

Cite this: *Chem. Sci.*, 2024, 15, 17642

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A photoinduced electron-transfer strategy for switchable fluorescence and phosphorescence in lanthanide-based coordination polymers†

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Smart optical materials with tunable fluorescence and room temperature phosphorescence (RTP) exhibit promising application prospects in the field of intelligent switches, information security, *etc.* Herein, a tetraimidazole derivative was grafted to one-dimensional lanthanum-diphosphonate through H-bonds, generating a coordination polymer (CP), $(\text{H}_4\text{-TIBP})\cdot[\text{La}_2\text{Li}(\text{H}_2\text{-HEDP})_4(\text{H-HEDP})]\cdot 3\text{H}_2\text{O}$ (termed **La**; TIBP = 3,3,5,5-tetra(imidazole-1-yl)-1,1-biphenyl; $\text{H}_4\text{-HEDP}$ = 1-hydroxyethylidene-1,1-diphosphonic acid) with a three-dimensional supramolecular structure. **La** shows dynamic fluorescence from blue to red and switchable monotonous yellowish-green RTP, which can be manipulated by reversible photochromism. It is worth noting that $\text{Eu}^{3+}/\text{Tb}^{3+}$ -doped CPs exhibit time-resolved (red to yellow) and monotonous green afterglow, respectively, which can be attributed to multiple emissions with different decay rates. The dynamic and multicolor luminescence endows these CPs with potential for application in the domains of optical communications, multi-step encryption, and anti-counterfeiting. This work not only integrates color-adjustable fluorescence, switchable RTP, and photochromism in one material, but also realizes the manipulation of the resultant optical performances *via* photochromism, paving the pathway for the design and synthesis of smart optical materials.

Received 12th July 2024
Accepted 1st October 2024

DOI: 10.1039/d4sc04632c
rsc.li/chemical-science

Introduction

Room temperature phosphorescence (RTP) materials, which have the characteristics of millisecond lifetimes and large Stokes shifts, have great promise for versatile applications in communications, security encryption, bioimaging, and information storage.^{1–5} Based on the composition, RTP materials can be divided into three categories: inorganics,^{6,7} organics,^{8–18} and hybrids.^{19–22} The emissions in organics and hybrids both originate from organic molecules, whereas hybrids were created to optimize the RTP properties of organics due to the heavy atom effect of metals/halogens and the rigid environment constructed by coordination bonds and intermolecular interactions, and both of them are collectively referred to as molecule-based RTP materials. To date, hundreds of molecule-based RTP materials have been reported, but most present static and

monotonous afterglow,² which limits their application in intelligent fields.

Smart stimulus-responsive RTP materials always exhibit adjustable afterglow (*i.e.*, color, lifetime, quantum yield, *etc.*) under external stimuli, including heat, light, force, and chemicals, which triggers dynamic RTP.^{23–28} Particularly, the light stimulus is of interest because of its low cost, cleanliness, remote control, and easy switching. However, most photo-responsive materials are based on the consumption of triplet oxygen, which is relatively uncontrollable and random.^{29,30} Photochromism with two stable forms has been employed to regulate RTP properties, and a molecule-based photochromic system is designable and accessible by (1) introducing photo-active functional groups, such as spiropyrans,³¹ azo,³² diarylethenes,^{33,34} *etc.*; (2) constructing feasible electron transfer (ET) pathways by selecting suitable electron donor/acceptor (ED/EA) pairs.^{35–38} For example, Ma *et al.* reported that the RTP colors could be remotely regulated by controlling the isomerization of the energy acceptor;³⁹ Yan and coworkers achieved a series of metal-organic frameworks (MOFs) with switchable long-lived RTP controlled by reversible ET-induced photochromism;⁴⁰ our group realized bidirectional regulation of photo-switchable RTP based on photo-generated radicals.⁴¹ However, research on similar materials is still scarce due to the competitive relationship between photochromism and luminescence^{42,43} but such a competitive relationship also provides opportunities to

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† Electronic supplementary information (ESI) available: Crystallographic data for **La** and **Laa** have been deposited. CCDC 2206527 and 2206528. For ESI and crystallographic data in CIF or other electronic format see DOI: <https://doi.org/10.1039/d4sc04632c>

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develop photo-controllable RTP based on reversible photochromism.

Considering the above factors, coordination polymers (CPs) can serve as promising candidates for the development of photo-controllable RTP, and two basic procedures are indispensable, (1) selecting an excellent phosphor containing heteroatoms to realize long-lived RTP in CPs by virtue of the heavy atom effect and anchoring phosphor in suitable CPs to mimic the freezing environment; (2) creating allowable ET paths between EDs and EAs to achieve ET-induced photochromism. Therefore, we chose 3,3',5,5'-tetra(imidazole-1-yl)-1,1'-biphenyl (TIBP) as the phosphor and EA, and 1-hydroxyethylidene-1,1-diphosphonic acid (H_4 -HEDP) as EDs, which are co-assembled with rare-earth metal and produced a new CP, $(H_4$ -TIBP)·[La₂Li(H_2 -HEDP)₄(H -HEDP)]·3H₂O (**La**). The heterometallic phosphonate chains act as EDs and also provide a rigid environment for phosphor, while protonated TIBP moieties serve as the phosphor and EAs. **La** is colorless and shows blue steady-state photoluminescence (PL) under UV light and yellowish-green afterglow after turning off the UV lamp. After continuous photo-stimulus, the photoactivated sample (termed **La**a) shows obvious ET-triggered photochromism from colorless to pale yellow, accompanied by fluorescence change from blue to red and afterglow disappearing. Substituting partial La³⁺ ions with other lanthanide ions, we successfully synthesized isostructural Eu³⁺/Tb³⁺-doped CPs, which were endowed with the characteristic emissions of Eu³⁺/Tb³⁺. Based on the photochromism-induced dynamic PL performances, as a proof of concept, we also demonstrated the

application prospects of these crystals in the fields of optical communications, multiple encryption, and anti-counterfeiting.

Results and discussion

Single-crystal X-ray diffraction (SCXRD) characterization shows that **La** crystallizes in the $P\bar{1}$ space group and its asymmetric unit contains two La³⁺ ions, one Li⁺ ion, five partially deprotonated diphosphonate units (four H_2 -HEDP units and one H -HEDP unit), two protonated TIBP (H_4 -TIBP) moieties with a site-occupancy factor of 1/2, and three guest water molecules (Fig. S1†). All metal ions are coordinated by O-atoms of phosphonate. La1 and La2 show octa/hepta-coordinated modes to generate biaugmented trigonal prism and capped trigonal prism configurations, respectively, whereas Li1 displays the [LiO₄] tetrahedron configuration (Fig. 1a, Table S2†). The distances of La–O and Li–O are in the range of 2.414(2)–2.625(3) Å and 1.918(7)–2.082(7) Å, respectively. The angles centered on La³⁺ and Li⁺ are between 64.37(8)°–150.90(9)° and 85.6(3)°–133.4(4)° (Tables S3 and S4†). The La³⁺ ions coordinate with diphosphonate ligands (H_2 -HEDP) forming a one-dimensional structure, while the Li⁺ ions binding a terminal-coordinated diphosphonate (H -HEDP) further connect with the former constructing the final anionic chain (Fig. 1b). The protonated TIBP balances the charges of anionic chains while serving as a mediator to connect four adjacent chains through H-bonds (N–H···O: 1.79–2.09 Å) to produce a three-dimensional (3D) supramolecular structure (Fig. 1c and d). Also, the lattice provides a rigid environment for TIBP which is beneficial for phosphorescence.^{19,40}



Fig. 1 (a) The geometric configurations of La1, La2, and Li1 (BTPR-8: biaugmented trigonal prism; CTPR-7: capped trigonal prism; vT-3: vacant tetrahedron); (b) heterometallic phosphonate chain; (c) H-bonds between protonated TIBP units and phosphonate chains (orange dashed); (d) 3D supermolecular structure.



La exhibits coloration from colorless to pale yellow within 3 min after photo-stimulus with a Xe-lamp (300 W). To investigate the photochromic process, the irradiation time-varied solid UV-vis absorption spectra were obtained. As shown in Fig. 2a and S2,[†] a new absorption peak at 250–450 nm can be observed after successive photo-stimulus, which can be ascribed to the yield of TIBP[•] radicals. The photo-induced radicals were further confirmed by solid-state electron paramagnetic resonance (EPR) measurements. The EPR spectra reveal a sharp signal at $g = 1.9931$ after photo-stimulus (Fig. 2b), while no signal is observed before, clearly indicating the generation of radicals. Detailed structural analysis was conducted to illustrate the photochromic mechanism. The rigid protonated TIBP guests are potential EAs, while the anionic heterometallic phosphonate hosts with O-abundant characteristics are potential EDs.^{35,38} The closest distance (2.602 Å) between N_{TIBP} and O_{HEDP} is advantageous to ET in the presence of H-bonds (N–H...O). Under UV light, **La** absorbs light energy and converts it into chemical energy, promoting the ET from O_{HEDP} to N_{TIBP} (Fig. 2c). Therefore, the photocoloration of **La** can be interpreted as photo-driven ET between EDs and EAs. The powder XRD and IR spectra (Fig. S3 and S4[†]) suggest that no apparent structural change is observable after coloration. Similar to ET-induced photochromic CPs, the **Laa** sample could relax to its original state after heating treatment at 120 °C for 3 h, which can be due to the oxidative quenching of TIBP[•] radicals at high temperatures.^{40,44–46} Also, thermogravimetric analysis (TGA) reveals that **La** can be stable up to 200 °C, which confirms the stability during the heating relaxation process (Fig. S5[†]).

The reversible photochromism induces dynamic dual fluorescence in **La**. The prompt PL spectra in Fig. 2d show one emission peak located at 334 nm with a shoulder in the range of 365–500 nm when excited by 280 nm UV light. The decay curves obtained at 334 nm suggest that the lifetime is 1.47 ns (Fig. 2e), indicating that the emission can be ascribable to fluorescence. After consecutive photoactivation, the emission peak at 334 nm is gradually quenched, while the shoulder peak centered at 422 nm appeared and gradually increased, showing the redshift of fluorescence from violet to blue (Fig. S6[†]). The decay curve obtained at 422 nm suggests the lifetime is 3.71 ns, which is also rooted in the singlet state. To understand the redshift of fluorescence, we obtained the PL spectrum of TIBP as shown in Fig. 2f, which shows a dominating emission band at 405 nm with a shoulder centered at 350 nm. The similar emissions between TIBP and **Laa** inspire us to speculate as follows, (1) the PL in **La** mainly originates from the TIBP ligand;^{19,40} (2) the light induces the weakening protonation of TIBP, and the increasing electron cloud density, which leads to the fluorescence of **Laa** tending towards the unprotonated TIBP.^{35,38}

When excited by 365 nm UV light, **La** exhibits blue emission while **Laa** changes to pink, so a series of excitation wavelength-varied emission spectra were obtained. As shown in Fig. S7,[†] **La** and **Laa** both exhibit wide excitation wavelength-dependent PL properties due to their multiple emissions involving singlet and triplet excitons. Surprisingly, the PL spectra of **Laa** show two new emissions at 580 nm and 631 nm when excited by 335–375 nm light, so a series of PL spectra with different irradiation times were obtained ($\lambda_{\text{ex}} = 370$ nm). As shown in Fig. 3a and S8,[†] one emission peak is located at 430 nm with a lifetime of



Fig. 2 (a) The irradiation time-dependent UV-vis absorption spectra; (b) EPR spectra before and after irradiation; (c) the schematic diagram of the ET process; (d) the irradiation time-dependent fluorescence spectra excited by 280 nm UV light; (e) the decay curves monitored in 334 nm and 422 nm; and (f) the prompt excitation and emission spectra of TIBP.



Fig. 3 (a and b) The irradiation time-dependent fluorescent spectra excited by 370 nm UV light and their corresponding CIE coordinates; (c) the irradiation time-dependent delayed PL spectra of **La** (inset: photographs of **La** and **La** after turning off the 365 nm UV lamp); (d) long-lived decay curves of **La**; (e) calculated molecular orbitals.

3.99 ns. As the irradiation time increases, the intensity at 430 nm decreases and the lifetime shortens to 2.79 ns, and two new peaks appear at 580 nm and 631 nm which increase gradually, which result in a red-shift in CIE coordinates (Fig. 3b). The decay curves prove that the lifetimes are 10.54 and 10.88 ns, respectively, suggesting that these two emissions may originate from intramolecular charge transfer (CT) rather than from rare earths.⁴⁷ The characteristic peaks of a series of $\text{Eu}^{3+}/\text{Tb}^{3+}$ -doped CPs in subsequent experiments are completely different from the two peaks mentioned above, which also confirms this point. In addition, the absorption band at 250–300 nm can be due to the π - π^* transition of aromatic rings, whereas the band at 300–400 nm is due to the n - π^* transition involving the ligand to ligand CT process (Fig. 2a and S2†). This also proves that the enhanced ligand to ligand CT process in the photoactivated sample may induce two new emissions.

The reversible photochromism also endows **La** with photo-switchable afterglow. As shown in Fig. 3c and S9,† **La** exhibits yellowish green afterglow after ceasing the 365 nm UV light, which almost disappears after coloration, so the delayed PL emission spectra with different irradiation times were measured. Before irradiation, the emission is at around 545 nm with a lifetime of 252.31 ms (Fig. 3d), which is mainly donated by TIBP ligands and can be assigned to RTP due to its long-lived lifetime and large Stokes shift (Fig. S10†). The temperature-varied delayed PL spectra in Fig. S11† show that the emission intensity increases with the temperature decreasing, further suggesting that the long-lived emission can be assigned to phosphorescence. The RTP intensity is gradually decreased as

the irradiation time increases, corresponding to the diminishing afterglow until it disappears, which is consistent with previous findings.^{40,43}

To understand the photophysical process, we built a model based on **La** and calculated the highest occupied molecular orbitals (HOMOs) and the lowest unoccupied molecular orbitals (LUMOs) based on the density functional theory (DFT) calculations. As shown in Fig. 3e and S12,† the HOMOs are mainly localized on HEDP, while the LUMOs are mainly populated on the benzene or imidazole group of TIBP, suggesting the occurrence of charge transfer from HEDP to TIBP (LLCT), which mainly serves the photoemission process.

Utilizing the similarity in coordination configurations and differences in characteristic emissions of rare earths, we doped a series of amounts of Eu^{3+} and Tb^{3+} (molar ratio: 1%, 2%, 5%, 10%, and 20%) into **La**, and obtained a series of isostructural CPs to achieve multicolor fluorescence and RTP (Fig. S13†).^{48,49} For Eu-doped CPs, the fluorescence goes from blue to pink while afterglow changes from yellowish-green to orange, and orange-red with the doping proportion increasing (Fig. 4a and S14†). Employing **La_{0.9}Eu_{0.1}** as an example, scanning electron microscope (SEM) and elemental mapping images reveal a microrod morphology with a uniform distribution of La and Eu elements on the crystal surface (Fig. 4b and S15†), and the Eu^{3+} content is determined to be 12.91%. To further confirm the ratio between metal centers, we also performed X-ray fluorescence (XRF) spectroscopy, with the more precise results summarized in Table S7.† The experimental XRF data indicate that **La_{0.9}Eu_{0.1}** contains a lower Eu^{3+} content of 8.57%. The



Fig. 4 (a) The photographs of $\text{La}_{0.9}\text{Eu}_{0.1}$ and $\text{La}_{0.9}\text{Tb}_{0.1}$ before and after turning off the UV lamp; (b) SEM and elemental mapping images of $\text{La}_{0.9}\text{Eu}_{0.1}$; (c) the irradiation time-dependent delayed PL spectra of $\text{La}_{0.9}\text{Eu}_{0.1}$ excited by 365 nm UV light; (d) long-lived decay curves of $\text{La}_{0.9}\text{Eu}_{0.1}$; (e and f) the time-resolved emission spectra (TRES) and the corresponding emission spectra; (g) the photophysical processes of $\text{La}_{0.9}\text{Eu}_{0.1}$.

prompt and delayed PL spectra of $\text{La}_{0.9}\text{Eu}_{0.1}$ both exhibit four new sharp peaks at 592, 616, 655, and 701 nm, corresponding to the $^5\text{D}_0\text{--}^7\text{F}_j$ ($j = 1\text{--}4$) transitions of Eu^{3+} . When excited by 280 nm UV light, the emission intensity at 334 nm decreases while other emission intensities increase with prolonged irradiation (Fig. S16a†). However, the prompt and delayed spectra in Fig. 4c and S16b† show that all emission intensities decrease as the irradiation time increases, suggesting that its PL properties can also be switched by ET-induced photochromism (Fig. S17†). These different phenomena also confirm the simultaneous occurrence of energy transfer from S_1 to Eu^{3+} and from T_1 to Eu^{3+} . It is worth mentioning that $\text{La}_{0.9}\text{Eu}_{0.1}$ exhibits time-resolved afterglow change from orange to yellow. To clarify the multi-color afterglow, the decay curves monitored in main peaks were exported. As shown in Fig. 4d, the lifetime at 545 nm is 296.45 ms, while that at 592, 616, and 701 nm is 41.97, 60.06, and 50.44 ms, suggesting that (1) the lifetime of characteristic transitions is significantly longer than that of traditional rare-earth-based materials, which results in the orange afterglow; (2) the decay rate at 545 nm is slower than others, so the afterglow changes from orange to yellow with time evolution. To capture the time-resolved afterglow, a series of time-resolved emission spectra (TRES) excited by 365 nm UV light were recorded. The delayed spectrum and TRES show that Eu^{3+} -dominated emission (592, 616, 655, and 701 nm) is

significantly stronger than the emission at 545 nm (from TIBP) when delayed by 1 ms and 10 ms, corresponding to orange-red afterglow (Fig. 4c, e and f). As aforementioned, Eu^{3+} -dominated emission decays rapidly while TIBP-dominated emission decays slowly, so as the delay time increases (>100 ms), the intensities of TIBP- and Eu^{3+} -dominated peaks trend to contribute equally, corresponding to yellow afterglow (Fig. S18†).

Different from Eu^{3+} -doped CPs, the fluorescence of Tb^{3+} -doped CPs does not exhibit obvious changes, but their afterglows are brighter (Fig. 4a and S19†). For $\text{La}_{0.9}\text{Tb}_{0.1}$, SEM and elemental mapping images also reveal a microrod morphology with a uniform distribution of La and Tb elements on the crystal surface (Fig. S20†), and the Tb^{3+} content is determined to be 12.23%. The XRF analysis shows that $\text{La}_{0.9}\text{Tb}_{0.1}$ contains a higher Tb^{3+} content of 19.61% (Table S7†). The prompt and delayed PL spectra both show four new sharp peaks at 490, 545, 585, and 622 nm, corresponding to the $^5\text{D}_4\text{--}^7\text{F}_j$ ($j = 3\text{--}6$) transitions of Tb^{3+} . When excited by 280 nm UV light, the Tb^{3+} -dominated emissions remain unchanged with prolonged irradiation (Fig. S21a†). However, the prompt and delayed spectra in Fig. S21b and S22† show that all emission intensities decrease as the irradiation time increases, corresponding to photo-switchable PL properties (Fig. S23†). The long-lived decay curves suggest that their corresponding lifetimes are 49.28, 138.30, 47.15, and 62.27 ms (Fig. S24†). Therefore, it can be

inferred that the brighter second-level afterglow is the result of the combination of the triplet state of TIBP and the antenna effect of Tb^{3+} . However, a series of Tb^{3+} -doped CPs show a monotonous green afterglow, which can be due to the characteristic emission of Tb^{3+} being similar to TIBP-originated RTP.

Benefiting from the photochromism and photo-controllable fluorescence, **La** microrods show photo-modulated optical waveguide properties, which are of great significance for the information security of photonic communications. As depicted in Fig. 5a and c, both endpoints of the rod-shaped crystal exhibit stronger emission under unfocused UV light, which means the optical waveguide properties. Therefore, detailed spatial-resolved PL microscopy images were obtained by changing the local positions on this microrod. Under a 375 nm laser beam, **La** exhibits bright blue emission, and the intensity at the tip gradually decreases with increasing propagation distance, which indicates that some of the photons generated by the excitation are effectively locked in the microrod and propagated from the excitation position to the tip. The optical waveguide loss performance can be evaluated by fitting the optical loss coefficient (R): $I_{\text{tip}}/I_{\text{body}} = A \exp(-RD)$ ($I_{\text{tip}}/I_{\text{body}}$ = the intensity at the tips/excited positions; D = the distance between exciting positions and tips). The R of **La** is $4.83 \times 10^{-4} \text{ dB } \mu\text{m}^{-1}$, which is significantly lower than that of most of the optical waveguide materials reported so far, indicating that **La** possesses excellent optical waveguide performance. This superior performance can be due to **La**'s high crystallinity, distinctive shape and smooth surface, which can effectively minimize the optical loss caused by scattering from the domain boundary.

Since **La** shows dynamic photo-induced PL emission, the optical waveguide properties after irradiation were also detected (Fig. 5b and d). **Laa** exhibits similar optical waveguide performance while the emission changes from blue to red under the 375 nm laser beam. The emission intensity gradually decreases as the excitation spot moves along the crystal, and the R value of **Laa** is $2.24 \times 10^{-3} \text{ dB } \mu\text{m}^{-1}$. **Laa** exhibits inferior optical waveguide performance compared to **La**, which can be due to two primary factors: (1) **Laa** exhibits stronger self-absorption (Fig. 2a). During photonic propagation within the **Laa** microrod, a portion of photons is lost through self-absorption, resulting in higher optical loss. (2) As depicted in the irradiation time-dependent fluorescence spectra (Fig. 3a), the emission intensity of **Laa** is lower than that of **La**. Also, the main emission peak (580 and 631 nm) of **Laa** exhibits a longer fluorescence lifetime (Fig. S8†), which competes with the quantum yield. However, both the R value and the waveguide color show significant changes before and after irradiation, suggesting that the optical waveguide behavior can be regulated by reversible photochromism.

To elucidate the relationships between 1D photonic properties and morphologies, the morphology simulation and prediction for **La** was conducted on the basis of Bravais-Friedel-Donnay-Harker (BFDH) theory (Fig. 5e).^{50,51} The simulated morphology suggests four fast-growing crystal facets of $\{010\}$, $\{001\}$, $\{011\}$, and $\{100\}$ with surface areas of 27.27, 24.80, 13.42, and 16.16%, respectively (Table S5†), and **La** grows into a 1D microrod crystal along the direction of the black arrow (Fig. S25†).

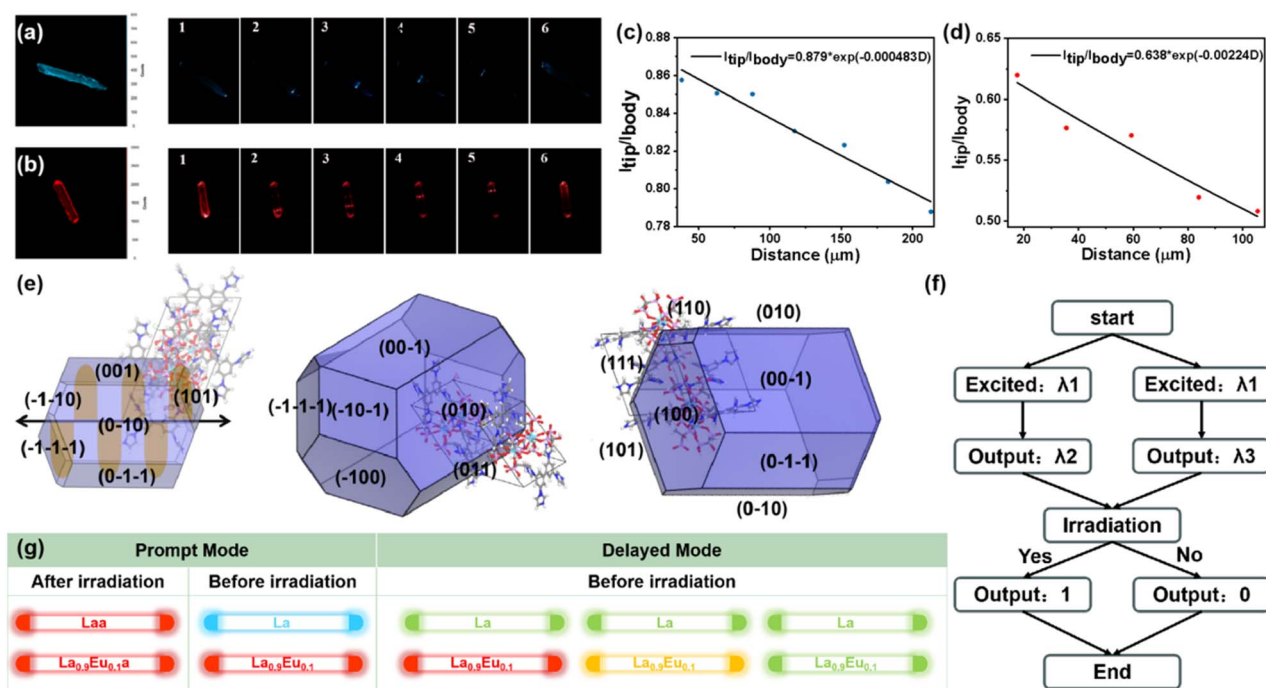


Fig. 5 PL images of microrods **La** (a) and **Laa** (b) excited with focused 370 nm light at different positions; the optical loss coefficients of **La** (c) and **Laa** (d); (e) the morphology simulation and corresponding crystal faces of **La** based on the BFDH method; (f) illustration of the optical logic gates; (g) predicted multi-mode 1D active waveguide properties.

By virtue of the dynamic optical waveguide behavior driven by photochromism, we designed an advanced secure optical logic gate prototype using **La** and **Laa** (Fig. 5f). We input an excitation wavelength (λ_1) into a flat waveguide coupler to activate the optical logic gate system, which can collect two wavelengths at the output port (λ_2, λ_3). If the emission in the tip is blue, the output port is “0”; otherwise the output is “1”. In this manner, a binary number output could be achieved within the 1D microcrystal. Based on the photo-controllable fluorescence and RTP in **La**, we reasonably speculate that the $\text{Eu}^{3+}/\text{Tb}^{3+}$ -doped CPs would exhibit optical waveguide behavior with multicolor emission, whether in prompt or delayed modes. As shown in Fig. 5g, **La** not only exhibits bi-mode optical waveguide properties (blue-red) in prompt mode, but is also predicted to present waveguide properties with yellowish-green emission in delayed mode. However, **La_{0.9}Eu_{0.1}** would exhibit completely different waveguide behaviors, which show red emissions before and after irradiation in prompt mode, and changes to orange and yellowish-green over time in delayed modes. Therefore, utilizing such smart and diverse waveguide behavior will be promising for designing more advanced optical logic gate systems.⁵²

The dynamic fluorescence and RTP properties provide **La** and $\text{Eu}^{3+}/\text{Tb}^{3+}$ -doped CPs with opportunities for applications in multi-step encryption and anti-counterfeiting fields. For example, we designed a house pattern that was filled with as-synthesized samples, in which the roof and door were filled with **La_{0.9}Eu_{0.1}**, the wall and tree trunk were filled with **La**, the chimney was filled with **Laa**, and leaves were filled with **La_{0.9}Tb_{0.1}** (Fig. 6a). When the pattern is exposed to daylight, all houses are white, which is easily imitated and forged. When the

pattern is exposed to UV light, and then turned off for 0.1 s, 1 s and 2 s, the pattern presents different emissions, which achieves temporal- and spatial-resolved multiple anti-counterfeiting. Meanwhile, the more emission states there are, the higher the difficulty for forging, confirming the application prospects of these CPs in the field of anti-counterfeiting.

Inspired by the time-resolved emission of **La_{0.9}Eu_{0.1}** and its similar emission to **La**, we also designed a Morse code application employing 2,6-naphthalene dicarboxylic acid (NDA) as a costar. As shown in Fig. 6b and S26,† all blocks are white under daylight, and show multi-color emissions under 365 nm UV light, so we cannot obtain any effective information. After turning off the UV light, the blocks filled with NDA disappear first, while **La** exhibits yellowish-green afterglow marked as “.” and Eu -doped CPs exhibit red-orange afterglow marked as “-”. The pattern can be horizontally decrypted as “SDQD”, which is the correct decoding letter representing “Shandong Qingdao”. Simultaneously, the “SFHA” in the vertical direction can confuse enemies. Also, the correct information “SDQD” can disappear in time. After turning off the UV lights for 1 s, the pattern changes to “SSHS” and “SHHI” in the horizontal and vertical directions, respectively. Based on the as-synthesized CPs, more diverse and useful Morse code can be designed and hopefully applied in practice.

These CPs are also promising for application in advanced data encryption and smart display. As depicted, an “8” pattern can also be designed as a digital encryption using **La**, **Laa**, and **La_{0.9}Eu_{0.1}** (Fig. 6c). Under daylight, the “8” pattern appears white. Under 365 nm UV light, all parts show indistinguishable emissions, so the pattern displays as “8”. When the UV light is extinguished, the emission of **Laa** disappears, while **La** and



Fig. 6 (a) The house pattern under daylight, UV light, and after turning off for 0.1 s, 1 s, and 2 s; (b) the model for Morse code application; (c) the digit model.

C₃₄H₅₇N₈O₃₈P₁₀La_{1.8}Tb_{0.2}Li (%). Calculated: C, 23.30; H, 3.54; N, 6.02. Found: C, 22.97; H, 3.50; N, 5.89. IR of **La_{0.9}Tb_{0.1}** (KBr pellets, cm⁻¹): 3435(s), 2977(w), 2927(w), 2854(w), 1618(s), 1468(w), 1422(w), 1125(m), 1072(m), 926(w), 804(w), 646(w), 557(w).

Experimental

Materials and measurements

IR spectra were obtained on a MAGNA-560 (Nicolet) FT-IR spectrometer. The room-temperature photoluminescence spectra were recorded on an F-4700 fluorescence spectrometer. The time-resolved emission spectra (TRES) were obtained on an FS5 fluorescence spectrometer. The temperature-dependent emission spectra and all decay curves were obtained on an FLS 1000 fluorescence spectrometer. The solid-state UV-vis spectra were measured on a Puxi Tu-1901 spectrophotometer with a BaSO₄ reference. Electron paramagnetic resonance (EPR) was recorded on a Bruker E500 spectrometer. A scanning electron microscope (SEM, Hitachi, S2400) equipped with an energy dispersive spectroscopy (EDS) detector was used to investigate the microstructures and elements. Chemical compositions of lanthanum oxide products were investigated by using a PANalytical Axios X-ray fluorescence spectrometer. The experimental powder X-ray diffraction (PXRD) analyses were conducted on a Rigaku D/max-2550 diffractometer with Cu-K α radiation ($\lambda = 1.5418 \text{ \AA}$). The simulated PXRD curve was derived from the SCXRD data and Mercury software. The Xe-lamp for time-dependent solid UV-vis absorption characterization is a Perfect Light PLS-SXE 300. The photosource for the luminescence characterization is the intrinsic light of the instrument.

Synthesis of La and Laa

A mixture of La_2O_3 (0.03 g, 0.092 mmol), TIBP (0.02 g, 0.05 mmol), $\text{H}_4\text{-HEDP}\cdot\text{H}_2\text{O}$ (0.10 g, 0.45 mmol), LiF (0.10 g, 4.00 mmol), 2 mL CH_3OH and 4 mL H_2O was sealed in a Teflon-lined autoclave and heated at 120 °C for 6 days, and colorless crystals (termed **La**) were obtained and washed with absolute ethanol. Elemental analysis for $\text{C}_{34}\text{H}_{57}\text{N}_8\text{O}_{38}\text{P}_{10}\text{La}_2\text{Li}$ (%). Calculated: C, 22.94; H, 3.23; N, 6.29. Found: C, 23.47; H, 3.58; N, 6.02. IR of **La** (KBr pellets, cm^{-1}): 3407(s), 3155(s), 2851(m), 2365(w), 1618(m), 1545(w), 1467(w), 1363(w), 1129(s), 1067(s), 920(s), 800(w), 650(w), 560(m). After sufficient illumination of the above crystals, light yellow crystals were obtained (termed **Laa**). IR of **Laa** (KBr pellets, cm^{-1}): 3395(s), 3155(s), 2970(m), 2932(w), 2859(m), 2744(w), 2645(w), 1618(m), 1543(w), 1468(w), 1365(w), 1125(s), 1076(s), 918(s), 857(w), 800(w), 646(w), 557(m).

Synthesis of Eu³⁺/Tb³⁺-doped CPs

Replace La_2O_3 with different doping proportions of Eu_2O_3 or Tb_2O_3 (1%, 2%, 5%, 10%, and 20%), and other processes are consistent with the synthesis of **La**. Elemental analysis for $\text{C}_{34}\text{H}_{57}\text{N}_8\text{O}_{38}\text{P}_{10}\text{La}_{1.8}\text{Eu}_{0.2}\text{Li}$ (%). Calculated: C, 22.90; H, 3.22; N, 6.28. Found: C, 23.01; H, 3.52; N, 6.33. IR of **La_{0.9}Eu_{0.1}** (KBr pellets, cm^{-1}): 3435(s), 2977(w), 2927(w), 2852(w), 1625(s), 1465(w), 1417(w), 1122(m), 1086(m), 1043(m), 926(w), 808(w), 643(w), 561(w). Elemental analysis for

X-ray crystallography

The single-crystal X-ray diffraction data of **La** and **Laa** were collected on a Rigaku Oxford Diffraction at 293(2) K with Mo-K α radiation ($\lambda = 0.71073$ Å). The SHELX-2016 software was employed to solve the structure.⁵³ Detailed crystallographic data for **La** and **Laa** are summarized in Table S1,[†] and the selected bond lengths and angles are listed in Tables S3 and S4.[†] Full crystallographic data have been deposited with CCDC 2206527 for **La** and 2206528 for **Laa**.

Calculation details

The calculations were conducted using the density functional theory (DFT) method in Gaussian16.⁵⁴ B3LYP-D3/def2-SVP was applied to get the Frontier Molecular Orbital.^{55–58} Wavefunction analyses were executed using Multiwfn.⁵⁹ Visualization of the frontier molecular orbitals was achieved using the VMD software.⁶⁰

Conclusions

In summary, a series of La^{3+} -based and $\text{Eu}^{3+}/\text{Tb}^{3+}$ -doped CPs were successfully assembled by using metal-diphosphonate and rigid tetraimidazole with the heterometallic phosphonate chains as electron-donating hosts and protonated tetraimidazole moieties as electron-accepting guests. Under photo-stimulus, these CPs characterized with an ED-EA skeleton not only show obvious ET-facilitated photochromism, but also display photochromism-induced color-adjustable fluorescence and switchable afterglow, which can be due to photo-generated radicals. It is worth mentioning that Eu^{3+} -doped CPs exhibit time-resolved afterglow, while Tb^{3+} -doped CPs exhibit brighter green afterglow, which inspires us that reasonable doping would be an effective strategy for constructing multi-color and efficient RTP materials. The variable fluorescence and RTP in these CPs provide these materials with more optical communication applications worth exploring, such as **La** microrods show photo-modulated bi-mode switchable optical waveguide properties (blue to red) with a low optical loss coefficient. In addition, a series of CPs also present promising application prospects in the fields of multi-step encryption and anti-counterfeiting. This work demonstrates photo-adjustable fluorescence and RTP by regulating photochromism, and provides a doping strategy to obtain phosphorescence with different surprising properties, affording a theoretical and experimental basis for the development of intelligent optical materials.

Data availability

The data supporting this article have been included as part of the Supplementary Information. Crystallographic data for

La and Laa has been deposited at the CCDC under 2206527 and 2206528.

Author contributions

Y.-J. Ma and F. Xu contributed equally to this work. Y.-J. Ma, J.-H. Li, S.-D. Han and G.-M. Wang conceived the main idea and designed the experiments. F. Xu, X.-Y. Ren and J. Pan synthesized and characterized the coordination polymers. Y.-J. Ma, F. Xu, X.-Y. Ren and F.-Y. Chen performed and analyzed the photophysical measurements. Y.-J. Ma, J.-H. Li, S.-D. Han and G.-M. Wang analyzed all data. Y.-J. Ma, F. Xu, J.-H. Li, S.-D. Han and G.-M. Wang wrote the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This study was supported by the National Natural Science Foundation of China (22071125 and 22071126), the Natural Science Foundation of Shandong Province (ZR2021MB064), the Youth Innovation Team Project of Shandong Provincial Education Department (No. 2023KJ234), and the Qingdao University discipline cluster interdisciplinary joint research project (FZ2024201).

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