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## Controlling through-space and through-bond intramolecular charge transfer in bridged D–D'–A TADF emitters†

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Donor–donor'–acceptor molecules where the donor' bridges the donor and acceptor have different possible interaction pathways for charge transfer. Here we study a series of donor–donor'–acceptor molecules, having the same acceptor and donor' but different donors, and donor'–phenyl spacer–acceptor to change the spatial separation and overlap between potential through-space donor–acceptor charge transfer (CT) in competition with donor'–acceptor through-bond CT. We determine that the charge transfer driving force plays a large role in dictating which charge transfer channel is favoured. Strong donors and acceptors with large driving force favour through-space CT. We also find that solid state host packing plays an important role, with small molecule hosts that pack tightly, distorting the guest molecules, reducing D–A separation to stabilise the through-space CT state over the through-bond state. Only the through-space CT states give fast reverse intersystem crossing and efficient TADF. These results give the first insight into the photophysics of through-space CT compared to through-bond states on the same molecule.

## Introduction

Organic molecules showing thermally-activated delayed fluorescence (TADF) are considered as the third generation of materials for organic light-emitting diodes (OLEDs).<sup>1</sup> Unlike phosphorescent emitters, TADF molecules harvest triplet excitons by a reverse intersystem crossing (rISC) mechanism

between the triplet (<sup>3</sup>CT) and the singlet (<sup>1</sup>CT) charge-transfer states, mediated by vibronic coupling between <sup>3</sup>CT and a local excited triplet state (<sup>3</sup>LE) that enables spin orbit coupling (SOC), when the energy gap between all three states becomes small, <100 meV. Mediation by the third state is required when <sup>3</sup>CT and <sup>1</sup>CT are so close in energy because direct spin orbit coupling (SOC) between them is forbidden because they have effectively the same orbital and no change in orbital angular momentum occurs during the direct transition <sup>3</sup>CT → <sup>1</sup>CT.<sup>2,3</sup> One of the simplest ways to achieve extremely small electron correlation energy, *i.e.* singlet–triplet splitting energy ( $\Delta E_{ST}$ ), is the use of conformational twisting between directly bridged donor (D) and acceptor (A) units in order to minimize the overlap between the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO).<sup>4,5</sup> Beyond the subtle manipulations of the degree of intramolecular through-bond charge transfer to control TADF properties, recent interest in through-space charge transfer between D and A, mediated by strong D A intramolecular  $\pi$ – $\pi$  dipolar interactions has been motivated by the potential of increasing photoluminescence quantum yield.<sup>6,7</sup> Such intramolecular through-space charge transfer is much like an exciplex state except that it offers far more control over the spatial separation and orientation of D and A compared to the random orientation of D and A in the intermolecular exciplex state. In a conjugated system, the D and A can be arranged co-facially, in close spatial proximity, using a non-coplanar molecular scaffold (bridge).<sup>8</sup> This approach has also been used in the design of non-conjugated polymers with spatially separated pendant D and A.<sup>9,10</sup> Despite the growing interest of intramolecular through-space charge transfer states as a means to TADF, very little photophysical study of the excited state dynamics and molecular conformations of such systems has been made.<sup>11</sup> Here we investigate the excited state dynamics of new molecules configured with asymmetrical donor–donor'–acceptor (D–D'–A) architectures, where the co-facial overlap between D and A is controlled by the introduction of a common

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Fig. 1 The chemical structures (a) and X-ray structures (b) of **Ph<sub>3</sub>TRZCzTPA**, **Ph<sub>2</sub>TRZCzTPA**, **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz**, thermal ellipsoids were drawn at a 50% probability level.

Table 1 The dihedral angles and distances of **Ph<sub>2</sub>TRZCzTPA**, **Ph<sub>3</sub>TRZCzTPA**, **Ph<sub>2</sub>TRZCzPhCz**, and **Ph<sub>3</sub>TRZCzPhCz**

	$\varphi_1$ (°)	$\varphi_2$ (°)	$\varphi_3$ (°)	$\varphi_4$ (°)	$\varphi_5$ (°)	$d_1$ (Å)	$d_2$ (Å)	$d_3$ (Å)	$d_4$ (Å)
<b>Ph<sub>2</sub>TRZCzTPA</b>	49.3	55.2				3.093	4.985		
<b>Ph<sub>2</sub>TRZCzPhCz</b>	36.5	47.1				3.373	4.926		
<b>Ph<sub>3</sub>TRZCzTPA</b>			75.1	70.9	0.6			3.150	5.295
<b>Ph<sub>3</sub>TRZCzPhCz</b>			65.4	57.8	2.5			3.201	4.924



weak (rigid) carbazole donor bridge (D') and by the introduction of phenyl spacer units between the acceptor and bridge, shown in Fig. 1. A common diphenyltriazine (dTRZ) acceptor is used and either a weak phenyl carbazole (PhCBZ) or strong triphenylamine (TPA) donor.

## Results

Synthetic methods and characterization of these new D–D'–A molecules and model compounds used in this study are reported in ESI† (Scheme S1–S3). Fig. 1b shows their single crystal structures and crystallographic data is summarized in Table S1 (ESI†). As shown, the pendant (spacer) aryl rings are

highly twisted from the carbazole bridge (D') due to congested steric interactions. All dihedral angles between the D, A and bridging D', selected closest atom-to-atom distances between the donor and the acceptor branches (3.09–3.37 Å), and distances between the donor N-atom and the centre of triazine acceptor ring (4.93–5.30 Å) are summarized in Table 1.

In general, **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** (triphenylamine donor) show larger donor-bridge dihedral angles as compared to those of the counterpart **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz** (PhCBZ donor). In addition, **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** show shorter closest atom-to-atom distances, but slightly longer distances between the donor N-atom and the centre of triazine ring. These distorted conformations and short distances imply that a through-space interaction between the donor and acceptor branches due to the stronger electron-donating ability of TPA may stabilize the twisted molecular structures. All of the molecules show excellent thermal stability required for stable film and device formation, and calculated HOMO/LUMO energy levels obtained from cyclic voltammetry (Fig. S1, ESI†) are given in Table 2. **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** show higher HOMO energy levels (–5.28/–5.27 V) as compared to **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz** (–5.85/–5.57 V) in accord with donor strength. Since the acceptor of all compounds is the same, the LUMO energy levels are similar (*ca.* –2.75 V) in all cases.

### DFT calculations

The ground state structures were first optimized at a B3LYP/6-311G(d) level. The calculated HOMO and LUMO distributions as well as energy levels are shown in Fig. S2 (ESI†). The HOMOs of **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** are mainly located at the



Table 2 The electrochemical properties and thermal properties of **Ph<sub>2</sub>TRZCzTPA**, **Ph<sub>3</sub>TRZCzTPA**, **Ph<sub>2</sub>TRZCzPhCz**, and **Ph<sub>3</sub>TRZCzPhCz**

	$E_{\text{onset}}^{\text{oxi}}$ <sup>a</sup> (V)	$E_{1/2}^{\text{red}}$ <sup>b</sup> (V)	HOMO <sup>c</sup> (eV)	LUMO <sup>c</sup> (eV)	$E_g^d$ (eV)	$T_d^e$ (°C)	$T_g^f$ (°C)	$T_m^f$ (°C)
<b>Ph<sub>2</sub>TRZCzTPA</b>	0.98 <sup>b</sup>	-1.45	-5.28	-2.73	2.55	324	111	212
<b>Ph<sub>3</sub>TRZCzTPA</b>	0.97 <sup>b</sup>	-1.53	-5.27	-2.74	2.53	389	111	279
<b>Ph<sub>2</sub>TRZCzPhCz</b>	1.21	-1.47	-5.85	-2.79	3.06	340	127	n.d. <sup>g</sup>
<b>Ph<sub>3</sub>TRZCzPhCz</b>	1.23	-1.50	-5.57	-2.77	2.80	419	129	338

<sup>a</sup> Calculated from the onset potential. <sup>b</sup>  $E = (E_{p,a} + E_{p,c})/2$ , where  $E_{p,a}$  and  $E_{p,c}$  stand for the peak potential at which anodic- and cathodic-direction scan, respectively. <sup>c</sup> HOMO and LUMO were determined from the electrochemical results in CH<sub>2</sub>Cl<sub>2</sub> and DMF solution, respectively. <sup>d</sup>  $E_g = \text{LUMO} - \text{HOMO}$  using electrochemical results. <sup>e</sup>  $T_d$  was analyzed using TGA (5% weight loss). <sup>f</sup>  $T_g$  and  $T_m$  were determined by DSC. <sup>g</sup> Not detected.

TPA moiety and slightly extend to the carbazole bridge. The HOMOs of **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz** are mostly positioned at the pendant PhCBZ donor with limited contribution from the carbazole bridge. Not surprisingly, the LUMOs of **Ph<sub>2</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzPhCz** are located at the dTRZ, which are well separated from their HOMOs. Whereas, the LUMOs of **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>3</sub>TRZCzPhCz** are located at the TRZ together with a small contribution on the carbazole bridge, giving weak but non zero HOMO–LUMO overlap. Obviously, the stronger donor, TPA, leads **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** to have higher HOMO levels as compared to those of **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz**. Whereas **Ph<sub>2</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzPhCz** without the aryl spacer between acceptor and bridge show slightly lower LUMOs. The calculated results are consistent with the observed CV data.

TD-DFT calculations were then made using a hybrid meta-generalized gradient-approximation functional, m06-2x/6-311(d), to optimize the S<sub>1</sub> and T<sub>1</sub> structure of these four molecules and calculate the natural transition orbitals (NTOs). The HONTO and LUNTO distributions are shown in Fig. S3 (ESI<sup>†</sup>). For the S<sub>1</sub> state of **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA**, the HONTOs distribute on the TPA moiety, while the LUNTOs delocalises at the dTRZ and TRZ, respectively. Since there is nearly no overlap between the HONTO and LUNTO, a through-space charge transfer character for the S<sub>1</sub> state is suggested. For the T<sub>1</sub> state configuration, the HONTO of the **Ph<sub>3</sub>TRZCzTPA** is delocalised on the TPA and TRZ, exhibiting both CT and LE transition character. On the other hand, the HONTO of the **Ph<sub>2</sub>TRZCzTPA** only distributes on the dTRZ, indicating the

<sup>3</sup>LE state is the lowest triplet state. Notably, a distinguishable change between the S<sub>1</sub> and T<sub>1</sub> orbitals could be observed for **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA**, which is a prerequisite for an allowed SOC spin–flip transition. Very interestingly, the different orbital distributions of S<sub>1</sub> and T<sub>1</sub> states could also be observed in the case of **Ph<sub>2</sub>TRZCzPhCz**. The HONTO of the S<sub>1</sub> state distributes on the PhCBZ donor and the carbazole bridge, while the LUNTO locates on the dTRZ. On the other hand, the HONTO of the T<sub>1</sub> state of **Ph<sub>2</sub>TRZCzPhCz** delocalises mainly on the dTRZ group and slightly on the phenylene of the pendant PhCBZ donor, whereas the LUNTO only locates on the dTRZ group. Thus, **Ph<sub>2</sub>TRZCzPhCz** is anticipated to possess both through-bond and through-space CT character due to the slight overlap of HONTO and LUNTO. For **Ph<sub>3</sub>TRZCzPhCz**, the HONTO of the S<sub>1</sub> state distributes on the pendant PhCBZ, whereas the LUNTO distributes on the TRZ group, leading to a through-space CT transition character. The HONTO and LUNTO distributions of the T<sub>1</sub> state are similarly delocalised on the PhCBZ and slightly coupled with the carbazole bridge, resulting in an apparent LE transition character. Although **Ph<sub>3</sub>TRZCzPhCz** shows considerable orbital deformation between S<sub>1</sub> and T<sub>1</sub> states, the larger  $\Delta E_{\text{ST}}$  undermines the probability of reverse intersystem crossing process.

### Steady state photophysics

Steady state absorption and emission of each compound was measured in methylcyclohexane (MCH) are shown in Fig. 2a and b, and in a range of different polarity solvents and solid state hosts in Fig. 3, for direct comparison.



Fig. 2 Extinction coefficients (a) and emission spectra (b) of the molecules in MCH solution (20  $\mu\text{M L}^{-1}$ ).





Fig. 3 Emission spectra of the compounds (a)  $\text{Ph}_3\text{TRZCzTPA}$ , (b)  $\text{Ph}_2\text{TRZCzTPA}$ , (c)  $\text{Ph}_2\text{TRZCzPhCz}$  and (d)  $\text{Ph}_3\text{TRZCzPhCz}$  in film and different solvents, methylcyclohexane (MCH), chlorobenzene (CB) and acetonitrile (MeCN).

The lowest energy absorption band, peaking at *ca.* 375 nm in  $\text{Ph}_3\text{TRZCzPhCz}$  and  $\text{Ph}_3\text{TRZCzTPA}$  is absent in  $\text{Ph}_2\text{TRZCzTPA}$  and  $\text{Ph}_2\text{TRZCzPhCz}$ . In  $\text{Ph}_3\text{TRZCzPhCz}$  this band correlates very well to the lowest energy absorption band in the model compound, 9-(4-(4,6-diphenyl-1,3,5-triazin-2-yl)phenyl)-9*N*-carbazole,  $\text{CzPhTrz}$ , as observed by Sharma *et al.*,<sup>12</sup> having strong  $\pi\pi^*$  character, and indicates a more delocalised acceptor unit, *i.e.* the phenyl spacer between A and bridge forms part of the acceptor unit itself in both cases, as suggested by our TD-DFT calculations. The magnitude of the extinction coefficients of these lowest energy bands is also  $> 2 \times 10^4 \text{ cm}^{-1} \text{ M}^{-1}$  indicating that they are not direct CT absorption bands as observed in typical D–A–D through bond TADF molecules.<sup>4</sup>

Emission spectra measured for each compound in MCH, Fig. 2b, show energy on-sets at; 2.89 eV, 2.79 eV, 3.59 eV and 3.32 eV for  $\text{Ph}_3\text{TRZCzTPA}$ ,  $\text{Ph}_2\text{TRZCzTPA}$ ,  $\text{Ph}_2\text{TRZCzPhCz}$  and  $\text{Ph}_3\text{TRZCzPhCz}$ , respectively.  $\text{Ph}_2\text{TRZCzPhCz}$  gives dual emission in MCH from a local donor at 3.59 eV and a charge transfer (CT) state at lower energy, in excellent agreement with the TD-DFT calculations.  $\text{Ph}_2\text{TRZCzTPA}$  emission shows some mixed local/CT character, indicated by the weak structure on the emission band. The stronger donor strength of  $\text{Ph}_3\text{TRZCzTPA}$  and  $\text{Ph}_2\text{TRZCzTPA}$  is clearly manifest by the large Stokes shift observed in their CT emission spectra. With increased solvent polarity, Fig. 3.  $\text{Ph}_3\text{TRZCzTPA}$ ,

$\text{Ph}_2\text{TRZCzTPA}$ , and  $\text{Ph}_3\text{TRZCzPhCz}$  all show strong solvatochromism, giving a positive Stokes shift indicative of CT emission. Emission of  $\text{Ph}_3\text{TRZCzTPA}$  and  $\text{Ph}_2\text{TRZCzTPA}$  in MeCN is highly red shifted and extremely weak indicating very low emission yields.

In general, the molecules with a Cz donor emit at higher energy than those having a TPA donor, in line with donor strength. This also dictates the driving force for charge transfer in each pair. The CT driving force may be estimated from the following equation;

$$\Delta G_0 = EA_A - EA_D - \Delta E_b$$

Where  $EA_x$  are the electron affinity of A and D fragment respectively, and  $\Delta E_b$  is the difference in binding energy between the initially created exciton and the relaxed charge separated state, see Scheme S3 (ESI<sup>†</sup>). In the case of these rigid orthogonal intramolecular D A molecules we assume that this difference in binding energy to be very small in non-polar media. Taking the measured (electrochemical) ionization potentials for the two families as that of the D units, the EA of the D units were estimated by adding the D optical excitation energy, yielding  $EA_{\text{Cz}} = -2.21 \text{ eV}$  and  $EA_{\text{TPA}} = -1.76 \text{ eV}$ . The measured EA of the A unit in all cases being  $-2.75 \text{ eV}$  from the CVs. For  $\text{Ph}_3\text{TRZCzTPA}$  and  $\text{Ph}_2\text{TRZCzTPA}$  we then estimate a CT driving force of *ca.* 1 eV, and for  $\text{Ph}_2\text{TRZCzPhCz}$  and



**Ph<sub>3</sub>TRZCzPhCz**, *ca.* 0.5 eV. We thus find a near double CT driving force for **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** compared to **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz**. It is interesting to note that for these rigid D–A intramolecular systems, in non-polar environment (minimising the Coulomb attraction energy) we also find that the difference between the (optical) D exciton energy and CT energy also gives very similar values, 0.8 eV for **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** and 0.5 eV for **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz**, yielding a potential very simple way to estimate the driving energy for photoexcited charge transfer in highly decoupled intramolecular D–A molecules.

The solution measurements for **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** are similar as can be seen in Fig. 3a and b, the bands shift by the same amount in different polarity solvents. Comparing the film spectra, we see that the zeonex films behave just like the MCH solutions for these materials with **Ph<sub>3</sub>TRZCzTPA** even showing a small blue shift compare to MCH. **Ph<sub>2</sub>TRZCzPhCz** also shows the same <sup>1</sup>LE and <sup>1</sup>CT dual emission bands, but the CT state has increased relative intensity in zeonex. However, in DPEPO films, no <sup>1</sup>LE component is observed from **Ph<sub>2</sub>TRZCzPhCz**, only the <sup>1</sup>CT emission is observed, at about the same peak wavelength as the corresponding **Ph<sub>2</sub>TRZCzPhCz** CT emission in toluene solution.

**Ph<sub>3</sub>TRZCzPhCz**, Fig. 3d, is seen to have a very similar spectrum with weakly structured emission in MCH compared to the model systems CzPhTrz<sup>12</sup> and *meta*-CzTRZ reported by Liu *et al.*<sup>13</sup> Thus, we ascribe its excited state as a through-bond CT state between the bridge Cz and TRZ D'–A pair. The CT state shows both weak solvatochromism and in solid state the emission has very similar band shape and width as in solution, mostly unaffected by host packing effects. Only in highly polar MeCN does the band become highly red shifted and very broad, which may point to a different excited state origin such as a through-space CT state.

**Ph<sub>2</sub>TRZCzPhCz** on the other hand is obviously different, being the only compound showing dual CT emission and very strong local emission, consistent with the X-ray data that shows a shallow 36.5° dihedral angle between D'–A. Cz donor units tend to have shallow ground state D–A dihedral angles giving strong LE character.<sup>14</sup> This is further emphasised by the localisation of the HOMO on the pendant Cz unit found in our DFT calculations. We therefore synthesised the model compound, 2-(carbazol-9-yl)-4,6-diphenyl-1,3,5-triazine (**CzTrz**), Fig. S4 (ESI†). The absorption spectrum of **CzTrz** indicates electronically decoupled D and A units, absorbing above 350 nm. Excitation into the A (TRZ) band yields both TRZ<sup>15</sup> and weak CT emission, whereas excitation into the D (Cz) band yields strong CT emission, onset at 3.26 eV. The CT emission on-set shifts to 3.1 eV in toluene and dominates the emission spectra. This shift is 35 nm more to the red than for the model **CzPhTrz** in toluene, which we ascribe to a smaller electron hole separation in **CzTrz** therefore larger Coulomb attraction energy, which is relaxed more strongly by the increasing solvent polarity red shifting the energy of the CT state to a greater extent.<sup>16</sup> This clearly corresponds with the CT state seen in **Ph<sub>2</sub>TRZCzPhCz** which can then be identified as a through-bond

CT state between the bridge Cz (D') and dTRZ unit. The higher intensity, narrow linewidth blue emission band with vibronic structure, also observed in **Ph<sub>2</sub>TRZCzPhCz**, onset *ca.* 3.54 eV, showing little or no solvatochromic shift, must be emission from a local excited state. Considering the individual D and A units in the molecule, PhTRZ is only very weakly emissive through excitation at 280 nm, emission onset at 3.44 eV, with slight vibronic structure.<sup>15</sup> Emission in our model **CzTrz** peaking at *ca.* 290 nm seems highly plausible to be from the dTRZ unit. This is not what we observe in **Ph<sub>2</sub>TRZCzPhCz**, however, carbazole shows strong emission, onset at 3.54 eV<sup>17</sup> and so we conclude that this local emission band in **Ph<sub>2</sub>TRZCzPhCz** comes from the pendent Cz D unit, consistent with the DFT calculations. This Cz must be strongly decoupled from the rest of the molecule as the emission is always present even in high polarity solvent, ruling out a through-space CT state. In highly polar MeCN, there is a small population of CT states in giving rise to the highly red shifted band which we believe to be through-space CT states, Fig. 3c.

With this in mind, we turn to **Ph<sub>2</sub>TRZCzPhCz** in zeonex (a branched polyolefin) and DPEPO (bis[2-(diphenylphosphino)phenyl]ether oxide) host films. In both cases the CT band is much stronger than in solution. In zeonex dual emission is still observed but the CT emission has equal intensity to the local emission, whereas in tightly packed DPEPO host, only a strong, red shifted and very broad CT emission is observed. As there can be no 'solid state solvatochromic effect' in a rigid solid host,<sup>18,19</sup> this must be due to a packing effect of the host. If the host forces the D and A units closer together, improving co-facial overlap, a through-space CT state can be stabilized and the increased charge transfer rate would effectively quench donor emission from the pendent Cz unit. Thus, in DPEPO we conclude that a through-space CT state is stabilized on **Ph<sub>2</sub>TRZCzPhCz**, in-line with TD-DFT calculations that strongly suggest both through-space and through-bond CT states are possible. We also note that this CT emission band in DPEPO is much broader and more red shifted than the CT emission observed from **Ph<sub>3</sub>TRZCzPhCz** in DPEPO. This we believe indicates a very inhomogeneous process stabilising the through-space CT state, clearly in-line with random host packing effects. The stable through-bond CT state in **Ph<sub>3</sub>TRZCzPhCz** remains narrow in DPEPO, which potentially indicates that the extra flexibility introduced by the spacer phenyl ring enables closer co-facial separation of D and A stabilising the through-space CT state so that host packing has little extra effect on this conformation. To confirm this hypothesis we measured **Ph<sub>2</sub>TRZCzPhCz** in both mCP (1,3-bis(*N*-carbazolyl)benzene) and CBP (4,4'-bis(*N*-carbazolyl)-1,1'-biphenyl) hosts, both having very low dielectric and polarizability values compared in DPEPO. In both cases clear CT emission is observed from **Ph<sub>2</sub>TRZCzPhCz**, Fig. S5 (ESI†), confirming that host packing plays a major role in stabilising through-space CT in these materials.

To understand **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** we recall an important observation from our previous work. A CT complex can form between TRZ and TPA in solution.<sup>11</sup> The



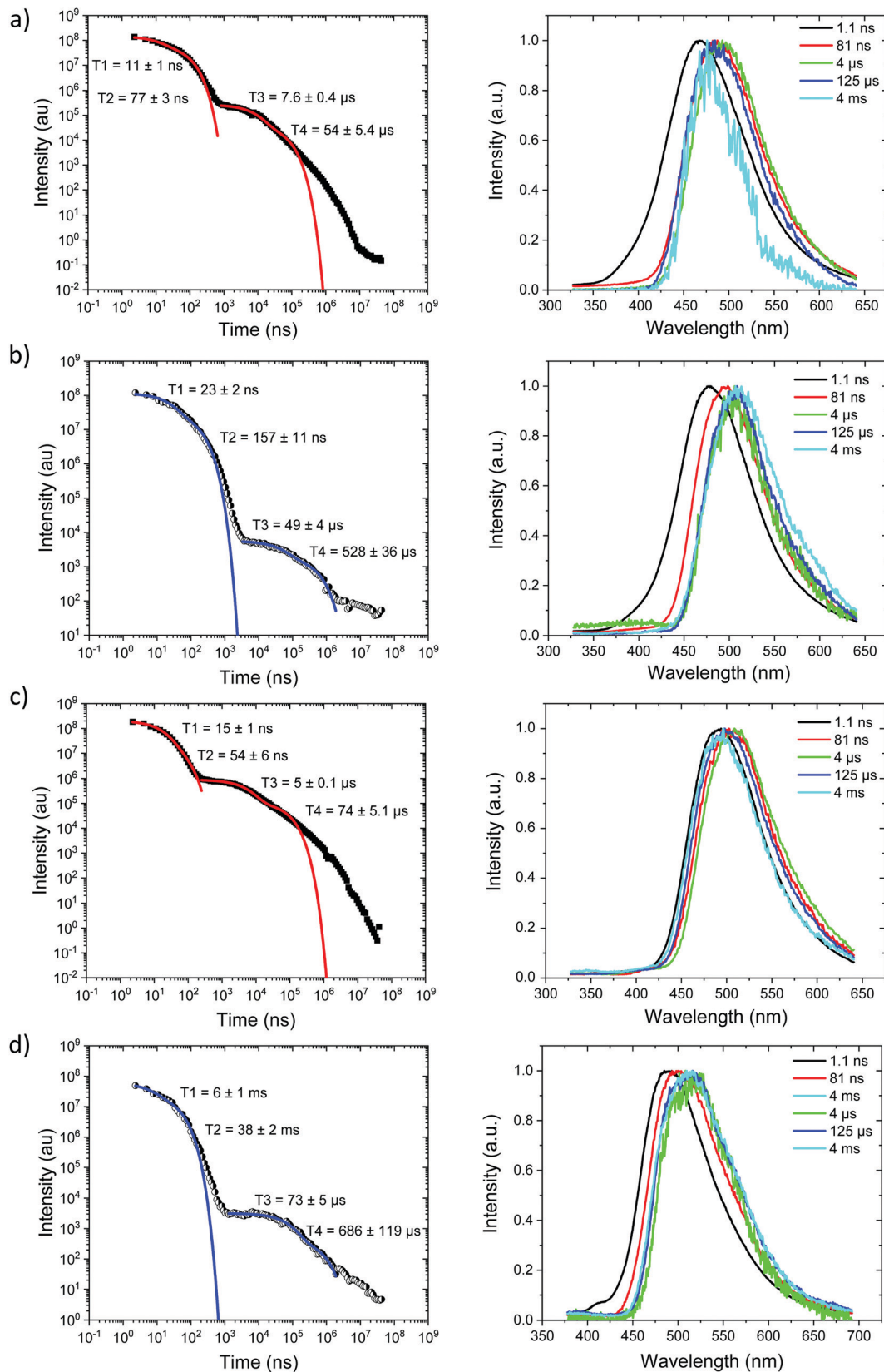


Fig. 4 Time-resolved spectrally integrated intensity decay (left) and representative time resolved spectra (right) of  $\text{Ph}_3\text{TRZCzTPA}$  (a) 300 K and (b) 80 K;  $\text{Ph}_2\text{TRZCzTPA}$  (c) 300 K and, (d) 80 K in zeonex matrix. Spectral times given in ns, and decay fitting data included in the decay plots.



resulting TRZ:TPA excited state emits at 492 nm (peak) in toluene (with solvatochromism being observed, Fig. S6, ESI<sup>†</sup>). This is at the same energy with the same solvatochromic behaviour as the band observed in both **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA**. We therefore identify the CT emission in both as coming from a through-space TRZ ··· TPA (strong D – strong A) CT excited state. No contribution from a through-bond, strong A – weak D' CT pair is observed. Concomitantly, emission is seen well to the red of either **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz** CT emission in line with a strong-D strong-A pair CT state. We note from X-ray measurements that this behaviour does not correlated simply to D–A co-facial separation, indicating the role of CT driving force. We also previously found that the intermolecular TRZ ··· TPA state has a ground state interaction, *i.e.* a heterodimer complex,<sup>11</sup> and we postulate that such a ground state interaction between D and A units may also help to drive and stabilise the through-space CT state over a through-bond interaction. Such ground state absorption is also seen in other intramolecular through-space CT systems.<sup>20</sup>

### Time-resolved spectroscopy

Starting with the compounds in zeonex host matrix. **Ph<sub>3</sub>TRZCzTPA** at room temperature, Fig. 4a, shows initial CT like emission (Gaussian band shape) in the first few ns, onset at 3.01 eV which relaxes in energy over 100 ns to give strong prompt (PF) CT emission at 2.88 eV and delayed fluorescence (DF) at 2.87 eV (on-sets). The DF then undergoes a monotonic blue shift of some 50 meV over the first 40 μs, Fig. S7 (ESI<sup>†</sup>). Phosphorescence, onset at 2.90 eV is seen at millisecond times. Fitting the emission decay indicates a fast 11 ns and a slow 77 ns prompt CT decays along with a 1 μs and 13 μs DF components. All parameters are given in Table 3. Here we note that the DF decays are highly non exponential at longer times, turning into a power law decay. This is very typical of TADF materials in the solid state. Thus, here we fit the early time DF with one or two exponential terms to model the majority of the DF emission and calculate rISC rates from a kinetic model as detailed previously.<sup>21</sup> The possible interpretation of the power law DF region will be discussed later.

At 80 K, Fig. 4b and Fig. S8 (ESI<sup>†</sup>), we see similar behaviour but the ns region prompt emission energy relaxation is slower. However, the emission band has a constant additional 60 meV red shift and exactly matches the band shape and energy position (in time) of the emission observed in **Ph<sub>2</sub>TRZCzTPA** (both at RT and 80 K, Fig. 4c and d). The large decrease in DF

intensity shows clear thermal activation of the DF mechanism. Phosphorescence at 80 K, Fig. 4b, also has a much lower onset of 2.76 eV. This large difference in CT state energy between RT and 80 K implies that thermal motion on average leads to a higher energy CT state in **Ph<sub>3</sub>TRZCzTPA**. As this behaviour is not observed in **Ph<sub>2</sub>TRZCzTPA** we conclude that it must be an effect due to the presence of the spacer phenyl ring between A and bridge. One explanation for this is that the thermal energy drives the spacer phenyl ring more planar (conjugated). So at 80 K the phenyl ring has a more orthogonal configuration with respect to the A and bridge, changing the coupling of the phenyl ring to the dTRZ and localising the LUNTO on the dTRZ unit analogous with **Ph<sub>2</sub>TRZCzTPA** as shown in our TDDFT calculations. This will also greatly affect the localisation of the lowest energy triplet state as well.

**Ph<sub>2</sub>TRZCzTPA**, Fig. 4c and d, shows constant <sup>1</sup>CT emission behaviour, onset at 2.82 eV both at RT and 80 K with a monotonic blue shift of 50 meV over the first 40 μs, Fig. S7 and S8 (ESI<sup>†</sup>). Phosphorescence, onset at 2.80 eV is seen at millisecond times. At RT, we see faster prompt decay, earlier time DF and faster DF decay compared to **Ph<sub>3</sub>TRZCzTPA**. As the  $\Delta E_{ST}$  are less than 30 meV (at RT) in both cases, this could indicate stronger magnetic coupling between D and A in **Ph<sub>2</sub>TRZCzTPA**, so more efficient SOC, given by the different spatial overlap and orientation of D and A units in the two compounds, again an effect that can be related to the spacer phenyl unit. As seen in the DFT calculations the D–A separation in **Ph<sub>2</sub>TRZCzTPA** (4.985 Å) is smaller compared to **Ph<sub>3</sub>TRZCzTPA** (5.295 Å) fully supporting this argument. Previously, Wada *et al.*<sup>20</sup> have proposed that the D A co-facial angle is critically in determining the SOC and rISC rate in a similar through-space CT system.

**Ph<sub>2</sub>TRZCzPhCz** at RT in zeonex, Fig. 5a, has a PF energy of 3.54 eV that decays within a few nanoseconds as expected for D emission. A simultaneous CT emission band (on our 1 ns timescale), onset at 3.10 eV, is also observed that decays very rapidly, within 100 ns, shown in detail in Fig. S18 (ESI<sup>†</sup>). At 100 ns we also see a weak transient CT emission band, onset at 2.95 eV, which also decays rapidly. This is similar to what we observe in DPEPO (see later) and may indicate a small population of weakly stabilised through-space CT states in **Ph<sub>2</sub>TRZCzPhCz**, as predicted by our calculations, and that are stabilised in high polarity MeCN solution, Fig. 3c. This band is better resolved at 80 K, Fig. 5b, with all prompt emission decayed by 200 ns. From 200 ns to 10 μs we observe no emission (above the noise floor of the iCCD), Fig. S7 (ESI<sup>†</sup>). At

Table 3 Time resolved data of zeonex film measurement at RT

Molecule	<sup>1</sup> LE <sup>a</sup> (eV)	$\tau_{LE}$ (ns)	<sup>1</sup> CT <sup>a</sup> (eV)	$\tau_{PF1}^b$ (ns)	$\tau_{PF2}^b$ (ns)	$\tau_{DF1}^b$ (μs)	$\tau_{DF2}^b$ (μs)	<sup>1</sup> CT <sup>a/c</sup> (eV)	<sup>3</sup> LE <sup>c</sup> (eV)	$\Delta E_{ST}^d$ (meV)
<b>Ph<sub>3</sub>TRZCzTPA</b>	3.01	~1	2.88	11	77	7.6	54	—	2.90	20
<b>Ph<sub>2</sub>TRZCzTPA</b>	—	—	2.82	15	54	5	74	—	2.80	20
<b>Ph<sub>2</sub>TRZCzPhCz</b>	3.54	3	3.10	3	9	342	4900	2.92	2.76	340
<b>Ph<sub>3</sub>TRZCzPhCz</b>	—	—	3.35	9	—	—	—	—	2.88	470

<sup>a</sup> Local (LE) and charge transfer (CT) emission energies calculated from the on-set of the emission band. <sup>b</sup> Prompt ( $\tau_{LE}$  and  $\tau_{PF}$ ) and delayed fluorescence ( $\tau_{DF}$ ) component lifetimes calculated from fits to the measured intensity decay curves. <sup>c</sup> Local triplet energy taken from the on-set of the phosphorescence measured at 80 K. <sup>d</sup>  $\Delta E_{ST}$  is the energy difference between the <sup>1</sup>CT and <sup>3</sup>LE energies.



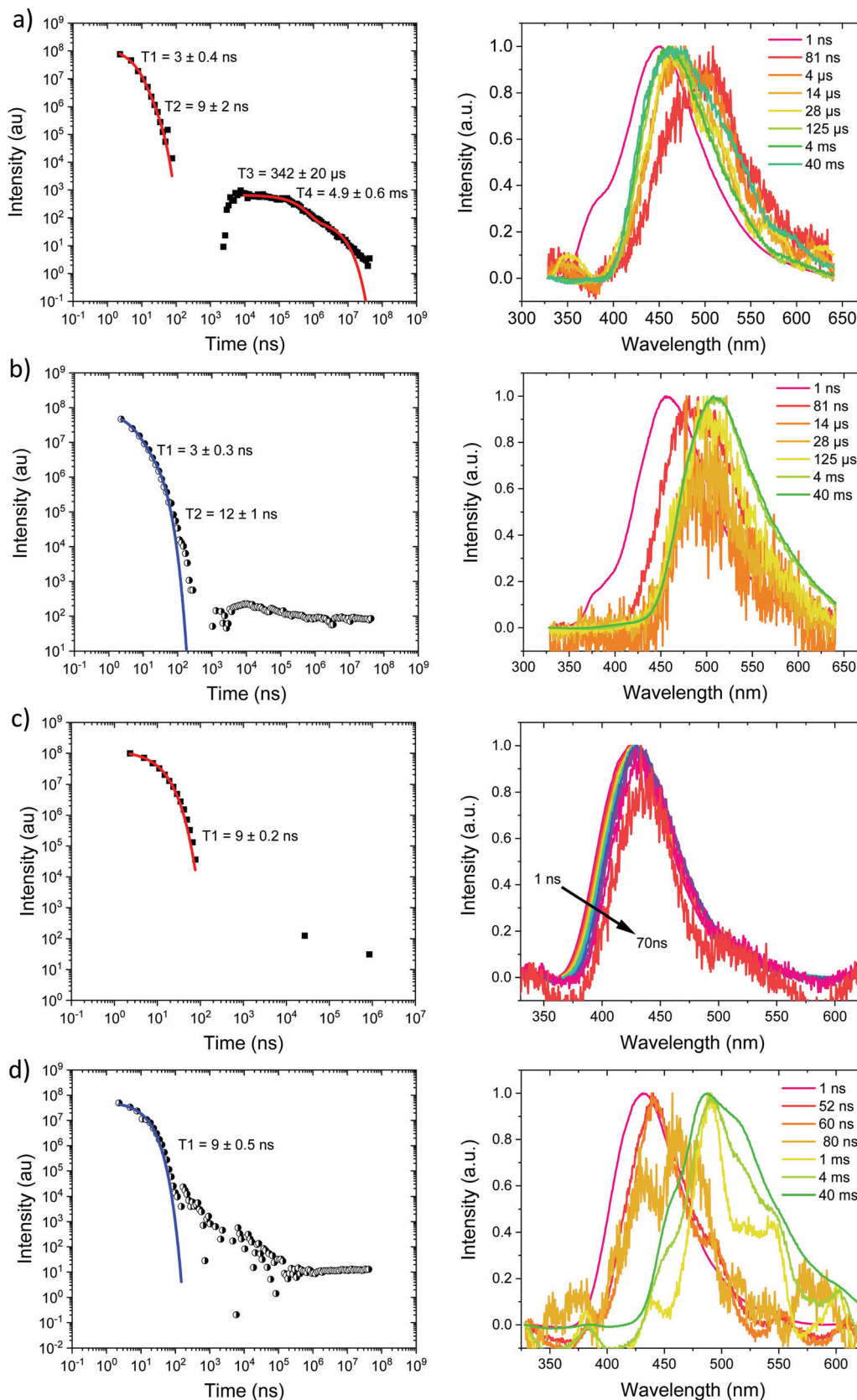


Fig. 5 Time-resolved spectrally integrated intensity decay (left) and representative time resolved spectra (right) of (a)  $\text{Ph}_2\text{TRZCzPhCz}$ , (a) 300 K and (b) 80 K;  $\text{Ph}_3\text{TRZCzPhCz}$ , (c) 300 K and, (d) 80 K in zeonex matrix. Spectral times given in ns, and decay fitting data included in the decay plots.





Fig. 6 Time-resolved spectrally integrated intensity decay (left) and representative time resolved spectra (right) of  $\text{Ph}_3\text{TRZCzTPA}$  (a) 300 K and (b) 80 K;  $\text{Ph}_2\text{TRZCzTPA}$  (c) 300 K and (d) 80 K in DPEPO host. Spectral times given in ns, and decay fitting data included in the decay plots.



millisecond times, we observe an emission with a new band shape and onset energy of 2.92 eV, which we assume to be phosphorescence. At 80 K the prompt spectra show minor shifts in energy of *ca.* 10 meV only. However, a much larger shift is seen in the phosphorescence which red shifts to an onset of 2.76 eV at 80 K. This is a large change in triplet energy, and looking critically at the band shape it is different. Careful inspection of the 40 ms emission at room temperature (compared to that at 4 ms) shows a shoulder has grown in on the red side of the band which we believe is a contribution from the low energy triplet species observed at 80 K. We have observed such dual phosphorescence before in D–A–D systems and this implies two coupled local triplet states in thermal equilibrium co-existing on the molecule.<sup>22</sup> As the lowest energy triplet state is at 2.76 eV, this yields a true  $\Delta E_{ST}$  of 340 meV and so it is not surprising that we observe little or no delayed emission.

**Ph<sub>3</sub>TRZCzPhCz** at RT and 80 K, Fig. 5c and d, behaves like a simple fluorescence emitter, having only a single emission band, onset 3.35 eV with mono exponential lifetime of 9 ns. Over its lifetime, the blue edge of this band decays faster than the red edge, a typical red edge effect. No DF is observed from this band at either temperature. At 80 K phosphorescence is observed in the 1–10 ms range, having an extremely well resolved spectrum, onset at 2.88 eV which is a clear match to a carbazole phosphorescence spectrum.<sup>23</sup> Further, by 40 ms the band shape has evolved, losing much of its structure, onset at 2.91 eV. The spectral band shape can be deconvolved into a linear combination of the carbazole phosphorescence and the phosphorescence band we observe in **Ph<sub>2</sub>TRZCzPhCz**, Fig. S9 (ESI<sup>†</sup>), indicating dual phosphorescence in **Ph<sub>3</sub>TRZCzPhCz** as well. This yields a  $\Delta E_{ST}$  of *ca.* 470 meV.

Measurements were also made at 10% by weight loading in DPEPO host, which has higher dielectric coefficient and polarizability than zeonex, also the DPEPO matrix greatly restricts the movement of the molecules because of tighter packing than the zeonex polymer matrix. Concomitantly, all CT emission is seen to be red shifted. In the case of **Ph<sub>3</sub>TRZCzTPA** the prompt emission at RT has onset at 3.0 eV, Fig. 6a. By 80 ns the emission has relaxed to 2.85 eV and from 500 ns until 4  $\mu$ s we see DF at 2.89 eV, Fig. S10 (ESI<sup>†</sup>). After 12  $\mu$ s until 4 ms weak emission is observed with onset at 2.85 eV. At 80 K, Fig. 6b, bi-exponential prompt emission, with lifetimes of 7 ns and 169 ns is observed, onset at 3.0 eV with a very weak blue knee ascribed to rapid D emission along with the longer lived prompt CT emission. The prompt and delayed CT emission behave similarly to room temperature but the DF intensity is lower, again showing thermal activation, Fig. S11 (ESI<sup>†</sup>). From 120  $\mu$ s to 40 ms emission is seen with slight structure and onset at 2.75 eV consistent with phosphorescence from the lowest triplet state. **Ph<sub>2</sub>TRZCzTPA** at RT shows dual PF, onset at 3.26 eV with lifetimes of 8.3 ns and 48 ns, Fig. 6c. From 80 ns until 4  $\mu$ s DF is observed at 2.64 eV, Fig. S10 (ESI<sup>†</sup>). From 12  $\mu$ s until 4 ms very weak emission onset *ca.* 2.79 eV is seen, ascribed to phosphorescence. At 80 K, Fig. 6d, the PF is again bi-exponential and the same delayed emission behaviour is

observed, Fig. S11 (ESI<sup>†</sup>). At 40 ms we again see phosphorescence at about 2.73 eV but the signal is very weak here. In both **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** the <sup>3</sup>LE energy from low temperature zeonex measurements at 2.76 and 2.80 eV respectively, is very consistent with the phosphorescence seen in DPEPO. This would then set  $\Delta E_{ST}$  at around 150–200 meV in DPEPO, consistent with the observed thermally activated delayed emission.<sup>24</sup> The calculated rISC rates of **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** in DPEPO films were calculated using a kinetic model. At RT, we estimated rates of  $6.3 \times 10^4 \text{ s}^{-1}$  and  $1 \times 10^5 \text{ s}^{-1}$  respectively, and at 80 K,  $1.4 \times 10^4 \text{ s}^{-1}$  and  $1 \times 10^4 \text{ s}^{-1}$  respectively, Fig. S12–S15 (ESI<sup>†</sup>).

For **Ph<sub>2</sub>TRZCzPhCz** in DPEPO at RT, Fig. 7a, emission is observed at 3.1 eV (onset) having a bi-exponential decay of 5.8 ns and 32 ns. From 500 ns to 35  $\mu$ s very weak emission is seen with an onset of 2.88 eV, Fig. S10 (ESI<sup>†</sup>), and at 4 ms there may be phosphorescence with onset at 2.92 eV but the signal is very weak. At 80 K, Fig. 7b, again we observe prompt emission at 3.1 eV and then very weak emission from 500 ns to 35  $\mu$ s at 2.88 eV, Fig. S11 (ESI<sup>†</sup>). At 40 ms emission is still seen at 2.83 eV which we ascribe to phosphorescence. The 3.1 eV CT band is red shifted by over 300 meV compared with that in zeonex and so we believe this shows that in DPEPO the through-space CT state is stabilised by the packing effect of the DPEPO forcing the D and A units closer together. In order to confirm this hypothesis, we measured **Ph<sub>2</sub>TRZCzPhCz** decays in both mCP and CBP hosts as well. In both cases, clear 3.1 eV CT emission is observed from **Ph<sub>2</sub>TRZCzPhCz** with long lifetime confirming DF, Fig. S5 (ESI<sup>†</sup>). These observations confirm that host packing plays a major role in the underlying physics of the CT states in **Ph<sub>2</sub>TRZCzPhCz**. All delayed emission in **Ph<sub>2</sub>TRZCzPhCz** is rather weak, and if we assume the same triplet energy as found in zeonex then  $\Delta E_{ST}$  will be of order 250 meV implying inefficient rISC in line with the very long lived weak DF. The CT band is also broaden which we assume implies a high degree of disorder within the host DPEPO, again consistent with an inhomogeneous, packing induced through-space CT state. **Ph<sub>3</sub>TRZCzPhCz** also shows similar behaviour to **Ph<sub>2</sub>TRZCzPhCz**, Fig. 7c and d. At RT we observe bi-exponential decay with lifetimes of 6 ns and 20 ns, emission onset at 3.1 eV. From 100 ns to 4  $\mu$ s this emission shifts to 2.91 eV, Fig. S10 (ESI<sup>†</sup>). Again the CT band is red shifted compared to zeonex pointing to DPEPO stabilising the through-space CT excited state and giving weak DF. At 80 K, prompt emission starts at 3.26 eV and shifts to 3.1 eV by 80 ns. The decay is bi-exponential, 6.3 ns and 31 ns, in line with a D emission as well as prompt through-space CT emission. There is little DF at 80 K, Fig. S11 (ESI<sup>†</sup>), in line with inefficient TADF, but at 40 ms clear phosphorescence emission with some structure is seen, onset at 2.95 eV in line with that observed in zeonex. All lifetimes are summarised in Table 4.

### Device results

**Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** were selected as the emitters to investigate electroluminescent (EL) characteristics. Four commonly used high-triplet energy hosts including;





Fig. 7 Time-resolved spectrally integrated intensity decay (left) and representative time resolved spectra (right) of  $\text{Ph}_2\text{TRZCzPhCz}$ , (a) 300 K and (b) 80 K;  $\text{Ph}_3\text{TRZCzPhCz}$ , (c) 300 K and (d) 80 K in DPEPO host. Spectral times given in ns, and decay fitting data included in the decay plots.



Table 4 Time resolved data of DPEPO film measurement at RT

Molecule	$^1\text{LE}^a$ (eV)	$\tau_{\text{LE}}$ (ns)	$^1\text{CT}^a$ (eV)	$\tau_{\text{PF1}}^b$ (ns)	$\tau_{\text{PF2}}^b$ (ns)	$\tau_{\text{DF1}}^b$ ( $\mu\text{s}$ )	$\tau_{\text{DF2}}^b$ ( $\mu\text{s}$ )	$^3\text{LE}^c$ (eV)	$\Delta E_{\text{ST}}^d$ (meV)
<b>Ph<sub>3</sub>TRZCzTPA</b>	3.00	~1	2.85	31	161	4	28	2.76	90
<b>Ph<sub>2</sub>TRZCzTPA</b>	2.95	~1	2.67	8.3	48	2.7	21	2.80	-130
<b>Ph<sub>2</sub>TRZCzPhCz</b>	3.10	~1	2.88	5.8	32	2	59	2.76	~250
<b>Ph<sub>3</sub>TRZCzPhCz</b>	3.10	~1	2.91	6	20	4	51	2.95	-40

<sup>a</sup> Local (LE) and charge transfer (CT) emission energies calculated from the on-set of the emission band. <sup>b</sup> Prompt ( $\tau_{\text{PF}}$ ) and delayed fluorescence ( $\tau_{\text{DF}}$ ) component lifetimes calculated from fits to the measured intensity decay curves. <sup>c</sup> Local triplet energy taken from the on-set of the phosphorescence measured at 80 K. <sup>d</sup>  $\Delta E_{\text{ST}}$  is the energy difference between the  $^1\text{CT}$  and  $^3\text{LE}$  energies.



Fig. 8 (a) Structural drawings of the materials used in OLEDs; (b) schematic structures of the fabricated OLEDs with different emitters.

4,4'-bis(*N*-carbazolyl)-1,1'-biphenyl (CBP) ( $E_{\text{T}} = 2.56$  eV), 2,6-bis(3-(9*H*-carbazol-9-yl)phenyl)pyridine (26DCzPPy) ( $E_{\text{T}} = 2.71$  eV), 1,3-bis(carbazol-9-yl)benzene (mCP) ( $E_{\text{T}} = 2.9$  eV), and 9-(4-*tert*-butylphenyl)-3,6-bis(triphenylsilyl)-9*H*-carbazole (CzSi) ( $E_{\text{T}} = 3.0$  eV) were tried. To match the HOMO and the LUMO levels of the emitting layer, di[4-(*N,N*-ditolyl-amino)-phenyl]cyclohexane (TAPC) and 1,3,5-tri[(3-pyridyl)-phen-3-yl]benzene (TmPyPB) were selected as the hole and electron transport layers respectively, see Fig. 8. Devices using mCP as the host gave the best performance in line with the photophysics, including relatively low operation voltages, higher maximum luminance, and improved efficiency. For doping concentration (2–20 wt%) of **Ph<sub>2</sub>TRZCzTPA** in the mCP further fine-tuning gave optimal performance, Fig. S16 (ESI<sup>†</sup>) depicts the EL characteristics and parameters are summarized in Table S4 (ESI<sup>†</sup>). From the EL spectra, no emission from mCP and/or carrier transport materials were observed when the doping concentration was greater than or equal to 2 wt%, implying effective charge transfer and exciton confinement on **Ph<sub>2</sub>TRZCzTPA**. The EL spectra of the devices were red-shifted from 505 nm to 529 nm as the doping concentration increased from 2 wt% to 20 wt%. The current density–voltage ( $J$ - $V$ ) curves shown in Fig. S16(b) (ESI<sup>†</sup>) indicate that current density increases with doping concentration whilst the turn-on voltages (at 1  $\text{cd m}^{-2}$ ) decrease from 3.3 V to 2.6 V, in line with **Ph<sub>2</sub>TRZCzTPA** being a bipolar charge transporting material. Fig. S16(d) (ESI<sup>†</sup>) depicts the external quantum efficiency (EQE) as a function of **Ph<sub>2</sub>TRZCzTPA** concentration, with

peak EQE of 16.3%, 49.7  $\text{cd A}^{-1}$ , and 59.3  $\text{lm W}^{-1}$  found at 12 wt% **Ph<sub>2</sub>TRZCzTPA**.

26DCzPPy was found to be the best host for **Ph<sub>3</sub>TRZCzTPA**. In addition, MoO<sub>3</sub>-doped TAPC was adopted as a hole injection layer to decrease the energy barrier between the ITO anode and the organic layer, Fig. 8. Fig. S17 (ESI<sup>†</sup>) depicts the EL characteristics while the corresponding parameters are summarized in Table S5 (ESI<sup>†</sup>). A weak emission from 26DCzPPy was observed in the EL spectra at doping concentrations of **Ph<sub>3</sub>TRZCzTPA** below 2 wt%, indicating the incomplete host-guest energy transfer. As the doping concentration increased from 2 wt% to 24 wt%, the devices showed the EL spectra red-shifted from 518 nm to 530 nm. The current density of the devices increases with the doping concentration, indicating the favourable carrier transport capability of **Ph<sub>3</sub>TRZCzTPA**. Given the inferior carrier mobility of 26DCzPPy; 2 wt% **Ph<sub>3</sub>TRZCzTPA** the turn-on voltage was 5.0 V but continuously decreased to 4.4 V with a higher doping concentration, to 24 wt%. The optimized concentration was found at 20 wt% **Ph<sub>3</sub>TRZCzTPA**, giving maximum EQE of 13.3%, 40.3  $\text{cd A}^{-1}$ , and 28.6  $\text{lm W}^{-1}$ . The corresponding EL characteristics of the optimized devices with **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** (*i.e.* devices A and B) are shown in Fig. 9 and the data are summarized in Table 5. Overall, the results demonstrate that both **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** give efficient TADF devices from through-space CT excited states.





Fig. 9 (a) Normalized EL spectra at a luminance of  $10^3$   $\text{cd m}^{-2}$ ; (b) current density–luminance–voltage ( $J$ – $V$ – $L$ ) characteristics; (c) external quantum efficiency versus luminance, (d) luminance/power efficiency versus luminance for devices A and B.

Table 5 EL characteristics of the devices with  $\text{Ph}_3\text{TRZCzTPA}$  and  $\text{Ph}_2\text{TRZCzTPA}$  emitters

Device	Host	Emitter	$V_{\text{on}}^a$ [V]	$L_{\text{max}}$ [ $\text{cd m}^{-2}$ ]	EQE [%]	CE [ $\text{cd A}^{-1}$ ]	PE [ $\text{lm W}^{-1}$ ]	$\lambda_{\text{Peak}}$ [nm]	CIE <sup>d</sup> [x, y]
A	CBP	$\text{Ph}_2\text{TRZCzTPA}$ (12 wt%)	2.9	27 158 (12.4 V)	16.3, <sup>b</sup> 14.7, <sup>c</sup> 10.9 <sup>d</sup>	49.7, <sup>b</sup> 44.9, <sup>c</sup> 33.4 <sup>d</sup>	59.3, <sup>b</sup> 28.2, <sup>c</sup> 15.0 <sup>d</sup>	522	0.32, 0.55
B	26DCzPPy	$\text{Ph}_3\text{TRZCzTPA}$ (20 wt%)	4.4	26 434 (11.0 V)	13.3, <sup>b</sup> 12.4, <sup>c</sup> 9.7 <sup>d</sup>	40.3, <sup>b</sup> 37.5, <sup>c</sup> 29.4 <sup>d</sup>	28.6, <sup>b</sup> 23.3, <sup>c</sup> 15.8 <sup>d</sup>	529	0.34, 0.54

<sup>a</sup> Turn-on voltage measured at 1  $\text{cd m}^{-2}$ . <sup>b</sup> Maximum efficiency. <sup>c</sup> Measured at  $10^2$   $\text{cd m}^{-2}$ . <sup>d</sup> Measured at  $10^3$   $\text{cd m}^{-2}$ .

## Discussion

We find  $\text{Ph}_2\text{TRZCzPhCz}$  and  $\text{Ph}_3\text{TRZCzPhCz}$  to have different photophysics compared to  $\text{Ph}_3\text{TRZCzTPA}$  and  $\text{Ph}_2\text{TRZCzTPA}$ , primarily because of their relative weak donor strength and smaller driving force for CT. A through-bond CT state is observed to be favoured (by comparison to model systems) and in  $\text{Ph}_2\text{TRZCzPhCz}$  stabilization of any CT state is hard to achieve so that D emission is always observed. In zeonex, the lowest energy triplet state is always found with onset at 2.76 eV, ascribed to the local triplet state of the pendent Cz donor, giving large  $^1\text{CT}^-\text{LE}$  energy gaps,  $>0.35$  eV, thus we observe little or no TADF from the through-bond CT states. In  $\text{Ph}_3\text{TRZCzPhCz}$  highly resolved Cz donor phosphorescence is observed indicative of a large local triplet population, remaining uncoupled from the rest of the molecule. In DPEPO films we observe a physical effect on the photophysics arising from the small molecule host packing, whereas in zeonex, which is an open polymer network having high free volume, the guest molecules behave much like in MCH solution, *i.e.* they are not constrained by the host. In DPEPO,  $\text{Ph}_2\text{TRZCzPhCz}$  and

$\text{Ph}_3\text{TRZCzPhCz}$  show no D emission but instead a highly red shifted, stable CT band. We propose that the packing forces of the DPEPO host distorts the guest molecules, forcing the D and A closer together, stabilizing a through-space CT excited state. This was confirmed measuring  $\text{Ph}_2\text{TRZCzPhCz}$  in both CBP and mCP hosts which also gave the same strong CT stabilisation as DPEPO does with long lived CT delayed emission. This DF, although weak, is seen to be temperature dependent in both compounds, *i.e.* TADF. In the case of the through-space CT state, with D and A still have a larger co-facial separation than an A–D' through-bond CT state (see X-ray data) and so the Coulomb attraction energy between separated electron and hole is reduced and thus the CT energy is red shifted. The  $^1\text{CT}$  through-space state being red shifted compared to the through-bond state, reduces the  $^1\text{CT}^-\text{LE}$  energy gap to *ca.* 120 meV and we thus observe DF in both cases.

In all cases where we observe a measurable DF signal, we observe a strongly non-exponential decay at long times. We believe this comes from the inhomogeneous environment of the doped films, with the hosts imposing various degrees of restriction and packing effects on the TADF molecules.



The through-space CT state systems seem to be particularly prevalent to this which is not surprising, showing a potentially high sensitivity to exact spatial separation and orientation of the D and A in a host environment. Recently, Suresh *et al.*<sup>25</sup> have discussed such a long time non-exponential DF tail in terms of combined TADF and TTA emission, however here we have only used 1% loading of the TADF molecules in zeonex and 10% in DPEPO which probably rules out TTA. However in devices were much higher loadings are used a TTA contribution along with TADF cannot be ruled out.

**Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA**, by comparison to the model TRZ-TPA excited state CT complex (exciplex), are seen to have stable through-space CT excited states. The stabilising of the through space CT state over a A-D' through bond CT state is ascribed to the larger CT driving force arising from their strong D, strong A, structure. From the time resolved decays and spectra, we do though see that in **Ph<sub>3</sub>TRZCzTPA**, an initial higher energy transient CT state which we tentatively ascribe to a short lived through-bond CT state (by comparison to model systems), potentially stabilised by the effect of the extra phenyl ring between acceptor and bridge. Relaxation of this phenyl ring configuration could then stabilise the through space CT state. Alternatively the through bond state may occur on a small population of **Ph<sub>3</sub>TRZCzTPA** molecules trapped in a non relaxed geometry in the film. This behaviour is observed in both zeonex and DPEPO which we believe favours the former explanation. Certainly we see no such behaviour in **Ph<sub>2</sub>TRZCzTPA** which lacks the spacing phenyl unit. **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA** show intense temperature dependent delayed emission, clearly TADF. The measurable differences between rISC rates of the two molecules we believe arises from differences in relative co-facial separation and orientation of D and A due to the phenyl spacer between A and D' (bridge) in **Ph<sub>3</sub>TRZCzTPA**, as seen in the X-ray data. This controls their magnetic coupling, *i.e.* spin orbit coupling matrix elements, that dictate rISC rate, given they have very similar, small singlet triplet energy gaps. By comparing the photophysics of these two systems, we see that strong acceptor, strong donor pairs favour through-space charge transfer whereas strong acceptor weak donor favours through-bond charge transfer, in line with both the driving force for charge transfer and the spatial separation and orientation of the non-conjugated D A pair. This clearly sets some basic design criteria for TADF intramolecular through-space charge transfer systems. The differences between the two molecules are also reflected in the device performance. We find that **Ph<sub>2</sub>TRZCzTPA** gives better device performance than **Ph<sub>3</sub>TRZCzTPA**, very much in line with the difference in rISC rates and the lack of possible residual through-bond CT states in **Ph<sub>3</sub>TRZCzTPA**. From these device results we again see sensitivity to D A orientation, flexibility of linkage to the bridge and also host packing effects. Much like exciplex emitters, these effects need to be carefully managed and controlled.<sup>26</sup>

In **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz**, the lowest local triplet state of the molecular resides on the pendant Cz unit which quenches higher energy D' (bridge) triplets leading to

negligible mediation of rISC for the through-bond CT state. All of our observations clearly align very well with ideas recently put forward by Wada *et al.* from their through-space charge transfer TADF molecule.<sup>20</sup> We also see that the effect of host packing has profound effects on the through-space CT states and these effects cannot be ignored, especially in devices.

## Conclusions

From this new series of D-D'-A materials we have gained a great deal of new understanding about through-space CT states, competition with through-bond CT formation and the role a host plays in controlling this equilibrium. In the two materials having strong D and strong A, **Ph<sub>3</sub>TRZCzTPA** and **Ph<sub>2</sub>TRZCzTPA**, we unambiguously identify stable through-space CT states *via* comparison to model compounds and solution state D A complexes. This stabilisation we ascribe to the large driving force for charge transfer (*ca.* 1 eV) and subsequent larger charge separation distance. For the weak D strong A pair, **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz**, a through-bond CT state between the A and the bridge D' is stabilised as the CT driving force is much less, *ca.* 0.5 eV. Little DF is observed unless an external perturbation further stabilises the CT state, *i.e.* solvent polarity or host packing forces. Indeed, in DPEPO, CBP and mCP with large packing forces we observe highly red shifted CT formation and subsequent TADF, which we ascribe to stabilization of a through-space CT state in **Ph<sub>2</sub>TRZCzPhCz** and **Ph<sub>3</sub>TRZCzPhCz**. This shows that such external forces can distort the molecule, forcing the D and A branches closer together to fully stabilise the through-space CT state over the through-bond state. Very satisfyingly, our TDDFT calculations identified the possibility of **Ph<sub>3</sub>TRZCzPhCz** having both through-space and through-bond CT excited states. In **Ph<sub>3</sub>TRZCzTPA** having an additional 'spacer' phenyl group between the bridge D' and A we observe a large temperature dependent red shift in the energies of both the <sup>1</sup>CT and <sup>3</sup>LE excited states. We ascribe this to thermal motion of this phenyl ring which reduces planarity between the A and the bridge, decoupling the phenyl ring from the dTRZ A moiety, making the **Ph<sub>3</sub>TRZCzTPA** act in a very similar fashion to **Ph<sub>2</sub>TRZCzTPA**. This decrease in conjugation also localises the <sup>3</sup>LE state on the dTRZ unit, increasing the phosphorescence energy as we observe. In view of these observations, having such a spacing phenyl unit that can rotate between the D and A units, might not be the optimum design strategy for TADF materials, as we find that **Ph<sub>2</sub>TRZCzTPA** gives better performing devices than **Ph<sub>3</sub>TRZCzTPA**. We also find that **Ph<sub>2</sub>TRZCzTPA** has faster rISC rates than **Ph<sub>3</sub>TRZCzTPA** which we believe indicates more optimal co-facial orientation (not separation) of D and A forming the through-space CT state, which is critical in controlling SOC properties and thus rISC, as proposed by Wada *et al.*<sup>27</sup> These differences in photophysics are also reflected in device performance with **Ph<sub>2</sub>TRZCzTPA** giving better performance in optimised devices. This first in-depth study of the physical properties and correlated device performance of through-space compared to



through-bond CT states and TADF, gives many new pointers and ideas to design more efficient and stable TADF emitters.

## Methods

Full details of the synthesis and physical characterisation of the materials reported here, along with photophysical measurements and data analysis are given in the ESI† file published alongside this paper.

## Author contributions

Hector Miranda-Salinas performed the majority of the optical and photoluminescence measurements and contributed to the analysis and helped in the preparation of the manuscript. Yi-Tzu Hung synthesized and characterize the physical property of titled compounds. Yi-Sheng Chen made the theoretical analysis of the titled compounds. Dian Luo and Hao-Che Kao fabricate the OLED device and collect the device data. Chih-Hao Chang (analyse the device data and helped to prepare the manuscript. Ken-Tsung Wong designed the project and molecules, and helped in the preparation of the manuscript. APM conceived and the model for the analysis of the photophysical results, lead the analysis of the photophysics and helped write the manuscript.

## Data

All data resulting from this work is given in the paper and ESI.†

## Conflicts of interest

There are no conflicts to declare.

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