


Cite this: *Mater. Adv.*, 2021,  
2, 6418

## Variation in solvato-, AIE- and mechano-fluorochromic behavior for furanyl and thiophenyl-substituted anthranil $\pi$ -conjugates: the role of tiny flanking donor groups†

Madhuparna Chakraborty and Manab Chakravarty \*

Only a few reports exist on thiophene or furan-linked typical tetraphenylethene or difluoroboron systems known as aggregation-induced emission-active fluorogens (AIEgens) with a substantial difference in solvato and mechano-fluorochromic (MFC) features. With a pressing demand for finding a smart strategy to achieve MFC-active materials, we herein report unsymmetrically substituted anthracene-vinyl-phenothiazine  $\pi$ -conjugates (**TAPTZ** and **FAPTZ**) linked with furan or thiophene as a tiny flanking donor group. Both the compounds are easily synthesized using an economic path by avoiding metal catalysts and harsh reaction conditions. In solvatofluorochromic studies, by varying polarity of solvents from hexane to acetonitrile, the bathochromic shifts of 118 nm (3082 cm<sup>-1</sup> Stokes shift) for **TAPTZ** and 112 nm (2840 cm<sup>-1</sup> Stokes shift) for **FAPTZ** are noticed. Such electron-rich but conformationally twisted molecules create a twisted intramolecular charge transfer (TICT) state responsible for the solvatofluorochromic feature. A relatively more significant change in the excited state dipole moment for **TAPTZ** causes the difference. Both these molecules exhibit blue-shifted AIE features (faint orange to intense yellow), where **TAPTZ** has appeared to be a relatively better AIEgen with a 65 nm average particle size. Interestingly, **FAPTZ** has emerged as a stronger emitter than **TAPTZ** in the solid state. By grinding in a mortar and pestle or by quick pressing (in an infrared pellet maker: 20 MPa), **TAPTZ** and **FAPTZ** display reversible MFC features with a 15 nm and 22 nm redshift, respectively. Single crystal study discloses the difference in the molecular twisting and packing in the solid-state for these analogs. A relatively large number of intramolecular interactions (dominated by S...S and C...S) make the crystal packing stronger for **TAPTZ** and make such a difference in the sensitivity. Thus, we found a decent impact of the flanking donor groups thiophene/furan on the emission behavior under different environments. The observed fact is further supported by powder X-ray diffraction and lifetime measurement studies. Also, the intermolecular interactions are quantified by Hirshfeld-surface analyses to validate the empirical facts further. Finally, **FAPTZ** is documented as a favorable platform for rewritable optical-recording/security-based applications.

Received 29th July 2021,  
Accepted 28th August 2021

DOI: 10.1039/d1ma00664a

rsc.li/materials-advances

## Introduction

The urge for the design and synthesis of stimuli-responsive small molecules offers ample demand in academics and research on security displays, sensors, and memory-based applications.<sup>1</sup> Amongst them, mechano-fluorochromic properties (MFC) have gained substantial attention due to the generation of admirable multiple colorations just by grinding or

rubbing the compound without performing cumbersome synthetic modifications.<sup>2</sup> There are multiple small organic molecular systems discovered with stimuli-responsive properties.<sup>2b,3a,b</sup> One of the essential needs for MFC-active solid-state fluorophores is to attain twisted conformation offering numerous intermolecular interactions.<sup>4</sup> In such a context, small heterocycles such as furan and thiophene exhibited a wide difference in MFC and electronic properties.<sup>5-8</sup> Nevertheless, thiophene and furan are generally considered weak electron-donors and are explored commonly in the donor- $\pi$ -acceptor like systems.<sup>6</sup> Mostly, thiophene has been established as a promising donor in semiconductor/transistor-based applications.<sup>7,8</sup> In contrast, the related investigations with furan are inadequate. However, recent studies revealed that

Department of Chemistry, BITS-Pilani Hyderabad Campus, Jawahar Nagar, Shameerpet, Hyderabad-500078, Telangana, India  
E-mail: manab@hyderabad.bits-pilani.ac.in

† Electronic supplementary information (ESI) available. CCDC 2100035 and 2100036. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/d1ma00664a



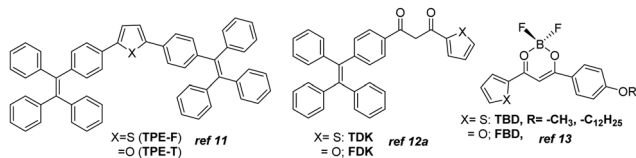


Fig. 1 Known furan/thiophene substituted MFC-active compounds.

the furan is also a favorable building block for generating semiconductors<sup>9a,b</sup> and has better charge carrier mobility.<sup>10</sup> The greater extent of biodegradation, smaller size, and high planarity make the choice of furan more favorable despite having relatively inferior aromaticity and polarizability features compared to thiophene. In particular, a household AIE-framework tetraphenylethene (TPE)-linked furan **TPE-F** (Fig. 1) was found to be superior in exhibiting chromic behavior to that of analogous **TPE-T** under external stimuli.<sup>11</sup> Additionally, the furan/thiophene-linked  $\beta$ -diketo building blocks were incorporated with the TPE system (**TDK/FDK**) and a variation in solvatofluorochromism, AIEE-behaviour, and (non)reversible MFC properties was identified.<sup>12</sup> Apart from the TPE system, the other difluoroboron  $\beta$ -diketonate was attached to thiophene or furan (**TBD** and **FBD**) and recognized as a reversible stimuli-responsive material with subtle differences between thiophene and furan analogs. The extent of molecular planarity, intermolecular interactions, and the crystal packing density were found to play a role behind the variation in the emission color.<sup>11–13</sup>

Thiophene or furan resides almost in the molecular plane with a slight twisting angle in all the reports above. However, the impact on emission color variation from the furan/thiophene analogs upon solvent or external pressure stimuli looks highly promising. Nevertheless, few molecules are only reported on the comparative solid-state and solvatofluorochromic features between furan and thiophene-conjugated analogs. In our current research on unsymmetrically substituted anthracene-based  $\pi$ -conjugated fluorophores, we noticed that a highly planar anthracene system could avoid disc-like molecular packing in the presence of flanking aryl/heteroaryl mainly due to the conformational variation (twisted at different torsion angles), resulting in the formation of AIEgens.<sup>14</sup> The van der Waals interaction between the anthracenyl peripheral H's and these groups creates such twisted stable conformation. In such a system, thiophene analogs appeared to be better AIEgens than furan.<sup>15</sup> However, solvato- and mechano-fluorochromic features were not promising for those molecules. Based on our interest in creating multi-stimuli-responsive materials,<sup>16</sup> we herein focus on introducing anthracene- $\pi$ -phenothiazine conjugates linked with furan or thiophene as flanking donor groups (Fig. 2).

The electron-rich and bulky phenothiazine unit is chosen due to its capability for forming weak C-H...S/S... $\pi$ /C-H... $\pi$  intermolecular interactions and the nonplanar structure of such a core would favor exhibiting AIE-features by avoiding the cofacial  $\pi$ ... $\pi$  stacking. Moreover, highly electron-rich phenothiazine can create a variation in the electronic

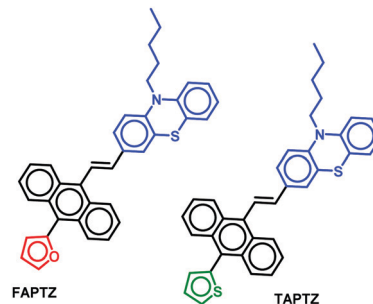


Fig. 2 Thiophene/furan-flanked anthracene- $\pi$ -phenothiazine molecules.

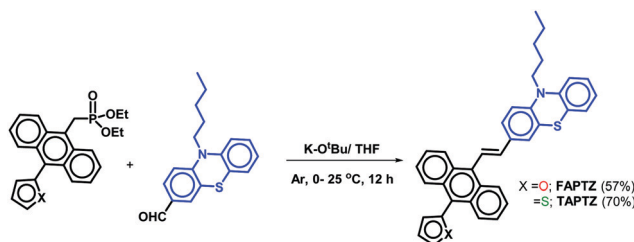
distribution within the system where the anthracenyl part would be relatively electron-poor. Thus, the system would display the solvatofluorochromic properties. The impact of flanking small heterocycles into this system would draw special attention. AIEgens can emit in the solid-state, and the possible conformational changes under mechanical force can offer MFC-feature. We found dissimilarity in terms of the emission behavior in the solution and aggregated and solid-state due to such a tiny difference in molecular structure (thiophene/furan). Thus, the variation in electronic feature, size factor and aromaticity between thiophene and furan can govern such a subtle disparity in stimuli-responsive emission feature for these unsymmetrically substituted  $\pi$ -conjugates. Besides, the difference in intermolecular interactions has also created a contrast in the solid-state emission features. All the observed properties are analyzed with the help of molecular structure analyses, powder X-ray diffraction (PXRD), and fluorescence lifetime studies.

## Results and discussion

### Synthesis and characterization

The established synthetic procedure finds a broad scope to tune the substitution effect; the planned molecules **FAPTZ** (furan-flanked) and **TAPTZ** (thiophene-flanked) are conveniently synthesized in a decent yield by simple, inexpensive, and efficient Horner–Wadsworth–Emmons (HWE) reactions at room temperature (*i.e.*, energetically economic) from easily accessible phosphonates (Scheme S1, ESI<sup>†</sup>) and phenothiazine-carbaldehyde (Scheme 1).

An expensive metal/ligand combination with high-temperature requirement is escaped to access such molecules



Scheme 1 Synthesis of **FAPTZ** and **TAPTZ**.



where thiophene/furan units are introduced *via* Friedel–Crafts arylation reaction.<sup>17</sup> The reaction can be scaled up on a g scale, indicating easy access to these materials for real-world applications. These compounds are soluble in most organic polar/nonpolar solvents and well-characterized by IR, NMR (<sup>1</sup>H/<sup>13</sup>C), and mass-spectroscopic studies. Finally, the molecules are crystallized by the room-temperature crystallization method, and the molecular structure with (*E*)-configuration is explicitly established by single-crystal X-ray diffraction studies (SCXRD; the details are discussed later). The phenothiazine unit's presence raises the thermal stability of the  $\pi$ -conjugates that show stability up to 400 °C and beyond as confirmed by thermogravimetric analysis (TGA, Fig. S1, ESI†).

### Solvatochromic studies

Primarily, the photophysical behaviors in the solution state were examined for these  $\pi$ -conjugates by changing the nature of the solvent with different polarities. A sharp maximum absorption at  $\lambda_{\text{abs}} \sim 405\text{--}410$  nm was detected for both the analogs in absorption spectroscopy (see Fig. S2, ESI† for all the absorption and emission spectra) due to  $\pi\text{--}\pi^*$  transition, and there was no impressive shift observed by changing the polarity of the solvent, indicating an unaffected ground state. A slightly higher  $\lambda_{\text{abs}}$  for the furan analog dictates a relatively better conjugation than that of thiophene. The emission spectra reveal a cleaner discernible coloration for **TAPTZ** than **FAPTZ** in most of the solvents of different polarities (Fig. 3). The emission at relatively longer  $\lambda_{\text{max}}$  for furan (**FAPTZ**) has further supported a more planar molecular structure for the furan analog. By varying  $E_{\text{T}}(30)^{18a}$  [a measure of microscopic solvent polarity] of different solvents, a promising positive solvatochromic effect (redshift) is noticed with both the compounds. The solvatochromic range between hexane (nonpolar) and MeCN (polar) is marginally better with a 118 nm redshift ( $3082\text{ cm}^{-1}$  Stokes shift) for **TAPTZ** than **FAPTZ** displaying a 112 nm redshift ( $2840\text{ cm}^{-1}$  Stokes shift). All the photophysical parameters, including the solvatochromic shifts for both the compounds, are listed in Table 1.

The overall difference of  $242\text{ cm}^{-1}$  between **TAPTZ** and **FAPTZ** in the Stokes shift (between hexane and MeCN) and

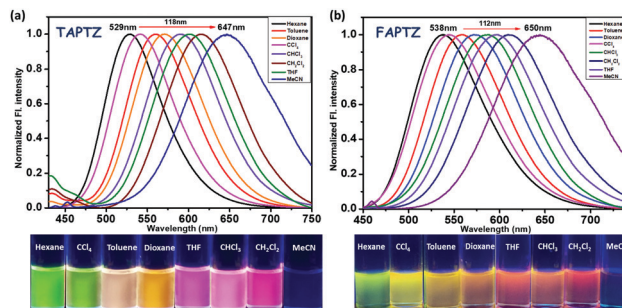


Fig. 3 Normalized emission spectra of (a) **TAPTZ** and (b) **FAPTZ**, in different solvents. Concentration of the probe:  $10\ \mu\text{M}$ ,  $\lambda_{\text{ex}} = 405$  nm. The image is taken under  $365$  nm UV light.

the considerable color variation are noteworthy (Fig. 3 and Table 1). The redshift may be attributed to the presence of the TICT (twisted intramolecular charge transfer) state, where the phenothiazine unit can be noted as an electron donor and anthracene as an electron acceptor. The role of significant twisting arises from the flanking thiophene and furan groups. The electronic structure is confirmed by performing a DFT study (Density Functional Theory) on these molecules. Ground-state optimization using DFT at the CAMB3LYP-6-31G(d,p) level identifies that the electron clouds in the highest occupied molecular orbital (HOMO) level are majorly located on the phenothiazine core (owing to the presence of two heteroatoms), while the lowest unoccupied molecular orbital (LUMO) level is predominantly placed over the anthracene ring with a slight contribution by flanking and phenothiazine groups (Fig. 4). Such electronic distribution can also support the ICT feature in the molecules. This is also consistent with the observed solvatochromic properties.

The LUMO and HOMO are relatively more stabilized for furan analogs, and the lower energy gap can cause longer wavelength absorption and emission for **FAPTZ**. The relatively lower quantum yields can be attributed to the presence of heavier sulfur atoms for **TAPTZ**. The DFT-optimized molecular structures demonstrate a more important  $\pi$ -conjugation for **FAPTZ** with a  $-67^\circ$  furan-anthracene torsion angle than **TAPTZ**

Table 1 Photophysical parameters of the probes in different solvents

Solvent	Abs $\lambda_{\text{max}}$ (nm) <b>TAPTZ</b> / <b>FAPTZ</b>	Emi $\lambda_{\text{max}}$ (nm) <b>TAPTZ</b> / <b>FAPTZ</b>	Stokes shift $\bar{\nu}_{\text{A}} - \bar{\nu}_{\text{F}}\text{ cm}^{-1}$		Quantum yield $\phi_{\text{f}}$ (%) <b>TAPTZ</b> / <b>FAPTZ</b>	Dipole moment	$E_{\text{T}}$ (30)
			<b>TAPTZ</b>	<b>FAPTZ</b>			
Hexane	402/404	529/538	5972/6165	14/13	0.08	31.0	
CCl <sub>4</sub>	409/413	541/553	5966/6130	8.8/13	0	32.5	
Toluene	406/412	560/566	6774/6604	10/10	0.43	33.9	
1,4-Dioxane	407/412	571/579	7057/7001	8.9/7.2	1.3	36.0	
THF	407/413	602/605	7959/7684	5.7/19	1.75	37.4	
CHCl <sub>3</sub>	408/412	590/595	7561/7465	5.7/22	1.15	39.1	
CH <sub>2</sub> Cl <sub>2</sub>	409/412	615/618	8190/8091	3.2/7.5	1.60	40.7	
Acetonitrile	408/410	647/650	9054/9005	0.32/6.4	3.45	45.6	

$\bar{\nu}_{\text{A}} - \bar{\nu}_{\text{F}}$  ( $\text{cm}^{-1}$ ): the wavenumber difference between absorbance and fluorescence. Relative quantum yields are measured using quinine sulfate as the standard (see the Experimental section).



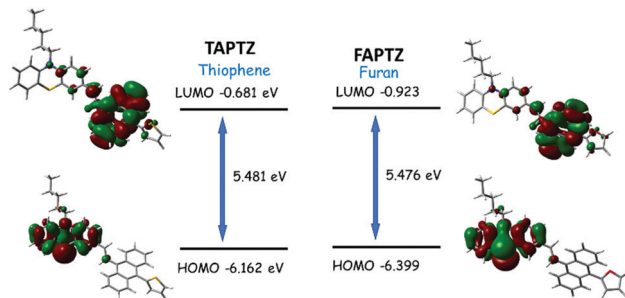


Fig. 4 Optimized [using the CAMB3LYP-6-31G(d,p) level] molecular structure of **TAPTZ** and **FAPTZ** with the electronic distribution in the HOMO and LUMO.

where thiophene is oriented in almost orthogonal ( $-88^\circ$ ) geometry with anthracene (Fig. S3, ESI<sup>†</sup>). The interplanar angles ( $58-66^\circ$ ) between anthracene and phenothiazine units are practically equal for **FAPTZ** and **TAPTZ** (Fig. S3, ESI<sup>†</sup>), indicating nearly identical coplanarity.

Moreover, a largely twisted conformational structure can create a more polar and energetically stabilized TICT state in the polar solvent, resulting in poor quantum yield with a high redshift.<sup>18b</sup> The plot of  $E_T(30)$  vs. Stokes shift and emission  $\lambda_{\max}$  (Fig. S4 and S5, ESI<sup>†</sup>) also supports the possible formation of the TICT state. The TICT state is typically more convincing in the typical D-A system, but the generation of the TICT state from such electron-rich and conformationally twisted systems is possible where phenothiazine acts as the donor and anthracene as an acceptor.<sup>19</sup> Moreover, anthracene and phenothiazine cores are coupled together through a *trans*-vinylidene spacer. Twisted conformation arises from the interactions between peripheral H-atoms of vinylidene and anthracenyl systems. However, the fluorescence (FL) intensity enhancement upon increasing the viscosity of the solvent medium (glycerol in methanol, Fig. S6, ESI<sup>†</sup>) indicates a substantial twisting in the molecular structures with such bulky rotors and creates the formation of a TICT state.<sup>18b</sup> The TICT state formation can also be favored due to the reversible first oxidation potential of phenothiazine upon photoexcitation and subsequent intramolecular rotation.<sup>20</sup> Although the conventional Lippert–Mataga equation might be inappropriate to apply for such a molecular shape, we have examined the Lippert–Mataga plot [Fig. S7 and Table S1, ESI<sup>†</sup>]. The Lippert–Mataga plot [Stokes shift against orientation polarizability ( $\Delta f$ )], reveals a quick rise in the Stokes shift with a polar solvent. It also confirms the enhanced dipole moment in the excited state,<sup>21</sup> showing 14.37 D for **TAPTZ** and 13.91 for **FAPTZ** [Table S1, ESI<sup>†</sup>]. This marginal change in dipole moment for **FAPTZ** validates a slightly better redshift for the thiophene analog.

### AIE-properties

The AIE properties are the most exciting photophysical behavior that has widened the application scope of the fluorophores in the solid/aggregated state and gained attention in the last two decades.<sup>22</sup> Acetonitrile was preferred as a good and

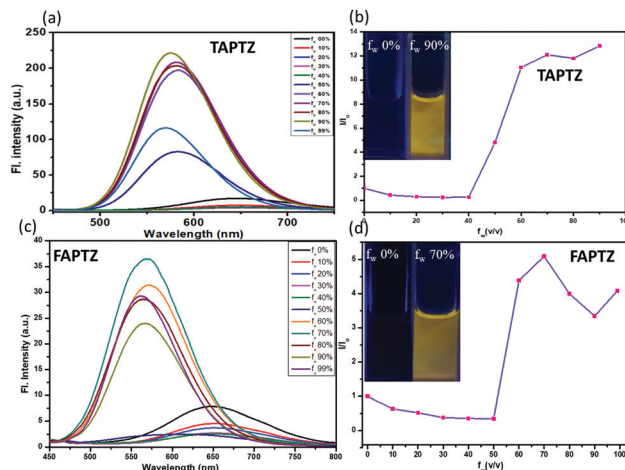


Fig. 5 Emission spectra of (a) **TAPTZ** and (c) **FAPTZ**. (b and d) The  $I/I_0$  plot ( $I_0$ : FL intensity before addition of water;  $I$ : FL intensity after addition of water) [concentration of the probe: 10  $\mu$ M,  $\lambda_{\text{ex}} = 405$  nm]. The image is taken at  $f_w = 0\%$  and  $f_w = 90\%$  (**TAPTZ**) and 70% (**FAPTZ**) under 365 nm UV light at different  $f_w$  in acetonitrile.

water-miscible solvent for these studies due to the weakest emission. The AIE properties were examined by measuring absorption (Fig. S8, ESI<sup>†</sup>) and emission spectra (Fig. 5) for each compound in 10  $\mu$ M acetonitrile solution upon slow addition of water fraction [a bad solvent  $f_w$ (v/v%)]. The reduction of FL intensity was observed initially due to the polarity effect until  $f_w = 40-50\%$  (Fig. 5b and d) for both cases. There was a sudden upturn in the FL intensity for **TAPTZ** from  $f_w = 40\%$ , and the intensity was maximum at  $f_w = 90\%$  with a 73 nm blue-shift (yellow emission at  $\lambda_{\max}$  574 nm). For **FAPTZ**, the FL enhancement started at  $f_w = 50\%$  and reached maximum until  $f_w = 70\%$  with an 81 nm blue-shift (yellow emission at  $\lambda_{\max}$  569 nm). Thus, both these compounds exhibit a similar AEE-effect (aggregation-enhanced emission) in terms of the emission color but more pronounced for **TAPTZ** (14 times enhancement) in comparison to **FAPTZ** (only five times enhancement). The ups and downs in the FL intensity at higher  $f_w$  for **FAPTZ** were observed earlier due to the formation of a nanoparticle suspension with both crystalline and amorphous characters that control the emission intensity in the solution.<sup>14</sup>

The aggregation at the highest FL intensity is recognized by the average size of the particles calculated from dynamic light scattering (DLS) studies that display 65 nm for **TAPTZ** and 85 nm for **FAPTZ** (Fig. S9, ESI<sup>†</sup>). The restricted intramolecular motion during the aggregation was earlier established as the primary cause for AIE behavior.<sup>22</sup> Such a highly twisted conformation with large rotors in an olefinic stator would mostly avoid the co-facial intermolecular aggregation. Thus, the intramolecular motion will be limited to fluoresce through a radiative channel. The torsion angle of  $67^\circ$  for thiophene would favor showing better AIE-properties than the furan-analog. The blue shift indicates the reduction in the molecular coplanarity upon aggregation. Blue-shifted AIE properties were also noticed earlier in a similar system.<sup>23,24</sup> A cross-aggregation (J-type) is found from the crystal packing (*vide infra*), and that causes a lack of



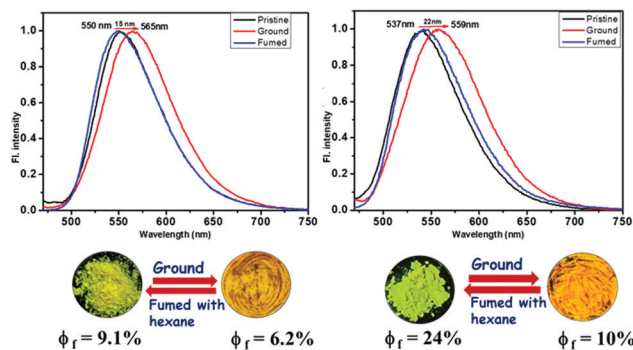


Fig. 6 Pristine and ground samples of **TAPTZ** and **FAPTZ** along with the corresponding (reversible) emission spectra [ $\lambda_{\text{ex}} = 420$  nm]. The images are taken under 365 nm UV-light.

conjugation to have a blue-shifted emission. The smaller particle size is possibly supporting to have more AIE-effect for **TAPTZ**.<sup>25</sup> The observed AIE properties are also competent enough to emit in the solid-state, and such a variously twisted conformation of the molecules will tend to exhibit MFC characteristics and thus, is explored.

### MFC-Studies

Initially, the solid-state photophysical properties are investigated before and after grinding both the pristine samples to examine the change in emission color. The solid-state UV-Vis spectrum displays a broad and substantial absorption in the region of 440 nm, mainly due to  $\pi-\pi^*$  transition for both the pristine samples (Fig. S10, ESI<sup>†</sup>). Upon grinding in a mortar and pestle or applying pressure (20 MPa) using an IR-pellet maker, the change in absorption spectrum is slightly blue-shifted ( $\sim 17$  nm) for **TAPTZ**, but almost no difference for **FAPTZ** (Fig. 6), indicating almost similar behaviour in the ground state. However, the solid-state FL spectrum for **TAPTZ** is noticeably red-shifted from 550 nm to 565 nm (15 nm) after grinding. The absolute quantum yield ( $\phi_f$ , %) for the pristine sample is 9.1, and that was weakened to 6.02 after grinding. Thus, one can visualize the sharp FL switching from light green to yellow emitting solid (15 nm contrast) through the naked eye under 365 nm UV-light after grinding the solid. On fumigation of the ground sample with hexane, the original green color returns. Notably, thermal energy could not bring back the actual green color. On the other hand, the furan analog **FAPTZ** emitted a relatively brighter green light (emission

Table 2 Solid-state photophysical parameters for compounds with absorption/emission wavelength  $\lambda_{\text{abs}}/\lambda_{\text{emi}}$  in nm, absolute  $\Phi_f$  in % (error:  $\pm 2\%$ ), an average lifetime ( $\tau$ ) and the ratio of radiative rate constant ( $K_r$ ) to non-radiative rate constant ( $K_{\text{nr}}$ ).  $K_r = \Phi_f/\tau$ ;  $K_{\text{nr}} = (1 - \Phi_f)/\tau$

Samples	States	$\lambda_{\text{abs}}^{\text{max}}$ (nm)	$\lambda_{\text{em}}^{\text{max}}$ (nm)	$\Phi_f$ (%)	$\tau$ (ns)	$K_r/K_{\text{nr}}$
<b>TAPTZ</b>	Pristine	440	550	9.1	0.65	0.100
	Ground	423	565	6.2	1.16	0.060
<b>FAPTZ</b>	Pristine	436	537	24.0	0.57	0.323
	Ground	438	559	10.0	0.54	0.121

$\lambda_{\text{max}} = 537$  nm) with  $\phi_f$  (%) 24 and switched to 559 nm upon grinding.

Additionally, the excited state's fluorescence lifetime (ns) was determined for solid **TAPTZ** and **FAPTZ** in both the pristine and ground states (Fig. S12 and Table S2, ESI<sup>†</sup>). All the related parameters are stated in Table 2. More  $\Phi_f$  (%) for **FAPTZ** can be elucidated with a relatively better ratio  $K_r/K_{\text{nr}}$  [radiative rate constant ( $K_r$ ) and non-radiative rate constant ( $K_{\text{nr}}$ )]. Grinding the compounds resulted in a decrease in the quantum yield and could be deciphered with lower  $K_r/K_{\text{nr}}$  values (Table 2).

### SCXRD studies

It is essential to examine the single-crystal X-ray structures for these  $\pi$ -conjugates to obtain a molecular-level understanding and investigate the cause for the FL-switching upon grinding. Typically, such FL switching is regulated by the change in the supramolecular assembly. The crystal structures were easily determined with suitably grown single crystals, obtained from 30% EtOAc in hexane through a room-temperature slow evaporation technique. Both the compounds **TAPTZ** and **FAPTZ** crystallized in a triclinic system with a  $P\bar{1}$  space group [see Table S3 (ESI<sup>†</sup>) for all the crystallographic parameters]. Predominantly, the molecular structures determined from SCXRD have presented the fundamental difference between these two molecules in the solid-state (Fig. 7).

Both the structures are significantly twisted in the solid-state and emit through a radiative channel to exhibit solid-state emission properties. Direct attachment of the thiophene or furan unit as a flanking donor causes the torsion angle of  $87.1^\circ$  and  $73.5^\circ$  with anthracene for **TAPTZ** and **FAPTZ**, respectively. A relatively smaller size of the furan ring enables a smaller

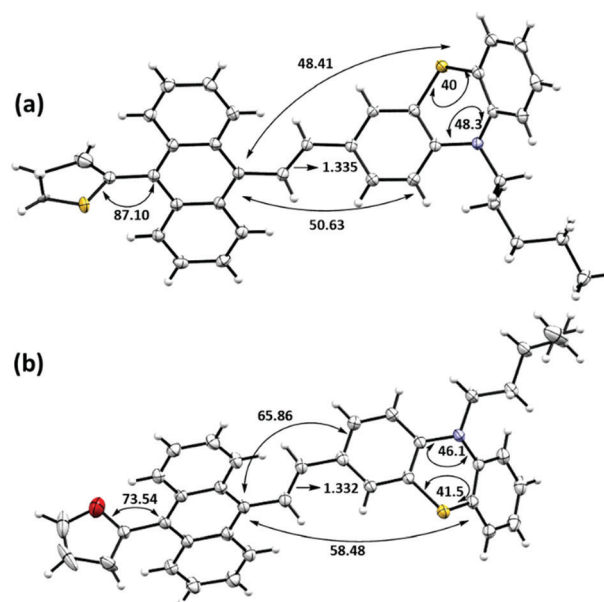


Fig. 7 Single-crystal X-ray structures (ORTEP diagram with 50% probability level) and few selected bond angles ( $^\circ$ ) and distances ( $\text{\AA}$ ): (a) **TAPTZ** (the disordered part is removed for clarity) and (b) **FAPTZ**. **TAPTZ** was solved with a slight disorder in the thiophene ring but refined successfully.



Table 3 Intermolecular interactions (Å) for both the  $\pi$ -conjugates

Compounds	C··H	O··H	S··S	S··H	C··S	H··H
<b>TAPTZ</b> (10 interactions)	2.887, 2.885 2.861, 2.829, 2.803, 2.816		3.524	2.789	3.335	2.399
<b>FAPTZ</b> (8 interactions)	2.833, 2.818 2.898, 2.747 2.865, 2.816	2.680				2.306

torsion angle but does not exhibit the emission at a longer wavelength in the solid state. A further detailed inspection indicates much smaller angles between anthracene and phenothiazine ( $\sim 50^\circ$ ) plane for **TAPTZ** than **FAPTZ** that attains an interplanar angle of  $\sim 66^\circ$ . Such a difference would facilitate more  $\pi$ -electron conjugation in **TAPTZ**, resulting in emission at a higher wavelength. A slightly longer (0.003 Å) olefinic C=C bond also supports better conjugation in **TAPTZ** than **FAPTZ**. It is worth noting that the DFT optimized molecular structures of **TAPTZ** and **FAPTZ** had almost equal coplanarity between anthracene and phenothiazine in the molecular state, and the differences are only observed in the solid-state due to the non-covalent interactions.

Next, we emphasized supramolecular interactions and mode of molecular crystal packing for **TAPTZ** and **FAPTZ**, which offer profound comparisons to recognize the switching of solid-state fluorescence behavior upon grinding.<sup>26,27</sup> The intermolecular interactions for these crystals are designated in Table 3 and Fig. 8. The differences in crystal packing from three different axes are presented in Fig. S13 (ESI<sup>†</sup>). The almost equal number of C··H interactions do not bring much variance in the MFC behavior. However, no  $\pi$ ·· $\pi$  interactions are observed because of such twisted conformations of these molecules, and cross J-type aggregation is perceived. The distances (Å) between two anthracenyl centroids are 9.743 and 5.010 for **TAPTZ** and **FAPTZ**, respectively. However, for **TAPTZ**, the flanking thiophene causes two additional interactions (C··S and S··S) that help to make the crystal packing more compact and can be further supported with a higher melting point for **TAPTZ** (165 °C) in comparison to **FAPTZ** (153 °C).

The **TAPTZ** molecules are oriented so that PTZ–PTZ units are closer, whereas PTZ–furan has appeared to be joined for **FAPTZ** in the crystal lattice (Fig. 8). The C··H interactions for both the compounds enforce more flexibility and are more helpful to exhibit MFC behavior.<sup>28a,b</sup> Nevertheless, a relatively more number of interactions for **TAPTZ** achieve little higher rigidity and higher crystal lattice energy<sup>29</sup> which thus becomes less sensitive. Notably, both these compounds display MFC-feature under grinding/pressure by improving the degree of planarization in molecular conformation that would elucidate the observed redshift.<sup>30</sup> The penetration of hexane vapor into the distorted crystal packing will permit molecules to develop flexibility and rearrange the molecular conformation to return to the pristine crystal packing to induce the authentic emission.

This repeatable fluorescence switching using **FAPTZ** is almost similar (change of  $\pm 10$  nm wavelength) even after multiple grinding/fuming (Fig. S11, ESI<sup>†</sup>) processes. Thus, the

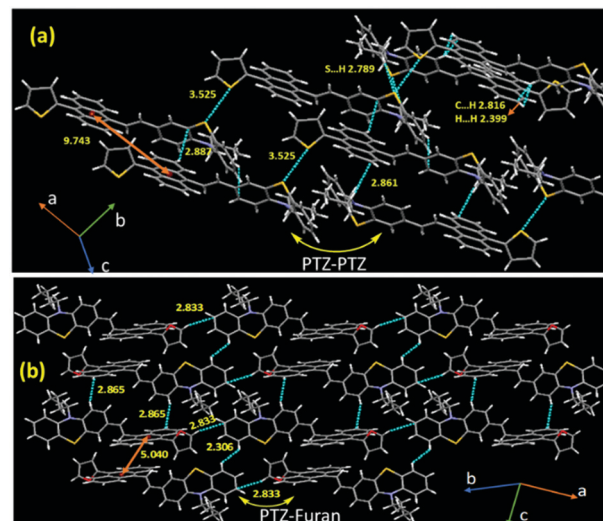


Fig. 8 Crystal packing for (a) **TAPTZ** and (b) **FAPTZ** showing a few intermolecular interactions ( $d/\text{Å}$ ).

material is supportive of numerous cycles of reversible color change upon solvent exposure.

### PXRD analysis

To gain a deeper insight into the above slight variation in MFC-feature, powder X-ray diffraction (PXRD) studies are performed. Both these analogs demonstrate many sharp and intense diffraction signals in the pristine state (Fig. 9), specifying a well-ordered crystalline structure. However, **TAPTZ** has appeared to be relatively more crystalline than **FAPTZ** as deciphered before. Some of the diffraction planes are designated based on the simulated diffraction patterns for both the compounds, and the diffractions are consistent with the

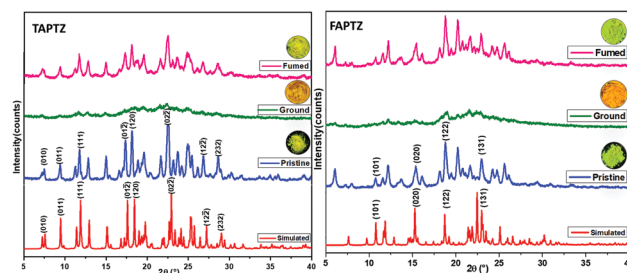


Fig. 9 The PXRD patterns of **TAPTZ** and **FAPTZ** in pristine, ground, and fumed states.



experimentally determined signals. The reversible MFC behavior for both the compounds is supported by the diffractogram for pristine, ground, and fumed samples. The crystal to amorphous state transformation is noticed after grinding because almost no diffraction peaks at the same  $2\theta$  are observed (Fig. 9). The original intense diffractions recur upon fuming with hexane. It depicts a considerable decrease in the crystallinity (transform towards the amorphous phase) after grinding the solid and the crystallinity is restored after fuming and fuming with hexane assists in rearranging the molecules to reach the original crystalline from the amorphous state. Thus, changes in the PXRD pattern support the difference in displaying MFC features for such a phase transition in these molecules.

### Hirshfeld surface analysis

The crystal packing showed the contribution of C···H interactions inside the crystal lattice. The quantitative analyses of intermolecular interactions are performed through the Hirshfeld surface generations (Fig. S14, ESI<sup>†</sup>). A significant amount of C···H type interactions (33.3%) is acting on the **FAPTZ** crystal compared to **TAPTZ** (30.5%). Notably, there are 4.5% C···C interactions for **TAPTZ**, whereas **FAPTZ** shows 3% of such interactions. The higher ratios of C···H/C···C interactions for **FAPTZ** indicate good MFC behavior in comparison to **TAPTZ**.<sup>31</sup> This Hirshfeld analysis is very well aligned with our experimentally observed features and the crystal packing analysis. Thus, the difference in the intermolecular interactions between these two molecules can elucidate the slight difference in MFC-feature for these two analogs.

### Application as an optical rewritable recorder

Further, such MFC-active materials are also recognised as a platform for rewritable optical recording. Such FL-switching is very much appropriate for real-world applications. An economical filter paper-based rewritable optical recorder is created by dispersing **TAPTZ** on the filter paper. Thus, the yellow-emitting surface becomes ready for the writing. The letters are suitably printed on this yellow-emitting surface by pressing with a metal spatula. The letters emit orange and are visible with bare eyes under 365 nm UV-torch (Fig. 10). The orange emitting part can further return to a yellow-emitting surface by fuming with hexane. Thus, this **TAPTZ**-coated filter paper is documented as an encouraging MFC material for repetitive usage in optical recording.

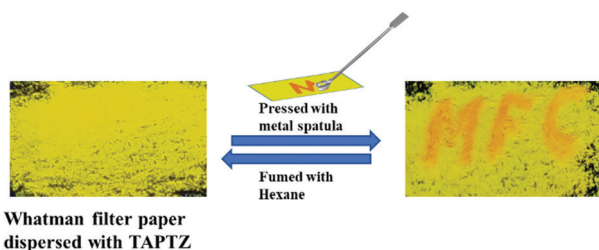


Fig. 10 Application of **TAPTZ** as a rewritable optical recorder.

## Experimental

### Materials

All of the chemicals were purchased from Merck, and the solvents for column chromatography were purchased from Finar. Spectroscopic grade solvents used for UV-vis and fluorescence spectroscopy were obtained from Sisco Research Laboratories (SRL). All the experiments were performed at room temperature,  $298 \pm 2$  K.  $^1\text{H}$ , and  $^{13}\text{C}$  NMR spectra were recorded on Bruker 400 MHz spectrometers with an operating frequency of 101 MHz for  $^{13}\text{C}$ . Chemical shifts ( $\delta$ ) are reported in ppm relative to the residual solvent signal ( $\delta$  7.26 for  $^1\text{H}$  NMR and  $\delta$  77.0 for  $^{13}\text{C}$  NMR).

### Methods and measurements

#### Steady-state absorption and fluorescence measurements.

The solution-state absorption spectra were recorded on a UV-vis-NIR spectrophotometer (Hitachi F7000, Japan), and solid-state absorption spectra were recorded on a JASCO-500 spectrophotometer. The solid-state emission spectra were recorded using a fluorimeter (Fluorolog, HORIBA) and solution-state (FP-6300, Jasco) using a 10 mm path length quartz cuvette. All the solution-state spectra were recorded, keeping the probe concentrations at 10  $\mu\text{M}$  (2 mL) in 3 mL cuvettes for all measurements. The emission spectra were recorded by excitation at their corresponding absorption wavelengths.

**Quantum yield measurement in the solution state.** The relative quantum yield of the sample was measured with a reference quinine sulfate (in 0.1 M  $\text{H}_2\text{SO}_4$ ) using the formula

$$\Phi_f = \Phi_{\text{ref}} \times \frac{a_R}{a_S} \times \frac{A_S}{A_R} \times \left( \frac{\eta_S}{\eta_R} \right)^2$$

where  $\Phi_f$  = quantum yield of the sample,  $\Phi_{\text{ref}}$  = quantum yield of quinine sulfate (0.54),  $a_R$  = area under the curve of emission spectra of quinine sulfate in 0.1 M  $\text{H}_2\text{SO}_4$ ,  $A_S$  = absorbance of the probe in different solvents,  $A_R$  = absorbance of quinine sulfate in 0.1 M  $\text{H}_2\text{SO}_4$ ,  $\eta_S$  = refractive index of the corresponding solvent, and  $\eta_R$  = refractive index of water. The error:  $\pm 10\%$ .

### Viscosity effect

The visochromic properties were studied by measuring absorption and emission spectra (Fig. S6, ESI<sup>†</sup>) for both **TAPTZ** and **FAPTZ** in 10  $\mu\text{M}$  MeOH solution upon incremental addition of glycerol fraction [highly viscous liquid  $f_g$  (v/v%): fraction of glycerol in methanol]. The FL intensities for **TAPTZ** and **FAPTZ** increased linearly with the gradual addition of glycerol ( $f_g$ , v/v%) in MeOH until  $f_g = 60\%$  and there was a sudden enhancement in the FL intensity at  $f_g > 60\%$ .

**Absolute quantum yield measurement.** The absolute QY was determined for the solid samples using a calibrated integrating sphere method with an absolute error of  $\pm 2\%$ .

**Powder X-ray diffraction and IR spectra.** The PXRD was recorded before and after grinding the samples using a Rigaku Ultima IV X-ray diffractometer by keeping the parameters constant for all samples, *i.e.*, step width 0.2, the scan rate of



$2^\circ \text{ min}^{-1}$  from  $5$  to  $45^\circ$  (Cu  $K_\alpha$  radiation  $\lambda = 1.54 \text{ \AA}$ ). The IR spectra were recorded using an FTIR spectrometer (FT/IR-4200, Jasco). Solid samples were mixed with KBr to record the spectra.

**Time resolved measurement.** Time-resolved fluorescence measurements were completed using a time-correlated single-photon counting (TCSPC) unit (Horiba Deltaflex). The pulse diode laser used was  $405 \text{ nm}$ , with a setup target of  $10000$  counts. The solid powder was made as a sandwich in between two quartz slides followed by keeping in a solid-sample holder. The instrument response function was measured before fluorescence lifetime measurements using aluminum foil. All of the decay curves were fitted using the supplied EZ Time software. All measurements were performed at room temperature ( $298 \text{ K}$ ). A magic angle ( $54.7^\circ$ ) configuration was used for all measurements. All fittings were done by keeping the  $\chi^2$  value nearer to  $1$ .

### Dynamic light scattering (DLS) measurement

The average particle size in the aggregated state was found using a Malvern particle size analyzer (zeta sizer nano-ZS), keeping a concentration of  $10 \mu\text{M}$ .

### Thermogravimetric analysis

The TGA thermograms were recorded using Themys One<sup>+</sup>, Setaram instrument keeping the temperature range  $30$ – $650^\circ \text{C}$  with  $10^\circ \text{C min}^{-1}$  under a  $\text{N}_2$  atmosphere.

### Single crystal X-ray measurement

All measurements were performed on a Rigaku XtaLAB P200 diffractometer using multi-layer mirror monochromated Cu- $K_\alpha$  radiation ( $\lambda = 1.54184 \text{ \AA}$ ). The data were collected at a temperature of  $-173 \pm 1^\circ \text{C}$  to a maximum  $2\theta$  value of  $149.8^\circ$ . Data were collected and processed using CrysAlisPro (Rigaku Oxford Diffraction). The linear absorption coefficient,  $\mu$ , for Cu- $K_\alpha$  radiation is  $18.391 \text{ cm}^{-1}$ . An empirical absorption correction was applied, which resulted in transmission factors ranging from  $0.227$  to  $0.593$ . The data were corrected for Lorentz and polarization effects. The structure was solved by direct methods (SIR2011)<sup>32</sup> and expanded using Fourier techniques. The non-hydrogen atoms were refined anisotropically. Hydrogen atoms were refined using the riding model. All calculations were performed using the Olex 2 crystallographic software package except for refinement, which was performed using SHELXL Version 2014/7.<sup>33</sup>

### Density functional theory

The DFT studies were performed on these molecules using a CAM-B3LYP 631-G(d,p) basis set. The HOMO and LUMO energies were calculated, and corresponding cubes were generated using the Gaussian 09 package.

### Hirshfeld surface analyses<sup>34</sup>

Hirshfeld surface analysis and void space calculation is a quantitative tool to understand non-covalent interactions. We have generated Hirshfeld surfaces for **TAPTZ** and **FAPTZ** with an iso-value of  $0.5 \text{ a.u.}$  The surface is generated, and the

interactions are shown in terms of  $d_e$  and  $d_i$ , where  $d_e$  and  $d_i$  are distances of an atom external or internal to the generated Hirshfeld surfaces; together, this pair ( $d_e$  and  $d_i$ ) generates a 2D fingerprint plot. The different colors on the fingerprint plot represent the frequency of occurrence of interaction. A red spot represents the direct interaction between two atoms. Ultimately it gives a normalized contact distance ( $d_{\text{norm}}$ ). The  $d_{\text{norm}}$  values are mapped onto the Hirshfeld surface using a red, white, and blue color scheme; red, white, and blue regions correspond to the strong, medium, and weak interactions, respectively. All the Hirshfeld surfaces were generated using Crystal Explorer 3.1 software.

### Synthesis and characterization

**(E)-10-Pentyl-3-(2-(10-(thiophen-2-yl)anthracen-9-yl)vinyl)-4a,10a-dihydro-10H-phenothiazine TAPTZ.** In a  $50 \text{ mL}$  round-bottomed flask, diethyl ((10-(thiophen-2-yl)anthracen-9-yl)methyl)phosphonate ( $0.50 \text{ g}$ ,  $1.218 \text{ mmol}$ ) was dissolved in  $30 \text{ mL}$  of dry THF under an argon atmosphere at room temperature.  $^t\text{BuOK}$  ( $0.410 \text{ g}$ ,  $3.65 \text{ mmol}$ ) was added and stirred for  $4$ – $5 \text{ min}$ . 10-Pentyl-10H-phenothiazine-3-carbaldehyde ( $0.398 \text{ g}$ ,  $1.34 \text{ mmol}$ ) was carefully added to the solution. The reaction was allowed to stir for  $12 \text{ h}$  and completion of the reaction was monitored by TLC. The resulting reaction mixture was quenched with water, extracted with ethyl acetate ( $20 \text{ mL} \times 3$ ), dried over anhydrous sodium sulphate, and concentrated under a rotary evaporator. The compound **TAPTZ** was purified by column chromatography ( $100$ – $200$  mesh-sized silica gel) using  $3\%$  ethyl acetate in petroleum ether.

**TAPTZ.** Yield:  $0.48 \text{ g}$ , yield  $70\%$ , m.p. –  $165$ – $167^\circ \text{C}$ . IR ( $\nu \text{ cm}^{-1}$ , in KBr):  $3066$ ,  $2928$ ,  $2861$ ,  $1704$ ,  $1603$ ,  $1460$ ,  $1334$ ,  $1296$ ,  $1242$ ,  $1126$ ,  $1037$ .  $^1\text{H NMR}$  ( $400 \text{ MHz}$ ,  $\text{CDCl}_3$ )  $\delta$   $8.37$  (dd,  $J = 7.8$ ,  $1.3 \text{ Hz}$ ,  $2\text{H}$ ),  $7.87$ – $7.76$  (m,  $3\text{H}$ ),  $7.60$  (dd,  $J = 5.2$ ,  $1.1 \text{ Hz}$ ,  $1\text{H}$ ),  $7.50$  (d,  $J = 2.0 \text{ Hz}$ ,  $1\text{H}$ ),  $7.39$ – $7.42$  (m,  $5\text{H}$ ),  $7.30$  (dd,  $J = 5.2$ ,  $3.4 \text{ Hz}$ ,  $1\text{H}$ ),  $7.19$ – $7.14$  (m,  $3\text{H}$ ),  $6.96$ – $6.87$  (m,  $3\text{H}$ ),  $6.83$  (d,  $J = 16.5 \text{ Hz}$ ,  $1\text{H}$ ),  $3.92$ – $3.86$  (m,  $2\text{H}$ ),  $1.92$ – $1.80$  (m,  $2\text{H}$ ),  $1.49$ – $1.31$  (m,  $4\text{H}$ ),  $0.92$  (t,  $J = 7.2 \text{ Hz}$ ,  $3\text{H}$ ).  $^{13}\text{C NMR}$  ( $101 \text{ MHz}$ ,  $\text{CDCl}_3$ )  $\delta$   $144.9$ ,  $136.5$ ,  $131.7$ ,  $131.7$ ,  $129.5$ ,  $129.3$ ,  $127.5$ ,  $127.3$ ,  $127.2$ ,  $127.0$ ,  $126.7$ ,  $126.2$ ,  $126.0$ ,  $125.22$ ,  $125.6$ ,  $125.4$ ,  $125.2$ ,  $125.0$ ,  $123.1$ ,  $122.5$ ,  $115.4$ ,  $47.6$ ,  $29.2$ ,  $26.6$ ,  $22.4$ ,  $14.1$  (some signals are merged together). HR-MS for  $\text{C}_{37}\text{H}_{32}\text{NS}_2$ , calc.  $554.1976$ , found to be  $554.1786$  [ $\text{M} + \text{H}$ ]<sup>+</sup>; X-ray structure is done for this sample (CCDC 2100036).<sup>†</sup>

**FAPTZ: (E)-3-(2-(10-(furan-2-yl)anthracen-9-yl)vinyl)-10-pentyl-4a,10a-dihydro-10H-phenothiazine.** Compound **FAPTZ** was prepared by following the synthetic procedure for compound **TAPTZ**. The compound **FAPTZ** was purified by column chromatography ( $100$ – $200$  mesh sized silica gel) using petroleum ether to obtain **FAPTZ** as a yellow solid. Yield:  $0.38 \text{ g}$ ,  $57\%$  yield m.p. –  $153$ – $155^\circ \text{C}$ . IR ( $\nu \text{ cm}^{-1}$ , in KBr):  $3396$ ,  $3064$ ,  $2925$ ,  $2860$ ,  $1597$ ,  $1481$ ,  $1459$ ,  $1332$ ,  $1251$ ,  $1140$ ,  $1105$ ,  $1020$ .  $^1\text{H NMR}$  ( $400 \text{ MHz}$ ,  $\text{CDCl}_3$ )  $\delta$   $8.43$ – $8.37$  (m,  $2\text{H}$ ),  $7.95$ – $7.90$  (m,  $2\text{H}$ ),  $7.86$ – $7.78$  (m,  $2\text{H}$ ),  $7.54$  (d,  $J = 2.0 \text{ Hz}$ ,  $1\text{H}$ ),  $7.51$ – $7.42$  (m,  $5\text{H}$ ),  $7.23$ – $7.18$  (m,  $2\text{H}$ ),  $6.99$ – $6.90$  (m,  $3\text{H}$ ),  $6.85$  (d,  $J = 16.5 \text{ Hz}$ ,  $1\text{H}$ ),  $6.78$ – $6.71$  (m,  $2\text{H}$ ),  $3.95$ – $3.89$  (m,  $2\text{H}$ ),  $1.94$ – $1.84$  (m,  $2\text{H}$ ),  $1.54$ – $1.36$  (m,  $4\text{H}$ ),  $0.96$  (t,  $J = 7.2 \text{ Hz}$ ,  $3\text{H}$ ).  $^{13}\text{C NMR}$  ( $101 \text{ MHz}$ ,  $\text{CDCl}_3$ )  $\delta$   $150.8$ ,  $145.1$ ,  $144.9$ ,  $142.9$ ,  $136.6$ ,  $135.2$ ,  $131.7$ ,  $131.6$ ,  $129.5$ ,  $127.5$ ,  $127.3$ ,  $126.6$ ,  $126.3$ ,  $126.1$ ,  $125.4$ ,  $125.2$ ,  $125.0$ ,





123.1, 122.5, 115.4, 112.3, 110.9, 47.6, 29.2, 26.6, 22.4, 14.1 (some signals are merged together). HR-MS for  $C_{37}H_{32}NOS$ , calc. 538.2205, found to be 538.2226  $[M + H]^+$ . X-ray structure is done for this sample (CCDC 2100035).†

## Conclusions

In conclusion, two new anthranil  $\pi$ -conjugates **TAPTZ** and **FAPTZ** linked with thiophene or furan as a flanking donor are easily synthesized. Thiophene analogs are found to be superior in displaying solvatofluorochromic features to the furan analog, with a slightly better variation in the excited state dipole moment. The emission wavelength in solution dictates a better conjugation in **FAPTZ** than **TAPTZ**, and that is interpreted by the DFT-optimized molecular structures, presenting better  $\pi$ -electronic conjugations in **FAPTZ** from a comparatively lesser torsion angle between furan and anthracene by keeping the rest almost similar. Contrarily, the solid-state emission appears at a lower wavelength for **FAPTZ** because of the higher torsion angle between anthracene (acceptor) and PTZ (donor). **TAPTZ** has appeared to be a better blue-shifted AIEgen with a relatively smaller particle size. The tilted conformation and a cross-packing enable these molecules to be emissive in the solid-state and exhibit reversible red-shifted MFC-feature. A relatively lower redshift and sensitivity for **TAPTZ** are described by more substantial crystal packing with two extra C $\cdots$ S and S $\cdots$ S interactions than **FAPTZ**. Such dissimilarity is deduced with SCXRD, PXRD, and fluorescence lifetime studies. Hirshfeld surface analyses indicate quantitatively more C $\cdots$ H interactions in **FAPTZ** than **TAPTZ**. Thus, this work has provided a thought on the effect of a tiny change in a flanking donor that can impact the solvato-, AIE- and MFC-features, which would develop new MFC-active materials. The practical application of reversible security writing demonstrates the efficacy of these  $\pi$ -conjugates.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

We thank DST-SERB (CRG/2018/000456) for financial support. MC thanks BITS-Pilani Hyderabad for NMR/X-ray facilities. We thank Prof Durba Roy for the DFT calculations.

## References

- (a) R. Gao, X. Fang and D. Yan, *J. Mater. Chem. C*, 2019, **7**, 3399–3412; (b) Q. Li and Z. Li, *Acc. Chem. Res.*, 2020, **53**, 962–973; (c) L. Bai, P. Bose, Q. Gao, Y. Li, R. Ganguly and Y. Zhao, *J. Am. Chem. Soc.*, 2017, **139**, 436–441; (d) Y. Gong, Y. Zhang, W. Z. Yuan, J. Z. Sun and Y. Zhang, *J. Phys. Chem. C*, 2014, **118**, 10998–11005.
- (a) Y. Sagara, S. Yamane, M. Mitani, C. Weder and T. Kato, *Adv. Mater.*, 2016, **28**, 1073–1095; (b) J. Xu and Z. Chi, *Mechanochromic Fluorescent Materials: Phenomena, Materials and Applications*, RSC Publishers, 2014.
- (a) G. Zhang, J. Sun, P. Xue, Z. Zhang, P. Gong, J. Peng and R. Lu, *J. Mater. Chem. C*, 2015, **3**, 2925–2932; (b) P. Xue, J. Ding and R. Lu, *J. Mater. Chem. C*, 2016, **4**, 6688–6706.
- (a) X. Huang, L. Qian, Y. Zhou, M. Liu, Y. Cheng and H. Wu, *J. Mater. Chem. C*, 2018, **6**, 5075–5096; (b) C. Wang and Z. Li, *Mater. Chem. Front.*, 2017, **1**, 2174–2194.
- Y. Yin, Z. Chen, Y. Yang, G. Liu, C. Fan and S. Pu, *RSC Adv.*, 2019, **9**, 24338–24343.
- (a) W. Sharmoukh, J. Cong, B. A. Ali, N. K. Allam and L. Kloo, *ACS Omega*, 2020, **5**(27), 16856–16864; (b) B. L. Reid, S. B. Briggs, L. E. Karagiannidis, S. Muzzioli, P. Raiteri, M. E. Light, S. Stagni, P. Brulatti, P. A. Gale, M. I. Ogden and M. Massi, *J. Mater. Chem. C*, 2013, **1**, 2209–2216.
- (a) G. Barbarella, M. Melucci and G. Sotgiu, *Adv. Mater.*, 2005, **17**, 1581; (b) A. Pachariyangkun, M. Suda, S. Hadsadee, S. Jungstutiwong, P. Nalaoh, P. Pattanasattayavong, T. Sudyoasuk, H. M. Yamamoto and V. Promarak, *J. Mater. Chem. C*, 2020, **8**, 17297–17306.
- M. El-Sayed, H. Müller, G. Rheinwald, H. Lang and S. Spange, *Chem. Mater.*, 2003, **15**, 746–754.
- (a) A. Pachariyangkun, M. Suda, S. Hadsadee, S. Jungstutiwong, P. Nalaoh, P. Pattanasattayavong, T. Sudyoasuk, H. M. Yamamoto and V. Promarak, *J. Mater. Chem. C*, 2020, **8**, 17297–17306; (b) O. Gidron, A. Dadvand, Y. Sheynin, M. Bendikov and D. F. Perepichka, *Chem. Commun.*, 2011, **47**, 1976–1978.
- (a) O. Gidron and M. Bendikov, *Angew. Chem., Int. Ed.*, 2014, **53**, 2546; (b) H. T. Turan, I. Yavuj and V. Aviyente, *J. Phys. Chem. C*, 2017, **121**, 25682–25690.
- Z. Zhao, H. Nie, C. Ge, Y. Cai, Y. Xiong, J. Qi, W. Wu, R. T. K. Kwok, X. Gao, A. Qin, J. W. Y. Lam and B. Z. Tang, *Adv. Sci.*, 2017, **4**, 1700005.
- (a) W. Liu, Y. Wang, J. Yang, X. Li, X. Wang and L. Ma, *Dyes Pigm.*, 2020, **175**, 108149; (b) N. Friebe, K. Schreiter, J. Kübel, B. Dietzek, N. Moszner, P. Burtscher, A. Oehlke and S. Spange, *New J. Chem.*, 2015, **39**, 5171.
- W. A. Morris, T. Butler, M. Kolpaczynska and C. L. Fraser, *Mater. Chem. Front.*, 2017, **1**, 158–166.
- M. Z. K. Baig, D. Majhi, R. N. P. Tulichala, M. Sarkar and M. Chakravarty, *J. Mater. Chem. C*, 2017, **5**, 2080–2387.
- W. Xu, M. M. S. Lee, Z. Zhang, H. H. Y. Sung, I. D. Williams, R. T. K. Kwok, J. W. Y. Lam, D. Wang and B. Z. Tang, *Chem. Sci.*, 2019, **10**, 3494–3501.
- T. Jadhav, B. Dhokale, Y. Patil, S. M. Mobin and R. Misra, *J. Phys. Chem. C*, 2016, **120**(42), 24030–24040.
- M. Z. K. Baig, G. Pallikonda, P. Trivedi, R. N. P. Tulichala, B. Ghosh and M. Chakravarty, *ChemistrySelect*, 2016, **1**, 4332.
- (a) C. Reichardt, *Chem. Rev.*, 1994, **94**, 2319–2358; (b) M. A. Haidekker, T. P. Brady, D. Lichlyter and E. A. Theodorakis, *Biorg. Chem.*, 2005, **33**, 415–425.



- 19 S. Sasaki, G. P. C. Drummen and G. Konishi, *J. Mater. Chem. C*, 2016, **4**, 2731–2743.
- 20 J. Zhang, B. Xu, J. Chen, L. Wang and W. Tian, *J. Phys. Chem. C*, 2013, **117**(44), 23117–23125.
- 21 U. Subuddhi, S. Haldar, S. Sankararaman and A. K. Mishra, *Photochem. Photobiol. Sci.*, 2006, **5**, 459–466.
- 22 (a) Y. Hong, J. W. Y. Lam and B. Z. Tang, *Chem. Soc. Rev.*, 2011, **40**, 5361–5388; (b) Y. Chen, J. W. Y. Lam, R. T. K. Kwok, B. Liu and B. Z. Tang, *Mater. Horiz.*, 2019, **6**, 428–433.
- 23 Z. Yang, W. Qin, J. W. Y. Lam, S. Chen, H. H. Y. Sung, I. D. Williams and B. Z. Tang, *Chem. Sci.*, 2013, **4**, 3725–3730.
- 24 B. Prusti, P. Sarkar, S. K. Pati and M. Chakravarty, *J. Mater. Chem. C*, 2021, **9**, 9555–9570.
- 25 N. Zhang, H. Chen, Y. Fan, L. Zhou, S. Trepout, J. Guo and M. H. Li, *ACS Nano*, 2018, **12**, 4025–4035.
- 26 B. Fang, M. Chu, Z. Wu, Y. Shi, Y. S. Zhao and M. Yin, *J. Mater. Chem. C*, 2019, **7**, 4434–4440.
- 27 B. Roy, M. C. Reddy and P. Hazra, *Chem. Sci.*, 2018, **9**, 3592–3606.
- 28 (a) A. Ekbote, S. M. Mobin and R. Misra, *J. Mater. Chem. C*, 2018, **6**, 10888–10901; (b) Y. Matsuo, Y. Wang, H. Ueno, T. Nakagawa and H. Okada, *Angew. Chem., Int. Ed.*, 2019, **58**, 8762–8767.
- 29 *Mechanochromic fluorescent materials phenomena, materials and applications*, ed. J. Xu and Z. Chi, Royal Society of Chemistry, 2014, p. 167.
- 30 Y. Ren and T. Baumgartner, *Inorg. Chem.*, 2012, **51**, 2669–2678.
- 31 B. Roy, M. C. Reddy and P. Hazra, *Chem. Sci.*, 2018, **9**, 3592–3606.
- 32 M. C. Burla, R. Caliendo, M. Camalli, B. Carrozzini, G. L. Cascarano, C. Giacobozzo, M. Mallamo, A. Mazzone, G. Polidori and R. Spagna, *J. Appl. Crystallogr.*, 2012, **45**, 357.
- 33 G. M. Sheldrick, *Acta Crystallogr., Sect. A: Found. Crystallogr.*, 2008, **64**, 112.
- 34 M. A. Spackman and D. Jayatilaka, *CrystEngComm*, 2009, **11**, 19–32.

