

RESEARCH ARTICLE

View Article Online
View Journal | View IssueCite this: *Mater. Chem. Front.*,
2020, 4, 3602

Exploiting trifluoromethyl substituents for tuning orbital character of singlet and triplet states to increase the rate of thermally activated delayed fluorescence†

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This work shows that trifluoromethyl (CF₃) substituents can be used to increase the rate of thermally activated delayed fluorescence (TADF) in conjugated organic molecules by tuning the excitonic character of the singlet and triplet excited states. The synthesis and detailed photophysical characterization of four new functionalized donor–acceptor–donor (D–A–D) compounds using CF₃-substituted-thioxanthene-*S,S*-dioxide and dimethylacridine units are presented. Several compounds are reported to exhibit rapid blue/green thermally activated delayed fluorescence with major delayed fluorescence lifetime components of ≈ 1 –2 μ s. Changes in the orbital character of singlet and triplet states (local vs. charge transfer) result in significant changes in rates of delayed fluorescence, despite similar ΔE_{ST} values. The change in orbital character is well supported by TD-DFT calculations. Electrochemical measurements reveal strong shifts in redox potentials that can be induced by σ -electron withdrawing CF₃ substituents. Trifluoromethyl substituted compounds with a wider ΔE_{ST} also perform more efficiently than might be expected due to the demonstrated changes in excited state character. This study reveals important photophysical and molecular design implications for the future development of TADF emitters.

Received 29th June 2020,
Accepted 22nd July 2020

DOI: 10.1039/d0qm00429d

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Introduction

Light emission from organic materials is revolutionizing the color-display and lighting industries.^{1–3} However, traditional fluorescent molecules have severe drawbacks in these applications, as only 25% of the excited states (singlets) that are generated under electrical excitation are harvested as emission; the remaining 75% of triplet excitons decay non-radiatively.⁴ Phosphorescent molecules can address this issue and achieve internal quantum efficiencies (IQEs) approaching 100% due to light emission from both singlet and triplet excited states.⁵ This has led to extensive studies on emitters containing heavy metal atoms, where strong spin–orbit coupling promotes intersystem crossing (ISC) and phosphorescence (PH) emission.^{6,7} The main disadvantages of organo-metallic emitters are their high cost due to the necessity of precious

and possibly toxic heavy metals (typically iridium and platinum) and the electrochemical and thermal instability of the blue and especially deep-blue emitting complexes.⁸

E-type delayed fluorescence,⁹ which has been re-named thermally activated delayed fluorescence (TADF) is an alternative mechanism that harvests triplet excited states in all-organic emitters for light emission.^{10,11} The main application of TADF materials is in organic light-emitting diodes (OLEDs).^{12–15} TADF materials have also been deployed in fluorescence imaging,^{16,17} optical temperature sensing^{18,19} and catalysis.²⁰ Specific molecular design features enable fast reverse intersystem crossing (rISC), which can convert up to 100% of triplet excited states to emissive singlet states, leading to TADF.²¹ Despite intense global research efforts, full understanding of the specific molecular features that enable rapid rISC remain elusive. Indeed, several different rISC mechanisms with different design rules have been discovered, including spin-vibronic coupling in donor–acceptor–donor (D–A–D) charge transfer (CT) emitters,²² upper triplet state crossings,²³ resonant non-overlapping orbital distributions in planar boron compounds,²⁴ exciplexes,^{25,26} and hybrid D–A compounds.²⁷

Ideal TADF materials possess both high photoluminescence quantum yields (PLQYs) and fast rISC rates, the latter of which are enabled by a small energy difference between the lowest

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† Electronic supplementary information (ESI) available: Synthesis and characterization details of the new molecules; methods and results for the photophysics, OLED devices and calculations (PDF). X-ray crystallographic data. CCDC 1887606 (3) and 1887607 (4). For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/d0qm00429d



singlet and triplet excited states (typically $\Delta E_{ST} < 0.25$ eV). These properties can be found in heteroaromatic donor–acceptor (D–A) molecules with charge-transfer (CT) excited states. In these D–A molecules, a large average dihedral angle between the planes of the D and A units ensures the highest occupied and lowest unoccupied molecular orbitals (HOMO and LUMO, respectively) are spatially separated. This minimizes electron exchange energy and thus also ΔE_{ST} . A detailed analysis of the mechanism of TADF in D–A–D materials has revealed a second-order spin–vibronic coupling process, in which a triplet state of local excitonic character (3LE) mediates rISC between the triplet and singlet charge transfer states (3CT and 1CT , respectively).^{28–30}

The key roles of D–A conjugation and dihedral angles, molecular rigidity, conformation, connectivity, and steric effects are also important factors that influence CT formation and hence TADF efficiency and emission color.^{31–37} Despite this level of understanding, stable blue emission, which is invaluable for high-quality full-color OLED displays, remains especially challenging to achieve. Only a limited number of blue/deep-blue TADF emitters possessing good color purity have been reported.^{36,38–44}

In this context a systematic series of TADF molecules that are functionalized at specific sites with trifluoromethyl substituents are herein presented. Trifluoromethyl (CF_3) is a strongly inductive electron-withdrawing group which makes it a useful substituent for emission colour tuning *via* donor and acceptor strength manipulation in D–A–D CT emitters. CF_3 is not π -conjugative, which makes it distinct from the electron-withdrawing groups that are typically used in TADF emitters, for example: cyano,^{10,45} sulfone,^{46,47} ketone,⁴⁸ triazine⁴⁹ and oxadiazole.⁵⁰ Trifluoromethyl groups can also be very stable to chemical, electrochemical, thermal and photochemical degradation, which are valuable properties to impart to TADF materials.^{51–54}

Extensive systematic studies of TADF emitters with CF_3 substituents are scarce. Current CF_3 functionality in TADF emitters are typically incorporated in the ‘Ar– CF_3 ’ type motif.⁵⁵ Examples include CF_3 –phenyl^{56–59} or CF_3 –pyridyl^{52,60} as an acceptor unit, or CF_3 –carbazole as a donor unit.^{61,62} This present study further diversifies the library of reported CF_3 functionalised TADF molecules and has led to new blue and green TADF emitters by manipulation of the electron-donating and electron-accepting abilities of the D and A subunits with CF_3 substituents. In particular, we have investigated the use of the CF_3 unit as part of the bridge of the acceptor unit as opposed to the typical ‘Ar– CF_3 ’ motif currently observed in the literature. There are few examples of CF_3 ‘bridges’ being utilized in TADF molecules.^{53,63} Swager and co-workers showed that geminal- CF_3 substituents on (non-TADF) polyfluorene derivatives were more photostable compared to geminal- CH_3 substituents.⁵³ Adachi and co-workers used a flexible bis(CF_3)-substituted methylene bridge directly between donor and acceptor in TADF emitters.⁶³

The synthetic routes, X-ray crystal structures, solution electrochemical data and a thorough photophysical investigation of the materials are presented here. These results show that

trifluoromethyl substituents can be used to impart large shifts in redox potentials and emission properties through inductive effects. Molecules **1** and **2** show highly similar ΔE_{ST} values, but **2** shows faster delayed fluorescence. Detailed analysis and supporting hybrid-DFT calculations reveal how the tuning of orbital character with trifluoromethyl substituents leads to an increase in TADF emission rates. OLEDs were also fabricated, although these devices showed unexpected instability.

Experimental section

The synthesis and structural characterization of the molecules are given in the ESI.[†]

X-ray diffraction experiments: full data (including structure factors) in CIF format is available as ESI[†] and have been deposited with the Cambridge Structural Database, CCDC 1887606 (**3**) and 1887607 (**4**).[†]

Optical and optoelectronic methods are given in the ESI.[†]

Results and discussion

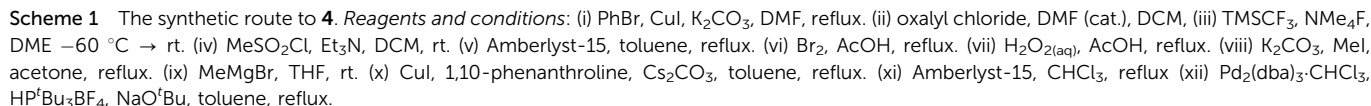
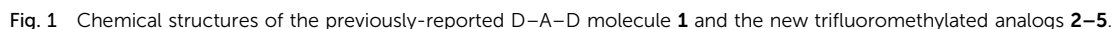
Molecular design and synthesis

Starting from the well-characterized blue TADF emitter 2,7-bis-(9,9-dimethylacridin-10-yl)-9,9-dimethylthioxanthene-*S,S*-dioxide (DDMA-TXO2) **1**, (Fig. 1), the motivation was to explore the effect on photophysical properties that arise from tuning the donor and acceptor abilities of the sub-units using systematic CF_3 substitution.^{64,65} Compound **1** was selected as it has efficient TADF properties and can be systematically modified. The DDMA-TXO2 triplet level (T_1) is a localized triplet (3LE) state and the energy splitting between 1CT and 3LE is $\Delta E_{ST} = (0.06 \pm 0.03)$ eV in evaporated DPEPO host.⁶⁵ Two independent studies have shown that DDMA-TXO2 affords blue OLEDs with EQEs > 20%.^{64,65}

Fig. 1 presents the chemical structures of **1** and the new analogs **2–5**. The methyl groups of the thioxanthene acceptor in **1** are replaced by CF_3 groups in **2–4** (Fig. 1), whereas CF_3 groups are added directly on to the aryl rings of the D unit in **5** (with both D and A modifications in **4**). The acceptor CF_3 groups in **2–4** are on the bridging sp^3 carbon atom and are not directly attached to the π -electron system. The potential ability of these CF_3 groups to alter the singlet and triplet levels (and ΔE_{ST}) are therefore of both fundamental and practical interest. The donor within the series **1–4** is systematically changed from the 9,9-dimethylacridine units (in **1** and **2**) to the 3,6-dimethoxycarbazole (in **3**) and 2,7-bis(trifluoromethyl)-9,9-dimethylacridine (in **4** and **5**).

The synthetic routes to **2–5** proceeded *via* palladium-catalysed Buchwald–Hartwig coupling of suitably functionalized D and A fragments.⁶⁶ The identity and high purity of **2–5** were confirmed by a combination of NMR spectroscopy, high resolution mass spectrometry, and X-ray crystallography (see the ESI[†]). The synthesis of molecule **4** is shown in Scheme 1, as a representative example.

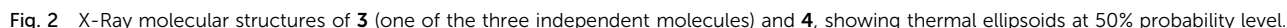




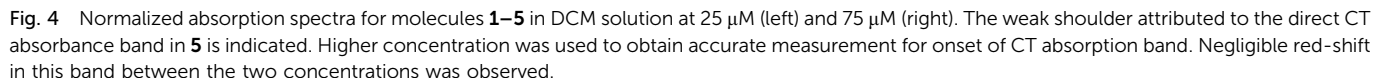
of **1**.⁶⁹ Typical oxidation of **12** gives the thioxanthene-*S,S*-dioxide acceptor precursor **13**.

To prepare the 2,7-bis-CF₃ substituted acridine donor **18**, the rings were constructed using starting materials with CF₃ functionality already installed. Ester formation yielded compound **15** from **14**. Installation of the dimethylmethylene bridge was achieved by reaction of MeMgBr with **15** to give alcohol **16**. Successful utilization of a recently reported protecting-group-free copper-catalyzed C–N coupling gave diarylamine compound **17**.⁷⁰ Compound **17** was cyclized to give new bis-CF₃-acridine donor **18** using Amberlyst-15. Compound **4** was synthesized from the coupling of **13** and **18** using typical Buchwald–Hartwig conditions, which generally tolerate a wide range of substituents for donor–acceptor couplings.^{47,65,71,72} Compounds **2**, **3** and **5** were also coupled using Buchwald–Hartwig conditions, with **2–5** prepared with yields ranging from 56 to 73%.

The crystal structures of **3** and **4** are shown in Fig. 2. Structure **3** is modulated along the $[1\ 1\ 1]$ direction: three symmetrically independent molecules are related roughly by pseudo-translations



All molecules in the series have π - π^* transitions in the 275–320 nm region. Molecule 2 has an additional shoulder at 325 nm. Compounds 1–5 have charge transfer (n - π^*) transitions at $\approx$ 350–375 nm. The dimethoxycarbazole moiety in 3 induces intense absorption bands in the 360–375 nm region,⁷⁴ which is due to a smaller torsion angle between the D–A units in carbazole functionalized systems (see Fig. 2: $<60^\circ$ dihedral angle for molecule 3 and $>70^\circ$ for molecule 4). The 360–375 nm band shape in 3 is not pure CT in character, *i.e.* not classically



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Table 2 Summary of key absorption and emission data in toluene and DCM solution

	Onset abs. (25 μ M)/nm ^a	Onset abs. (75 μ M)/nm ^a	Onset em./nm ^b	Onset em./nm ^a	λ_{max} em./nm ^b	λ_{max} em./nm ^a	Onset em. solvent shift/nm ^c	λ_{max} em. solvent shift/nm ^c
1	406	407	409	441	452	503	32	51
2	428	427	446	479	515	572	33	57
3	413	416	427	457	502	554	30	52
4	391	391	406	423	462	492	17	30
5	374	374	373	392	410	441	19	31

^a Measured in DCM solvent. ^b Measured in toluene solvent. ^c Shift in emission onset or λ_{max} from toluene to DCM solvent.

stronger CF₃ substituted acceptor (as in 2) is offset by the blue-shift associated with weaker donor (as in 5), leading to a similar emission profile as in 1. Unsubstituted 1 has slightly bluer emission in toluene and the emission of 4 is slightly bluer in dichloromethane, suggesting weaker CT emission character in 4 compared to 1. This difference is reflected in the solvatochromic shifts from toluene to dichloromethane in molecules 1 and 4.

Steady-state solid state photoluminescence

Compounds 1–5 were also characterized in three solid state host materials of differing polarities: zeonex, 1,3-bis(triphenylsilyl)benzene (UGH3), and (bis[2-(diphenylphosphino)phenyl]ether oxide) (DPEPO). UGH3 and DPEPO were chosen due to their high triplet energies.

All molecules 1–5 show some degree of red-shift in emission from apolar zeonex to DPEPO and UGH3 hosts. This red-shift is mainly due to packing effects in the solid state, as can be seen through the consistent similarity between UGH3 and DPEPO film spectra. Molecule 4 shows a significantly smaller shift in emission color between hosts. It is suggested that the weak CT strength makes 4 less sensitive to polarity of the surrounding medium, as found in solution state emission measurements (Fig. 5). Interestingly, the high degree of hydrophobicity imparted by the more extensive trifluoromethylation on molecule 4 may limit spectral shifts associated with modifications in molecular packing, considering that 5 (with a weaker acceptor than 4) and subsequently weaker charge transfer still significantly shifts from zeonex to DPEPO or UGH3 hosts.

Time-resolved photoluminescence

Time-resolved emission spectroscopy was performed on films of 1–5 in DPEPO host to determine the effect of trifluoromethyl substituents on the TADF properties of 2–5 in comparison to 1. Fig. 7 shows the time resolved emission decays and Fig. 8 shows snapshots of the emission at various time delays during the time resolved decay.

Compounds 1–4 all show charge transfer (CT) emission in the early ns region with a slight red-shift of the emission during in the first 80 ns of decay (Fig. 8 and Table 4). The CT emission at 80 ns representing the prompt fluorescence (PF) closely resembles the delayed fluorescence (DF) emission at 2 μ s. This is typical of emission originating from a TADF process, where intersystem crossing to the triplet excited state, and reverse intersystem crossing back to the ¹CT state has occurred. Molecules 1–4 have ΔE_{ST} gaps ≤ 0.22 eV with varying DF emission lifetimes. Temperature dependence measurements on 2–4 show increased emission intensity in the μ s delayed fluorescence region (Fig. S5a and b, ESI[†]). Power dependence measurements (Fig. S5c, ESI[†]) rule out multi-exciton emission processes with a slope close to one for 2–4, confirming that vibronically coupled TADF processes are occurring.

The lifetimes for PF and DF emission for 1–4 were determined by fitting with dual exponential decay curves (Fig. 7). Dual exponential curves gave good R_2 values (>0.98 , ESI[†]) for all fittings with $<10\%$ standard error in all cases (Table S1, ESI[†]). The emission for 3 and 5 was very weak and consequently the error in fitting the emission is larger. The start of the fitting for the DF was carefully considered to minimize interference

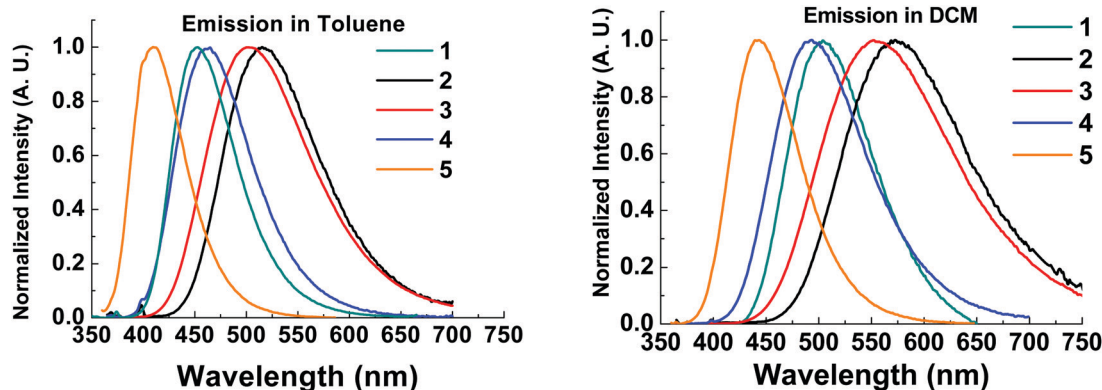


Fig. 5 Solution steady state emission spectra in toluene (left) and dichloromethane (right) of compounds 1–5. $\lambda_{\text{exc}} = 355$ nm.

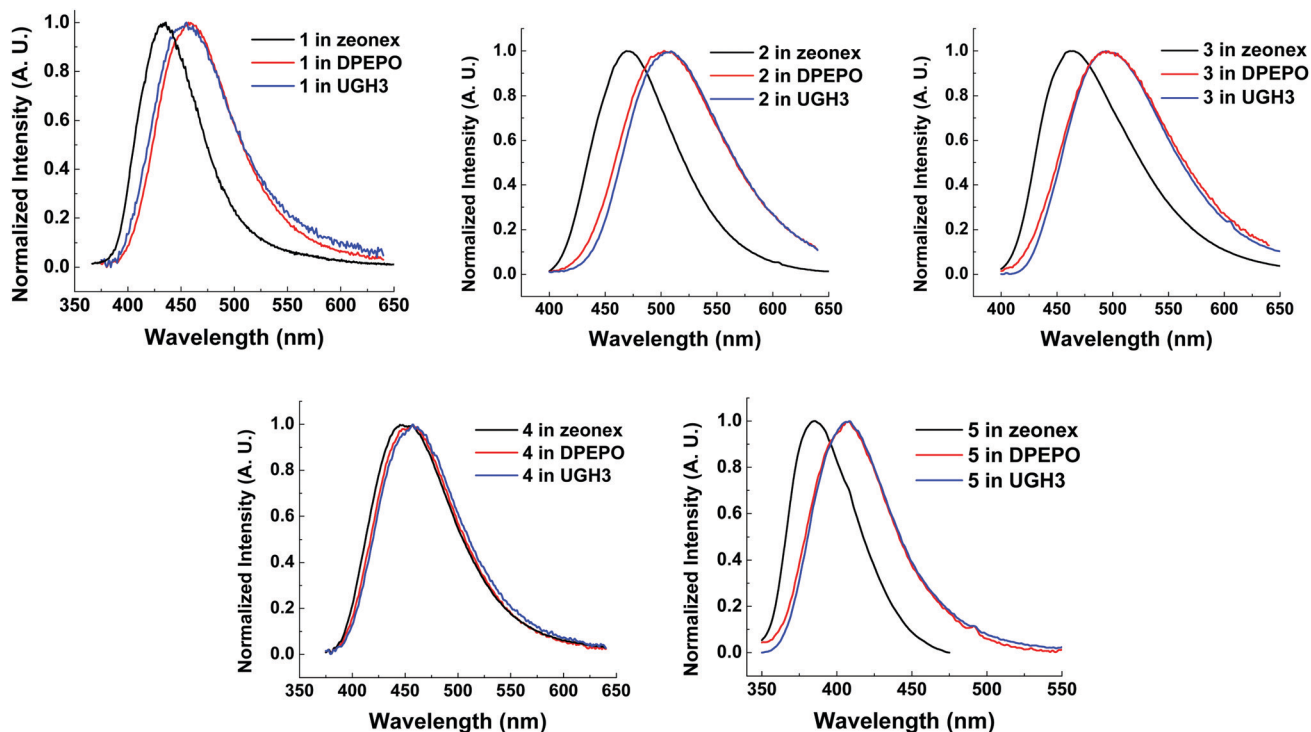


Fig. 6 Normalized steady state emission spectra from drop cast films of molecules **1–5** at 10 wt% in solid state hosts at room temperature. $\lambda_{\text{exc}} = 355$ nm.

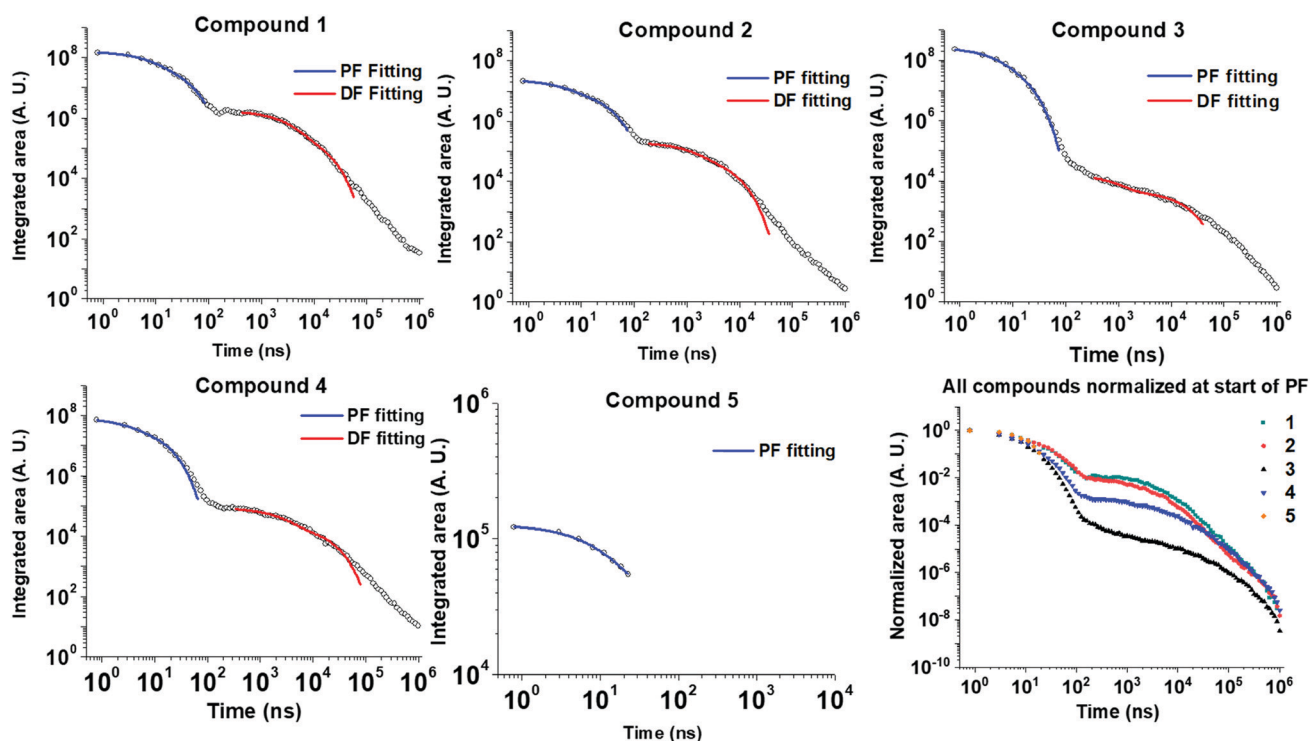


Fig. 7 Time-resolved photoluminescence decays of drop-cast films of **1–5** at 10 wt% in DPEPO host matrix. Dual exponential fittings performed for the PF (prompt fluorescence) in the early ns region and for the DF (delayed fluorescence) in the late ns/early μ s region. Fitting parameters are shown in Table S1 (ESI[†]). Lifetimes and corresponding weightings are shown in Table 3.

from the overlapping PF signal as much as possible at approx. 100 ns. For **1**, **2** and **4** both components of the dual exponential

decays contribute significantly to the total PF and DF emission (Table 3). This kinetic analysis for **1**, **2** and **4** suggests that there

Table 4 Summary of emission data for **1–5** in drop cast films at 10 wt.% in DPEPO host matrix

#	$E\ S_1/\text{eV (nm)}^a$	$E\ S_1/\text{eV (nm)}^b$	$E\ T_1/\text{eV (nm)}^c$	$\Delta E_{ST}/\text{eV}^d$	$\Delta E_{ST}/\text{eV}^e$	$\lambda_{\text{MAX all Em.}}/\text{nm}^f$	$\Phi_{\text{em}} \pm 5\%$
1	3.08 (403)	3.00 (413)	3.01 (412)	0.07	−0.01	460	80 ^g
2	2.86 (433)	2.77 (448)	2.78 (446)	0.08	−0.02	504	74
3	2.93 (423)	2.81 (442)	2.71 (458)	0.22	0.10	495	46
4	3.14 (395)	3.05 (406)	2.92 (425)	0.22	0.13	454	50 ^h
5	3.42 (363)	3.30 (376)	3.07 (404)	0.35	0.22	407	5 ⁱ

^a Onset of steady state emission in DPEPO host (Fig. 6). ^b Onset of relaxed CT emission at approx. 75 ns as indicated in Fig. 8 for **1–4**. For **5** the relaxed onset was taken at 11 ns due to little signal beyond this time. ^c Onset of phosphorescence emission at 80 K with a time delay of 80 ms (Fig. 8). ^d Calculated from steady state emission in DPEPO as in *a*. ^e T_1 energy taken from *c*. ^f Calculated from relaxed CT emission as in *b*. ^g T_1 energy taken from *c*. ^h Taken from steady state emission in DPEPO host (Fig. 6). ⁱ Taken from evaporated DPEPO film data from literature.⁷³ ^h Measured at 25 wt% in DPEPO evaporated film. Evaporated film at higher emitter concentration required in the blue region to obtain strong signal due to scattering and lower detector sensitivity in the blue region of the spectrum. ⁱ 1 wt% in zeonex matrix. Signal was too low to be measured in DPEPO film as shown by the signal to noise ratio in Fig. 7.

Computations

Geometry optimizations on the ground state (S_0) geometries of **1–5** at the hybrid DFT functional CAM-B3LYP/6-31G(d) revealed D–A orientations to be orthogonal in all molecules except for **3** where the averaged D–A dihedral angle of approximately 50° is obtained (Table 5). Good agreements between computed and experimental geometries were found for **3** and **4** (Fig. S7a, ESI†) so the optimized geometries for **1**, **2** and **5** are likely to be present as major conformers in solutions. Simulated absorption spectra from time-dependent DFT (TD-DFT) data (Fig. S7b, ESI†) on **1–5** are in accord with experimental absorption spectra with the intense mixed $^1\text{CT}/^1\text{LE}$ band for **3** present in the 370 nm region (oscillator strength, $f = 0.7$). The much greater intensity and strong CT/LE mixing in **3** compared to CT bands in **1**, **2**, **4** and **5** is due to the different D–A orientations. Constraining the D–A orientations to 90° in **3** results in a CT band with near zero intensity ($f = 0.0003$) (Fig. S7c and Table S5, ESI†). The perpendicular D–A energy barrier in **3** is estimated to be 2 kcal mol^{−1} thus a range of conformers of **3** in solution would be present.

Excited state geometries of **1–5** were also optimized here by TD-DFT for S_1 states and by unrestricted DFT (UDFT) for T_1 states with D–A dihedral angles of the optimized geometries listed in Table 5. For all molecules except **3**, the D–A orientations remain orthogonal on going from S_0 to S_1 so a relatively small geometrical rearrangement would take place after excitation and geometry relaxation to the most stable S_1 state for each geometry. To gain insight into the observed different emission data for **1–5**, results from TD-DFT data carried out on the optimized S_1 geometries are analyzed here with the assumption that these S_1 geometries are responsible for the charge-transfer emissions and are summarized (Table S8, ESI†).

Table S8 (ESI†) and Fig. 9 show the energies of the S_1 , T_1 and T_2 states and their orbital/excitonic characters for **1** and **2**. The ^3LE state is necessary to allow ISC/rISC processes to occur since the ^1CT and ^3CT exchange is formally forbidden.^{28,29} Large spin–orbit couplings (SOC) between ^1CT and ^3LE (Table S9, ESI†) with vibronic coupling (VC) between ^3CT and ^3LE do take place if the ^3LE state, ^3CT and ^1CT are close together. Compound **2** has a fast DF rate and T_1 with ^3CT character, whereas compound **4** has a slower observed DF rate and T_1 with ^3LE character like **1** (Table 3, and Fig. S7d, ESI†). Comparing compounds **1**, **2** and **4**, where very small systematic substitutions are made to the structure, it appears that the different T_1 character predicted is related to the DF rate where a faster rate occurs when T_1 has ^3CT character.

Fig. 9 highlights the differences between **1** where the ^1CT – ^3LE gap dominates whereas in **2** the ^3CT – ^3LE gap dominates. Thus in **1** rIC to populate T_2 is the rate limiting step, whereas rIC is not rate limiting in **2**, and **2** has faster rISC than **1**.⁷⁶

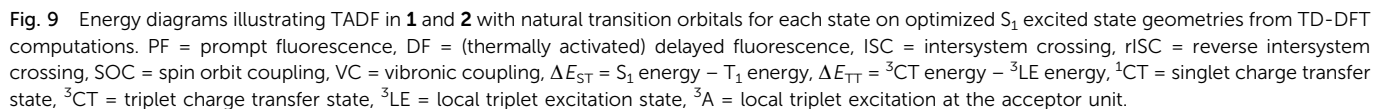
In the case of **3**, there is considerable early prompt fluorescence ^1CT (Fig. 8) and relatively low DF intensity, making it difficult to make quantitative comparisons in rates between **3** and the other compounds (Fig. 7). If the geometry in **3** contains D–A dihedral angles constrained at 50° in the S_1 state, the TD-DFT data provide an entirely different picture where the S_1 , T_1 and T_2 states are up to 0.59 eV apart and no rISC process would be expected (Fig. S7e and Table S4, ESI†). For **4**, the S_1 , T_1 and T_2 states are all within 0.17 eV and the larger energy gap suggests a less efficient rISC process in **4**. The nature of the ^1CT character in **4** with only 74% CT (and thus 26% local) is much less than the ^1CT characters in the other molecules (**1–3** and **5**)

Table 5 Relative D–A dihedral angles in degrees in optimized S_0 , S_1 and T_1 geometries.^a Experimental mean values in parentheses

	1	2	3	4	5
S_0	89.6, 89.6	84.1, 87.1	47.9, 49.1 (46.5)	84.4, 88.5 (81.3)	89.6, 89.6
S_1	85.8, 90.0	87.3, 87.8	52.2, 87.6	88.2, 89.0	82.3, 89.5
T_1	57.7, 89.4	55.8, 87.1	45.2, 49.3	54.7, 85.8	54.4, 89.3

^a Two averaged dihedral angles are reported because of the D–A–D architecture. When **1–5** are excited and charge transfer occurs, only one electron is transferred from a single donor to the central acceptor. In this excited state the other donor is acting as a ‘pendant’ substituent that will play a role in the electronics, but in a more passive fashion compared to the donor that transfers an electron. The ‘pendant’ substituents in the excited state geometries (S_1 , T_1) contain similar dihedral angles to those in the ground state geometries (S_0).





produces more stable devices,⁶⁵ it is likely that the acceptor is responsible for the instability in the OLED devices of **2** and **4**. The instability is likely to be due to the very high acceptor strength which results in cleavage of a C–C or C–S bond of the acceptor on reduction, rather than instability of the CF₃ functionality itself.

Conclusions and outlook

The CF₃-substituted acceptor in 2–4 has been an invaluable structural tool for probing the effects of orbital/exciton character on the rates of TADF emission to further understand the TADF mechanism and improve future molecular design. A key finding is that changing the character of the TADF-active singlet and triplet states through tuning with trifluoromethyl substituents improves the rate of TADF with rapid delayed emission components in the early microsecond regime. This work highlights how changes in the orbital character of singlet and triplet states

(local vs. charge transfer) lead to significant changes in rates of delayed fluorescence, despite the molecules possessing similar ΔE_{ST} values. The change in orbital character is supported by TD-DFT calculations. These results open up new important considerations for the design of TADF emitters where the ordering of the excited states has an impact on the rates of delayed fluorescence. To further enhance the rate of delayed fluorescence, and thereby reduce the probability of triplet interactions which cause detrimental annihilation effects, it is necessary to gain a deeper understanding of how singlet and triplet state character and orbital distribution affect emission rates. This will be a focus of future studies.

Author contributions

The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript. J. S. W. wrote the main body of the manuscript with assistance from all authors. J. S. W. designed the molecules and performed all the chemical synthesis and characterization under the supervision of M. R. B. A. D. performed the majority of the photophysical measurements with assistance from P. S. under the supervision of A. P. M. Processing of the photophysical data was performed by J. S. W., A. D. and P. S. M. A. F. performed and reported the DFT calculations. A. S. B. performed X-ray diffraction experiments and analysis on crystals grown by J. S. W. OLED devices were fabricated by A. D. Solution electrochemistry measurements were performed by J. S. W. and P. S. The funding for this work was obtained jointly by M. R. B. and A. P. M.

Conflicts of interest

The authors declare no competing financial interest.

Acknowledgements

We thank EPSRC grant EP/L02621X/1 and EU Horizon 2020 grant agreement number 732013 (HyperOLED) for funding. Dr Paloma Lays dos Santos is thanked for technical assistance. Dr Luke O' Driscoll is thanked for useful synthetic discussions. Dr Kleitos Stavrou and Dr Nils Haase are thanked for useful photophysical discussions.

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