rao, N. Sivaraman and S. K. Sahoo, *Inorg. Chem.*, 2018, 57, 15270–15279.
- 60 S. Seenan, S. Manickam, S. Sawminathan, D. Jothi and S. K. Iyer, *J. Photochem. Photobiol. Chem.*, 2022, **430**, 113952.
- 61 S. Sawminathan and S. K. Iyer, *New J. Chem.*, 2021, **45**, 6033–6041.
- 62 Y. Shen, Y. Wei, C. Zhu, J. Cao and D.-M. Han, Coord. Chem. Rev., 2022, 458, 214442.
- 63 S. M. Kumar, S. Munusamy, D. Jothi, S. Enbanathan and S. K. I. Iyer, *J. Mol. Liq.*, 2023, 373, 121235.
- 64 S. M. Kumar, S. Munusamy, D. Jothi, S. Enbanathan, J. Haribabu and S. K. Iyer, *Opt. Mater.*, 2023, **144**, 114382.