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Strong Circularly Polarized Luminescence from the Supramolecular Gels of an Achiral Gelator: Tunable Intensity and Handedness

Zhaocun Shen, Tianyu Wang, $*^a$ Lin Shi, Zhiyong Tang and Minghua Liu $*^{ab}$

Although the importance of circularly polarized luminescence (CPL) materials has been widely recognized, the CPL responses of supramolecular gels are still rarely studied. Moreover, developing CPL materials based on supramolecular gels is of great significance, due to their special advantages and important applications. Herein, we report the first circularly polarized supramolecular gels self-assembled exclusively from a simple achiral C_3 -symmetric molecule. Most importantly, the excellent tunability of these novel CPL materials, which benefits from the achiral molecular building blocks as well as the nature of supramolecular gels, has been investigated. Thus, the CPL intensity of these supramolecular gels is easily enhanced by mechanical stirring or doping chiral amines. And the handedness of CPL signals is controlled by the chirality of organic amines.

Introduction

Recently, developing the materials exhibiting circularly polarized luminescence (CPL) has attracted more and more interest, not only for understanding the mysteries of chirality, 2 but also due to the important potential applications of CPL materials. Chiral molecular systems showing differential emission of right-handed and left-handed circularly polarized light have been explored to be used as display devices,³ optical storage devices, 4 CPL sensors, 5 biological probes 6 and even catalysis for asymmetric photochemical synthesis. Although various chiral molecular systems, such as chiral metal complexes, 8, 1h chiral polymers, 9 liquid crystals 10, 1e and few chiral organic molecules in solution^{11, 1b} or in solid state¹² have been found to be CPL candidates, the supramolecular gels with the ability to emit circularly polarized light are still rarely constructed. Actually, the supramolecular gels based on the self-assembly of small organic molecules are very important soft matters with many exceptional advantages, such as flexibility, biocompatibility and easy processing. 13 Therefore, for the application of CPL systems, the supramolecular gels emitting circularly polarized light could be of special significance.

In addition, the known organic molecules showing CPL are usually chiral molecules with complicated but specific structures, for example chiral helicenes, the liberature of th

capability of emitting strong CPL signals becomes extremely important for constructing novel CPL devices.

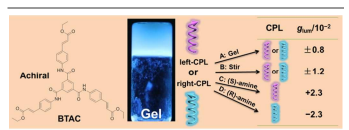


Fig. 1 First circularly polarized supramolecular gels self-assembled exclusively from a simple achiral C_3 -symmetric molecule (BTAC). The amplitude of CPL signals can be further enhanced upon stirring or adding chiral amines, and the handedness of CPL signals can be controlled well by the chirality of organic amines

We also notice that almost all of the CPL materials are based on chiral molecular systems. The only known exception from the literature is the co-assembly of an achiral ionic polymer and Rhodamine B, which shows CPL under mechanical stirring. Our previous study has demonstrated that the simple achiral C_3 -symmetric molecules (Tris(ethyl cinnamate) Benzene-1,3,5-tricarboxamides , BTAC) can self-assemble into supramolecular gels with intense circular dichroism (CD) signals (Fig. 1). In this paper, we present supramolecular gels that show excellent CPL performance (luminescence dissymmetry factor, $g_{lum} = \pm 0.8 \times 10^{-2}$), even though these gels are composed of identical achiral organic molecules. Most importantly, both the intensity and handedness of CPL signals of these gels can be easily modulated. Upon

Chemical Science Page 2 of 6

ARTICLE Journal Name

mechanical stirring, the CPL intensity can be enhanced; while adding some simple chiral dopants not only can enhance the amplitude of CPL signals but also can readily control the handedness of CPL signals (Fig. 1).

Results and discussion

CPL activity of BTAC gels

BTAC is a C_3 -symmetric molecule with three aromatic ethyl cinnamate groups connected to a benzene ring via amide bond (Fig. 1). For the self-assembly of BTAC, we propose that both π - π interactions from aromatic rings and hydrogen bonding from the amide groups play very important roles. 16 Achiral BTAC molecules could form supramolecular gels in DMF/H₂O mixture (v/v: 5/2) with spontaneous symmetry breaking, as proved by the unequal amount of left- and righthanded twists and the strong circular dichroism (CD) signals.¹⁶ In this case, supramolecular chirality can be obtained from the gelation of achiral BTAC molecules. While the handedness of these chiral assemblies can be left-handed or right-handed by chance. Moreover, when BTAC gels in DMF/H2O mixture (v/v:5/2) are irradiated by 330 nm light, the strong fluorescence with emission maximum at 448 nm is detected (Fig. 2b), even though BTAC molecules themselves have very week fluorescence in DMF solution. The absolute fluorescence quantum yield ($\Phi_{\rm F}$) of BTAC solution in DMF is 0.003, while the BTAC gels have very strong fluorescence with $\Phi_{\rm F}$ equal to 0.105. significant gelation-induced fluorescence enhancement encouraged us to investigate CPL response of these supramolecular gels which consist of only achiral molecular building blocks.

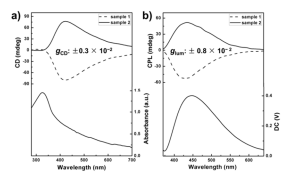


Fig. 2 (a) CD (left axis) and UV-Vis (right axis) spectra of BTAC gels (1% wt/vol) in DMF/ H_2O (v/v: 5/2); (b) CPL spectra (left axis) and fluorescence spectra (right axis) of BTAC gels (1% wt/vol) in DMF/ H_2O (v/v: 5/2) excited at 330 nm. The sample with negative Cotton effect displays right-handed CPL (dash curves), while the sample with positive Cotton effect exhibits left-handed CPL (solid curves).

Amazingly, from different batches of BTAC gels in DMF/H₂O mixture (v/v: 5/2), strong CPL signals with different handedness and maximum at 427 nm can be observed (Fig. 2b and Fig. S1), highlighting excited-state supramolecular chirality of these assemblies formed by achiral molecular building blocks. In contrast, no CPL signals can be detected from BTAC solution (Fig. S2). For understanding the relationship between

the ground-state supramolecular chirality and excited-state supramolecular chirality of BTAC gels, the correlation between the CD signs and CPL signs was studied. The results reveal that the samples with positive Cotton effect display left-handed CPL, while the samples with negative Cotton effect display right-handed CPL (Fig. 2).

The magnitude of the circular polarisation at the ground state is defined as $g_{CD} = 2 \times (\varepsilon_L - \varepsilon_R)/(\varepsilon_L + \varepsilon_R)$, where ε_L and ε_R refer to the extinction coefficients for left- and right- handed circularly polarised light, respectively. Experimentally, the value of g_{CD} is defined as $g_{CD} = [ellipticity/32,980]/absorbance$ at the CD extremum. The magnitude of CPL can be evaluated by the luminescence dissymmetry factor (g_{lum}) , which is defined as $g_{\text{lum}} = 2 \times (I_L - I_R)/(I_L + I_R)$, where I_L and I_R refer to the intensity of left- and right-handed CPL, respectively. The maximum g_{lum} value ranges from +2 for an ideal left CPL to -2 for an ideal right CPL, while $g_{lum} = 0$ corresponds to no circular polarization of the luminescence. Experimentally, the CPL were measured from JASCO CPL-200 spectrometer, and the value of is defined g_{lum} g_{lum} [ellipticity/(32,980/ln10)]/total fluorescence intensity at the CPL extremum.

Although BTAC gels were formed by identical achiral molecules, the absolute value of dissymmetry factor ($|g_{lum}|$) of their CPL signals can be about 0.80×10^{-2} (Fig. 2b), which is even much larger than those $|g_{lum}|$ values of some chiral organic molecules in solution or in solid state. To the best of our knowledge, this is the first example that a supramolecular gel, prepared by only small achiral molecules, exhibits CPL feature without any chiral additives.

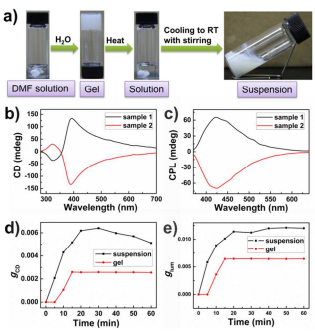
Stirring-induced CPL enhancement

It has been reported that the vortex stirring would lead to symmetry breaking and produce supramolecular chirality for some supramolecular assemblies formed by achiral molecules.17 More importantly, the handedness supramolecular chirality of some systems, such as the porphyrin assembly in solution, can be controlled by changing the direction of stirring.17 In contrast, for BTAC gels, the change from achiral molecules to chiral supramolecular assemblies is established simply upon gelation, not necessary to external mechanical force. In this context, it is also interesting to evaluate the effect of vortex stirring on the chirality of BTAC assemblies. Can mechanical force manipulate the symmetry breaking from gelation and finally decide the handedness of supramolecular chirality?

The BTAC gels formed in DMF/ H_2O (v/v: 5/2) are thermally reversible, and the vortex stirring can be introduced into the system during the gelation process (Fig. 3a). The self-assembly of BTAC under stirring was performed in a 5 mL vial containing a 5.0×10.0 mm Teflon-coated magnetic stirring bar at the bottom, and the clockwise (CW) or counterclockwise (CCW) stirring at 900 rpm was applied during the sol-gel process. The CD and CPL spectra of BTAC assemblies before and after vortex stirring with different directions were studied in detail.

ARTICLE

Page 3 of 6 Chemical Science



Journal Name

Fig. 3 (a) Photographs showing BTAC assemblies (1% wt/vol) in DMF/H₂O (v/v: 5/2) upon 900 rpm stirring during gelation; (b, c) CD spectra (b) and CPL spectra (c) of BTAC assemblies (1% wt/vol) after 900 rpm clockwise stirring during gelation; the sample with negative Cotton effect displays right-handed CPL (red curve), while the sample with positive Cotton effect shows left-handed CPL (black curve); (d, e) changes of $g_{\rm CD}$ (d) and $g_{\rm lum}$ (e) of BTAC gels (red curves) (1% wt/vol) and suspensions under 900 rpm stirring during gelation (black curves) in DMF/H₂O (v/v: 5/2); suspensions under 900 rpm stirring during gelation show much larger dissymmetry factor than that of BTAC gels without stirring.

Interestingly, although the gels are broken into suspensions upon long time stirring during gelation, the CD and CPL signals of BTAC assemblies could always be detected readily. The results show that BTAC suspensions show stronger CD and CPL signals after stirring (Fig. 3b, 3c). The strong CD signals with negligible linear dichroism (LD) artefacts are also confirmed by LD spectrum (Fig. S3). The SEM image of the BTAC assemblies after stirring also shows chiral twists, which are similar to the nanostructures of as-prepared BTAC gels (Fig. 4).

Most impressively, compared with the case of BTAC gels without stirring, both the CD signals and CPL activity of BTAC assemblies are found to be enhanced after stirring, as shown in Fig. 3 and Fig. S4. Thus, if as-prepared BTAC gels have positive Cotton effect and left-handed CPL, respectively, they will have more intensive positive Cotton effect and left-handed CPL after stirring, whether the corresponding stirring is clockwise (CW) or counterclockwise (CCW) (Fig. S5). Thus, it is interesting to note that in our case, the handedness of the supramolecular chirality cannot be controlled by stirring direction. Once the handedness has been decided by spontaneous symmetry breaking on the earlier stage of selfassembly, it would not be changed by macroscopic mechanical force from outside. The vortex stirring cannot change the interactions between different molecules. However, the stirring is able to enhance the certain signals. Presumably it is because that vortex stirring could promote the orderly arrangement of supramolecular assemblies around nanoscale. This case can be proved by the scanning electron microscopy (SEM) images, in which many large assemblies can be detected after stirring (Fig. 4). Moreover, the unchanged molecular packing mode of BTAC assemblies before and after stirring can be demonstrated by the X-ray diffraction (XRD) patterns (Fig. S6).

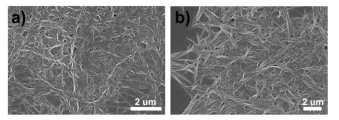


Fig. 4 SEM images of BTAC gels (a) and the corresponding BTAC suspensions after 900 rpm clockwise stirring during gelation (b) in DMF/H₂O (v/v: 5/2).

It is worth mentioning that stirring has to be applied during the sol-gel process to enlarge the supramolecular chirality of BTAC assemblies. Stirring after gelation cannot increase the intensity of the corresponding CD and CPL signals (Fig. S7).

further understanding the stirring-enhanced supramolecular chirality, we altered the stirring speeds during the self-assembly process, and the CD spectra of different samples after diverse stirring speeds were recorded. Interestingly, very slow stirring rates (300 rpm) could decrease the g-factor (Fig. S8. Perhaps, slow stirring rates cannot promote the orderly arrangement of nanostructures. However, the enhancement of supramolecular chirality can be clearly detected when the stirring speed is further increased (Fig. S8). Thus, the 900 rpm stirring has been selected for general investigation. The influence from the stirring time is also explored. Upon long time stirring, BTAC assemblies would become suspensions with good dispersion (Fig. S9). The results show that both the CPL and CD signals come to the maximum after about 30 minutes stirring at 900 rpm. However, further prolonging the stirring time does not improve the supramolecular chirality further (Fig. 3d, 3e). Compared with the change of CD intensity of BTAC gels without stirring (Fig. 3d), the enhancement of supramolecular chirality by mechanical force is remarkable. It is worth mentioning that very high luminescence dissymmetry factor $(g_{lum} = 1.2 \times 10^{-2})$ is achieved upon long time stirring (Fig. 3e). Therefore, the CPL materials constructed by supramolecular gels have a very nice tunable property. Altogether, CPL supramolecular gels are fabricated by simple achiral C_3 -symmetric molecules, and the intensity of the corresponding CPL signals can be further enhanced via mechanical stirring.

Chiral amine-induced enhancement and control of CPL

It has been reported that the chirality of supramolecular assemblies can be controlled by chiral additives.¹⁹ We have found that chiral organic amines have the capability to inherit their molecular chirality to the supramolecular assemblies

Chemical Science Page 4 of 6

ARTICLE Journal Name

formed by achiral BTAC molecules and therefore control the handedness of these chiral assemblies.¹⁶

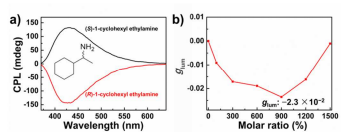


Fig. 5 (a) CPL spectra of BTAC gels (1% wt/vol) containing 900 mol% chiral 1-cyclohexyl ethylamine in DMF/H₂O (v/v: 5/2); (b) the dissymmetry factor (g_{lum}) of BTAC gels with different molar ratios of (R)-1-cyclohexyl ethylamine/BTAC.

With regard to the supramolecular chirality of BTAC assemblies on the excited-state optical activity, addition of the chiral amines not only controls the handedness of CPL but also enhances the amplitude of CPL. For the BTAC gels containing (R)-1-cyclohexyl ethylamine with a molar ratio of BTAC/amine = 1:9, a strong right-handed CPL signal was detected (Fig. 5a). In contrast, in the case of BTAC gels containing (S)-1cyclohexyl ethylamine, the mirror-imaged CPL spectrum was discerned (Fig. 5a). Remarkably, the dissymmetry factor (g_{lum}) increases greatly to $\pm 2.3 \times 10^{-2}$ with doping chiral organic amines. Therefore, both the intensity and handedness of CPL signals can be easily modulated. On the other hand, although a higher concentration of the chiral amine in the system could result in higher CPL activity, overloaded chiral components could also destroy the assembly and decrease the intensity of CPL peaks (Fig. 5b).

Conclusions

In conclusion, the supramolecular gels with strong CPL responses have been fabricated from very simple achiral C_3 -symmetric gelator. Moreover, the excited-state supramolecular chirality of these materials can be controlled perfectly. Thus, the luminescence dissymmetry factor (g_{lum}) can be enhanced largely by simply using mechanical stirring or adding chiral amine dopants, which is also useful for controlling the handedness of CPL signals. Considering the explosive growth of demand for developing novel CPL materials, our work opens new strategies to achieve low-cost CPL supramolecular gels, which is easy to be fabricated into different devices with desirable flexibility and biocompatibility. Most of all, our results also firstly demonstrate that the excited-state supramolecular chirality can also be obtained from the self-assembly of identical small achiral molecules.

Experimental section

Instruments and methods

Scanning electron microscopy (SEM) was performed on a Hitachi S-4800 FE-SEM with an accelerating voltage of 10 kV. Before SEM measurements, the samples on silicon wafers were

coated with a thin layer of Pt to increase the contrast. UV-Vis, CD and LD spectra were obtained using JASCO UV-550 and JASCO J-810 spectrometers, respectively. CPL measurements were performed with a JASCO CPL-200 spectrometer. 0.1 mm cuvettes were used for measuring the UV-Vis, CD, LD and CPL spectra of samples. For the measurement of CD spectra, the cuvette was placed perpendicularly to the light path of CD spectrometer and rotated within the cuvette plane, in order to rule out the possibility of the birefringency phenomena and eliminate the possible angle dependence of the CD signals. Fluorescence spectra were recorded on a Hitachi F-4500 fluorescence spectrophotometer. The absolute fluorescence quantum yield was measured by using an absolute PL quantum yield spectrometer (Hamamatsu Photonics) with a calibrated integrating sphere. X-ray diffraction (XRD) analysis was performed on a Rigaku D/Max-2500 X-ray diffractometer (Japan) with $Cu_{K\alpha}$ radiation (λ =1.5406Å), which was operated at a voltage of 40 kV and a current of 200 mA.

Materials

All the starting materials and solvents were obtained from commercial suppliers and used as received. 1,3,5-Benzenetricarbonyl trichloride and ethyl 4-aminocinnamate were purchased from Alfa Aesa. The C_3 -symmetric gelator (BTAC) was synthesized by following the method reported previously. Milli-Q water (18.2 M Ω cm) was used in all cases. (R)-1-cyclohexyl ethylamine and (S)-1-cyclohexyl ethylamine were purchased from TCI and used as received.

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