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A tunable full-color lanthanide noncovalent polymer based on cucurbituril-mediated supramolecular dimerization†

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The construction of lanthanide multicolor luminescent materials with tunable photoluminescence properties has been developed as one of the increasingly significant topics and shown inventive applications in miscellaneous fields. However, fabricating such materials based on synergistically assembly-induced emission rather than simple blending of different fluorescent dyes together still remains a challenge. Herein, we report a europium-based noncovalent polymer with tunable full-color emission, which is constructed from the 2,6-pyridinedicarboxylic acid-bearing bromophenylpyridinium salt. This rationally designed bifunctional component can concurrently serve as a guest molecule and a chelating ligand to associate with cucurbit[8]uril and europium ions, thus leading to the formation of a trichromatic (red-green-blue, RGB) photoluminescent polypseudorotaxane-type noncovalent polymer in aqueous solution. Meanwhile, the full-color emission enclosed within the RGB color triangle could be readily produced by simply tuning the molar ratio of cucurbit[8]uril and europium ions. The lanthanide supramolecular polymer featuring tricolor emission, long lifetime, high photoluminescence efficiency and low cytotoxicity could be further applied in multicolor imaging in a cellular environment. These results provide a new and feasible strategy for the construction of full-color single lanthanide self-assembled nanoconstructs.

Introduction

Multicolor photoluminescent materials, especially white lightemitting devices with an emission spectrum spanning the whole visible region, have attracted considerable attention¹⁻³ due to their promising applications in optical sensors,4-6 bioimaging agents,7-10 flexible full-color displays,11-14 and lightemitting diodes.¹⁵⁻¹⁷ Common methods to obtain multicolor emitting materials rely on simply mixing three primary redgreen-blue (RGB) colors or covalent attachment of complementary chromophores in a precise ratio, which often involves tedious synthetic procedures and undesired cross-color interference among the multicomponent emitters. 18-20 To alleviate these shortcomings, macrocycle-involved host-guest interactions have been developed as a powerful approach to design multicolor luminescent materials, in which individual lumiwell-ordered nophores self-assemble

Lanthanide complexes have been widely utilized in the design of luminescent materials and imaging agents due to their excellent photophysical properties, such as large Stokes shifts, narrow emission bands, and long-lived excited states. 33-35 Although multicolor emission by blending discrete lanthanide metal ions and organic ligands has been known in some cases, 36-39 full-color emission solely based on a single lanthanide metal ion still remains a challenge in aqueous media, to the best of our knowledge. This is mainly due to the vulnerable excited state of lanthanide complexes that can be severely quenched by the O–H group of water molecules and the established chelating ligands around lanthanide metal ions that lack tunable photoluminescence characteristics. 40,41

nanoarchitectures with distinct changes in photophysical properties. ^{21–24} Among the most frequently encountered macrocycles, cucurbiturils (CBs), a family of macrocyclic receptors possessing hydrophobic cavities and carbonyl-laced portals, have been explored as versatile building blocks to fabricate supramolecular nanoconstructs with elaborate structures and functions. ^{25–27} In this context, cucurbit[8]uril (CB[8]), an eightmembered CB homologue, has been drawn into the limelight because it can enable two specific guest molecules to stack in close proximity within the rigid cavity and often leads to dramatic photoluminescence enhancement in both inanimate milieus and biological fluids. ^{28–32}

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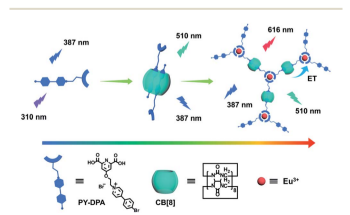
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Recently, we have developed several types of CB-confined supramolecular nanosystems with a wide color range and purely organic room-temperature phosphorescence (RTP).42-44 This motivated us to integrate the host-guest-binding-induced phosphorescence and intrinsic lanthanide photoluminescence into a single self-assembled entity, with the aim of attaining a supramolecular lanthanide assembly possessing tunable fullcolor emission properties in aqueous solution. In this work, a heteroditopic guest, 2,6-pyridinedicarboxylic acid-modified bromophenylpyridinium salt (PY-DPA) could form a stable biaxial pseudorotaxane with CB[8], accompanied by a conversion from blue fluorescence to green phosphorescence in water. Meanwhile, the coordination of terminal carboxylic groups with europium ions (Eu³⁺) could further lead to the formation of a lanthanide supramolecular polymer, which would significantly enhance the characteristic luminescence of Eu³⁺ via the complexation-triggered intramolecular transfer energy pathway. As a result, tunable full-color luminescence (including the white-light emission) enclosed within the RGB triangle could be conveniently achieved by varying the proportion of Eu3+ and CB[8]. Furthermore, this nanoassembly proved to be highly biocompatible and capable of multicolor cellular imaging upon excitation at a single wavelength. Therefore, it can be envisioned that the resultant Eu³⁺@PY-DPA⊂CB[8] assembly featuring full color (red-green-blue), different photophysical origins (photoluminescence from singlet and triplet excited states), and diverse scales in lifetimes (ns-level fluorescence, us-level phosphorescence, and ms-level lanthanide emission) will offer a new and elegant strategy for the fabrication of multicolor smart light-emitting materials.

Results and discussion

The construction of a full-color emissive lanthanide non-covalent polymer is depicted in Scheme 1. The dual-functional guest PY-DPA was synthesized by the reaction between a 2,6-pyridinedicarboxylic acid derivative and 4-(4-bromophenyl)-pyridine. The structural characterization is shown in the ESI (Chart S1 and Fig. S1–S6).† To investigate the binding behavior



Scheme 1 Schematic illustration of a lanthanide noncovalent supramolecular polymer (Eu³⁺@PY-DPA⊂CB[8] assembly) for tunable full-color emission.

between PY-DPA and CB[8], ¹H NMR titration experiments were preliminarily conducted. As shown in Fig. 1a, the aromatic protons assigned to the PY moiety (Ha-d) underwent a dramatic upfield shift upon gradual addition of CB[8], while the protons of the DPA unit (H_e) shifted to downfield. These phenomena suggested that the PY moiety was deeply encapsulated in the cavity of CB[8] and the DPA part was located outside the cavity of CB[8]. Besides, the changes of chemical shifts of all aromatic protons achieved the equilibrium state with 0.5 equivalents of CB[8], implying a 1: 2 binding stoichiometry between CB[8] and PY-DPA. In addition, 2D rotating-frame Overhauser effect spectroscopy (ROSEY) was employed to gain more insight into the binding mode. Strong correlated signals between H_b and H_d protons were observed, indicative of the head-to-tail stacking mode of two pyridinium units in the cavity of CB[8] (Fig. 1b). Moreover, the formation of the ternary PY-DPA ⊂ CB[8] inclusion complex was also evaluated by UV-vis spectroscopy. As shown in Fig. 2a, free PY-DPA exhibited the main absorption peak at 308 nm, which experienced a slight bathochromic shift to 312 nm and a decrease in intensity upon the stepwise addition of CB[8], accompanied by the emergence of two isosbestic points at 247 nm and 330 nm. The binding stoichiometry between CB[8] and PY-DPA was further validated as 1:2 from the Job plot, and the association constant (K_a) in the PY-DPA \subset CB[8] complexation was calculated to be 7.46 \times 10¹¹ M⁻² by the nonlinear least-squares method (Fig. S7†).

Subsequently, the photoluminescence properties of the PY-DPA CB[8] complex in aqueous solution were further explored. As can be seen from Fig. 2b, free PY-DPA showed blue emission centered at around 387 nm. The appearance of a new emission peak at 510 nm with a decrease in emission at 387 nm was clearly observed upon stepwise addition of CB[8] into the aqueous solution of PY-DPA. The luminescent color changes from blue to green could also be recognized by the naked eye. With a delay time of 50 μs, the emission at 387 nm completely disappeared and only the peak at 510 nm was maintained (Fig. 2c). As revealed by the time-resolved decay curves, the lifetime measured at 387 nm was on a nanosecond scale and the lifetimes were assigned as 0.43 and 0.46 ns for free PY-DPA and the PY-DPA CB[8] complex, respectively (Fig. S8†). In comparison, the lifetime measured at 510 nm for the PY-

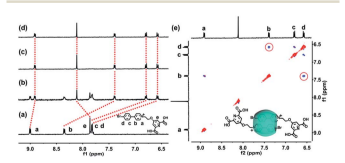


Fig. 1 1 H NMR titration spectra (400 MHz, D₂O, 298 K) of PY-DPA (5.0 \times 10⁻⁴ M) upon addition of (a) 0.00, (b) 0.25, (c) 0.50, and (d) 1.0 equivalent of CB[8]; (e) 2D-ROSEY spectrum of the PY-DPA \subset CB[8] complex in D₂O ([PY-DPA] = [CB[8]] = 5.0 \times 10⁻⁴ M).

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(a) 0.4 (b) 1.6x10³ (c) 1.2x10³ (d) 1.6x10⁴ (d) 1.6x10⁴ (e) 1.6x10⁴ (e) 1.6x10⁴ (e) 1.6x10⁴ (f) 1.6x

Fig. 2 (a) UV-vis absorption spectra ([CB[8]] = $0-1.3 \times 10^{-5}$ M) and (b) prompt photoluminescence spectra of PY-DPA (1.0×10^{-5} M) upon gradual addition of CB[8] ([CB[8]] = $0-1.0 \times 10^{-5}$ M) in aqueous solution. Inset: photograph showing the changes in luminescent color under light irradiation at 302 nm; (c) prompt and gated photoluminescence spectra (delayed time = $50 \mu s$) of the PY-DPACB[8] complex; (d) phosphorescence lifetime decay curve of the PY-DPACCB[8] complex at 510 nm ([PY-DPA] = [CB[8]] = $1.0 \times 10^{-5} \text{ M}$, $\lambda_{ex} = 310 \text{ nm}$, 298 K).

elength (nm)

Time (ms)

DPA \subset CB[8] complex was on a microsecond scale (650 µs, Fig. 2d). Moreover, the intensity at 510 nm showed an apparent enhancement after deoxygenation treatment, and the lifetime increased to 2748 µs. Meanwhile, no obvious change was found with respect to the intensity at 387 nm (Fig. S9 \dagger). Thus, it could be reasonably inferred that the emission at 510 nm with a relatively longer lifetime came from host–guest-binding-induced RTP.

It is documented that 2,6-pyridinedicarboxylic acid could strongly coordinate with lanthanide metal ions (i.e. Eu3+ and Tb³⁺) and harvest energy for the sensitization of the characteristic emission.45 Therefore, a UV-vis titration experiment was performed to study the metal-chelating behaviors between PY-DPA and Eu³⁺. With the addition of Eu³⁺ to the PY-DPA solution, the absorbance of PY-DPA gradually decreased, as a result of the coordination of Eu3+ with the carboxylic groups of PY-DPA (Fig. S10†). At the same time, the coordination stoichiometry was determined to be 1:3 by analyzing the absorbance at 300 nm versus [Eu3+]/[PY-DPA] molar ratios. In addition, the characteristic emission of Eu³⁺ with four sharp peaks at 595 nm (5D₀ \rightarrow 7F₁), 616 nm (5D₀ \rightarrow 7F₂), 650 nm (5D₀ \rightarrow 7F₃) and 695 nm $(5D_0 \rightarrow 7F_4)$ was observed in the Eu³⁺@PY-DPA and Eu³⁺@PY-DPA CB[8] complexes, on account of the efficient energy transfer from the DPA moiety to Eu³⁺ (Fig. 3c). It was also found that two sets of emission peaks, especially the phosphorescence one, showed an obvious decline by adding Eu³⁺ to the PY-DPA CB[8] complex (Fig. 3d and e). Thus, we inferred that an energy transfer process might occur between the PY ⊂ CB[8] part and Eu³⁺@DPA center. The more direct evidence was provided by the time-resolved curves and excitation spectra. With the gradual addition of Eu³⁺ to the PY-DPA ⊂ CB[8] complex, the

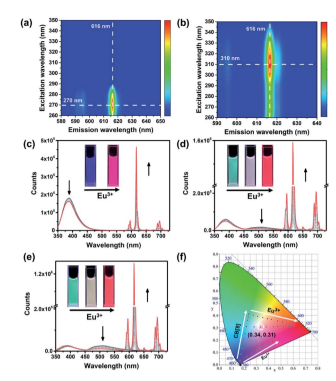


Fig. 3 Excitation–emission mapping of (a) Eu³+@DPA ([DPA] = 1.0×10^{-4} M and [Eu³+] = 3.3×10^{-5} M) and (b) Eu³+@PY-DPA \subset CB[8] complex in H2O ([PY-DPA] = 1.0×10^{-5} M, CB[8] = 5.0×10^{-6} M, and [Eu³+] = 3.3×10^{-6} M); prompt photoluminescence spectra of (c) PY-DPA and (d) PY-DPA with 0.5 equivalents of CB[8] and (e) 1.0 equivalent of CB[8] upon gradual addition 1/3 equivalents of Eu³+ in H2O. Inset: photograph showing the changes in luminescent color under light irradiation at 302 nm. (f) The 1931 CIE chromaticity diagram illustrating the luminescent color changes of DPA-PY with the addition of Eu³+ and CB[8] ($\lambda_{\rm ex}=310$ nm, 298 K).

lifetime at 510 nm remarkably decreased from 544 to 220 µs (Fig. S11†). Meanwhile, the excitation spectra of the Eu³⁺@PY-DPA⊂CB[8] complex monitored at 616 and 510 nm substantially coincided in the region above 290 nm, and the maximum peak was located at 311 nm, which was identical to the main absorption peak of the PY-DPA CB[8] complex. In clear contrast, the effective excitation of Eu³⁺@DPA monitored at 616 nm appeared in the region below 290 nm (Fig. S12†). Moreover, the excitation-emission mapping showed that the optimum excitation wavelengths for Eu³⁺@DPA and the Eu³⁺@PY-DPA CB[8] complex were 270 and 310 nm, respectively (Fig. 3a and b). The distinct redshift of the excitation wavelength validated the occurrence of energy transfer, which could facilitate the luminescence of the Eu³⁺ center at the longer excitation wavelength. It was noteworthy that the luminescence intensity of the Eu³⁺@PY-DPA⊂CB[8] complex at 616 nm was 3.6 times higher than that of Eu³⁺@PY-DPA under the same experimental conditions (Fig. 3c and d), and the lifetime and quantum yield of the Eu³⁺@PY-DPA \subset CB[8] complex ($\tau = 1.64$ ms and $\Phi =$ 21.1%) were significantly higher than those of Eu³⁺@PY-DPA (τ = 1.22 ms and Φ = 2.85%, Table S1†). These spectroscopic results jointly demonstrate that the complexation with CB[8] can not only induce the phosphorescence emission but also

enhance the photoluminescence efficiency of the lanthanide center *via* the energy transfer pathway.

To deepen the understanding of CB[8]-induced luminescence enhancement behaviors, we estimated the number of water molecules that coordinate to Eu³⁺ in the first coordination domain for the Eu³⁺(a)PY-DPA complex and Eu³⁺(a)PY-DPA⊂CB [8] assembly. The excited-state lifetime of Eu³⁺@PY-DPA was measured to be 1.22 ms in H₂O and 2.59 ms in D₂O, while the lifetime of Eu³⁺@PY-DPA \subset CB[8] was 1.64 ms in H₂O and 2.20 ms in D₂O (Fig. S13†). Accordingly, the coordination numbers of water molecules (q) were calculated to be -0.11 and 0.22 with and without CB[8], respectively, indicating that the coordination of water molecules was significantly inhibited in the Eu³⁺@PY-DPA⊂CB[8] assembly (Table S1†).46 The protection of Eu³⁺ from the quenching of water molecules was attributed to the formation of a close-packed supramolecular nanostructure in the case of the Eu³⁺@PY-DPA CB[8] assembly. Indeed, compared to the binary Eu³⁺@PY-DPA or PY-DPA⊂CB[8] complex, the Eu^{3+} @PY-DPA \subset CB[8] assembly showed an obvious Tyndall effect and a clear decline in transmittance, implying the existence of large-sized species (Fig. S14†). In addition, the intuitive morphology of the assembly was revealed by scanning electron microscopy (SEM) and transmission electron microscopy (TEM). The SEM and TEM images consistently showed that the Eu³⁺@PY-DPA CB[8] assembly exhibited a flake-like morphology, which probably contributed to the extensive intermolecular self-assembly of the supramolecular polymer (Fig. S15†).

Given that the Eu³⁺@PY-DPA⊂CB[8] assembly could emit three-primary-color (RGB) luminescence including blue fluorescence, green phosphorescence and red emission of lanthanide metal ions, multicolor photoluminescence outputs enclosed within the RGB color triangle could be obtained by rationally manipulating the host-guest and metal-ligand interactions. As depicted in the CIE 1931 chromaticity diagram (Fig. 3f), the luminescent color changed linearly from blue to pink with the stepwise addition of Eu³⁺ to free PY-DPA aqueous solution. Unexpectedly, when Eu³⁺ was continually added to the solution of the PY-DPA CB[8] complex, a linear change of luminescent color from cyan to red including the white emission was achieved with the chromaticity coordinate (0.34, 0.31). Moreover, benefitting from the CB[8]-induced fluorescencephosphorescence transition, the luminescent color range could be further expanded by changing the amount of CB[8]. For instance, when the molar ratio of CB[8] was fixed at 1 equivalent, the luminescent color of the PY-DPA CB[8] complex could be tuned linearly from green to red with the addition of Eu³⁺. On the foundation of these results, a multicolor emission hydrogel was conveniently fabricated by dispersion of agarose into the specifically emissive solution. As shown in Fig. 4a, four luminescence hydrogels with blue, green, red, and white emission could be intuitively recognized by the naked eye under UV light irradiation at 302 nm.

Considering that long-lived luminescent materials can significantly distinguish between background fluorescence and autofluorescence in living organisms, we wondered whether the Eu^{3+} @PY-DPA \subset CB[8] assembly with superior photophysical

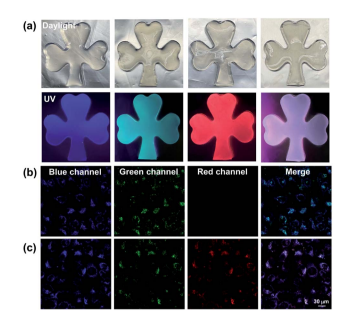


Fig. 4 (a) Photographs of multicolor hydrogels under daylight and 302 nm UV light irradiation; confocal microscopy images of HeLa cells treated with the (b) PY-DPA \subset CB[8] complex and (c) Eu $^{3+}$ @PY-DPA \subset CB[8] assembly. The excitation wavelength was set at 405 nm ([PY-DPA] = CB[8] $=1.0\times10^{-5}$ M, [Eu $^{3+}$] $=6.7\times10^{-7}$ M and scale bar $=30~\mu\text{m}$).

properties could be further applied in cell imaging. Then, HeLa cells were incubated with the PY-DPA CB[8] complex and Eu³⁺@PY-DPA⊂CB[8] assembly for 12 h, and then observed by confocal laser scanning microscopy. As shown in Fig. 4b, the cells treated with the PY-DPA CB[8] complex showed blue fluorescence and green phosphorescence upon excitation with a 405 nm laser. Besides, benefitting from the efficient energy transfer from PY⊂CB[8] to the Eu³⁺@DPA center, brighter red emission assigned to the millisecond-level Eu3+ complex was immediately found in the case of the Eu³+@PY-DPA⊂CB[8] assembly at the same excitation wavelength (Fig. 4c). In addition, the cell cytotoxicity of the Eu³⁺@PY-DPA⊂CB[8] assembly was evaluated by the cell counting kit-8 (CCK8) assay. It was found that more than 90% of the cells survived after being treated with the ternary assembly for 24 h, indicative of the excellent biocompatibility of the assembly (Fig. S16†). As we know, the potential cytotoxicity of metal complexes is considered one of the major obstacles limiting their practical applications. It is necessary to develop novel labelling methods using highly biocompatible metal complexes. Therefore, it is believed that our obtained Eu³⁺@PY-DPA⊂CB[8] assembly with tricolor emission and multidimensional excited-state lifetimes (ns-level blue, µs-level green, and ms-level red emission) will have promising potential in time-resolved multicolor bio-imaging.

Conclusions

In conclusion, a CB[8]-mediated lanthanide noncovalent polymer was successfully constructed *via* synergistic host-guest complexation and metal-ligand coordination, which was able to

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show RGB photoluminescence and cell imaging under a single wavelength excitation. The encapsulation by CB[8] could induce PY-DPA to form a stable homoternary inclusion complex, thus giving rise to dual fluorescence-phosphorescence emission in aqueous solution. Meanwhile, the resultant complex could further assemble with Eu³⁺ to generate the polypseudorotaxanetype supramolecular aggregate, accompanied by the significant enhancement of red emission via the complexation-assisted energy transfer process. Interestingly, multicolor emission including the unique white light could be produced in aqueous solution and the gel phase through simply adjusting the amounts of CB[8] and Eu3+. More significantly, the obtained supramolecular lanthanide assembly with a long lifetime and full-color emission could also be applied for multicolor cell imaging. Thus, we believe that the present work will bring about new enlightenment for the fabrication of multicolor lightemitting materials and high-resolution reliable bio-imaging probes.

Data availability

The data that support the findings of this study are available in the ESI.†

Author contributions

H.-J. Yu, Y.-M. Zhang and Y. Liu conceived and directed this study. H.-J. Yu conducted the synthesis, characterization, and analysis of all materials described with assistance from X.-L Zhou, F.-F. Shen, and Q. Zhou. The cell imaging experiment was conducted by Y. Dai. H.-J. Yu wrote this manuscript. Y.-M. Zhang, X. Xu and Y. Liu supervised the work and edited the manuscript. All authors discussed and reviewed the manuscript.

Conflicts of interest

There are no conflicts to declare.

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