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Combined Aggregation Induced Emission (AIE), Photochromism and Photoresponsive Wettability in Simple Dichloro-substituted Triphenylethylene Derivatives

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A dichloro-substituted triphenylethylene derivative (TrPEC2) with aggregation-induced emission (AIE), photochromism and photoresponsive wettability has been synthesized. The new compound shows fast-response photochromic behaviour, with good ON/OFF repeatability by utilizing a proposed stilbene-type intramolecular photocyclization in the solid state. Compared with the more usual diphenylethylene derivatives, the photochromic properties of the triphenylethylene derivative is much more striking and easier to achieve. The triphenylethylene derivative also displays AIE properties leading to strong fluorescence in the solid state. Therefore, both the ultraviolet-visible absorption and fluorescence emission are drastically changed during the photochromic processes. Furthermore, the morphology of the TrPEC2 microcrystalline surface could be controlled by irradiation. The wettability of the surface could be drastically decreased with contact angles of a water droplet changing from 73° to 118°. The triphenylethylene derivative with a simple molecular structure is, therefore, attractive for multifunctional materials.

Introduction

Photochromic materials are a type of photoresponsive compounds which undergo reversible chemical transformations between two (or more) well-defined states which have different optical properties. In recent years they have attracted much attention for their fundamental properties and for their emerging applications in security markings, optical shutters, photo-switchable molecular devices and optical memory storage systems.1,2 Accordingly, a wide range of photochromic systems have been developed including stilbene- and azo-containing compounds,3 spiropyran,4 spiroxazine,5 spiroxazine6 and diarylethylene5 (especially dithienylethene) and dihydroazulene derivatives.7

Photochromic materials are also investigated as topographical change materials.8 Furthermore, the wettability of the surfaces could be changed with morphology control by irradiation.8,9 In previous studies, topographical change properties of diarylethene derivatives have been systematically investigated by Uchida,10 Kobatake,11 and other groups.12 Spiropyran and azo-containing compounds have also been used as topographical change materials.13 Widespread applications are still restricted by the unstable surfaces properties, the demands of long irradiation times and complex synthetic procedures. Therefore, fast-response photochromic systems with simple molecular structures obtained from readily-available starting materials are in high demand. In this regard, diphenylethylene derivatives have been investigated; however, their photochromic properties are difficult to observe as the key cyclization reaction occurs only in the cis-isomer and the ring-closed states are quite unstable.1 Triphenylethylene derivatives can overcome these problems and might, therefore, be ideal fast-response photochromic systems. In addition, introducing halogen atoms into triphenylethylene derivatives can be expected thereby enhance the ring-closing photochromic process.6a-e,6f

Triphenylethylene derivatives are also well known as aggregation-induced emission (AIE) molecules. In contrast to the aggregation-caused quenching (ACQ) process, the emission intensities of AIE molecules are significantly increased by aggregation where the restriction of intramolecular rotation can efficiently avoid the usual emission quenching in the solid state. Since the pioneering work of Tang et al in 2001,14 several AIE systems have been developed including silole,15...
tetrphenylethenes,\textsuperscript{16} triphenylethylene derivatives\textsuperscript{17} and others.\textsuperscript{18} AIE has applications in OLED materials, piezochromic materials, chemosensors, photosensitizers and biolabels.\textsuperscript{14-18} We are aware of only one report of molecules which show both photochromic and AIE properties. During the course of our present studies, Tang et al described fast responsive, photo-reversible and thermo-irreversible photochromic properties of three triarylethylene-based polycyclic hydrocarbons which are AIE active.\textsuperscript{19}

Triphenylethylene derivatives were first reported to be AIE active by Chi et al\textsuperscript{17a} and the piezochromic properties of these molecules have been widely investigated.\textsuperscript{17b} We now report the first example of triphenylethylene derivatives which possess combined AIE, photochromic and photoresponsive wettability properties, namely TrPECl\textsubscript{2} (Scheme S1). As in Tang’s recent report,\textsuperscript{15} photochromism arises due to the photocyclization of stilbene moieties in the chromophores. Compared with previous AIE photochromic system, the response times of the triphenylethylene derivatives are shortened from minutes to seconds. Therefore, these materials have advantages in the areas of optical shutters and photo-switchable molecular devices.\textsuperscript{20} To the best of our knowledge there has been no previous report of an AIE-photochromic-photoresponsive wettability system with such simple structures and short response times. The straightforward synthetic route to TrPE and TrPECl\textsubscript{2} is shown in Scheme S1 in Supporting Information.

Results and discussion

Triphenylethylene (TrPE) is a typical AIE unit for the construction of AIE-active molecules\textsuperscript{17} and compound TrPECl\textsubscript{2} also shows AIE properties. AIE behavior in solution is commonly proven by the enhancement of emission resulting from fluorescent nanoparticle formation by increasing the ratio of a poor solvent.\textsuperscript{14-18} The data for TrPECl\textsubscript{2} in THF/water mixed solvent system at room temperature is shown in Figure 1. In the pure THF solution (good solvent), the photoluminescence (PL) of the compound is not detectable. This can be attributed to the fact that intramolecular rotation, which serves as the relaxation channel for the excited state to decay, is active. However, when the water fraction reached 90% (v/v) (water is a poor solvent for TrPECl\textsubscript{2}), the PL intensity was boosted to many fold with an emission peak at $\lambda_{\text{max}}$ 425 nm. This dramatic enhancement in luminescence indicates that the nanoparticles are formed in the proportion of 90:10 (v/v) water/THF, and that intramolecular rotation is restricted due to aggregation of the molecules, as is typically observed in AIE-active molecules. The emission band mainly originates from the $\pi-\pi^*$ transition of the triphenylethylene moieties according to the previous reports.\textsuperscript{17a} UV-vis spectra of TrPECl\textsubscript{2} in THF/water mixed solutions containing 0% and 90% water fractions are shown in Figure S2. It is obvious that the baseline is enhanced when the water fraction reached 90% which indicates the formation of nanoparticles.\textsuperscript{21}

For compound TrPECl\textsubscript{2}, blue emission is observed in the solid state upon excitation with UV light (365 nm), which is similar to that in AIE studies. Meanwhile the color of the solid rapidly changes from white into bright red under the UV irradiation as shown in Figure 2(b) and the blue emission is quenched subsequently as shown in Figure 2(d). The red solid reverts into a white solid after several seconds, and the photochromic bleaching process is described in Figure 3. After turning to white, the emission can be observed under UV irradiation again. This cycle can be repeated many times without significant fatigue. The repeatability of the photochromic process will be discussed later. TrPE shows similar photochromic properties with shorter irradiation saturated times and photochromic blenching times under the same conditions.

Figure 1. Emission spectra of TrPECl\textsubscript{2} in THF/water mixed solutions containing 0% and 90% water fraction; the inset shows the photographs of the TrPECl\textsubscript{2} in THF/water mixed solutions containing 0% and 90% water fractions.

Figure 2. Photographs of the crystals of TrPECl\textsubscript{2}; a) Before irradiation, the picture was taken under room light, b) after irradiation, the picture was taken under room light, c) before irradiation, the picture was taken under UV light, and d) after irradiation, the picture was taken under UV light.
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Figure 3. Photochromic bleaching process of TrPECl$_2$ at room temperature after irradiation with UV light stopped for: (a) 0 s, (b) 20 s, (c) 40 s, and (d) 60 s.

Figure 4. (a) Time dependent UV-vis reflectance spectra of TrPECl$_2$ during the irradiation process. (b) Time dependent UV-vis reflectance spectra of TrPECl$_2$ during the photochromic bleaching process.

To further evaluate the photochromic properties of TrPECl$_2$ and TrPE time dependent UV-vis reflectance spectroscopy studies were carried out: the spectra of TrPECl$_2$ under irradiation are shown in Figure 4(a). In the UV-vis reflectance spectra, the absorption bands below 400 nm are assigned as the $\pi-\pi^*$ transitions, in accordance with previous studies of the triphenylethylene derivatives. Before irradiation, bleaching signals with $\lambda_{max}$ 429 nm are also detected in the UV-vis reflectance spectra which are mainly due the emission of TrPECl$_2$. As shown in Figure 4(a), the low-energy absorption bands at around 514 nm increase with the irradiation time, suggesting increased conjugation in the molecule. Meanwhile the bleaching signals at 429 nm gradually decrease to zero with the quenching of emission which is in accordance with the phenomena described in Figure 2. After 30 s, the absorption bands at 514 nm reach their maximum intensity and no increase is observed with further irradiation under a deuterium light source (irradiation saturation time). Figure 4(b) shows the time dependent UV-vis reflectance spectra for the photochromic bleaching process of over-irradiated TrPECl$_2$. The absorption bands at 514 nm decreased as soon as the irradiation stopped and disappeared after 60 s (photochromic bleaching time). It could be detected that the absorption spectra in Figure 4(b) are broader than in Figure 4(a). This is mainly due to the bleaching signals (emission bands of TrPECl$_2$) under UV irradiation during the photochromic process. The compound TrPE shows shorter irradiation saturation time (<1 s) and photochromic bleaching time within 1 s at ambient room temperature and lighting. The much shorter irradiation and photochromic bleaching processes are mainly due to the instability of the ring-closed structures of TrPE, which will be discussed later. It should be noted that the photochromic processes of TrPE and TrPECl$_2$ are generally much faster than the previously reported AIE-photochromic systems (ca. 60 s). The increase and decrease of the reflectance of TrPECl$_2$ and TrPE at 515 nm as a function of time are shown in Figure 4(a-b) and Figure S5, respectively. The repeated photochromic switching processes of TrPECl$_2$ are shown in Figure S6 for 20 cycles without fatigue. Good repeatability of these photochromic materials indicates their potential applications as rewritable materials. It is obvious from these data that triphenylethylene derivatives show both fast-response properties and high stabilities in the photochromic processes.

In the triphenylethylene derivatives, the distance between the phenyl rings and the ethene-1,1-diyldibenzene moieties are short enough for the ring-closure reactions. The photochromic response after irradiation is assigned to a stilbene-type 6-π electron ring-closure as depicted in Figure 5. The ring-closure of similar structures has been reported in previous studies. In contrast to the previous report on AIE-photochromic materials, the structure of TrPECl$_2$(B) is unstable and quickly reverted to structure TrPECl$_2$(A) even in the dark at room temperature. The instability of TrPECl$_2$(B) structure indicates a lower energy barrier during the photochromic process, which is in accordance with the less extended π structure compared with the previously reported system. After irradiation for many cycles in air, the white solid TrPECl$_2$(A) became a pale yellow color. This solid is weakly emissive in solution which is different from pure compounds TrPECl$_2$(A) and TrPECl$_2$(B) which are non-emissive in solution. According to a previous study, the yellow colour after irradiation is mainly due to the dehydrogenation process of compound TrPECl$_2$(B) in the presence of oxygen to afford TrPECl$_2$(C) (Figure 5). However, the photochromic processes of these compounds were not observed in solution states. This is mainly attributed to the twisted structures of these AIE molecules in solution states which resist the ring-closure reaction. Similar results were also reported in AIE photochromic materials in previous studies.

Figure 5. Proposed mechanism of photochromism of TrPECl$_2$. 

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In an attempt to provide evidence for structure TrPECl$_2$(C) a photo-oxidation reaction was performed: compound TrPECl$_2$ was irradiated for 48 h in the solid state in the presence of air. The pale yellow solid is formed, which is proposed to be dehydrogenated product TrPECl$_2$(C). The $^1$H NMR spectrum of this solid showed additional peaks in the aromatic region compared to compound TrPECl$_2$(A), consistent with the formation of dehydrogenated phenanthrene derivative TrPECl$_2$(C) as shown in Figure S7 and S8 in supporting information. To further confirm the structure of TrPECl$_2$(C), a small amount of TrPECl$_2$(C) was purified from the irradiated product by silica gel column chromatography with hexane as eluent. High resolution EI mass spectrum of TrPECl$_2$(C) was shown in Figure S9 and obtained validation.

The TrPECl$_2$ microcrystalline surfaces were initially prepared by spin-coating a TrPECl$_2$ dichloromethane solution (5 mg mL$^{-1}$) onto two identical pieces of SiO$_2$ substrate. After the solvent was evaporated in vacuum at room temperature, one TrPECl$_2$ microcrystalline surface (surface A) was formed by storage in the dark for 3 days at ambient temperature (around 303 K). The other microcrystalline surface (surface B) was formed by irradiation under UV light (365 nm) for 3 minutes and followed by storage in the dark for 3 days at room temperature. Scanning electron microscopy (SEM) and contact angle analysis studies were carried out to establish the morphology of TrPECl$_2$ microcrystalline surfaces without or with irradiation. The SEM images without or with irradiation are displayed in Figure 6(a, b) and Figure 6(d, e) respectively. On surface A without the 3 minutes irradiation, microcrystals of TrPECl$_2$ with rod-shape-crystals gradually grow in the dark. A branched pattern is formed on the surface by the rod-shape-crystals as shown in Figure 6(a). The contact angle of a water droplet on surface A is 73$^\circ$ (Figure 6c). However, the microcrystals of TrPECl$_2$ on surface B grow into a different pattern of scale-shape crystals (Figure 6d and 6e) and the contact angle of a water droplet on surface B increases from 73$^\circ$ to 118$^\circ$ (Figure 6f). These results demonstrate that irradiation strongly affects the patterns of TrPECl$_2$ microcrystals and further controls the wettability of surfaces. The topographical change of TrPECl$_2$ microcrystals on the surface is mainly attributed to the formation of closed-ring isomer TrPECl$_2$(B) as nuclei at the beginning of crystal growth; a similar mechanism of the topographical change has been reported previously. The TrPECl$_2$(B) could be changed back to TrPECl$_2$(A) quickly in the crystal-growth process in the dark as mentioned above. It should be noted that these two surfaces with different patterns are formed with the same isomer [TrPECl$_2$(A)] of the photochromic material which is very different from the topographical change and wettability-controllable systems based on diarylethene derivatives. These new topographically-changeable and wettability-controllable surfaces are much more stable in light because the different microcrystalline patterns are all formed by the stable isomer of the photochromic process. In addition, the photoresponsive patterning process requires much shorter irradiation time compared with the same process for diarylethene derivatives (usually several hours). These advantages could promote these topographically-changeable and wettability-controllable materials based on triphenylethylene to industrial applications.

**Conclusions**

In summary, a structurally-simple triphenylethylene derivative TrPECl$_2$ has been successfully designed and synthesized and it provides a rare example of combined AIE, fast-responsive photochromism, topographically-changeable and wettability-controllable properties in a single compound. To study the photochromic properties of these materials in the solid state, UV-vis reflectance spectroscopy studies have been performed. SEM and contact angle analysis have been used to investigate the morphology and wettability control of the TrPECl$_2$ microcrystalline surfaces by irradiation. These new compounds enlarge our understanding of the photophysical and photochemical properties of triphenylethylene derivatives and present a new strategy to design multifunctional materials with simple structures.

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**Notes and references**


