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Effect of Ionic Interaction on the Mechanochromic Property of Pyridinium Modified Tetraphenylethene

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A pyridinium modified tetraphenylethene-based salt shows aggregation-induced emission enhancement property and irreversible mechanochromic behaviours.

As typical soft materials, molecular solids tend to show more pronounced responses to external stimuli when compared to the covalent and ionic solids. For conjugated organic compounds, the responses reflect not only on the macroscopic change in shape and the microscopic change in molecular packing, but also on the optic and electronic properties. A representative example is the mechano-chromic effect observed for many organic dyes that possess aggregation-induced emission (AIE) or aggregation-induced emission enhancement (AIEE) property.^{1,2} The term of mechano-chromism refers as to the phenomenon that a solid changes its absorption and/or emission spectrum under a certain kind of mechanical force applied to it.^{1b,3} For organic AIE/AIEE solids, the incentives are largely classified into two categories. One is the changes of physical factors and the other is rooted in the chemical structures.

The physical factors that result in mechanochromism can be further divided into intramolecular and intermolecular changes, which are conformation (e.g. torsional angle) and molecular packing (e.g. J-aggregate), respectively. Regardless to the formation and fracture of chemical bond, the potential barrier of the mechanochromism induced by physical factors is relatively low and the process is usually reversible. For instance, most of the AIE/AIEE materials show higher emission efficiency in crystal state but the efficiency becomes lower accompanying with red-shifted emission spectrum when crystals are transformed into amorphous solids by mechanical forces such as grounding and shearing. These feature changes can fully recovered by appropriate thermal annealing and/or vapor fuming.^{1b,4} This essence allows organic AIE/AIEE solids to be used as smart action elements such as mechano-sensors, security papers, and data storage devices. To this end, the softness and reversibility are highly desirable. But the evident reversibility and

softness make such kind of organic solids not stable enough sustain long term storage or to endure stronger external stress. In fact, the amorphous state of organic solids is metastable and it is destine to enter the most stable crystal state in proper conditions or in period that is long enough. Therefore, it is a challenging task postpone the spontaneous transition from amorphous to crystal states

Static attraction between cationic and anionic species is strong r than van der Waals interaction between uncharged species. As the two opposite poles, most inorganic salts show high hardness ar melting point, while some organic salts can be ionic liquid at root temperature. These features suggest that, with rationally designed size and symmetry, the ionic compounds constructed by larg organic cations and small inorganic anions are expected to keep the balance between the rigidity of ionic solid and the flexibility covalent solid.

The strategy to take the advantages of ionic interaction to enhance the emission of organic luminogens has been reported by F. Fery-Forgues and colleagues. 4-Hydroxy-7-nitrobenzoxadiazole phenola anions and tetrabutylammonium bromide cations formed **Chart 1.** Molecular structures of (*E*)-BPHTATPE, $[TPE-Py-Me]^+[PF_6]^2$, ar (E)-TPE-DPy-DMe]²⁺[PF₆]².



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hydrophobic salt, which readily crystallized in the medium and was accompanied by a strong increase in fluorescence intensity.⁵ Lately, this strategy was carried forwarded and expanded by different research groups in fluorescent turn-on detection of ionic chemicals and charged biomacromolecules, such as heparin and DNA.⁶⁻⁸

Recently, we reported an ion pair containing tetraphenylethene (TPE) derivative, [TPE-Py-Me]⁺[PF₆]⁻ (Chart 1).^{9, 10} It is a typical AIEE molecule and showed evident mechanochromic effect. Like other uncharged TPE-derivatives, the solid-state emission of [TPE- $Py-Me]^{+}[PF_{6}]^{-}$ was reversibly switched between green and yellow color by grinding-fuming and grinding-heating processes with a high contrast due to the transformation from the crystalline to the amorphous state and vice versa. Despite the similarity, the chromic switching occurred more difficult than those uncharged TPEderivatives. For example, the emission of the ground powder of $[TPE-Py-Me]^+[PF_6]^-$ recovered by fuming with acetone vapor for 10 min or heating at 150 °C for 10 min. Another charge-free TPE derivative BPHTATPE displayed excellently reversible mechanochromic behaviours. Heating of the ground or pressurized sample at 120 °C for a short while (1 min) or fuming of the ground sample with the vapor of polar solvents in 3 min could recover its emission color, although BPHTATPE has larger molecular weight than [TPE- $Py-Me]^{+}[PF_{6}]^{-11}$ Based on these comparative results, we realize that the introduction of ionic interaction is helpful to acquire more robust mechanochromic effect and non-volatile emission colour change.

Herein, we report our attempt to enhance the robustness of the mechanochromism by introducing more ionic species into an AIEactive molecule. As shown in Chart 1, it is a homolog of [TPE-Py- $Me^{+}[PF_{6}]$, in which the AIE-gen, cationic and anionic moieties are TPE, pyridinium and hexafluorophosphate ([PF₆]), respectively. This compound ($[TPE-DPy-DMe]^{2+}[PF_6]_2^{2-}$) demonstrates typical AIEE and mechanochromic behaviors as most TPE derivatives have shown. But the experimental data reveal that the transition between the highly efficient yellow emission of the crystalline and the moderate red emission of amorphous $[TPE-DPy-DMe]^{2+}[PF_6]_2^2$ solid is irreversible by simple treatments of thermal annealing and/or solvent vapor fuming, whereas the ground sample can recover its emission only by recrystallization from dissolving the amorphous solid to proper solvents. These behaviours are evidently distinct from the phenomena observed for [TPE-Py-Me]⁺[PF₆]⁻ and other uncharged TPE derivatives.

The synthetic route towards $[TPE-DPy-DMe]^{2+}[PF_6]_2^{2-}$, the experimental procedures and the structure characterization data are described in Electronic Supplementary Information (ESI, Schemes S1, S2 and Fig. S1-S9). In brief, the dibromo-modified TPE was derived from a McMurry coupling reaction and the resultant was a mixture of E/Z isomers. The mixture was directly used in the successive Heck reaction, through which the pyridine moieties were linked to the TPE core via C=C double bonds. Heck reaction allowed both of the C=C bridges to be in sole trans-configuration, which has been confirmed by the characteristic chemical shifts and coupling constant in ¹H NMR spectrum (Fig. S1). The E and Zisomers of the pyridine-modified TPE (TPE-DPy) could be readily separated by using silicon gel column chromatography with the mixture of petroleum (60-90 °C) and tetrahydrofuran (5:1 by volume) as eluent, and the E-isomer was the predominant resultant in a total yield of 37%. Then, the pure E-isomer of TPE-DPy was used to react with iodiomethane at room temperature to give rise to [TPE-DPy- $DMe]^{2+}[I]_2^{2-}$, which is an organic salt. This salt emits yellow fluorescence in crystalline state and has a relative low quantum efficiency ($\Phi = 0.8\%$), due to the presence of iodine anions, which partially quenched the fluorescence. Therefore, we replaced the iodine anions by [PF₆]⁻ anions and got the target compound [TPE-DPy-DMe]²⁺[PF₆]₂²⁻ in a good yield of 96%. The as-prepared

crystalline powder of $[TPE-DPy-DMe]^{2+}[PF_6]_2^{2-}$ emits strong yellow-greenish fluorescence with a emission peak (λ_{em}) at 558 n p and an evidently enhanced quantum efficiency (Φ) of 43% (Fig. S10). The characterization data indicate 10 that ${}^{39/2}$ [TFFE-DFFC] DMe]²⁺[PF_6]_2²⁻ molecule takes a *trans*-configuration and the following experimental results are based on this *E*-isomer [(*E*)-TPF DPy-DMe]²⁺[PF_6]_2²⁻.



Fig. 1 (A) Fluorescence (FL) spectra of [(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ In acetone/hexane mixtures with different hexane fractions ($f_{\rm H}$). (B) Plot c dependent peak FL intensity, where I_0 and I are the peak FL intensity at $f_{\rm H} = v$ and other fractions, respectively; and the data are extracted from (A). [4], TPE-DPy-DMe]²⁺[PF₆]₂²⁻ concentration : 10⁻⁵ M. Excitation wavelength (κ_{ex}). 395 nm. Inset of (B): FL images of [(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ in acetone/hexane mixtures taken at $f_{\rm H} = 0\%$ and $f_{\rm H} = 90\%$. λ_{ex} : 395 nm.

[(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ is soluble in acetone but insoluble in hexane. Thus we used acetone and hexane as solvent and norsolvent to examine the AIE behaviours of the salt and the results a shown in Fig. 1. In dilute acetone solution $(1 \times 10^{-5} \text{ mol/L})$, it shows faint emission when excited at a wavelength of 395 nm (λ_{exs} , u =maximum of the salt's UV-vis absorption spectrum, see Fig. S11). keeps weakly emissive in the acetone/hexane mixtures with increasing the hexane fraction (f_{H} , by volume) to 60%, and λ_{em} appears at around 600 nm. When the f_{H} is up to 70%, the mixture becomes evidently emissive and λ_{em} blue-shifts to about 586 nm. Further increasing f_{H} up to 80% and 90%, the emission intensit grows higher and λ_{em} localizes at around 586 nm. The f_{H} dependent fluorescent behavior is displayed in Fig. 1B, which indicates the satisfies a typical AIEE-active molecule.

The blue-shifted fluorescent spectrum can be ascribed to the solvatochromic effect of the [(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ molecu¹ This effect usually occurs to fluorogens bearing electron donor (2) and acceptor (A) moieties.¹² Such fluorogens usually show red-shifted fluorescence spectrum and concomitantly decreased fluorescence intensity as the polarity of the environment enhance . In a [(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ molecule, the TPE core and th



Fig. 2 (A) Variations of the emission wavelengths of [(E)-TPE-DP DMe]²⁺[PF₆]₂²⁻ solids in grinding-fuming (blue trace) and grinding-recrysta - lization (red trace) cycles. (B) X-ray diffraction patterns for [(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ of as-prepared microcrystals (upper), ground powders (midd', and re-crystalline sample (lower).

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pyridinium units play the roles of D and A, respectively. Thus it should have typical solvatochromic behaviour. Indeed, this behaviour has been confirmed by monitoring the absorption and emission spectra of [(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ in a series of organic solvents with different polarities. As displayed in Fig. S12, the emission maximum shifts from 565 to 613 nm when the solvent changes from non-polar dioxane to highly polar ethanol.

Lumpy solid was obtained after the solvents were fully evaporated, and it was used to investigate the mechanochromic behaviours. The as-synthesized [(E)-TPE-DPy-DMe]²⁺[PF₆]²⁻ is a light vellow solid with a strong yellow-greenish emission peaked at ~560 nm (Fig. 2A). After the lumpy solid was ground into fine powder with a pestle or shearing with a spatula, the emission peak shifts to around 605 nm, showing a red-shift of 45 nm. The emission peak exhibits a trend of monotonous blue-shift in grinding-fuming cycles. This phenomenon can be associated with the accumulation of fuming effect.¹³ The fluorescence intensity of the ground $[(E)-TPE-DPy-DMe]^{2+}[PF_6]_2^{2-2}$ powder is much weaker than the untreated sample (ref the images in Fig. 3). Quantitatively, the quantum efficiency drops from 43% to 18% by grinding treatment. This is a typical mechanochromic behaviour that has been observed for a variety of AIE/AIEE compounds.¹⁻³ The change of emission colour is associated with the changes of the molecular packing in the solids before and after grinding. X-ray diffraction patterns of [(E)-TPE-DPy-DMe]²⁺[PF₆]². samples indicate that the as-prepared and ground solids are in polycrystalline and amorphous states, respectively (Fig. 2B).



Fig. 3 (A) Emission peak (colour) changes of [(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ samples upon grinding the as-prepared microcrystals, heating the ground powders at different temperatures ($(120 \circ C, 1 h; (140 \circ C, 1 h; (160 \circ C, 1h; (180 \circ C, 2h)))$ and recrystallization of the ground powders in DCM solution. (B) Photographs of the light emission (λ_{ex} = 365 nm) from the samples of as-prepared crystalline, ground powder, DCM vapor fuming, thermal annealing, and recrystallization from DCM.

It has been found that a variety of uncharged TPE-based molecules exhibit a solvent-dependent vapochromic effect.²⁻⁶ Their ground samples are sensitive to volatile polar solvents. After exposure to the solvent vapor for a while, the red-shifted emission of the ground samples are largely reverted back to the original emission of the as-prepared sample, owing to the fuming-induced recrystallization process. We investigated the vapochromic effect of [(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ by exposure the ground sample to dichloromethane (DCM) vapor for ~5 min. But the emission had not been recovered to the original yellow-greenish. Using the first cycle as an example, the grinding resulted in a red-shift of emission peak from 558 to 605 nm. The DCM vapor fuming for 5 min could only lead to a blue-shift of emission peak from 605 to 598 nm. Extending the fuming time to 10 min brought about little change in the emission features. However, when the amorphous ground powder was redissolved in DCM and dried, the emission intensity and colour reversed almost to the solid before grinding. In the first cycle, the emission peak shifted from 605 to 565 nm. The XRD-pattern recorded from the recrystallization sample (Fig. 2B) indicated a crystalline state that is similar to the as-prepared solid.

According to the thermodynamic principle, the amorphous ground sample is in a metastable state, thus it will transforms back to the

thermo-dynamically stable crystalline state. This transformation can be achieved by thermal annealing. To fix the annealing temperatur the thermal transition property was evaluated by takinguithermad gravimetric analysis (TGA) and differential scanning colorimet (DSC) measurements at a scan rate of 10 °C/min under nitrogen flo (Fig. S13 and S14). The decomposition temperature of $[(E)-T^{D}F]$ DPy-DMe]²⁺[PF₆]₂²⁻ powder is determined to be 340 °C. The a prepared crystalline solid shows an endo-thermal transition peak 265 °C and immediately follows an exo-thermal transition peak around 274 °C. The first peak can be assigned to the melting point the microcrystals and the second may be associated with the recrystallization of the melt just formed. After grinding, the fire powder only has an exo-thermal transition at around 263 °C, which is tentatively assigned to amorphous to crystal transition. The samp... recrystallized from the ground powder in DCM shows thermal transition behaviour similar to the as-prepared microcrystals, having an endo-thermal transition peak at 252.7 °C and immediately follov s an exo-thermal transition peak at 268.5 °C. The small decrease in the melting point can be explained by the fact that the recrystallize sample has a relatively lower degree of crystallization, which is accordance with what is revealed by XRD-patterns (Fig. 2B).

Based on the thermal evaluation results, we investigated the end of thermal annealing on the mechanochromic behaviours of the ground [(E)-TPE-DPy-DMe]²⁺[PF₆]2²⁻ sample. As shown in Fig. 5, the emission peak undergoes a blue-shift of about only 2 nm after annealing at 120 °C for 1 h. In a typical run of elevating the annealing time for 1 h, the blue-shift of the emission peak is recorded to be 6, 3, and 6 nm, respectively. Extending the thermal annealing at 180 °C to 2h, the emission peak shift is about 12 nm. The larger spectral shift at higher temperature and for longer time the reasonable because of the metastable state of the ground sample. All of these spectral shifts are quite small if compared with that to be compared with the transformation of the spectral shift between the ground and recrystallize samples (> 40 nm).



Fig. 4 Schematic illustration of the irreversible mechanochromic process occurred to [(E)-TPE-DPy-DMe]²⁺[PF₆]₂⁻² salt. The ionic interaction holds the luminogens together and bestows the amorphous solid robustness to endure the solvent vapor fuming and thermal annealing treatments.

The above experimental results indicate that the mechanochrom. behaviour of [(E)-TPE-DPy-DMe]²⁺[PF₆]²⁻ is irreversible by simple solvent vapor fuming and thermal annealing treatments. But tl 2 emission features can only be reverted from the mechanically treated (ground) sample to the original (as-prepared) one by recrystallizatio treatments. This behaviour is evidently distinct from the reversible mechanochromism observed for many uncharged TPE-bas(1 molecules reported in recent literature. Considering that the basic difference between the previously reported molecules and he present [(E)-TPE-DPy-DMe $]^{2+}$ [PF₆] $_{2}^{2-}$ focuses on the existenc or absence of the sufficient static charges in molecules, we ascribe lowered reversibility and enhanced robust mechanochromic effect of [(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ to the static interaction between the charged species. On one hand, the static interaction has 1 directionality and saturability, thus each of the negatively charged $[PF_6]^-$ is an attractive center to surrounding positively charged $[(E)^-]$ TPE-DPy-DMe]²⁺ species and vice versa. The ionic interaction

between net charges, on the other hand, is stronger than the van der Waals interaction. In addition, the unmatched sizes between the positively and negatively charged species make the amorphous solid more stable, as demonstrated by ionic liquids. Consequently, the amorphous ground sample of [(E)-TPE-DPy-DMe]²⁺[PF₆]₂²⁻ exhibits extraordinary stability.

In summary, we have shown that the modification of TPE fluorogen with ionic groups allows the derived TPE-based salt to hold the AIEE activity and to achieve effectively enhanced mechanochromic robustness. With only weak van der Waals interaction, the molecular solid of TPE-derivative exhibits highly reversible mechanochromic behaviours under moderate thermal annealing temperature and shorter solvent vapor fuming time. Introducing ionic species into TPE-derivative, [TPE-Py-Me]⁺[PF₆]⁻ exhibits reversible mechano-chromic behaviours under elevated annealing temperature and longer fuming time. Enhancement of the ionic strength, $[TPE-DPy-DMe]^{2+}[PF_6]_2^{2-}$ exhibits irreversible mechanochromic behaviours under elevated annealing temperature and longer fuming time. The emission features can only be reverted from the mechanically treated (ground) sample to the original (asprepared) one by recrystallization treatment. The distinct behaviours are reasonably ascribed to the static interaction between the cationic and anionic species, which bestows the solid with excessive stability in comparison with uncharged TPE-derivatives. The present work gives a hint to rationally design organic photo/electronic materials and to acquire the ingenious balance between the robustness of ionic solids and the flexibility of covalent solids.

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Electronic Supplementary Information (ESI) available: Synthesis and characterization, materials, procedures of all experiments; additional absorption/emission spectra; photographs of fluorescence emission; TGA and DSC curves. See DOI: 10.1039/c000000x/

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Introducing ionic interaction into soft tetraphenylethenebased AIE-molecule furnished the compound with robust mechanochromic effect while holding aggregation-induced emission enhancement property.

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