Materials Advances



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Cite this: *Mater. Adv.*, 2021, 2, 6075

Received 10th August 2021, Accepted 13th August 2021 DOI: 10.1039/d1ma00709b

rsc.li/materials-advances

1. Introduction

The phenomenon and concept of the aggregation-induced emission (AIE) of organic compounds, proposed by Tang and co-workers in 2001,¹ have been attracting considerable research interest and it has shown extensive application prospects in many fields, including chemical sensing,² fluorescent sensors,³ and bioimaging.⁴ In the AIE process, AIEgens exhibit strong fluorescence emission in the aggregated state,^{5,6} which is different from the aggregation-caused quenching (ACQ) effect. Supramolecular materials,⁷ where the component molecules are combined together through noncovalent bonds, have been attracting great interest. In these systems, AIE has provoked much wholesale academic interest for innovative practical applications.

Regulation of the bi-color fluorescence changes of AIE supramolecular self-assembly gels by interaction with Al³⁺ and energy transfer[†]

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Supramolecular fluorescent materials have attracted considerable attention in recent years since they endow specific and unique properties to materials. Nevertheless, the utilization of photoresponsive characteristics to modulate their fluorescence emission behaviors and functions remain rarely explored. Here a facile fabrication strategy for producing dual-emissive materials based on supramolecular gels is proposed. A bi-acylhydrazone supramolecular gelator BD was designed and synthesized by a Schiff base reaction. Interestingly, the gelator BD could self-assemble into the stable supramolecular gel BDG with a strong aggregation-induced emission (AIE) in DMF-H₂O binary solutions *via* π - π stacking interactions. On the one hand, the BDG could selectively identify Al³⁺ in the gel state. Upon addition of Al³⁺, AIE BDG showed an obvious blue-shift (85 nm, from yellow-green to sky-blue). On the other hand, artificial light-harvesting systems were successfully fabricated in a gel environment based on the supramolecular strategy. In these systems, efficient energy transfer occurs between the BD assembly and the loaded acceptors. For instance, the transition from yellow-green to red light could be accomplished in the BDG/SR 101 system. Based on these, the manipulation of bi-color fluorescence emission was realized by interaction with Al³⁺ and energy transfer.

Supramolecular gels can be constructed by single noncovalent interactions or by the combination of multiple noncovalent interactions.⁸ By comparison, the dynamic and reversible properties of multiple noncovalent interactions endow supramolecular gels with an excellent response to various external stimuli, such as heat, light, pH, and metal cations.⁸ Moreover, the gel is a soft substance between a liquid state and a solid state, and a large amount of solvent molecules are fixed/wrapped by gelators, in which the solvent can contribute also. These characteristics provide a broad scope for the fabrication of smart materials and devices.

The recognition and sensing of metal ions have become the focus of considerable attention in biological, chemical, material, and environmental fields.^{9,10} Aluminum is the third most abundant of all the elements (after oxygen and silicon) in the earth's crust,^{11,12} accounting for 8.3% of the earth's total mineral components. However, a high aluminum intake can damage the central nervous system and is involved in Alzheimer's disease, Parkinson's disease, bone softening, chronic renal failure, and smoking-related diseases.¹³ Accordingly, the development of convenient and efficient testing methods for Al³⁺ is of great significance for environmental protection and human health. In this way, developing a fluorescent chemosensor for the selective recognition and monitoring of Al³⁺ in environmental systems or in living cells¹⁴ is essential.

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[†] Electronic supplementary information (ESI) available. See DOI: 10.1039/d1ma00709b

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Photosynthesis plays a vital role in the survival foundation of organisms, in which the large number of closely packed antenna pigments (*ca.* 200) around the reaction center is applied in the energy transfer and accumulation of sunlight.^{15–18} Several scaffolds have shown extraordinary results in mimicking the natural light-harvesting process by achieving efficient energy transfer from donors to acceptors through a Förster resonance energy transfer (FRET) process,^{19,20} such as protein assemblies,²¹ dendrimers,²² metal complex polymers, and porphyrin assemblies. These artificial light-harvesting systems/materials are of significant importance for practical applications, which are still less emphasized in the gel state to date.²³

Fluorophores with aggregation-induced emission properties are good candidates for the construction of fluorescent supramolecular systems. However, supramolecular gels with tunable emission have been rarely reported, although some progress has recently been made on the construction of discrete fluorescent supramolecular assemblies.²⁴ Traditionally, more than one kind of fluorophores should be incorporated to construct this kind of gel, thus resulting in complexity in the designed systems. Besides this method, the emission behavior could also be tuned by the introduction of interactions with ions, hostguest interactions, or other forms of interaction,^{9,10,23} resulting in a change in the emission wavelength. Herein, we report a biacylhydrazone compound, which was found to exhibit competitive guest stimuli responsiveness in the gel state and AIE properties. The stable organogel (BDG) showed a selective and sensitive stimuli-response for Al³⁺ but the BD in solution could not detect Al³⁺ in our fluorescence analysis experiments. Moreover, efficient light-harvesting systems (LHSs) were successfully fabricated using the organogel BDG as an energy donor, and a series of dyes, such as sulforhodamine 101, acridine red, rhodamine B, and rhodamine 6G, were utilized as energy acceptors, respectively. Additionally, the fluorescence color and emission spectrum of BD could be effectively tuned by metal ion coordination or by fabricating light-harvesting systems. As far as we know, this is the first report that connects self-assembly, AIE activity, light-harvesting, and ion recognition through a multifunctional bi-acylhydrazone derivative (Scheme 1).

2. Experimental

2.1. Materials and general methods

For the materials, 2-(2-hexyl-1H-benzimidazole-1-yl)acetohydrazide, terephthalaldehyde, organic solvent and metal salt were purchased from commercially available sources and were used without further purification. Deionized water was used for all the experiments. The measurements of steady-state luminescence were performed with a Shimadzu RF-5301PC spectrometer and spectrofluorophotometer (F-4500, Japan). Ultraviolet-visible (UV-vis) spectra were recorded on a Shimadzu UV-1750 spectrometer. The infrared spectra were recorded on a Thermo Scientific Nicolet iS5 FT-IR spectrophotometer. Fluorescence micrographs (FOM) of the samples were obtained with an Olympus IX 71 system. The ¹H and ¹³C NMR spectra were recorded on a Bruker 400 MHz spectrometer. The internal standard was TMS and the solvent was dimethylsulfoxide $(DMSO-d_6)$. Time-resolved photoluminescence decay measurements were carried out using a time-correlated single-photon counting (TCSPC) spectrometer (Edinburgh, FLS 980). Mass spectra were obtained on a Bruker solanX 70 FT-MS mass spectrometer equipped with an ESI interface and ion-trap analyzer. The morphologies of the as-synthesized samples were characterized with an SM-74190UEC SEM system using an accelerating voltage of 10 kV. The quantum yield was measured on an FLS980 fluorescence spectrometer. X-ray diffraction patterns (XRD) were determined with a X'Pert PRO diffractometer using Cu-K_{α} radiation over the 2 θ range of 5–90°.



Scheme 1 Schematic representation for the bi-color fluorescence changes of a bi-acylhydrazone compound.

2.2. Synthesis of the compounds

As shown in Scheme 2, 2-(2-hexyl-1*H*-benzimidazole-1-yl)acetohydrazide (6.855 g, 25 mmol) and terephthalaldehyde (1.341 g, 10 mmol) with a molar ratio of 2.5:1 were dissolved in DMF (100 mL). The mixture was stirred under reflux at 80 °C for 8 h with acetic acid added (2–3 drops). After the reaction was finished and the solution cooled to room temperature, the precipitate was collected by filtration, washed three times with hot absolute ethanol, and then dried in a vacuum; giving a faint yellow powder compound BD (5.030 g, yield: 61%). ¹H NMR (400 MHz, DMSO-*d*₆, Fig. S1, ESI†) (ppm): 11.88 (d, 2H, N–H), 8.31–8.11 (d, 2H, N=CH), 7.94–7.15 (m, 12H, ArH), 5.49–5.03 (d, 1H). ESI-MS: *m/z* 647.4 ([C₃₈H₄₆N₈O₂] + H)⁺ (Fig. S3, ESI†).

2.3. Fluorescence measurements

Various nitrate solutions containing Na⁺, K⁺, Mn²⁺, Ni²⁺, Zn²⁺, Pb²⁺, Cu²⁺, Cr³⁺, Fe³⁺, Hg²⁺, Cd²⁺, Co²⁺, Mg²⁺, and Al³⁺ of 1 mol L⁻¹ were used for testing the fluorescence performance of the gel BDG toward metal ions. All the fluorescence spectra experiments were implemented in a DMF-H₂O (1:4, v/v) binary system. Any fluorescence changes of the compounds were recorded upon the addition of nitrate salt, while the gelation concentration of the supramolecular gel was 0.5% in all the experiments. To further study the sensitivity of the gel BDG for Al³⁺ detection, the gelator BD in DMF and various concentrations of Al³⁺ aqueous solution were mixed intensively before measurement.

3. Results and discussions

3.1. Formation of the gel BDG

First of all, to exam the assembly ability and gelation properties of the gelator BD, we configured BD in a variety of polar, nonpolar organic, and component solvents (organic solvents– H_2O), respectively, in the proportion of 5% and heated it completely. After cooling to room temperature, we found that BD could





Fig. 1 (a) FE-SEM micrographs of the xerogel of BD. The gel was two days old before the images were taken. (b) FOM images of the BDG (0.5%, V_{DMF} : $V_{\text{water}} = 1:4$).

self-assemble into a stable gel only in DMF, DMF-H₂O binary solution, DMSO, and DMSO-H2O binary solution. The successful preparation of the gel BDG was verified via a "stabilized reverse tube" method and the results are summarized in Table S1 (ESI[†]). As a general procedure, the compound BD (0.015 mmol) was completely dissolved in DMF (0.4 mL), then H₂O (1.6 mL) was added, dissolved completely by mixing as a hot solution, and finally the mixture was cooled to ambient temperature. As a vital parameter of supramolecular gel, we investigated the effect of different water contents on the assembly ability of the gel (Fig. S4, ESI⁺), which was to play an important role when we were choosing the solvent composition of BD gel (V_{DMF} : $V_{\text{water}} = 1:4$, Fig. S5, ESI[†]). In the DMF– H₂O binary system, with a water content less than 67.5%, the gel of BD could not be obtained. Meanwhile, we also conducted partial tests for DMSO (Table S1, Fig. S6 and S7, ESI⁺). By comparison, BD dissolved more quickly in DMF than in DMSO. The lowest critical gelation concentration (CGC) in DMF was 2.5% (w/v%, 10 mg mL⁻¹ = 1.0%) and the gel-sol transition temperature $(T_{
m gel})$ in DMF was 74 °C. More interestingly, the CGC of the BD in DMF-H₂O (1:4, v/v) was 0.5% and the T_{gel} was 95 °C in DMF-H₂O (1:4, v/v). Hence, the DMF-H₂O (1:4, v/v) binary system was determined to be optimal for the formation of the gel BDG. As shown in Fig. 1a, the morphology structure of the BDG was further investigated by scanning electron microscopy (SEM), and was observed to be in a homogeneous uniform state. We could find that the large fibers exhibited some fine structure and appeared to be multi-layered. 3D entangled network structures comprising fibers with high aspect ratios were obtained, revealing well-ordered molecular packing properties. The fibers lacking twisting and regular shapes existed with widths of 200-500 nm and lengths of dozens of micrometers. As the FOM (fluorescent optical microscopy) images show, the morphology of BDG comprised dense aggregated bundles of fibers (Fig. 1b). Interestingly, a significantly yellow-green light could be achieved in the BDG.

3.2. AIE behaviors

As shown in Fig. 2, the solution of BD had inappreciable fluorescence. Also, while the emission intensity emerged rapidly, and reached a steady state finally at $\lambda = 520$ nm, in the process, the temperature of the heated DMF/water solution declined under that of the T_{gel} ($T < T_{gel}$) of BDG, which



Fig. 2 Temperature-dependent fluorescent spectra of BDG (V_{DMF} : $V_{water} = 1:4, 0.5\%$) during the gelation process ($\lambda_{ex} = 372$ nm).

indicated that the fluorescence of the supramolecular gel BDG was aggregation-induced emission (AIE), accompanied by vellow-green fluorescence. In order to indicate that this was not a purely thermal effect, we did another experiment as supplementary proof of the AIE properties of the BD gel (Fig. S8, ESI[†]). From the experiment, we could see that BD powder (25 °C) showed a yellow-green light under UV light, corresponding to the description in Fig. 2. However, this phenomenon did not exist in the BD powder (25 °C) under daylight or BD in DMF $(1 \times 10^{-4} \text{ M}, 25 ^{\circ}\text{C})$ under UV light. This means that the molecule BD presented an interesting aggregation-induced emission phenomenon. BD was insoluble in water, and the fluorescence of BD in the aggregation state was also studied in DMF/water mixtures with different water fractions to investigate its solvent-dependent aggregation behavior (Fig. S9, ESI⁺). The emission peak at 430 nm of BD (3 \times 10⁻⁴ M) was very weak in DMF and almost kept constant until the water fraction reached 30%. The fluorescence intensity of BD showed a remarkable enhancement caused by the intriguing AIE effect owing to the formation of aggregates when the water volume reached 40%. Meanwhile, it exhibited an apparent red-shift (from λ = 430 nm to 520 nm) with the increment of the water content. At about 70% water, the maximal emission intensity was achieved, which was almost 7-fold greater than that in dilute solution. In the DMSO/water mixtures with different water fractions, BD showed a similar phenomenon (Fig. S10, ESI[†]).

3.3. Interaction of Al³⁺

Investigations of BDG to assess the influence of metal ions on it in mixed solutions were performed (Fig. S11, ESI[†]). The addition and diffusion of 10.0 equiv. of various metal ions (Na⁺, K⁺, Mn^{2+} , Ni^{2+} , Zn^{2+} , Pb^{2+} , Cu^{2+} , Cr^{3+} , Fe^{3+} , Hg^{2+} , Cd^{2+} , Co^{2+} , Mg^{2+} , and Al^{3+} using their nitrate salts as solutions) to BD generated the corresponding metallogels (such as BDG-Al). In the fluorescence spectrum, upon the addition of 10.0 equiv. Al^{3+} to the BDG in the process of gel formation, the emission of BDG showed an obvious blue-shift (Fig. 3), which indicated that Al^{3+} could be clearly detected using BDG. Considering the above interesting blue-shift in the fluorescence spectrum, the effects of various concentrations of Al^{3+} on the fluorescence spectrum



Fig. 3 Fluorescence spectra (λ_{ex} = 372 nm) of BDG (0.5%, V_{DMF} : V_{water} = 1:4) before and after the addition of Al³⁺ (10.0 equiv.). Inset: Photographs showing the fluorescence change of BDG and BDG-Al³⁺ under illumination at 365 nm.

of BDG were investigated in the gel state (Fig. S12, ESI[†]). When the concentration of Al³⁺ was further increased, a blue-shift of 85 nm emerged; meanwhile, the original peak disappeared. Then the fluorescence intensity of BDG at 435 nm was enhanced gradually with the continuing increase in the Al³⁺ concentration. The fluorescent selectivity of BDG toward Al³⁺ may be attributed to the smaller ionic radius (0.5 Å), which allowed a suitable coordination geometry with BD and a higher charge density that made Al³⁺ coordinate to BD strongly.²⁵ In the presence of the Al³⁺ ion, the carbonyl O and acylhydrazone N could coordinate with the Al³⁺ central atom, which increased the energies of the n-p* transitions compared to the corresponding p-p* transitions, and finally the PET process was interrupted, and the fluorescence changed from yellow-green to blue emission. As the Job's plot experiments showed, we could see an obvious peak at 3:7 of BDG–Al³⁺, assignable to the best composition ratio of BDG-Al³⁺ (1:2) (Fig. 4a). In terms of the performance of identifying Al³⁺, for example, Fig. 4b shows an investigation in which various metal cations (including Na⁺, K⁺, Mn²⁺, Ni²⁺, Zn²⁺, Pb²⁺, Cu²⁺, Cr³⁺, Fe³⁺, Hg²⁺, Cd²⁺, Co²⁺, Mg²⁺, and Al³⁺) were put into dot arrays of BD-gel, but only Al³⁺ gave BDG a bright sky-blue color under UV light at 365 nm.

In addition, the response mechanism of the BDG for Al³⁺ was further investigated by ¹H NMR titration, IR, and XRD. In the ¹H NMR titrations of Al³⁺ for BD (Fig. 5) with the increasing concentration of Al³⁺, the N-H (H¹) signals showed a downfield shift. When adding proper Al^{3+} , the weakness of the N-H(H¹) signal at δ 11.9 ppm indicated that the deprotonation process had occurred. In the meanwhile, the hydrogen bond existing in the acylhydrazone group was destroyed during the coordination of Al³⁺ with the carbonyl group of BD. Additionally, at low concentration of BD, =CH (H²) on BD appeared and its signals showed a downfield shift with the increasing concentration of Al³⁺. Therefore, a possible sensing mechanism based on deprotonation was proposed, as given in Fig. S13 (ESI[†]). The XRD patterns of free BDG and BDG-Al (Fig. S14, ESI⁺) showed *d*-spacings of 3.52 Å and 11.45 Å at $2\theta = 25.23^{\circ}$ and 7.71°, supporting the presence of π - π stacking and hydrophobic



Fig. 4 (a) Job's plots according to the method for continuous variations, indicating the 1:2 stoichiometry for BDG-Al³⁺ (the total concentration of BD and Al³⁺ was 1×10^{-3} mol L⁻¹). (b) Fluorescence responses of the BD-gel-based sensor array to the presence of various cations under UV at 365 nm. In gels, the concentration of BD in the sample was 0.5 wt%.



Fig. 5 Partial ¹H NMR spectra (400 Hz, 298 K) of BD (2.0×10^{-2} M) in DMSO- d_6 with increasing amounts of Al³⁺ (0.1 M, DMSO- d_6), (a) free BD; (b) 1.0; (c) 2.0; (d) 5.0 equiv.

interaction in the BD. In the meantime, *d*-spacings of 3.51 Å and 3.80 Å at $2\theta = 25.37^{\circ}$ and 23.34° suggested that the π - π stacking remained in the metallogel BDG-Al. FT-IR spectra of the powder gelator BD and the xerogel BDG-Al were obtained to deeply understand the coordination of Al³⁺ and the gelator BD (Fig. S15, ESI[†]). The bands at 1695 and 1596 cm⁻¹ corresponding to V (C=O) and V (C=N) were revealed to be present in the IR spectrum of the gelator BD. However, with the addition of 2.0 equiv. of Al³⁺ and the formation of the Al³⁺ coordinated metallogel BDG-Al, the V (C=N) band was shifted

to a lower wavenumber (1562 cm^{-1}), respectively, due to coordination with the metal center.

3.4. Light-harvesting ability

Due to the unique AIE effect, compound BD in the aggregated form exhibited a broad emission at 520 nm, which enables BD to act as a remarkable donor in gels. The potential fluorescence resonance energy transfer (FRET) between AIE (BDG) and various acceptors was next investigated. Sulforhodamine 101 (SR 101), acridine red, rhodamine 6G (Rh 6G), and rhodamine B (Rh B) were selected as energy acceptors, respectively, since they show strong absorption in the visible region and overlap with the emission from BDG. As shown in Fig. 6a, and Fig S16a, S16c, and S16e (ESI⁺), the absorption band of these dve molecules overlapped well with the fluorescence emission of the BD assembly. On the other hand, a key parameter in facilitating an efficient energy-transfer process is spatially well-organized chromophores. This could be realized in between the BD assembly and the dyes. As shown in Fig. 6b, with the gradual addition of dye SR 101, the fluorescence intensity of BDG decreased significantly, while the fluorescence emission band of SR 101 appeared and increased when excited at 372 nm, accompanied with a change in the emission color from green/yellow to red under a UV lamp.

The fluorescence quantum yield of the BDG/SR 101 system was estimated to be 13.28% (Fig. S17a, ESI†). According to



Fig. 6 (a) Normalized emission spectrum of BDG (7.7 mM), absorption spectrum of SR 101 (1 \times 10⁻⁵ M). (b) Fluorescence spectra of BDG (7.7 mM) in the gel with different concentrations of SR 101 (38.5 μ M, λ_{ex} = 372 nm, Inset: photographs of BDG, BDG/SR 101 under UV light).

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fluorescence titration spectrum, the energy-transfer efficiency was calculated to be 80% (Fig. S18a, ESI[†]) at a donor/acceptor ratio of 200:1. Upon the gradual addition of acridine red, Rh 6G, and Rh B as acceptors to the BD assembly, respectively, the emission peak of BDG began to decay while the emissive intensity of the acceptors kept increasing (Fig S16b, S16d, and S16f, ESI[†]). According to fluorescence titration spectrum, the energy-transfer efficiencies ($\Phi_{\rm ET}$) of acridine red, Rh 6G, and Rh B as acceptors were calculated to be 98%, 99%, and 98% (Fig S18b, S18c, and S18d, ESI[†]). The fluorescence quantum yields of the acridine red, Rh 6G, and 7.51% (Fig. S17b, S17c, and S17d, ESI[†]).

Furthermore, to clearly show the energy transfer (ET) from BDG to the acceptors during the gelation process, we measured the fluorescence spectra of gelation in energy-transfer systems (Fig. S19, ESI[†]). In view of the rapid gelation of BD, we performed the experiment at lower concentrations. Primarily, when adding SR 101 into BDG (molar ratio of BDG: SR 101 was 3000:1) in DMF/H₂O solution, the fluorescence spectrum of BDG/SR 101 showed a continuous significant increase at 610 nm during the gelation process (Fig. S19a, ESI[†]). In addition, compared with the emission spectrum of BDG (Fig. S20, ESI[†]) at the same concentration, the fluorescence intensity of BDG/SR 101 at 515 nm after 10 min was only half as much as the intensity of BDG. Finally, the morphologies of BDG/SR 101, BDG/acridine red, BDG/Rh 6G and BDG/Rh B were investigated by scanning electron microscopy (SEM) (Fig. S21, ESI⁺). The SEM images of the light-harvesting systems showed a number of micron rod-like structures. The transformation from network structures into stick structures indicated that the dye molecules could create an interaction with BDG to change the previous intramolecular interactions; thereby the formation of a rod-like supramolecular structure was observed. Subsequently, the fluorescent optical microscopy (FOM) images further truly reflected the morphology and luminescence phenomenon of the light-harvesting systems (Fig. S22, ESI⁺).

4. Conclusion

In conclusion, we designed a novel organogelator BD that could be self-assembled into an organogel BDG showing strong AIE. The supramolecular gel BDG can serve both as a selective fluorescence sensing system for Al^{3+} and light-harvesting energy donor with several kinds of dyes through the FRET mechanism. Notably, the supramolecular coordination complexes BDG- Al^{3+} formed by the metal-coordination-driven self-assembly emitted from yellow-green to sky-blue as the concentration of Al^{3+} increased. The fluorescence emission from yellow-green to red was also obtained *via* the host-guest interactions in artificial light-harvesting systems. This simple bi-acylhydrazone demonstrates an example of a multifunctional light-emitting material including AIE activity, lightharvesting abilities, sensing properties, and so on.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by the National Natural Science Foundation of China (No. 21961029), Key Research Foundation of Department of Science and Technology of Ningxia, China (No. 2019BDE03019, 2018BEG03020), The Natural Science Foundation of Ningxia, China (No. 2020AAC02031); CAS "Light of West China" (No. 2017-32), The Science and Technology Foundation of Guyuan, China (No. 2019GKGY025), Research Award Fund for First-class Discipline Construction (Education Discipline) in Higher Education Institutions of Ningxia, China (No. NXYLXK2017B11), The Foundation of Ningxia Normal University (No. NXSFZDA1905), Engineering Research Center of Liupanshan, School Chemistry and Chemical Engineering of Ningxia Normal University (No. HGZD21-02), are gratefully acknowledged.

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