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## ARTICLE

# Highly bright mechanoluminescence and remarkable mechanochromism based on a tetraphenylethene derivative with aggregation-induced emission†

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Organic materials exhibiting mechanoluminescence (ML) are promising for usages in displays, lighting and sensing. However, the mechanism for ML generation remains unclear, and the light-emitting performance of organic ML materials in solid state has been severely limited by the aggregation-caused quenching (ACQ) effect. Herein, we present two strongly photoluminescent polymorphs (*i.e.*, C<sub>g</sub> and C<sub>b</sub>) with distinctly different ML activities based on a tetraphenylethene derivative P<sub>4</sub>TA. As an aggregation-induced emission (AIE) emitter, P<sub>4</sub>TA perfectly surmounted the ACQ, making the resultant block-like crystals in the C<sub>g</sub> phase exhibit brilliant green ML under daylight at room temperature. The ML-inactive prism-like crystals C<sub>b</sub> can also turn on its ML by transiting toward C<sub>g</sub> with the aid of dichloromethane vapors. Moreover, the C<sub>g</sub> polymorph shows ML and mechanochromism simultaneously and respectively without and with UV irradiation under force stimulus, thus suggesting a feasible design direction for developing efficient and multifunctional ML materials.

## Introduction

The mechanoluminescence (ML) phenomenon has been first found by Francis Bacon in 1605.<sup>1</sup> Heretofore, research on exploiting advanced ML materials has not yet been a major focus.<sup>2</sup> The materials with brilliant ML are actually of great importance from both fundamental and practical viewpoints because they are promising for usages in displays, as well as light source and sensors.<sup>2,3</sup> However, the comprehensive understanding on the crystal properties required for ML activity and the corresponding mechanisms is less well demonstrated.<sup>4</sup>

This lack in understanding leads to the feasible design principles of these emitters, particularly those with satisfactory ML brightness, being rarely found.<sup>5</sup> As reported previously, the performance of organic ML compounds can be related to both their molecular and molecular-assembly structures.<sup>6</sup> Therefore, controlling the molecular arrangements in the solid state and achieving a molecular-level understanding of the relationship between the molecular conformations and packing characteristics and the resulting optical properties are the essential issues in obtaining efficient ML materials.

Notably, non-covalent intermolecular interactions, such as  $\pi$ - $\pi$  stacking and hydrogen bonding, are important in constructing the supramolecular systems.<sup>7</sup> These interactions are able to influence the final packing structure strongly, thereby making polymorphism with different ML activities more probable.<sup>6a,8</sup> Nevertheless, in most cases, typical  $\pi$ - $\pi$  stacking interactions often lead to aggregation-caused quenching, which poses

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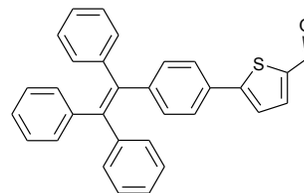


Fig. 1 Molecular structure of P<sub>4</sub>TA.

significant difficulties in developing high-performance ML materials.<sup>4b,9</sup> By contrast, a diametrically opposed effect is recently operative in a class of chromophores with twisted conformations (*e. g.*, tetraphenylethene derivatives), which exhibit enhanced emission in the solid state with respect to the fluid solution.<sup>10</sup> The discovery of this abnormal phenomenon, known as the aggregation-induced emission (AIE), has sparked a rapid expansion in the field of photoluminescent sensors and electroluminescent devices.<sup>11</sup> AIE also provides new possibilities for designing highly mechanoluminescent materials.

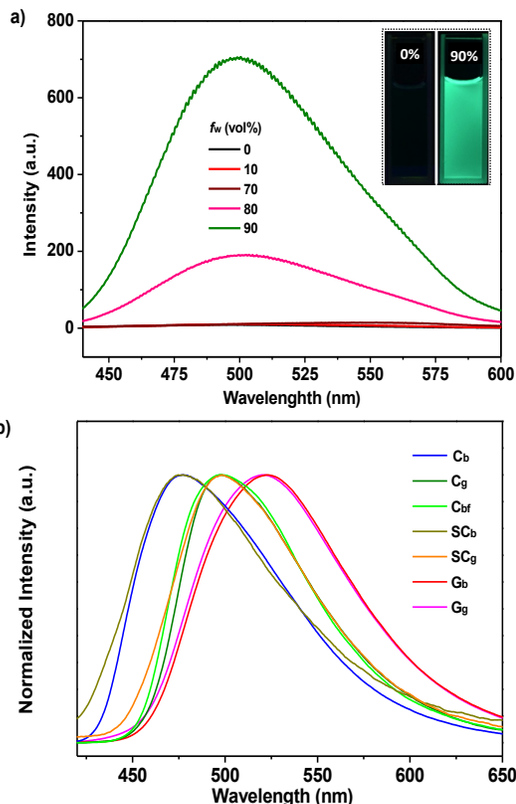
This study presents a new polymorphic system that can be facilely and controllably constructed using a tetraphenylethene derivative [*i. e.*, 5-(4-(1,2,2-triphenylvinyl)phenyl)thiophene-2-carbaldehyde (P<sub>4</sub>TA)] as the building blocks (Fig. 1). The two crystalline polymorphs of P<sub>4</sub>TA show strong blue- and green-colored photoluminescence (PL). The blue-light crystals are significantly ML inactive, whereas the green-light ones are highly mechanoluminescent because of its distinctly different molecular packing mode and unique AIE character. The existence of polymorphs from the same molecule with exactly opposite properties provides a unique prototype to investigate the crystalline structures required for ML activity and the effect of AIE property on ML enhancement. The relationship between the ML and the mechanofluorochromism of P<sub>4</sub>TA is also presented.

## Results and discussion

P<sub>4</sub>TA was straightforwardly prepared through a palladium-catalyzed coupling reaction by introducing 2-thiophenylaldehyde to the tetraphenylethene moiety (Scheme S1). The purified material was then characterized using nuclear magnetic resonance (NMR) spectroscopy and X-ray crystallography. The satisfactory data obtained fully confirmed its expected molecular structure (ESI).

The UV-visible absorption spectrum of P<sub>4</sub>TA was measured in a dichloromethane (DCM) solution. Two absorption bands centered at 317 and 366 nm were observed. These two bands were associated with the  $\pi$ - $\pi^*$  transition and the intramolecular charge transfer, respectively (Fig. S1). The large Stokes shift (68,027 cm<sup>-1</sup>) between the absorption and the fluorescence spectra ( $\lambda_{\text{em-max}}=513$  nm) of P<sub>4</sub>TA in DCM was indicative of the structural difference between the ground and excited states. The resulting P<sub>4</sub>TA compound would probably also be AIE-active in considering that tetraphenylethene is the most frequently used AIE unit. To identify this probability, a tetrahydrofuran (THF) solution of P<sub>4</sub>TA was titrated with water and the change in fluorescence emission was monitored. P<sub>4</sub>TA exhibited an extremely weak emission in the THF solution, where it was well dissolved (Fig. 2a). Hence, almost no PL signal was recorded. However, when 90% (*v/v*) of water was added, a strong green emission peaked at 499 nm was observed and the corresponding intensity (~705 a. u.) dramatically increased by up to ~100 times as compared to that at 0% water fraction (~7 a. u.). Adding water to the THF solution of P<sub>4</sub>TA significantly induced the formation of nanoparticle because P<sub>4</sub>TA molecules contained highly hydrophobic aromatic rings. In other words, the emission enhancement was caused by molecule aggregation which suggested that P<sub>4</sub>TA was AIE-active.<sup>12</sup>

The prominent AIE character of P<sub>4</sub>TA motivated the application of material in the solid state. Accordingly, block-like crystals (C<sub>g</sub>-form) were achieved through solvent evaporation of P<sub>4</sub>TA in the mixture solvent of *n*-hexane and



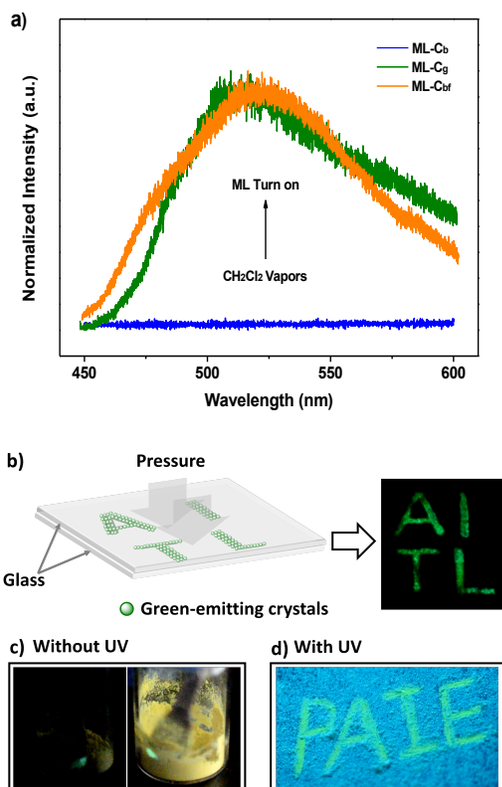
**Fig. 2** a) PL spectra of the dilute solutions of P<sub>4</sub>TA in water/THF mixtures with different water fractions ( $f_w$ ). The inset depicts the emission images of the compound in pure THF and 90% water fraction mixtures under 365 nm UV illumination (10  $\mu$ M). b) PL spectra of P<sub>4</sub>TA in different phases.

DCM (Fig. S2a). Compared to the emissive nanoaggregates in THF/H<sub>2</sub>O, the as-prepared sample C<sub>g</sub> exhibited even a stronger green-light emission centered at 498 nm [ $\Phi_{\text{F,S}}=52\%$ ] (Fig. 2b). By grinding the C<sub>g</sub> crystals with a pestle or shearing them with a spatula, a highly bright green light emission peaking at 517 nm was observed in the dark without UV irradiation (Figs. 3a and 3c, and Video S1). This experiment unambiguously illustrated that P<sub>4</sub>TA in C<sub>g</sub>-form was ML-active. The strong ML of C<sub>g</sub> was indeed clearly seen even under daylight at the room temperature and maintained while the crystals were crushed (Fig. 3c and Video S2). Certain organic materials, such as coumarin, phenanthrene, *N*-acetyl anthranilic acid, *N*-isopropyl carbazole and *N*-phenyl imides, were also reported to show mechanoluminescent activities. However, none of these materials could emit a ML strong enough to be observed with the naked eye under daylight at the room temperature.<sup>5,13</sup> The poor performance of the conventional organic ML materials should be attributed to their intrinsic ACQ property, which led to the low emitting efficiency in the solid state. By contrast, the unique AIE feature of P<sub>4</sub>TA perfectly surmounted the ACQ effect and exhibited a positive effect on luminescence enhancement, thereby giving a brilliant ML of C<sub>g</sub>. To further demonstrate the ML characteristic of C<sub>g</sub>, a simple device was made by sandwiching the sample between two pieces of pre-sculptured glasses. The capital letters 'AITL' were clearly displayed when pressure was used as the driven force, which suggested the ML 'display capability' of C<sub>g</sub> (Fig. 3b). As such,

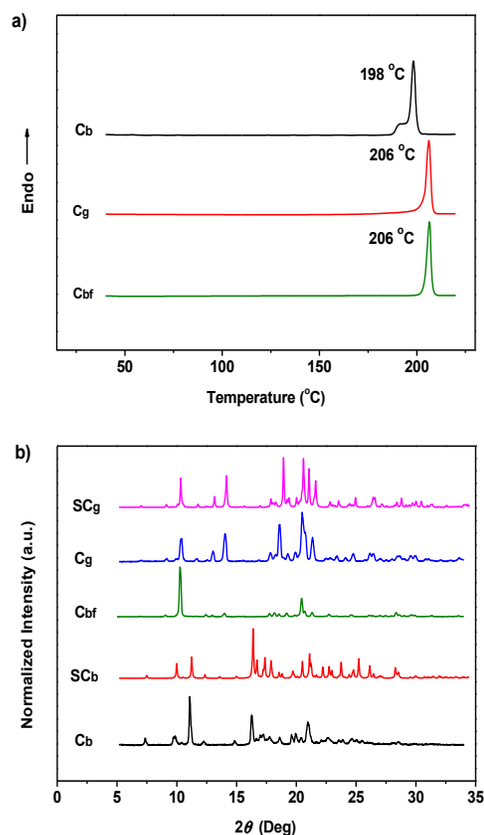
the extraordinary AIE-active ML material as P<sub>4</sub>TA will be a promising candidate for displays and optical recording.

Interestingly, another type of prism-like crystals (C<sub>b</sub>-form) was observed while exploring different processing conditions. The C<sub>b</sub>-form crystals, which showed an intense blue-light emission peaked at 476 nm ( $\Phi_{F,s}=36\%$ ), could be obtained by adding ethanol into a P<sub>4</sub>TA/DCM solution under the action of ultrasonic (Fig. S2b). However, in contrast to the vivid C<sub>g</sub> phenomenon, the P<sub>4</sub>TA sample completely lost its ML activity when aggregated in the C<sub>b</sub>-form (Fig. 3a). The C<sub>b</sub> crystals showed a small melting endothermic shoulder peak at 191 °C and a sharp peak at 198 °C in the first heating curve of differential scanning calorimetry (DSC) (Fig. 4a), indicating that C<sub>b</sub> was mainly composed by the microcrystals melted at 198 °C. This result was different from that of C<sub>g</sub> melted at 206 °C. Moreover, the powder X-ray diffraction (XRD) spectra also exhibited distinctly different patterns for the two samples (Fig. 4b). These results implied that the different ML activities between C<sub>g</sub> and C<sub>b</sub> might be attributed to their dissimilar molecular packing modes. A single crystal X-ray analysis was thus performed for the P<sub>4</sub>TA crystals to obtain more insight into this aspect. The single crystals with two polymorphs (*i. e.*, SC<sub>g</sub> and SC<sub>b</sub>) suitable for the X-ray structural analysis were isolated through the slow solvent evaporation of P<sub>4</sub>TA in the mixtures of ethanol and CHCl<sub>3</sub> with different concentrations.

The SC<sub>g</sub> and SC<sub>b</sub> samples emitted intense green and blue lights peaked at 499 and 476 nm, respectively (Figs. 2b and 5a-5b). These lights were similar to those of the as-prepared

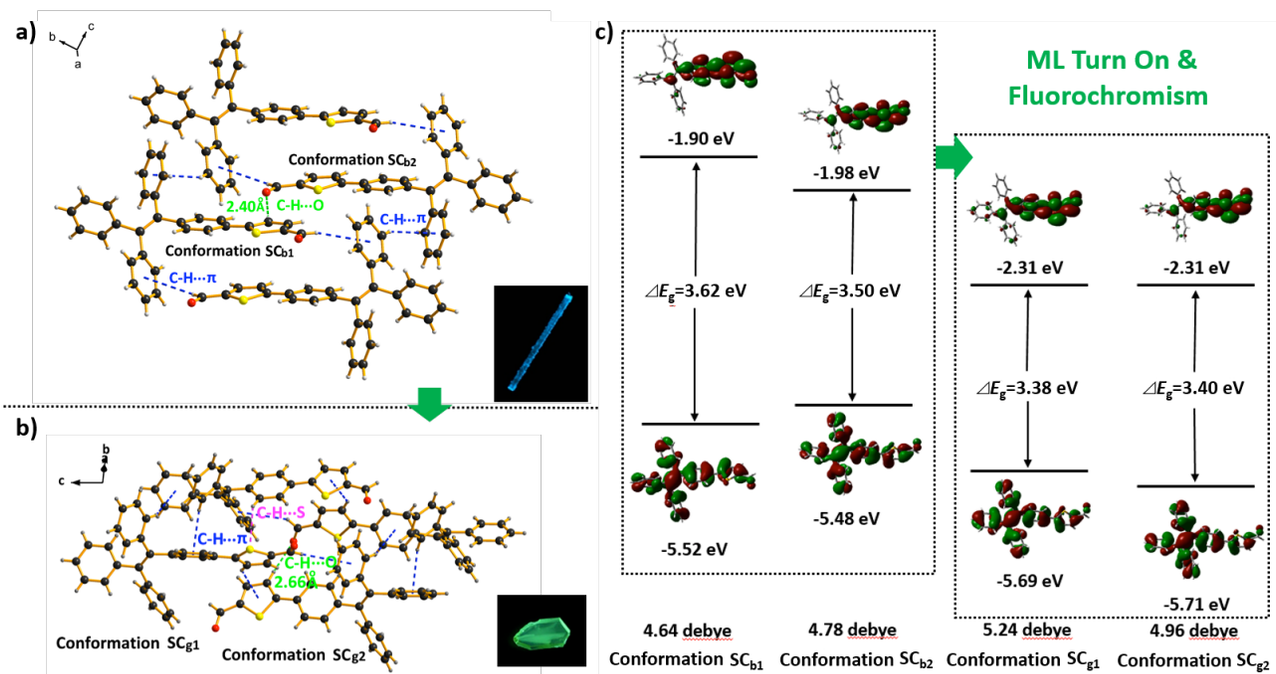


**Fig. 3** a) ML spectra of P<sub>4</sub>TA in different phases. b) The image of capital letters 'AITL' shown through ML of P<sub>4</sub>TA in the dark under the pressure stimulus at room temperature. c) ML images of P<sub>4</sub>TA in the dark (left) and under daylight (right) at room temperature. d) Writable mechanochromic fluorescence of P<sub>4</sub>TA demonstrated by capital letters 'PAIE' generated with a metal rod.



**Fig. 4** DSC curves a) and XRD patterns b) of P<sub>4</sub>TA in different phases.

crystals of C<sub>g</sub> and C<sub>b</sub>. The main peaks of the simulated XRD patterns of SC<sub>g</sub> and SC<sub>b</sub> also agreed well with those in the patterns obtained from P<sub>4</sub>TA in the C<sub>g</sub> and C<sub>b</sub> phases, which suggested that the initial powders were mainly composed of P<sub>4</sub>TA microcrystals in the polymorphs of SC<sub>g</sub> and SC<sub>b</sub> (Fig. 4b). Further systematic analysis revealed that both SC<sub>g</sub> and SC<sub>b</sub> belonged to the non-centrosymmetric polar space group of *P*(2)*1* (Table S1). Some previous reports had showed that the dipolar structures and the non-centrosymmetric molecular arrangements were in favor of obtaining the piezoelectric properties, which were closely pertinent to the ML activities of the crystals.<sup>14</sup> In principle, the fracture of crystals with a strong piezoelectric effect will lead to electronic discharge at the crack surface, which would result in dye excitation and generation of ML for the crystals.<sup>5,15</sup> The molecular structure of P<sub>4</sub>TA and the crystalline symmetry of SC<sub>g</sub> and SC<sub>b</sub> also met the requirements of piezoelectric properties, thus making the as-prepared crystals of C<sub>g</sub> and C<sub>b</sub> more impetus and more possibility to achieve the ML character. However, the dipole moments and the HOMO-LUMO band gaps ( $\Delta E_g$ ) of the molecules in SC<sub>g</sub> and SC<sub>b</sub> polymorphs were different. These differences were caused by their distinct molecular conformations and packing characteristics. The asymmetric units in both polymorphs (*i. e.*, SC<sub>g1</sub> and SC<sub>g2</sub> for SC<sub>g</sub>, and SC<sub>b1</sub> and SC<sub>b2</sub> for SC<sub>b</sub>). Each molecule showed the formation of a C-H...O intermolecular hydrogen bond (Figs. 5a-5b, Figs. S3 and S4). Compared with SC<sub>b</sub>, the most notable conformational difference of P<sub>4</sub>TA in the SC<sub>g</sub> polymorph was the dihedral



**Fig. 5** Stacking modes and intermolecular interactions of the molecules in polymorphs SC<sub>b</sub> a) and SC<sub>g</sub> b); the insets show the fluorescence images of SC<sub>b</sub> and SC<sub>g</sub> under an excitation of 365 nm UV light. c) HOMO (lower images) and LUMO (upper images) of the four conformations of P<sub>4</sub>TA in polymorphs SC<sub>b</sub> and SC<sub>g</sub> calculated at the B3LYP/6-31G (d, p) level.

angle  $\theta$  between the thiophene and the adjacent phenyl ring. While the conformations of P<sub>4</sub>TA were twisted ( $\theta=18.9^\circ$  for SC<sub>g1</sub> and  $6.1^\circ$  for SC<sub>g2</sub>) in polymorph SC<sub>g</sub>, the two aromatic rings were nearly coplanar in polymorph SC<sub>b</sub> ( $\theta=2.5^\circ$  for SC<sub>b1</sub> and  $4.7^\circ$  for SC<sub>b2</sub>) (Table S2). In the case of SC<sub>g</sub>, the two thiophene rings of SC<sub>g1</sub> and SC<sub>g2</sub> were almost vertical to each other, showing a dihedral angle of  $85.8^\circ$ . By contrast, an anti-parallel packing mode was observed between the thiophene rings of SC<sub>g1</sub> and SC<sub>g2</sub> ( $\theta=3.7^\circ$ ) in SC<sub>b</sub>.

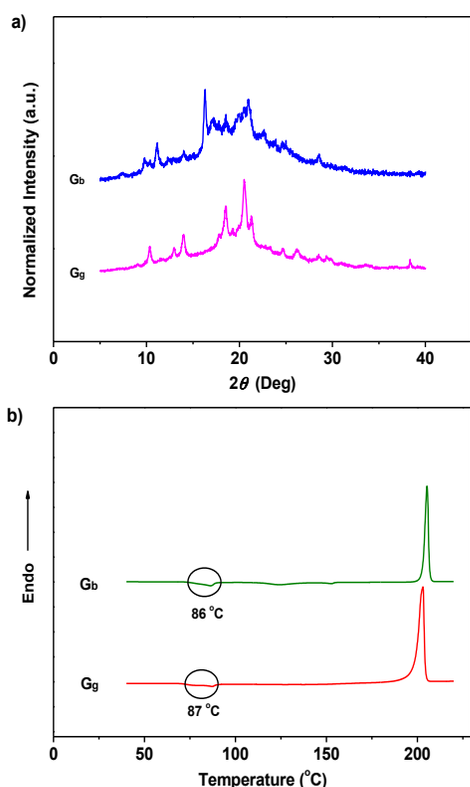
The most popular B3LYP density functional theory was then used to calculate the dipole moments and the  $\Delta E_g$  of P<sub>4</sub>TA in the four conformations at the 6-31G (d, p) level based on their ground state geometries in the single crystals. Fig. 5c presents the results. The dipole moments of SC<sub>g1</sub> and SC<sub>g2</sub> in the SC<sub>g</sub> polymorph were 5.24 and 4.96 debye (D), respectively. Both values were larger than those of SC<sub>b1</sub> (4.64 D) and SC<sub>b2</sub> (4.78 D) in SC<sub>b</sub>. The larger dipole moments of the molecules combining the non-centrosymmetric molecular arrangement may result in a larger net-dipole moment of the crystalline structure, and would subsequently lead to a stronger piezoelectric effect in the SC<sub>g</sub> polymorph when breaking the crystals. The theoretical calculation results also suggested that the molecules in both SC<sub>g</sub> and SC<sub>b</sub> had ICT characteristics: the electronic transitions (mainly from HOMO to LUMO for all the four conformations in SC<sub>g</sub> and SC<sub>b</sub>) from the occupied orbitals delocalized over the TPE (donor) moiety to the thiophenylaldehyde (acceptor) moiety made major contributions to the excited states (Fig. 5c and Table S4).<sup>16</sup> SC<sub>g1</sub> and SC<sub>g2</sub> showed even smaller  $\Delta E_g$  (HOMO→LUMO) values at 3.38 and 3.40 eV, respectively, as compared to those of SC<sub>b1</sub> (3.62 eV) and SC<sub>b2</sub> (3.50 eV) in the SC<sub>b</sub> polymorph. The calculations showed good agreement with the solid state UV-Visible spectra of C<sub>g</sub> and C<sub>b</sub> which absorbed

at 373 nm and 367 nm, respectively (Fig. S5). The preceding results thus indicate that the electrons of the molecules in SC<sub>g</sub> can be excited with a lower energy. Hence, the stronger piezoelectric effect and the lower electronic transition energy made for the excitation of P<sub>4</sub>TA molecules and the generation of ML in the SC<sub>g</sub> phase by breaking the crystals. Meanwhile, all the molecules adopted a highly twisted propeller-like conformation in the SC<sub>g</sub> polymorph, which prevented the formation of detrimental species, such as excimers or exciplexes, caused by  $\pi$ - $\pi$  stacking interactions. Furthermore, numerous intermolecular interactions of C-H... $\pi$  and C-H...S might also exist in the crystals aside the C-H...O hydrogen bonding (Fig. 5b and Table S3). These multiple interactions had rigidified the molecular conformations and impeded the intramolecular rotations, which largely reduced the energy loss via non-radiative relaxation channels, and subsequently resulted in a notable AIE effect and high  $\Phi_{F,S}$  value for P<sub>4</sub>TA. The preceding factors consequently made the ML of sample C<sub>g</sub>, which was mainly composed of P<sub>4</sub>TA microcrystals in SC<sub>g</sub> polymorph, highly emissive under the stimulus of mechanical force. By contrast, the weaker piezoelectric effect in the SC<sub>b</sub> polymorph seemed not to reach the higher energy requirement for the electronic excitation although SC<sub>b</sub> also exhibited strong photoluminescence. Consequently, P<sub>4</sub>TA lost its ML activity when aggregated in the C<sub>b</sub> phase. These results also suggested a feasible design direction for the development of efficient ML materials by combining the prominent piezoelectric property for molecular excitation and the abnormal AIE character for emission.

The PL of C<sub>b</sub> has been remarkably changed when the sample was exposed to DCM or acetone vapors for about 10 min, passing from an initial blue to green light at 499 nm (C<sub>b</sub>-form).

The resulting spectrum of  $C_{bf}$  was superimposable on that of  $C_g$  (Fig. 2b). The coincidence of PL emissions suggested that the fumed sample of  $C_{bf}$  probably took the same molecular arrangement with that of the  $C_g$  polymorph. Further evidence for this standpoint was provided by their similar XRD patterns and their overlapped DSC curves with the same melting point at 206 °C (Figs. 4a and 4b). As mentioned in the preceding discussion, sample  $C_g$  was ML-active. And expectantly, the ML activity of  $C_b$  could be tuned by simply altering the molecular packing mode upon fumigation. To verify this hypothesis, the ML spectrum of  $C_b$  was collected after exposure to DCM vapors ( $C_{bf}$ ). As anticipated,  $C_{bf}$  also exhibited a strong green light emission without UV irradiation by pressure, which revealed that the ML of  $C_b$  was facile to be turned on with the aid of DCM vapors. The ML emission maximum of  $C_{bf}$  located at 520 nm, which was close to that of  $C_g$  (Fig. 3a).

Noticeably, the ML maxima of  $C_g$  and  $C_{bf}$  were both significantly red-shifted ( $\Delta\lambda_{em,max}\approx 21$  nm) as compared to their



**Fig. 6** XRD patterns a) and DSC curves b) of the ground samples of  $P_4TA$ : ( $G_b$ ) ground sample from blue-light crystals; ( $G_g$ ) ground sample from green-light crystals.

PL spectra. This result shows a special mechanofluorochromic effect. To gain an understanding of this, the influence of applied pressure on the luminescence was investigated. The PL maximum of the pristine  $P_4TA$  in the  $C_g$ -form shifted from 498 nm to 521 nm ( $G_g$ ) after pressing or grinding (Fig. 2a), agreed well with its ML emission (Fig. S6), thereby confirming that the bathochromic shift between ML and PL of  $C_g$  was caused by its intrinsic mechanochromic property. The phase characteristic of the ground sample  $G_g$  was determined by XRD

to decipher further the relationship between the ML and the mechanochromism of  $C_g$ . Most of the diffraction peaks were diffuse or even disappeared although some resolvable peaks of  $G_g$  were consistent with those of their original crystals (Fig. 6a). This revealed that the ground sample was partially in a metastable amorphous state. Accordingly, DSC was performed for the sample after grinding (Fig. 6b). Compared with  $C_g$ , an additional exothermic peak around 86 °C was observed in the DSC thermogram of  $G_g$ , which demonstrated that the  $C_g$  crystals were partially destroyed and converted to an amorphous state by the grinding or pressing treatment. The  $P_4TA$  molecules in the  $C_g$  phase adopted twisted conformations in the crystalline state to fit into the crystalline lattice, and the crystalline lattices may collapse when triggered with mechanical force. The dye molecules then also relaxed to a more planar conformation, thereby emitting the redder ML and PL. In other words, the bathochromic shift of ML for  $C_g$  is originated from the microcrystal amorphization and the extension of molecular conjugation, which were believed to be the main reasons for the  $C_g$  mechanochromism.<sup>16,17</sup> Unlike other conventional stimuli-responsive materials, the  $C_g$ -form of  $P_4TA$  can show luminescence response and luminescence color change simultaneously and respectively without and with UV irradiation under force stimulus. This new kind of force-responsive material with AIE property had not yet been achieved before, and would facilitate applications of ML materials in the field of sensors.<sup>11c</sup> In addition, the fluorescence spectroscopy was also performed to evaluate the mechanochromic behavior of  $P_4TA$  in the  $C_b$  phase, and the  $C_b$  sample exhibited a more remarkable emission wavelength change of 47 nm upon grinding. Furthermore, the corresponding PL spectrum ( $G_b$ ) with  $\lambda_{em,max}=523$  nm fitted well with that for the powder ground from  $C_g$  (Fig. 2b), which indicated that  $P_4TA$  could switch to the same emission under force stimulus regardless of its initial state. Also the fluorescence ‘writability’ of  $C_b$  can be verified when being written on a piece of filter paper as shown in Fig. 3d. The mechanofluorochromism of  $C_b$  should undergo the similar mechanism proposed for  $C_g$ .

## Conclusions

Based on the AIE-active  $P_4TA$  molecules, two photoluminescent polymorphs (*i.e.*,  $C_g$  and  $C_b$ ) with multiple molecular conformations were achieved. They can show opposite mechanoluminescent activities by tuning the molecular assembly structures in the crystals. The block-like crystals of  $C_g$  exhibited highly bright green color ML upon pressing or grinding under daylight at room temperature. This unique property should be attributed to the strong piezoelectric effect of the crystals and the positive effect of the AIE property on luminescence enhancement. Moreover, with UV irradiation, the  $C_g$  of  $P_4TA$  showed mechanofluorochromism under mechanical stimulus. This new kind of force-responsive compound with the AIE property would facilitate applications of ML materials in the display and sensor fields. This work may provide a feasible design direction for developing more efficient ML materials by combining the prominent piezoelectric property for molecular excitation and the unique AIE character for emission.

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