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1. Introduction

The fluorescence mechanism of rare earth compounds closely resembles atomic emission.¹⁻³ These compounds exhibit many advantages in fluorescence, including narrow emission bandwidths, long fluorescence lifetimes, and a wide range of colors. Due to their unique luminescent properties, rare earth compound-based luminescent materials have found extensive applications in fields such as luminescent devices,⁴⁻⁷ chemical sensing, $8-10$ biological imaging, $11-13$ commodity anticounterfeiting¹⁴⁻¹⁶ over the past few decades. However, due to the parity forbidden of $f \rightarrow f$ transition, the direct absorption intensity of rare earth ions is weak, which leads to the application bottleneck associated with the low fluorescence quantum efficiency of many rare earth compounds.17,18 To address this issue, the incorporation of specific organic ligands for coordinating with rare earth ions represents an efficacious strategy.¹⁹⁻²¹ The presence of these ligands can enhance the light absorption coefficient of rare earth compounds in the ultraviolet range, thereby effectively exciting the rare earth ions through a non-radiative energy transfer process and significantly boosting their fluorescence performance. The phenomenon is commonly referred to as the "antenna effect". The

Synthesis and fluorescence properties of europium complex functionalized fiberglass paper†

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The development of novel rare earth fluorescent materials and the exploration of their applications have consistently been focal points of research in the fields of materials science and chemistry. In this work, a novel rare earth composite material with good photo-fluorescence properties and self-supporting has been prepared via a simple ultrasonic solvent reaction method. Initially, the Phen moieties is immobilized onto the surface of a self-supporting fiberglass paper using ICPTES, followed by the coordination of Eu(TTA)₃ moieties with Phen moieties through a convenient ultrasonic solvent reaction. The resulting GF-Phen-Eu(TTA)₃ has been characterized using FTIR, UV-Vis DRS, fluorescence measurements, and so on. The results indicate that the composite material exhibits strong fluorescent emission and presents a vivid red color under ultraviolet light. Further research has shown that the fluorescence of GF–Phen– Eu(TTA)₃ strips demonstrated a pronounced quenching effect in response to some transition metal ions (1 mM). Hence, the rare earth composite materials presented here can be utilized not only for the production of optical materials, but also for the development of fluorescence sensing strips. PAPER
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ligands commonly employed for sensitizing the luminescence of rare earth ions include derivatives of carboxylic acids, $22-24$ β diketones,^{25–27} N-heterocyclic compounds,^{28–30} etc. The incorporation of these organic ligands not only significantly enhances the fluorescence quantum efficiency, fluorescence emission intensity, and prolongs the fluorescence lifetime but also facilitates the covalent assembly of rare earth complexes onto diverse matrix materials through ligand modification, thereby achieving the construction of multi-component luminescent materials based on rare earth elements.

In recent decades, there has been a significant surge in attention towards this multi-component composite, which is based on the covalent assembly of rare earth complexes.³¹–³⁶ The commonly employed matrices for fabricating rare earth multicomponent hybrid materials encompass silicon oxide,³⁷⁻³⁹ carbon, $40-42$ metal oxide, $43-45$ polymer, $46-48$ and zeolite $49-51$ materials. The incorporation of these matrices signicantly enhances the stability of rare earth complexes and broadens the application fields of rare earth-based luminescent materials. However, with the exception of polymer-based rare earth multicomponent hybrid materials that can be directly shaped, most other types of rare earth multi-component hybrid materials exist in powdered form and cannot undergo direct processing for molding. Therefore, researchers have redirected their focus towards the development of rare earth multi-component hybrid materials based on substrates possessing inherent selfsupporting properties such as polymer matrix,⁵² glass,⁵³ fiber, 54 etc. For instance, the successful integration of the Europium complex onto ITO Glass by Santos et al.⁵⁵ resulted in

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the fabrication of remarkably efficient Organic Light-Emitting Diodes (OLEDs). Shang et al ⁵⁶ successfully achieved the grafting of europium complexes onto carbon fiber, resulting in the development of a composite material exhibiting strong red fluorescence. Tong et al ⁵⁷ fabricated a lanthanide Metal-Organic Frameworks functionalized carbon cloth by in situ growth method, followed by modification of poly vinyl terephthalic acid through in situ polymerization, which resulted in the production of a novel fluorescent composite thin film with exceptional sensitivity towards UO_2^{2+} ions. Currently, the fabrication of rare earth fluorescent composites using selfsupporting matrices and their subsequent applications in sensing present a significant challenge.

The glass fiber paper is an environmentally friendly, highly stable, cost-effective, and easily modifiable material that serves as a well self-supporting matrix for fabricating composites.^{58,59} Herein, a very simple strategy was proposed to prepare the highly photo-fluorescent rare earth composite by directly surface-modifying glass fiber filter paper. We successfully immobilized a europium complex onto the surface of a glass fiber filter paper using a molecular bridge, resulting in the formation of a composite luminescent material. Finally, the resulting material has been thoroughly investigated in terms of their photophysical properties and their potential application for metal ion sensing.

2. Experimental details

2.1 Materials and methods

The $1,10$ -phenanthrolin-5-amine (Phen-NH₂, 97%), isocyanatopropyltriethoxysilane (ICPTES, 95%), 2-thenoyltri fluoroacetone (TTA, 98%) and $EuCl₃·6H₂O$ (99.99% metals basis) were purchased from Shanghai Aladdin Biochemical Technology Co., Ltd. The glass fiber (GF) filter was purchased from Weifang Kaisheng Electronic Technology Co., Ltd. All other reagents and solvents were obtained from Sinopharm Chemical Reagent Co., Ltd. The Fourier transform infrared spectroscopy (FTIR) were determined Agilent Cary 630. The fluorescence excitation and emission data were tested with SHIMADZU RF-6000 fluorescence spectrophotometer. NMR spectra were recorded on AVANCE NEO 400MHZ. Elemental analysis were tested with Elementar UNICUBE. The ultravioletvisible diffuse reflection spectrum (UV-Vis DRS) were recorded by Shimadzu UV-3600Plus spectrophotometer. The powder Xray diffraction patterns (XRD) were acquired on Rigaku SmartLab SE equipped with Cu anode. The fluorescence lifetime was measured on an Edinburgh Instruments FLS 920 fluorescence spectrometer.

2.2 Synthesis of europium complex

The lanthanide complexes were synthesized by typical methods. For Eu(TTA)₃ \cdot 2H₂O, typically: At room temperature, 6 mmol TTA was dissolved in ethanol, followed by 3 mL 1 mol per L NaOH aqueous solution for deprotonation; And then an ethanol solution containing 2 mmol $EuCl_3 \cdot 6H_2O$ was slowly introduced into the solution. The resulting mixture was stirred for 30 min

until it is slowly transformed into the $Eu(TTA)₃·2H₂O$ solution. The solution was not further processed and was directly reserved for further use. For elemental analysis and ¹H NMR spectroscopy, the solution was initially concentrated to induce the precipitation of $Eu(TTA)_3 \tcdot 2H_2O$. The resulting precipitate was subsequently washed with water and petroleum ether, dried, yielding a crude product that was characterized as Eu(TTA)₃ · 2H₂O. ¹H NMR (CDCl₃, 400 MHz, δ ppm): 7.07 (s, 3H, H of thiophene moiety), 6.51 (s, 3H, H of moiety), 6.07 (s, 3H, H of moiety), 0.88 (s, 3H, H of β -diketonate). Elemental analysis was performed for C, H, S, atoms. Calc. for $\text{EuC}_{24}H_{16}F_9O_8S_3$: C, 33.85; H, 1.89; S, 11.30. Found: C, 33.43; H, 2.34; S, 11.32.

2.3 Synthesis of europium complex functionalized glass fiber

First, the cut glass fiber was soaked in 25% NaOH solution and activated for about 30 min, then washed with deionized water and dried at 60 °C, denoted GF. Subsequently, the as-prepared GF was immersed in an ethanol solution containing 1 mmol Phen-NH₂ and 7 mmol ICPTES. The reaction system was then transferred to an ultrasonic cleaning machine (40 kHz, 50 W) for ultrasonic treatment for 60 min. After treatment, the glass fiber was took out and washed with anhydrous ethanol, and dried at 60 \degree C to obtain the Phen-loaded glass fiber, which is referred to as GF–Phen. The GF–Phen was immersed in the asprepared $Eu(TTA)₃·2H₂O$ solution and treated in the ultrasonic cleaning machine again for 60 min. Finally, the europium complex functionalized glass fiber was took out and washed with anhydrous ethanol, and dried at 60 °C to obtain the GF– Phen-Eu(TTA)₃ (Scheme 1). RSC Advances

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2.4 Sensing experiments

The GF-Phen-Eu(TTA) $_3$ was cut into strips and immersed in a series of solutions containing different metal ions for 10 min under ultrasonic treatment. Subsequently, the resulting strip

Scheme 1 Synthesis strategy of the europium complex functionalized glass fiber and the photos of the resultant product under daylight and 365 nm UV light.

3. Results and discussion

The main composition of glass fiber and the other glass is essentially identical, consisting primarily of a mixture of various silicate compounds. Upon activation, the surface of glass fiber becomes enriched with numerous active silicon hydroxyl groups (Si–OH), thereby endowing it with exceptional potential for surface modification and rendering it highly suitable as a matrix material for preparing composite materials. On the other hand, the bridge-linking method is widely recognized as the primary approach for fabricating hybrid materials with covalent bonds, and it has been extensively employed in the synthesis of rare earth composite luminescent materials. Based on these findings, a self-supporting glass fiber filter paper was chosen as the substrate. Its surface was activated with NaOH aqueous solution and subsequently modified with Phen-NH₂ and ICPTES in combination to establish a molecular bridge, the obtained materials are marked as GF–Phen. The superior chelating properties of the Phen moiety enable GF–Phen to effectively adsorb unoccupied rare earth complexes on its surface. Ultimately, the rare earth europium complex was immobilized onto the surface of glass fiber through the immersion of GF–Phen in Eu(TTA)₃ \cdot 2H₂O solution followed by ultrasonic treatment, resulting an innovative composite luminescent material (GF–Phen–Eu(TTA)₃). The synthesis strategy is shown in Fig. 1. As illustrated, the obtained GF-Phen-Eu(TTA)₃ exhibits negligible coloration under daylight; however, it manifests a vivid red color upon irradiation with 365 nm UV light. This transformation can be attributed to the characteristic emission of the europium complex when excited by UV light. It can also be regarded as evidence of the successful immobilization of rare earth complexes onto glass fiber filter paper. At the same time, in order to investigate the impact of surface modification on the microstructure of glass fiber paper, the Puper

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XRD tests on the glass fiber paper before and after modification were conducted (Fig. S1, ESI†). The results indicate that the surface complexation did not alter the microstructure of the glass fiber paper. Whether before or after modification, the material exhibits a broad peak centered around approximately 25°, indicative of the characteristic diffraction pattern of the amorphous silicon framework material. Furthermore, there are no discernible crystalline regions in the XRD patterns, suggesting that the europium complex introduced during material modification should be at a mesoscale level. Therefore, no relevant crystallographic data appeared in the XRD pattern (Fig. S1, ESI†).

To further investigate the assembly of europium complexes on the surface of glass fiber, the Fourier transform infrared spectroscopy (FTIR) was employed to characterize the relevant materials. As shown in Fig. 1, the spectrum of GF is characterized by a simple profile and exhibits typical features of silicate infrared spectra; it shows a prominent absorption at 1060 cm^{-1} , which derives from the asymmetric stretching vibration of Si– O–Si moieties and consistents with the spectra of GF–Phen and $GF-Phen-Eu(TTA)₃$. The spectrum of $GF-Phen$ closely resembles that of GF, with a more pronounced absorption peak at 1609 cm⁻¹, attributed to the stretching vibration of C=O and bending vibration of N–H in amide group. For the GF–Phen– Eu(TTA)₃, the peak observed around 1609 cm⁻¹ exhibits a notable increase in intensity. This augmentation is primarily ascribed to the stretching vibration of C=O in β -diketone moieties, which shifts to lower wavenumber after coordinating with europium(m) ions. The peak at 1544 cm⁻¹ of GF-Phen- $Eu(TTA)$ ₃ also arises from the downward shift stretching vibration of $C=C$ within the ring-shaped β -diketone coordination structure. Furthermore, the peaks observed at 1308 cm^{-1} and 1356 cm^{-1} are indicative of the symmetric and asymmetric stretching vibrations of the CF_3 group, while the peak at 715 cm^{-1} can be attributed to the stretching vibrations of the C–S bond. The presence of these absorption peaks in spectrum of GF–Phen–Eu(TTA)₃ provides compelling evidence that the Eu complex has been successfully deposited onto the surface of the GF.

As a down-conversion phosphor material, the absorption capacity of GF–Phen–Eu(TTA) $_3$ for ultraviolet-visible light plays a crucial role in its fluorescent properties. The ultraviolet-visible diffuse reflection spectra (UV-Vis DRS) of the materials are depicted in Fig. 2. As shown in figure, the GF exhibits the characteristic absorption spectrum of glass materials in response to ultraviolet light. After chemical coupling treatment with Phen, the glass fiber demonstrated enhanced light absorption in the 320–420 nm, albeit with a subtle effect. For europium complexes-loaded glass fibers, a prominent absorption band in the 305–410 nm range is observed for GF–Phen– $Eu(TTA)₃$, attributed to the exceptional absorption capability of TTA. According to this figure, GF–Phen–Eu(TTA)₃ demonstrates the highest absorption capacity in the ultraviolet region at approximately 350 nm which closely aligns with the maximum fluorescence excitation wavelength of $GF-Phen-Eu(TTA)₃$ shown in Fig. 3. This result indicates that GF-Phen-Eu(TTA)₃ Fig. 1 FTIR spectra of GF, GF–Phen and GF–Phen–Eu(TTA)₃. can absorb abundant energy in the ultraviolet range and then

UV-Vis DRS spectra of GF, GF–Phen and GF–Phen–Eu(TTA)3.

Fig. 3 Fluorescence excitation and emission spectra of GF–Phen– $Eu(TTA)_{3}$

transfer it through the "antenna effect" to sensitize the fluorescence emmission properties of the europium (m) ions loaded on glass fiber. Therefore, we can expect that GF-Phen-Eu(TTA)₃ will have excellent photoluminescence performance under ultraviolet light, which has been confirmed by their corresponding fluorescence emission spectra (Fig. 3) and their photo under ultraviolet light (Fig. 1).

As previously mentioned, the europium complexfunctionalized glass fiber exhibits a vivid red fluorescence when exposed to 365 nm UV light, attributed to the characteristic emission of europium (m) ions. In order to further investigate its fluorescence properties, Fig. 3 illustrates the fluorescence excitation and emission spectra of GF-Phen- $Eu(TTA)₃$. By monitoring the strongest emission peak of euro $pium(m)$ ions at 613 nm, the fluorescence excitation spectrum can be obtained. The spectrum reveals that the peak of

fluorescence excitation spectrum is situated at about 332 nm and exhibits a broad band-like structure. As previously discussed, the excitation band closely aligns with the UV absorption range of the β -diketone (TTA). Based on this, we selected the 332 nm as the excitation wavelength and performed the fluorescence emission measurement of the GF–Phen–Eu(TTA)₃. As shown on the right side of Fig. 3, the fluorescence emission spectrum of GF–Phen–Eu(TTA)₃ displays the characteristic narrow band emission associated with rare earth elements. There are four distinct emission peaks can be observed from le to right, with their centers located at 592, 613, 652, and 703 nm, which can be aligned to the transitions of europium (m) ions from ${}^5D_0 \rightarrow {}^7F_1$, ${}^5D_0 \rightarrow {}^7F_2$, ${}^5D_0 \rightarrow {}^7F_3$ and ${}^5D_0 \rightarrow {}^7F_4$ transitions, respectively. It is evident that the overwhelming majority of these emissions lie within the red light spectrum, thereby accounting for its appearance as red under a UV light. It is also evident that the narrowband emission peak corresponding to ${}^{5}D_{0}$ \rightarrow ${}^{7}F_{2}$ transition overwhelmingly dominates, resulting in its exceptionally high purity in the red color. Furthermore, it is widely recognized that the electric dipole transition ${}^5\text{D}_0 \rightarrow {}^7\text{F}_2$ of europium (m) ions demonstrates a high sensitivity to local spatial symmetry, while the magnetic dipole transition ${}^{5}D_{0} \rightarrow {}^{7}F_{0}$ of europium(m) ions exhibits relatively low sensitivity to ${}^{7}F_{1}$ of europium(III) ions exhibits relatively low sensitivity to local spatial symmetry. As a result, the ratio (R) of emission intensities from the ${}^{5}D_0 \rightarrow {}^{7}F_2$ and ${}^{5}D_0 \rightarrow {}^{7}F_1$ transitions of europium (m) ions $(i.e. I⁽⁵D₀ \rightarrow {}^{7}F₂)/I⁽⁵D₀ \rightarrow {}^{7}F₁))$ can be seen as a crucial indicator for assessing the radial symmetry around europium(m) ions. Here, the R value of GF–Phen–Eu(TTA)₃ is as high as 18.1, indicating that europium (m) ions is situated within an environment of low symmetry, which is in line with the hypothesis that europium (m) ions occupy the coordination center of [Eu(TTA)₃Phen] moieties. PSC Advances Article 2023. This article is a september 2024. The common article is a september 2024. The common article is licensed under a creative Creative Commons Article is a september 2024. At a september 2024. The c

The fluorescence decay curve of GF–Phen–Eu(TTA)₃ was measured using an excitation wavelength of 332 nm at room temperature (Fig. S2, ESI†). The decay data can be well fitted to a double exponential function, and the decay time was calculated to be about 4.7 µs. Moreover, the $^5\mathrm{D}_0$ quantum efficiency of the GF–Phen–Eu(TTA)₃ was estimated based on the emission spectra and lifetime (τ) , in accordance with the Judd-Ofelt theory (further details are listed in ESI†). The fluorescent quantum efficiency of GF-Phen-Eu(TTA) $_3$ was calculated to be about 0.5%. The relatively low value may be attributed to the limited capacity of the rapid assembly method to accommodate a substantial quantity of europium complex on the glass fiber, thereby resulting in a reduced theoretical quantum efficiency. In short, these optical properties primarily stem from the unique coordination structure of $[Eu(TTA)_{3}(Phen)]$ and its relatively low loading content.⁶⁰⁻⁶²

As a self-supporting material, one of the most important advantages of glass fiber is that it can be freely cut into a certain shape. In order to further expand the applications of europium complex-loaded glass fiber, the GF–Phen–Eu(TTA)₃ was cutted into strips and tested its potential performance in metal ion sensing. The fluorescence response of GF-Phen-Eu(TTA) $_3$ to a range of common metal ions $(L⁺, Na⁺, K⁺, Ca²⁺, Mg²⁺, Zn²⁺,$ Cu^{2+} , Fe³⁺ and Co²⁺) at a concentration of 1 mM was assessed. As shown in Fig. 4, the fluorescence intensity of GF-Phen-

 $Eu(TTA)$ ₃ did not show significant change after immersed in solutions containing Li $^+$, Na $^+$, K $^+$, Ca $^{2+}$ or Mg $^{2+}$; meanwhile, the corresponding strips exhibited negligible color changes when examined under a 365 nm lamp following immersion. However, upon immersion in the solution containing Zn^{2+} , Cu^{2+} , Fe^{3+} and $Co²⁺$ ions, the fluorescence intensity of the strips exhibited a significant decrease. This phenomenon is also observable through their fluorescent expression when exposed to a 365 nm light. As shown in the figure, the fluorescence of the strip immersed in solutions containing Cu^{2+} , Fe³⁺ and Co^{2+} was significantly quenched. Only the strip immersed in the Zn^{2+} solution exhibited residual weak fluorescence. It is evident that the fluorescence quenching of GF–Phen–Eu(TTA)₃ by Cu²⁺, Fe³⁺ and $Co²⁺$ can be attributed to their strong coordination ability, enabling them to effectively compete with $Eu³⁺$ for chelating with the Phen moieties. These ions have the capability to displace the europium complex from the surface of glass fiber, consequently leading to the loss of characteristic fluorescence of europium(III) ions for those soaked strips. While for Zn^{2+} ions or other ions that belongs to main group, owing to limited coordination ability, they are unable to effectively displace the europium (m) ions on the paper. Consequently, complete quenching of their fluorescence is unattainable, or their impact on the fluorescence performance of the strip is minimal. These results indicate that the GF-Phen-Eu(TTA)₃ exhibits specific selectivity and high sensitivity to some transition metal ions, rendering it a promising metal ion sensor of broad-spectrum. Paper

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Regrettably, upon further reduction of the solution concentration, for instance by decreasing the metal ion concentration to 100 µmol, only few transition metal ions, such as Cu^{2+} and $Fe³⁺$, can effectively quench the fluorescence of the test paper. However, in addition to the cost-effective glass fiber paper and ethanol, the other primary reagents utilized in this study are relatively economical (Table S2, ESI†), and the experimental procedures are straightforward with a high success rate. The

Fig. 4 Photographs of the GF-Phen-Eu(TTA)₃ strips under 365 nm UV lamp after treated with different metal ion solutions for 10 min, and comparison of their corresponding fluorescence intensity.

production cost for creating the fluorescent test paper described in this study is minimal. Furthermore, the preparation process for this test paper does not necessitate stringent requirements; it can even be conducted in chemistry laboratories at primary and secondary schools. As long as standard safety goggles and gloves are worn during experimentation, it can be safely performed. The resulted GF-Phen-Eu(TTA)₃ displays certain stability. A vivid red fluorescence remained detectable under 365 nm UV light even after prolonged storage in a standard ziplock bag within the laboratory for a year. Despite the decrease in fluorescence intensity, it remained suitable for transition metal ion sensing tests. To evaluate the detection capability of the fluorescent paper in real water samples, we collected multiple water samples from a local river and observed that the natural water samples did not quench the fluorescence of our paper. However, introduction of ions such as Cu^{2+} and Fe³⁺ resulted in quenching, indicating potential practical applications for GF–Phen–Eu(TTA)₃. Thus, the sensing performance of GF–Phen–Eu(TTA)₃ towards transition metal ions demonstrates remarkable environmental stability and resistance to interference.

4. Conclusions

In this paper, a novel rare earth composite material was synthesized through a simple ultrasonic solvent reaction method using glass fiber as the matrix. The prepared GF-Phen- $Eu(TTA)₃$ have outstanding fluorescent performance and exhibits intense red fluorescence when exposed to 365 nm light. The developed composite material also exhibit good photostability, mechanical properties, suggesting that the GF–Phen– Eu(TTA)₃ are a very promising fluorescent materials. Moreover, the inherent self-supporting nature of glass fiber enables GF-Phen–Eu(TTA)₃ to be tailored into diverse structures. Here, GF– Phen–Eu(TTA)₃ has been fabricated into strip format and observed its specific sensing capability towards some transition metal ions, offering a novel avenue for the development of test strip that can be used for the detecting of transition metal ion.

Data availability

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

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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