Research Article
Barbituric Derivative Nanoaggregates with Aggregation-Induced Emission and Mechanofluorochromism

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Three new nonplanar barbituric derivatives, named as TTB, TTTB, and TOB, were synthesized. The D-π-A type conjugated compounds showed obvious intramolecular charge transfer (ICT) property, which was evidenced by theoretical calculations and spectral analyses. All of them exhibited aggregation induced emission (AIE) when formed nanoaggregates. These nanoaggregates also showed reversible mechanofluorochromism (MFC). Their red light emission became deep red after grinding and then recovered with dichloromethane fuming. Hence, a strategy to fabricate mechanofluorochromic nanoaggregate phosphors via nonplanar π-skeleton and steric effect was demonstrated, and these nanophosphors possess potentials for mechanosensors and anticounterfeiting technology.

1. Introduction

Conventional organic luminescent materials have little fluorescence due to aggregation (aggregation-caused quenching (ACQ) effect), which severely limits the applications in solid state [1, 2]. Fortunately, Hong and coworkers discovered a series of compounds and opened a new door for aggregation-induced emission (AIE) [3, 4]. Many research groups have developed strong interest in further designing and synthesizing some AIE organic luminescent materials and have discovered and summarized the interesting properties of AIE materials [5–12]. Though many AIE materials have been reported, however, the relationship between molecule structure and the AIE effect still remains ambiguous. Therefore, it is of great significance to design and synthesize more AIE compounds and explore the structure-property relationship.

In our former work, a series of thiophene derivatives with one, two, and four branches attaching on thiophene, respectively, were synthesized [13]. In these compounds, intramolecular charge transfer (ICT) occurred from each branch to thiophene, producing multidirectional polarization body. The relationship between the multidirectional polarization and the AIE effect was studied. It was found that simple one-branched compound with dipolar ICT showed advantage in AIE effect over the multibranched one.

Based on the former work, we further design and synthesize several dipolar compounds constituted by thiophene and barbituric acid in the current submission. In this work, the monomer, dimer, and trimer of thiophenyl serve as electron donor. The emphasis of this work is put on the relationship between the degree of oligomerization in the electron donor and the AIE effect. The research results show that the compound with the trimer of thiophenyl exhibits the strongest AIE effect. Additionally, these barbituric derivative nanoaggregates exhibited strong mechanofluorochromism (MFC) effects. The detailed results are reported below.

In this paper, three barbituric derivatives were synthesized and exhibited very good AIE property. Piezochromatic
luminescent materials are a type of force-stimulated responsive materials that change the luminescence color and intensity by varying the physical packing of the molecules [14–17].

2. Experimental Section

2.1. Materials and Characterizations. All the reagents were obtained commercially and used without further purification: 1,3-dimethylbarbituric acid (Aladdin, 98%), 2,2'-bithiophene-5-carboxaldehyde (Macklin, 98%), and 2,2':5',2'-ter thiophene-5-carboxaldehyde (Macklin, 98%). All the solvents used in this study were purified according to standard methods prior to use. The fluorescence quantum yield of TTB, TTTB, and TOB. Theoretical calculations at the B3LYP/6-31G (d) basis were used to determine the HOMO and LUMO energies for the three compounds.

2.2. Syntheses. The compound of 1a was synthesized according to literature methods [19]. The target luminogens (TTB, TTTB, TOB) were synthesized by the Knoevenagel condensation of aldehyde-based intermediates with 1,3-dimethyl barbituric acid (Scheme 1).

2.2.1. Synthesis of 2,2'-Bithiophene-1,3-Dimethylbarbituric Acid (TTB). A mixture of 1,3-dimethylbarbituric acid (0.960 g, 6.18 mmol) and 2,2'-bithiophene-5-carboxaldehyde (1.00 g, 5.15 mmol) in acetic (5 mL) and acetic anhydride (5 mL) was refluxed for 2 h. The reaction was cooled to room temperature and solid particles were filtered. The product was then purified by recrystallization with acetic acid to give 0.58 g (61% yield): 1H NMR (400 MHz, CDCl3): δ 8.65 (s, 1H), 3.80 (s, 1H), 7.55 (s, 1H), 7.45 (d, J = 5.4 Hz, 1H), 7.30 (d, J = 6.6 Hz, 1H), 7.12 (s, 1H), 3.43 (s, 6H) (Figure S1); 13C NMR (151 MHz, CDCl3): δ 162.18, 157.65, 134.69, 130.97, 130.33, 127.52, 125.40, 124.62, 124.45, 124.30, 123.78, 28.62, 27.97 (Figure S4).

2.2.2. Synthesis of 2,2':5',2''-Ter thiophene-1,3-Dimethylbarbituric Acid (TTTB). A mixture of 1,3-dimethylbarbituric acid (0.670 g, 4.35 mmol) and 2,2':5',2''-ter thiophene-5-carboxaldehyde (1.00 g, 3.62 mmol) in ethanol (10 mL) and two drops of acetic acid was refluxed for 4 h. The reaction was cooled to room temperature, and the solid particles were filtered. The product was then purified by recrystallization with acetic acid to give 0.85 g (83% yield): 1H NMR (400 MHz, CDCl3): δ 8.63 (d, J = 6.6 Hz, 1H), 7.80 (d, J = 4.6 Hz, 1H), 7.47 (d, J = 4.2 Hz, 1H), 7.36 (d, J = 4.2 Hz, 1H), 7.40 (d, J = 3.4 Hz, 1H), 3.43 (s, 6H) (Figure S2); 13C NMR (101 MHz, CDCl3): δ 162.22, 161.56, 153.11, 150.90, 147.99, 146.47, 135.79, 134.99, 128.14, 127.59, 126.37, 124.14, 108.50, 28.39, 27.63 (Figure S3).

2.2.3. Synthesis of 5-Thiophen-2-yl-Furan-2-Carbaldehyde, 1,3-Dimethylbarbituric Acid (TOB). A mixture of 1,3-dimethylbarbituric acid (1.05 g, 6.74 mmol) and 1a (1.00 g, 5.62 mmol) in ethanol (10 mL) and two drops of acetic acid was refluxed for 6 h. The reaction was cooled to room temperature and the solid particles were filtered. The product was then purified by recrystallization with acetic acid to give 0.89 g (90% yield): 1H NMR (400 MHz, CDCl3): δ 8.65 (s, 1H), 7.80 (s, 1H), 7.55 (s, 1H), 7.44 (d, J = 5.4 Hz, 1H), 7.37 (d, J = 4.6 Hz, 1H), 7.12 (s, 1H), and 3.43 (s, 6H) (Figure S5).
for TTB and TOB, as well as obvious that the fluorescence intensity increases as \( f_w \) is increasing, accompanied by a red shift of the emission wavelength (\( \lambda_{em} \)). This may be caused by intramolecular charge transfer (ICT). As the water content increases, the fluorescence intensity first increases and then decreases, and the emission wavelength is always red-shifted. To verify the ICT mechanism, emission spectra were recorded in different solvents with various polarities. As can be seen from Figure 3, the emission wavelengths of the barbituric acid derivatives TTB, TTTB, and TOB were red-shifted with the increase of the solvent polarity. This phenomenon is typical behavior of molecules with ICT nature. The theoretical calculation results are consistent with this phenomenon (Figure 4). When \( f_w \) increases, the molecules begin to accumulate in a limited space and, to some extent, limit the free rotation of the intramolecular rotatable groups, which makes the radiation transition pathway active and leads to enhanced fluorescence emission [20–22]. However, when \( f_w > 80\% \) for TTB and TOB, as well as \( f_w > 70\% \) for TTTB, molecules began to precipitate, resulting in weak emission and blue shifts of the \( \lambda_{em} \) [23, 24]. Similarly, it can be observed from the photographs of TTB, TTTB, and TOB (Figure 5), with the \( f_w \) ranging from 0 to 90\%, all the three compounds are AIE active.

The AIE index \( I_m/I_0 \), where \( I_m \) is the maximum fluorescence intensity of the nanoaggregate solution \( (0 < f_w < 100\%) \), and \( I_0 \) is the fluorescence intensity when \( f_w = 0\% \), is generally used to quantify the AIE activity [25]. From Figure 2(b), obviously, TTTB exhibits enhanced AIE effect compared to TTB and TOB: in aggregation solution, the luminescence intensity of TTTB increased by 20-fold \( (f_w = 70\%/f_w = 0\%) \), with the fluorescence quantum yield \( (\Phi_F) \) increasing from 1.5\% to 14.9\%. However, the maximum luminescence enhancement of TTB \( (f_w = 80\%/f_w = 0\%) \) is only 4-fold \( (\Phi_F : 0.3\% \text{ to } 1.6\%) \) and the maximum emission enhancement of TOB \( (f_w = 80\%/f_w = 0\%) \) is only 6-fold \( (\Phi_F : 0.4\% \text{ to } 2.1\%) \). TTTB shows the highest AIE index. This may be attributed to two facts: (i) the energy gap between HOMO and LUMO of TTTB is the smallest of the three compounds, resulting in the strongest emission; (ii) oligothiophene is a stronger electron donor than the other two compounds, which may lead to a stronger ICT in TTTB.

### 3. Results and Discussion

#### 3.1. Optical Properties

As shown in Figure 1, the absorption peak wavelength (\( \lambda_a \)) and emission wavelength (\( \lambda_{em} \)) of TTB and TOB are similar in THF solution, indicating that both of the molecules have similar distribution of electronic levels. The \( \lambda_a \) and \( \lambda_{em} \) for TTB are much longer than those of TTB and TOB, which may be due to TTTB being a larger conjugated molecule with an additional thiényl in the electron donor (Figures 1(a) and 1(b)). As shown in Figure 1(b), the emission wavelengths of the three compounds are 585, 654, and 640 nm, respectively. Obviously, in the solid state, the emission wavelength of TTB shows a blue shift compared to TTTB and TOB. These results have been proved by theoretically calculated.

#### 3.2. AIE Activities

The photoluminescence (PL) behaviors of compounds TTB, TTTB, and TOB were investigated in THF/H\(_2\)O mixed solvent (Figure 2) with the concentration of all the three compounds of \( 1.0 \times 10^{-5} \text{ mol/L} \). From Figure 2(a), it can be clearly seen that the three compounds have obvious AIE properties which show weak fluorescence in pure THF solution and strong emission in the aggregate state. For Figure 2(b), when \( f_w \leq 80\% \) for TTB and TOB, as well as \( f_w \leq 70\% \) for TTTB, it is obvious that the fluorescence intensity increases as \( f_w \) is increasing, accompanied by a red shift of the emission wavelength (\( \lambda_{em} \)) which may be caused by intramolecular charge transfer (ICT).

**Figure 1:** (a) Absorption and (b) emission spectra of three compounds in THF solution (concentration: \( 1.0 \times 10^{-5} \text{ mol/L} \); excitation wavelength, TTB: 360 nm, TTTB: 480 nm, and TOB: 360 nm) and solid state (excitation wavelength, TTB: 410 nm, TTTB: 530 nm, and TOB: 440 nm).
Figure 2: (a) FL spectra of TTB, TTTB, and TOB in THF/H$_2$O with different water fractions ($f_w$). (b) FL peak location (blue) and emission intensity (red) of TTB, TTTB, and TOB versus $f_w$ in THF/H$_2$O (concentration: 1.0 × 10$^{-5}$ mol/L; excitation wavelength, TTB: 360 nm, TTTB: 480 nm, and TOB: 360 nm).
meaning that enhanced electrostatic interaction between TTTB molecules that limits the intramolecular rotation of some bonds in the nanoaggregate state [26–28].

The morphology and size of the nanoaggregates in the THF/H2O with different f_w were observed by scanning electron microscopy (SEM) and dynamic light scattering (DLS) (Figure 6), which show the formation of nanoaggregates with relatively narrow sized distributions. As shown in Figure 6, the average diameters (d) of the nanoaggregates for TTB, TTTB, and TOB with their strongest emission intensities (when f_w = 80%, 70%, and 80%) are 140, 200, 130 nm, respectively.

3.3. Theoretical Calculations. As shown in Figure 7, the HOMO-LUMO gaps for TTB, TTTB, and TOB are 3.07, 2.71, and 3.07 eV, respectively. The HOMO mainly localizes in the donor furan or thiophene units, and the LUMO distributes over all the molecules; it was clearly confirmed that a charge transfer effect occurred in the molecules [29]. As shown in Figure 7, the visualized electrostatic potential exhibited intramolecular charge transfer with a positive region (blue) on the donor moiety and a negative region (red) on the acceptor moiety. This theoretical result illustrates the transfer of electrons from the donor to the acceptor moiety, corresponding to the observed optical properties.

Figure 3: PL spectra of TTB, TTTB, and TOB in different solvents with varying polarities (concentration: 1.0×10⁻⁵ mol/L; excitation wavelength, TTB: 360 nm, TTTB: 480 nm, and TOB: 360 nm).

Figure 4: Plots of the electrostatic potential for TTB (a), TTTB (b), and TOB (c) with B3LYP/6-31G (d) on the isodensity surface of 0.001. The color is coded as red for strong negative and blue for strong positive.

Figure 5: Photographs of TTB, TTTB, and TOB at f_w = 0-90% under 365 nm UV light illumination.
3.4. Mechanofluorochromic Properties. Based on the MFC characteristics of nonplanar AIE molecules, the MFC performances of TTB, TTTB, and TOB with helical structure were investigated. As shown in Figures 8 and 9, TTB, TTTB, and TOB are red powder (nanoaggregates) with red fluorescence ($\lambda_{\text{max}} = 585$, 654, and 640, respectively) under 365 nm UV excitation. After grinding, the color of TTB and TOB only changed a little ($\lambda_{\text{max}} = 593$ and 645, respectively), indicating that the MFC activities for both TTB and TOB are very weak and can be negligible. However, the color of TTTB became deep red with a red shift of 26 nm. The color change of TTTB before and after grinding can be easily observed with naked eyes.

Figure 6: SEM images (a) and DLS sizes (b) of TTB, TTTB, and TOB in THF/H$_2$O (nanoaggregate state) at a concentration of $1 \times 10^{-5}$ mM.
After grinding, the three ground powders were exposed to CH$_2$Cl$_2$ vapor at room temperature. As can be seen from Figure 8, the fluorescent color of the compound after the fume almost recovered to its original state. In particular, the color change of TTB is more obvious, its fluorescent color changes from a ground deep-red to orange-red after fuming, and the process is completely reversible.

Powder X-ray diffraction (XRD) were used to further explore the MFC mechanism of these nanoaggregates. As shown in Figure 10, all of the original nanoaggregate samples had sharp and strong peaks, but the intensity of the peaks decreased or even disappeared after grinding. The result implies that the molecular morphology changed from crystalline to amorphous [30–32], and red shift of $\lambda_{em}$ was attributed to the planarization induced by grinding. AIE molecules usually adopt a twisted configuration due to the steric hindrance between moieties and the formation of intermolecular cavities with a loose packing mode. The binding energy released after grinding and accompanied by the destruction of crystal lattice. After fumed with methylene chloride gas,
the XRD patterns gained their initial characteristic peaks, indicating that the crystal structures were restored (Figure 10).

4. Conclusions

In summary, we reported three compounds (TTB, TTTB, and TOB) with typical AIE characteristics and highly reversible piezochromic and vapochromic properties. Among them, TTTB displayed the largest MFC spectra shift (26 nm) with visible color changes. It was confirmed that the emission bands of these compounds and their intensity dramatically changed by switching the nanoaggregates between crystalline and amorphous. These nanomaterials can be used in color emission and security [33].

Data Availability

The data used to support the findings of this study are available from the corresponding authors upon request.

Conflicts of Interest

The authors declare that they have no conflicts of interest.

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Supplementary Materials

Figure S1: the 1H-NMR spectra of TTB. Figure S2: the 1H-NMR spectra of TTTB. Figure S3: the 1H-NMR spectra of TOB. Figure S4: the 13C-NMR spectra of TTB. Figure S5: the 13C-NMR spectra of TTTB. Figure S6: the 13C-NMR spectra of TOB. Figure S7: the mass spectra of TTB. Figure S8: the mass spectra of TTTB. Figure S9: the mass spectra of TOB. (Supplementary Materials)
References


[29] X. Q. Zhang, Z. G. Chi, B. J. Xu et al., "Comparison of responsive behaviors of two cinnamic acid derivatives containing..."


