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The interplay of thermally activated delayed fluorescence (TADF) and room temperature organic phosphorescence in sterically-constrained donor—acceptor charge-transfer molecules†

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A series of phenothiazine-dibenzothiophene-S,S-dioxide charge-transfer molecules have been synthesized. Increasing steric restriction around the donor-acceptor bond significantly alters contributions from TADF and phosphorescence. Bulky substituents on the 1-(and 9) position(s) of the phenothiazine result in no TADF in the solid state; instead strong phosphorescence is observed at ambient temperature.

The study of new fluorescent and phosphorescent emitters is central to the development of a wide range of optoelectronic technologies, including organic light-emitting diodes (OLEDs), electrochromic devices, sensors and fluorescence imaging.¹ Spin statistics dictate that the efficiency of traditional fluorescent emitters which utilise only singlet excited states for light emission is limited to 25%.² Phosphorescent emitters based on transition metal complexes, especially third row ions such as Ir(III) and Pt(II), are popular because the heavy metal facilitates spin-orbit coupling and harvesting of both singlet and triplet excitons by enhanced intersystem crossing (ISC). Internal quantum efficiencies (IQEs) approaching 100% can be achieved by this mechanism.³ However, these phosphors suffer from limited stability in practical applications, especially for blue and white emission, and the scarcity and expense of precious metals restricts their commercial applications.

Triplet fusion is an alternative way to convert triplets into emissive singlet states although the IQE in this process is limited to a theoretical maximum of 62.5%. Currently, a leading strategy is to exploit thermally activated delayed fluorescence (TADF) to convert triplets to singlet states with a potential IQE of up to 100% using all-organic (metal-free) materials. The mechanism of TADF

is fundamentally different from traditional fluorescence. TADF uses thermal energy to assist reverse intersystem crossing (RISC) of lower-energy triplet states into emissive singlets. Two main parameters control this mechanism: (i) the energy splitting between the lowest singlet (S_1) and triplet (T_1) states should be minimised, and (ii) the non-radiative pathways by which the excited singlet and triplet states can decay need to be suppressed in order to achieve high fluorescence quantum yields and long triplet excited state lifetimes.⁶

Based on the TADF properties of a wide range of donoracceptor (D-A) molecules, 7-15 it is clear that a prerequisite for efficient TADF is the spatial separation of the highest occupied molecular orbital (HOMO) on the electron donor and the lowest unoccupied molecular orbital (LUMO) on the electron acceptor moieties of the emitter molecule. This can be achieved by enforcing a twist between the planes of the D and A moieties to restrict the conjugation leading to an intramolecular charge transfer (ICT) state with a small (ΔE_{S1T1}) energy gap.⁷ For example, tuning of TADF in the green-orange region has been achieved by stabilisation of the ICT state between a phenoxazine donor unit and 2,4,6-triphenyl-1,3,5-triazine acceptor unit separated by a large twist angle.8 A 9,9'-spirobifluorene unit has been used to enforce orthogonality between D and A units in a TADF emitter. Systematically engineering the twist angles within a series of D-A structures is, therefore, an important strategy for understanding the parameters which control the overall emission, including TADF, in these types of molecules.

We now report a series of D–A–D and D–A molecules based on phenothiazine donor and dibenzothiophene-*S*,*S*-dioxide acceptor units in which the steric bulk of alkyl groups on the donor restricts rotation around the C–N bond linking the D and A units (Fig. 1). Remarkable effects of this functionalization on the structural and photophysical properties of the molecules are observed. Compounds 1^{16a} and 6^{16b} exhibit efficient TADF in zeonex films whereas the more sterically hindered analogues 3 and 4, which have less conformational flexibility, show strong contributions from phosphorescence, even at RT.

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Fig. 1 Structures of D-A-D and D-A molecules.

Functionalization of the phenothiazine unit(s) with alkyl substituents at the 1 (and 9) position(s) affords the new compounds 2-5, 7, 8. The synthetic details are reported in the ESI,† S2. The structures of 2.THF, 3, 5.2CH₂Cl₂ and 7 were determined by singlecrystal X-ray diffraction (Fig. 2 and ESI,† S8).

The dibenzothiophene moiety in 2 and 5 is practically planar; in 3 it is slightly twisted and in 7 folded, with the two arene rings forming a dihedral angle of 6.9° and 12.5°, respectively.

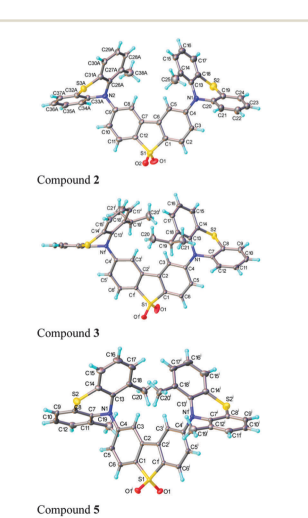


Fig. 2 X-ray molecular structures of compounds 2 (major conformer), 3 and 5 displayed with thermal ellipsoids at 50% probability.

The molecules of 3 and 5 possess a crystallographic two-fold axis, hence the phenothiazine substituents have transoid orientation with respect to the dibenzothiophene plane, i.e. their S atoms lie on opposite sides of this plane. 2 has a similar conformation, although the molecule has no crystallographic symmetry. Interestingly, one of the phenothiazine moieties [at C(9)] is disordered in a 0.6:0.4 ratio between two conformations, with the methyl substituent on opposite sides of N(2) and the tricyclic system librating around the N(2) as the pivot, to provide room for the methyl group. The phenothiazine moiety is always folded along the N···S vector, forming a dihedral angle of 128.7° (3), 135.5° (5) and 135.3° (7). In 2, the ordered phenothiazine moiety is folded by 131.1° and the disordered one by 133.6° or 135.0°. In every case, the N atom is substantially pyramidalised; its lone pair is oriented favourably for conjugation with the dibenzothiophene π -system rather than with the arene rings of phenothiazine itself. Therefore, the N-C(dibenzothiophene) bond is shorter than N-C(phenothiazine), on average by 0.035 Å in 3 and 5 and 0.039 Å in 7 (s.u. 0.001–0.002 Å); the similar difference (0.038 Å) in 2 is not statistically significant due to disorder and high s.u. The packing of 2 and 5 is characterised by stacking $(\pi - \pi)$ interactions between phenothiazine arene rings (related via inversion centres and therefore parallel) with interplanar separations of 3.50 and 3.60 Å (intermittently) in 2, or 3.66 Å (uniformly) in 5. These interactions link molecules into infinite chains (ESI,† S8). No stacking is evident in the structures of 3 and 7.

The ¹H NMR spectra of 1, ^{16a} 2, 5-8 at 298 K display sharp peaks showing that there is one species in solution at this temperature on the NMR timescale. In contrast, compounds 3 and 4 which have a bulkier iPr or tBu substituent on each phenothiazine ring exhibit restricted rotation. The spectrum of compound 3 at 298 K clearly shows two species in solution: upon heating the peaks coalescence to reveal only one species at 373 K (Fig. 3). 1H NOESY and ROESY NMR experiments strongly suggest that the two species are interconverting with each other (see ESI,† S4). For the t-butyl analogue 4, at 298 K selected peaks are extremely broad and the spectra at 348 K and 373 K are comparable to 3 at 298 K. Rotation in compound 4 is still partially restricted even at 373 K (Fig. 4). ¹H NOESY NMR experiments with 4 also demonstrate there is some exchange occurring between the two very broad peaks at 298 K (see ESI,† S4). The presence of these two species is likely to be due to up-up

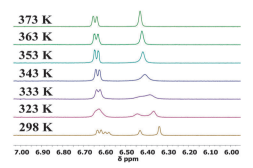


Fig. 3 500 MHz variable temperature ¹H NMR experiments for **3** from 298-373 K in DMSO-d₆.

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373 K
348 K
323 K
298 K
7.0 6.9 6.8 6.7 6.6 6.5 6.4 6.3 6.2 6.1 6.0 5.9 5.8 5.7 5.6 5.5 5.4 5.3

Fig. 4 $\,$ 500 MHz variable temperature 1 H NMR experiments for **4** from 298-373 K in DMSO-d₆.

and up-down configurations of the i Pr or t Bu groups with respect to the acceptor.

Photophysical measurements on 2–8 reveal how the increasingly bulky phenothiazine donors affect their optical properties, primarily by restricting rotation around the N–C bond that links the D and A units. Consistent with previous work, ¹⁰ it has been shown that **1** and **6** emit strongly *via* a TADF mechanism ¹⁶ (see ESI,† S6).

The D-A compound 7 has an additional methyl group in the 1-position on phenothiazine compared to 6, and this has a dramatic effect on the emission of this molecule. The same difference is also observed in the D-A-D molecule 2 (Fig. 5).

Introduction of the methyl group into molecules 2 and 7 increases the lifetime of the DF emission in toluene solution (compared with molecules 1 and 6, ESI,† S6.3). Temperature and power dependence data confirm that the delayed emission still occurs *via* a TADF mechanism in solution (Fig. 6).

There is also a reduction in intensity of the delayed fluorescence on introduction of the methyl group into 2. We propose that the methyl group slows molecular reorganization processes resulting in a slower initial electron transfer step as well as increasing the time taken for the ¹CT state to emit. These effects increase the amount of non-radiative decay and direct decay from ¹D-¹A states, as well as reducing the number of CT states initially created. This is evidenced by the increasing contribution of emission around 400 nm in the steady-state data. The pronounced influence of the side groups in the TADF emission is clearly seen in the case of compound 4, where even in toluene solution no ¹CT emission is observed.

The most noticeable difference between unsubstituted **1** and methyl substituted **2** is that in the solid state in a zeonex film, the TADF emission is no longer present. A large time gap with

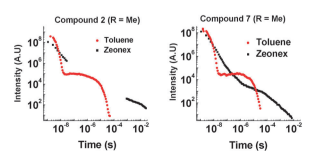


Fig. 5 Time-resolved emission decay for 2 and 7 at 290 K (355 nm excitation).

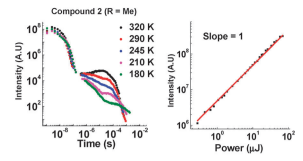


Fig. 6 Left: Temperature dependence of emission decay in toluene for $\bf 2$. Right: Power dependence of DF emission in toluene for $\bf 2$ (400 ns delay and 50 μs integration time).

no emission is observed with 2 after fast donor fluorescence decays. It is suggested that the methyl group's ability to restrict reorganization is enhanced in the solid state, resulting in such slow electron transfer that the donor states have all decayed before appreciable electron transfer occurs. The unusual feature in the solid state is that 2 emits significantly more *via* phosphorescence compared with 1. Quenching experiments of the emission with oxygen show that 2 and 7 have increasing phosphorescent character (Fig. 7). This is consistent with the efficient ISC to the donor triplet state instead of CT formation.

D–A–D structure 2 shows more phosphorescence than its D–A analogue 7. More fine structure is observed with 2 and significantly more quenching is observed in the presence of oxygen. This room temperature phosphorescence is unusually strong for an organic molecule, ^{17–20} and the methyl group plays a key role. With increasing steric hindrance the phosphorescence increases even further (Fig. 8). The distinction between TADF and phosphorescence is discussed in more detail in ESI,† S6.8.

With a bulkier ⁱPr group in the D-A system **8**, TADF is still observed in solution (ESI,† S6.5). In zeonex no TADF is observed, but strong phosphorescence is observed at RT. For the D-A-D system **3**, the amount of TADF is lowered significantly, even in toluene solution. This correlates with the solution NMR studies showing restricted rotation. With the two ⁱPr systems **3** and **8**, the extent of phosphorescence increases significantly (Fig. 9).

The emission of 3 and 8 in zeonex shows even more fine structure compared with 2 and 7 due to the increased ratio of phosphorescence to ¹CT fluorescence. D-A-D 3 with two ⁱPr groups, prevents molecular movement more than in the D-A

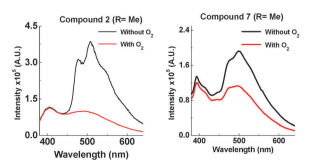


Fig. 7 Emission of 2 and 7 at 290 K in zeonex (excited at 355 nm)

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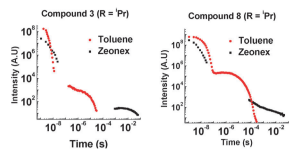
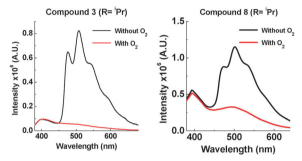


Fig. 8 Time-resolved emission decay for 3 and 8 at 290 K (355 nm excitation)

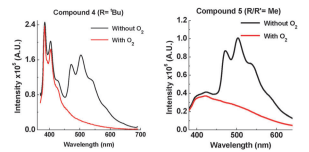


Emission of 3 and 8 at 290 K in zeonex (excited at 355 nm).

analogue 8 (see Fig. 3 and ESI,† S3). The emission from 3 in the solid state consists almost entirely of phosphorescence as shown by the large decrease in emission on introduction of oxygen. It is unusual for an organic molecule to decay almost completely by phosphorescent pathways. Temperature dependence studies in solid state show that 8 does not emit via a TADF mechanism (ESI,† S6.6). The introduction of these bulkier groups acts like a switch, turning off electron transfer and turning on phosphorescence.

Having observed the drastic change caused by an ¹Pr group, ^tBu groups were introduced into 4, and two methyl groups were attached onto each phenothiazine in 5. A large contribution from phosphorescence is observed in the solid state of 4 (Fig. 10), and there is also a large contribution of fast fluorescence from the excited S₁ state. It is proposed that the motion in 4 in the solid state is so restricted that electron transfer from the ¹D-¹A states to the ¹CT state is prevented. This results in fast fluorescence decay from S₁ (¹A and ¹D) and long lived phosphorescence (³A and ³D) as the main radiative pathways for the excited state to decay. Supporting evidence for extended ¹D emission in 4 is seen in time-resolved studies (see ESI,† S6.3). Compound 5 also shows strong phosphorescence at RT, but less ¹A and ¹D fluorescence than 4, which correlates with the less bulky substituents.

In conclusion, suppressing vibrational quenching in a series of D-A-D and D-A molecules strongly activates phenothiazine phosphorescence. By sterically engineering near-orthogonal orientation and restricting rotation between the D and A fragments, in the solid state the main non-radiative deactivation pathway that affects the lowest triplet excited state is thus suppressed and so phosphorescence from phenothiazine is not quenched.



Emission of 4 and 5 at 290 K in zeonex (excited at 355 nm)

This provides valuable mechanistic information on TADF and has led to strongly phosphorescent molecules at room temperature.

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