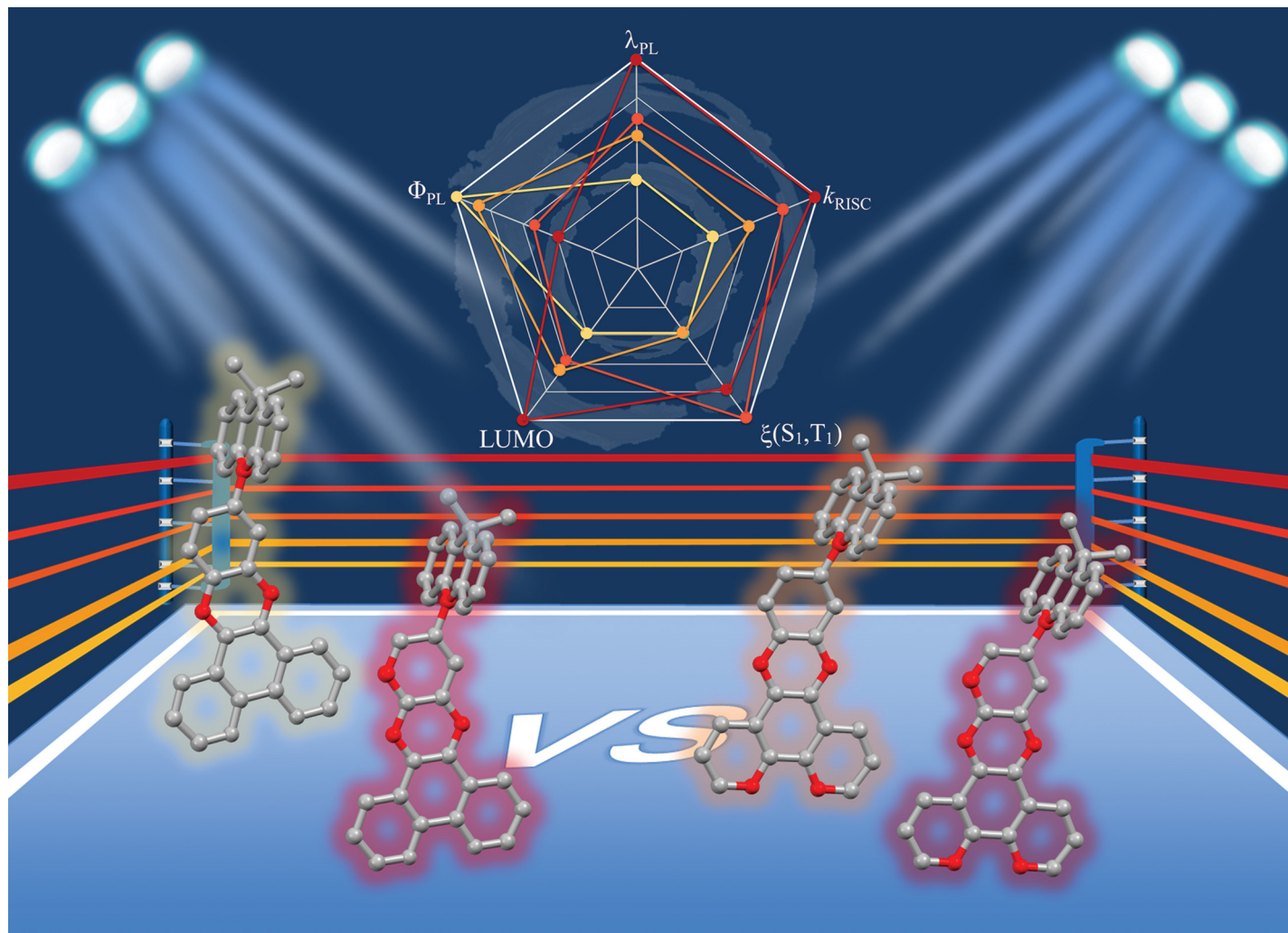


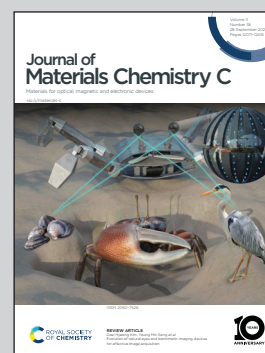


Highlighting research from the Zysman-Colman Group,  
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The influence of nitrogen doping of the acceptor in  
orange-red thermally activated delayed fluorescence  
emitters and OLEDs

A family of four orange-to-red TADF compounds  
containing different numbers of nitrogen atoms within  
the conjugated acceptor core has been systematically  
investigated. The findings indicate that increasing the  
nitrogen atom content in the acceptor of the D-A  
compounds results in a more stabilized LUMO,  
smaller  $\Delta E_{ST}$  and faster  $k_{RISC}$ .

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# The influence of nitrogen doping of the acceptor in orange-red thermally activated delayed fluorescence emitters and OLEDs†‡

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Nitrogen-containing polycyclic aromatic hydrocarbons (N-PAH) have been widely used as deep lowest unoccupied molecular orbital (LUMO) acceptors in donor–acceptor (D–A) red thermally activated delayed fluorescence (TADF) emitters and in organic light-emitting diodes (OLEDs). However, most of the studies have focused disparately on donor/acceptor combinations to yield efficient emitters, while a methodological study investigating the influence of nitrogen (N) doping ratios on the ground and excited states of PAH acceptors is rare. Here, we report a family of four different N-PAH acceptors containing different numbers of nitrogen atoms within the N-PAH and their use in D–A TADF emitters, **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN**, when coupled to the same donor, 9,9-dimethyl-9,10-dihydroacridine (DMAC). As the nitrogen content in the acceptor increases the LUMO becomes progressively more stabilized while the singlet–triplet energy gap ( $\Delta E_{ST}$ ) decreases and the rate constant for reverse intersystem crossing ( $k_{RISC}$ ) increases. In particular, introducing nitrogen at the 10-position of dibenzo[a,c]phenazine (BP) leads to a more than ten-fold enhancement in  $k_{RISC}$  in **DMACPyBP** and **DMACPyBPN** compared to **DMACBP** and **DMACBPN**. Among the OLEDs with all four emitters that with **DMACBPN** demonstrates the highest EQE<sub>max</sub> of 19.4% at an emission peak of 588 nm, while the deepest red emitting device employed **DMACPyBPN** ( $\lambda_{EL}$  = 640 nm) with an EQE<sub>max</sub> of 5.4%.

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## Introduction

Thermally activated delayed fluorescence (TADF) is a photophysical mechanism that manifests itself in dual emission, a rapid prompt fluorescence from singlet excitons that decay radiatively, and a slow delayed fluorescence that occurs as a result of the thermal up-conversion of triplet excitons into singlets *via* reverse intersystem crossing (RISC) prior to light generation.<sup>1–3</sup> The efficient harvesting of both singlet and triplet excitons to produce light emission makes TADF materials particularly

attractive as emitters for electroluminescent devices as, like phosphorescent devices, they can achieve 100% internal quantum efficiency (IQE).<sup>4,5</sup> Most of the reported TADF emitters employ a strongly twisted donor (D) and acceptor (A) structure, which leads to localization of the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) on the D and A, respectively, and thus to a small singlet–triplet energy gap ( $\Delta E_{ST}$ ). A sufficiently small  $\Delta E_{ST}$  is necessary to enable endothermic upconversion of triplet excitons to singlets by RISC.<sup>6–8</sup> The apparent paradox in TADF emitter design is that the design requirements for efficient RISC are in contrast to those to attain high photoluminescence quantum yield ( $\Phi_{PL}$ ), which require significant orbital overlap to maximize the oscillator strength of the transition.<sup>9,10</sup> Balancing these two criteria in the design of high-performance orange-red emitters becomes that much more difficult as radiative decay decreases while non-radiative decay increases due to the energy-gap law.<sup>11–13</sup> The general molecular design principles for red/deep red D–A TADF emitters are: (1) use of strong donor and acceptor motifs is a common strategy, affording a shallow HOMO and a deep LUMO for D and A, respectively; and (2) construction of a rigid and typically planar fused ring donor and/or acceptor for the suppression of nonradiative transitions. Within this model, most of the acceptors are

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derivatives of either acenaphtho[1,2-*b*]pyrazine-8,9-dicarbonitrile (APDC),<sup>14</sup> heterocyclic quinoxaline-6,7-dicarbonitrile (QCN),<sup>15</sup> 11,12-dicyanodibenzo[*a,c*]phenazine (CNBPz),<sup>16</sup> 2,3-dicyanodibenzo[*f,h*]quinoxaline (CNBQx),<sup>17,18</sup> or dibenzo[*a,c*]phenazine-3,6-dicarbonitrile (PZCN).<sup>19</sup> Nitrogen-doped polycyclic aromatic hydrocarbons (N-PAH), like dibenzo[*a,c*]phenazine (BP), have proven to be particularly popular as moieties in orange-red TADF emitters (Fig. S1, ESI†).<sup>20–26</sup> Zhao *et al.* first reported BP-based TADF compounds, containing one-to-three 9,9-dimethyl-9,10-dihydroacridine (DMAC) donor moieties (Fig. S1, ESI†). The orange-red OLEDs with **3DMAC-BP**, an emitter with three DMAC donors situated at the 3-, 6-, and 11-positions of the BP, showed a maximum external quantum efficiency (EQE<sub>max</sub>) of 22.0% with an electroluminescence peaking at  $\lambda_{\text{EL}}$  = 606 nm.<sup>23</sup> Yang *et al.* reported a related compound using the BP acceptor, **DMAC-11-DPPZ** with the DMAC donor attached at the 11-position of the BP acceptor, which emits at  $\lambda_{\text{PL}}$  of 576 nm, has a photoluminescence quantum yield,  $\Phi_{\text{PL}}$ , of 57% and a delayed lifetime,  $\tau_{\text{d}}$ , of 1.53  $\mu\text{s}$  in 10 wt% doped films in CBP. The optimized OLED with **DMAC-11-DPPZ** displays an orange emission at 588 nm with an EQE<sub>max</sub> of 23.8%.<sup>24</sup> Tang *et al.* introduced one more nitrogen at the 10-position of the BP core to generate a new acceptor unit, BPQ. Star-shaped isomers **3,6,11-triAC-BPQ** and **3,6,12-triAC-BPQ** contain three DMAC donors at either the 3,6,11-positions or 3,6,12-positions of BPQ. **3,6,12-triAC-BPQ** showed a small red-shifted emission at 611 nm in toluene and a slightly shorter  $\tau_{\text{d}}$  of 2.25  $\mu\text{s}$  in 15 wt% doped films in mCBP compared to **3DMAC-BP** ( $\lambda_{\text{PL}}$  = 605 nm in toluene,  $\tau_{\text{d}}$  = 2.9  $\mu\text{s}$  in 20 wt% doped films in mCBP), which is a compound that does not have an N-atom at the 10-position of the acceptor. In contrast, **3,6,11-triAC-BPQ** exhibited a blue-shifted emission at  $\lambda_{\text{PL}}$  of 565 nm compared to **3,6,12-triAC-BPQ**. The devices with **3,6,11-triAC-BPQ** and **3,6,12-triAC-BPQ** showed an EQE<sub>max</sub> of 22.0% [ $\lambda_{\text{EL}}$  = 581 nm, Commission International de l'Éclairage, CIE, coordinates of (0.51,0.48)] and 16.5% [ $\lambda_{\text{EL}}$  = 616 nm, CIE coordinates of (0.58,0.39)], respectively.<sup>25</sup> The same group reported an orange-yellow TADF material **DPPZ-DMAC**, where the related DPPZ acceptor contains two additional nitrogen atoms compared to BP. This compound emits at  $\sim$ 590 nm, has a smaller  $\Delta E_{\text{ST}}$  of 0.01 eV, and a higher  $\Phi_{\text{PL}}$  of 91.6% in 6 wt% doped films in CBP, compared to that of **DMAC-11-DPPZ**, a compound without the two additional nitrogen atoms in the acceptor. The **DPPZ-DMAC**-based device showed an EQE<sub>max</sub> of 27.8% with  $\lambda_{\text{EL}}$  at 598 nm. Zhang *et al.* reported a similar emitter **SAF-2NP** that employs a structurally related 10H-spiro(acridine-9,9'-fluorene) (SAF) donor instead of DMAC in combination with the same acceptor as **DPPZ-DMAC**. As expected, **SAF-2NP** shows similar photophysics as **DPPZ-DMAC** but has a slightly blue-shifted emission at  $\lambda_{\text{PL}}$  of  $\sim$ 580 nm and a much higher  $\Phi_{\text{PL}}$  of 99%. The **SAF-2NP**-based OLED demonstrated a very high EQE<sub>max</sub> of 32.5% with an  $\lambda_{\text{EL}}$  at 576 nm, an efficiency linked to the high  $\Phi_{\text{PL}}$  of 99% and horizontally oriented transition dipole ( $\theta_{\parallel}$ ) of 85% of the emitter in the host matrix. When DPPZ is coupled with a triphenylamine (TPA) donor, as in **oTPA-DPPZ**, the emission shifts to the red at 605 nm in the 30 wt% doped films ( $\Phi_{\text{PL}}$  = 75%;  $\Delta E_{\text{ST}}$  of 0.07 eV).<sup>26</sup> The OLEDs

with **oTPA-DPPZ** showed an EQE<sub>max</sub> of 18.5% at  $\lambda_{\text{EL}}$  of 600 nm. By adjusting the position of the TPA groups, a “T-shape” deep-red TADF emitter, **pTPA-DPPZ**, was obtained.<sup>26</sup> Compared to **oTPA-DPPZ**, the rational spatial arrangement of D and A groups in **pTPA-DPPZ** accelerated radiative decay from the singlet state by 90-fold, giving rise to a  $\Phi_{\text{PL}}$  of 87% in the neat film. The corresponding OLEDs, having a simplified bilayer non-doped structure, showed an EQE<sub>max</sub> of 12.3% at  $\lambda_{\text{EL}}$  of 652 nm.

Examples of D–A TADF emitters have illustrated that the use of N-PAHs like BP or DPPZ accesses orange-to-red emission. Although numerous molecules have been reported, to the best of our knowledge, there is no one study that has been conducted to correlate the number of nitrogen atoms in the N-PAH acceptor (Fig. 1(a)) to the photophysical properties of the emitter. To address this question, we designed and synthesized four DMAC-containing TADF emitters containing different numbers of nitrogen atoms in the N-PAH acceptor: 11-(9,9-dimethylacridin-10(9H)-yl)dibenzo[*a,c*]phenazine (**DMACBP**) (2 nitrogen atoms), 12-(9,9-dimethylacridin-10(9H)-yl)dibenzo[*f,h*]pyrido[2,3-*b*]quinoxaline (**DMACPyBP**) (3 nitrogen atoms), 11-(9,9-dimethylacridin-10(9H)-yl)dipyrido[3,2-*a*:2',3'-*c*]phenazine (**DMACBPN**) (4 nitrogen atoms) and 12-(9,9-dimethylacridin-10(9H)-yl)pyrido[2',3':5,6]pyrazino[2,3-*f*][1,10]phenanthroline (**DMACPyBPN**) (5 nitrogen atoms). The variations in key photophysical properties of these four emitters are summarized in Fig. 1(b). It becomes clear that the rate of reverse intersystem crossing,  $k_{\text{RISC}}$ , can be enhanced when the BP acceptor is modified to contain an additional nitrogen atom at the 10-position. The OLEDs with **DMACBPN** demonstrated an EQE<sub>max</sub> of 19.4% at  $\lambda_{\text{EL}}$  of 588 nm. The EL spectra shift to red when there is an increasing nitrogen content in the acceptor, with the reddest device using **DMACPyBPN** emitting at 640 nm.

## Results and discussion

### Synthesis and characterization

The synthesis of **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** is outlined in Scheme 1. The intermediates **DMACPhSN** and **DMACPySN** were obtained by a nucleophilic substitution reaction between 5-fluorobenzo[*c*][1,2,5]thiadiazole and 9,9-dimethyl-9,10-dihydroacridine (DMAC), and Buchwald–Hartwig coupling between 6-bromo-[1,2,5]thiadiazolo[3,4-*b*]pyridine and DMAC, respectively, followed by ring-opening of the benzothiadiazole (BTD) with lithium aluminum hydride (LiAlH<sub>4</sub>). Intermediate **1** was then reacted with phenanthrene-9,10-dione and 1,10-phenanthroline-5,6-dione in 1-butanol to afford **DMACBP** and **DMACBPN**, respectively, in good yield. **DMACPyBP** and **DMACPyBPN** were obtained from the reaction of **2** with phenanthrene-9,10-dione and 1,10-phenanthroline-5,6-dione in 1-butanol, respectively. The structural identity and purity of the four emitters were verified by <sup>1</sup>H and <sup>13</sup>C nuclear magnetic resonance (NMR) spectroscopy, melting point determination, high-resolution mass spectrometry, elemental analysis, and high-performance liquid chromatography (HPLC) (Fig. S2–S24, ESI†).





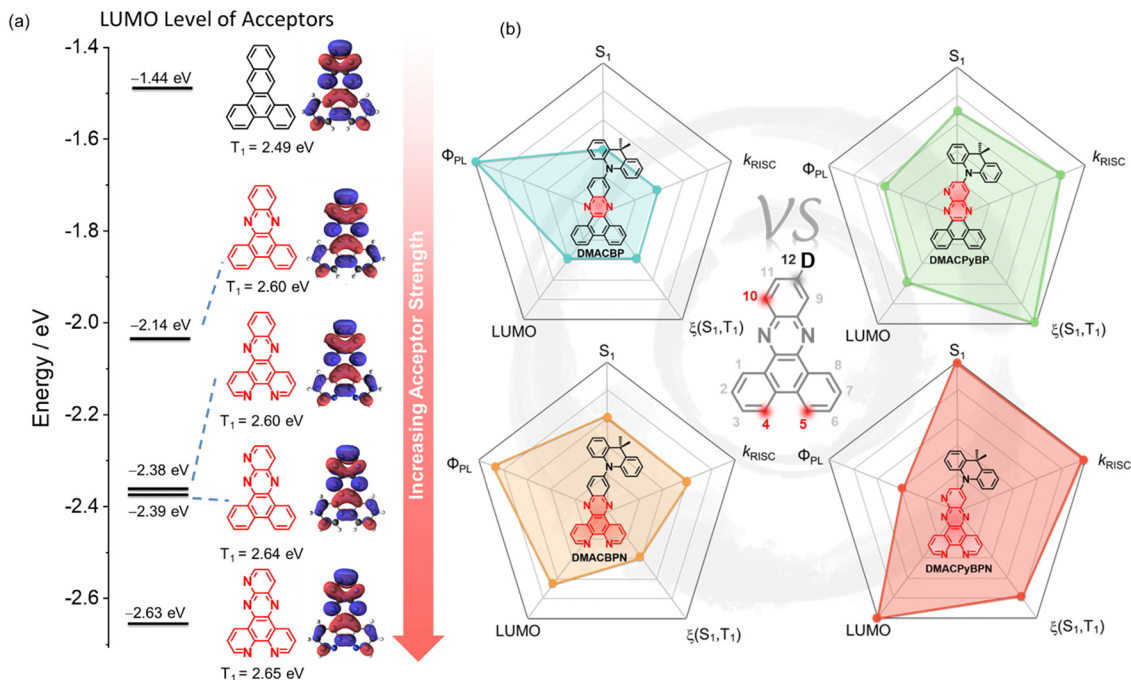
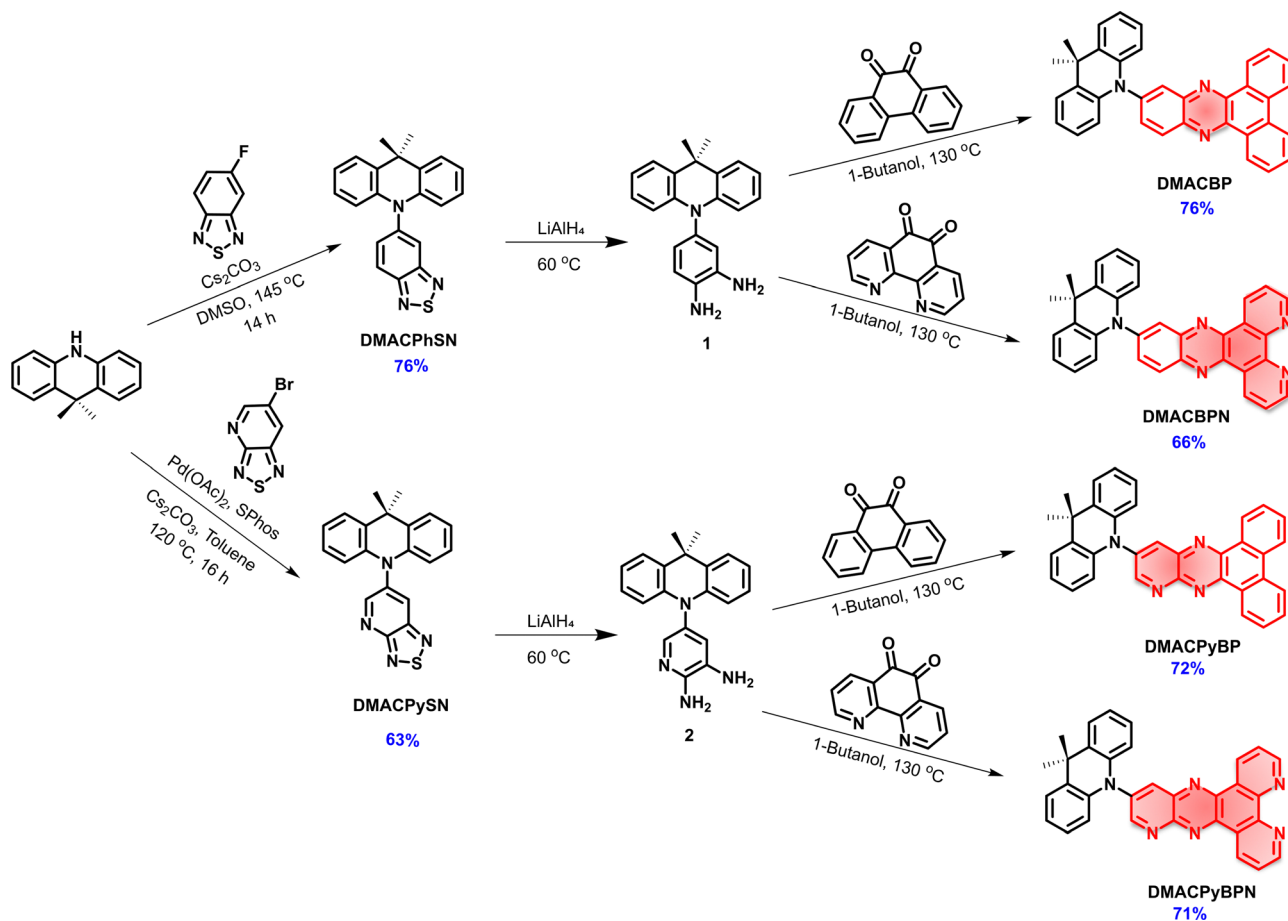


Fig. 1 (a) Electron-accepting capability (the LUMO level) of nitrogen-doped PAH acceptor units, estimated at the PBE0/6-31G(d,p) level. (b) The main parameters ( $\Phi_{PL}$ ,  $S_1$ , LUMO,  $k_{RISC}$ , and  $\xi(S_1, T_1)$ ) values vs. four emitters with different N/C ratio acceptors in this work.



Scheme 1 Synthetic routes for DMACBP, DMACPyBP, DMACBPN and DMACPyBPN emitters.



To predict their optoelectronic properties, we first modelled the ground-state structure of the four emitters in the gas phase using density functional theory (DFT) at the PBE0/6-31G(d,p) level of theory (Fig. 2). The calculated energy levels of the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO) are shown in Fig. 2(a) and Fig. S25 (ESI<sup>†</sup>). As expected, the HOMO is localized on the DMAC donor while the LUMO is localized on the nitrogen-rich electron acceptors. The gradually stabilized LUMO from **DMACBP** to **DMACPyBP**, **DMACBPN** and **DMACPyBPN** originates from the enhanced electron-withdrawing strength of the acceptor with increasing nitrogen content. However, due to the enhanced electronic coupling between the donor and the acceptor, the HOMO levels of **DMACPyBP** and **DMACPyBPN** are more stabilized as these are delocalized over both DMAC and the pyridine (Py) rings, as compared to the DMAC-localized orbitals in **DMACBP** and **DMACBPN** (Fig. 2(a)). The HOMO–LUMO gap,  $\Delta E_{\text{HOMO-LUMO}}$ , decreases from 2.84 eV for **DMACBP**, to 2.75 eV for **DMACPyBP**, 2.72 eV for **DMACBPN** and 2.63 eV for **DMACPyBPN**, following a trend similar to that observed for the LUMO and governed by the strength of the electron-acceptor moiety.

The excited-state properties were then calculated using the time-dependent DFT (TD-DFT) within the Tamm–Dancoff approximation (TDA) based on the optimized ground-state geometry.<sup>27</sup> The  $S_1$  energies are 2.17 eV for **DMACBP**, 2.04 eV for **DMACPyBP**, 2.05 eV for **DMACBPN** and 1.91 eV for **DMACPyBPN**, while the  $T_1$  energies likewise decrease from 2.15 eV, to 2.00 eV, 2.03 eV and 1.89 eV, respectively, following a trend of increasingly stabilized excited states with increasing N atom content in the acceptor, although **DMACPyBP** and **DMACBPN** are close in energy. The corresponding  $\Delta E_{\text{ST}}$  values are all relatively small at 0.02, 0.04, 0.02 and 0.03 eV, due to the large dihedral angles between the DMAC and the acceptor that effectively minimize the electronic coupling [**DMACBP** (82.2°), **DMACPyBP** (82.7°), **DMACBPN** (83.8°) and **DMACPyBPN** (86.1°) based on the optimized  $S_0$  geometries] (Fig. S26, ESI<sup>†</sup>). To gain insight into the photophysical properties of the  $S_1$  excited states, we performed natural transition orbital (NTO) analyses at the  $S_0$  geometry (Fig. 2(b)). The  $S_1$  state of each of the compounds possesses an evident intramolecular charge transfer (ICT) character. Hole and electron distributions mirror HOMO/LUMO localization, respectively. Root-mean-square displacements (RMSDs) between  $S_0$ ,  $S_1$  and  $T_1$  were calculated (Fig. 2(c) and Fig. S27, ESI<sup>†</sup>) using the VMD program to evaluate the structural changes that occur upon excitation and intersystem crossing.<sup>28</sup> The RMSD values of **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** between  $S_0$  and  $S_1$  were simulated to be 0.17, 0.14, 0.13 and 0.07 Å, respectively. The smaller RMSD of **DMACPyBPN** indicates that the introduction of N at the 10-position or 4/5-position reduces the magnitude of structural relaxation. In the relaxed  $S_1$  geometry, there is a larger  $S_1$ – $T_1$  spin-orbit coupling (SOC) matrix element of 0.05 cm<sup>−1</sup> in **DMACPyBP** and of 0.04 cm<sup>−1</sup> in **DMACPyBPN** than that in **DMACBP** (0.01 cm<sup>−1</sup>) and **DMACBPN** (0.01 cm<sup>−1</sup>) (Fig. 2(b)), indicating that the rate constants of intersystem crossing ( $k_{\text{isc}}$ ) in **DMACPyBP** and

**DMACPyBPN** will be faster than in the latter two compounds. As shown in the electrostatic potential maps (Fig. S38, ESI<sup>†</sup>), the nitrogen-rich components have a more negative electrostatic potential while the other parts have a neutral and a positive potential, which correlates with the trend in increasing electron-withdrawing ability of the acceptor with the inclusion of an increasing number of nitrogen atoms within the acceptor moiety.

### Photophysical investigations

The energies of the frontier molecular orbitals (FMOs) were inferred from the electrochemical behavior of **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** by cyclic voltammetry (CV) and differential pulse voltammetry (DPV) in degassed dichloromethane (DCM) with tetra-*n*-butylammonium hexafluorophosphate ([<sup>n</sup>Bu<sub>4</sub>N][PF<sub>6</sub>]) as the supporting electrolyte (Fig. 3(a)). Electrochemical potentials are reported *versus* a saturated calomel electrode. The reduction potentials ( $E_{\text{red}}$ ), determined from the DPV peak values, are −1.39 V (**DMACBP**), −1.22 V (**DMACPyBP**), −1.33 V (**DMACBPN**) and −1.11 V (**DMACPyBPN**), respectively, reflecting the expected anodic shift associated with increasing acceptor strength that is mirrored in the trend of the calculated LUMO levels from BP < BPN < PyBP < PyBPN (Fig. 1(a)). The calculated LUMO energies are −2.94 eV, −3.12 eV, −3.00 eV and −3.20 eV for **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN**, respectively. **DMACPyBP** (1.04 V) and **DMACPyBPN** (1.04 V) have more positive oxidation potentials than those of **DMACBP** (1.00 V) and **DMACBPN** (0.98 V), in line with the DFT calculations. The HOMO levels of **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** are −5.34, −5.39, −5.33 and −5.39 eV, respectively. The HOMO–LUMO gaps for **DMACBP**, **DMACBPN** and **DMACPyBPN** are 2.40, 2.33 and 2.19 eV, respectively, which mirror the trend in the DFT calculated values of 2.84, 2.72, and 2.63 eV; however, **DMACPyBP** exhibits a much smaller HOMO–LUMO gap (2.27 V) inferred from its electrochemistry than **DMACBPN** (2.33 V), which is opposite to the DFT calculated results in the gas phase. However, the inferred LUMO energies obtained from CV exactly mirror the trend from the DFT calculated LUMO of the acceptors as shown in Fig. 1(a). The inconsistency of the data for **DMACPyBP** to its corresponding acceptor likely indicates a deviation of the D–A dihedral angles between the DFT calculated structure in the gas phase and that exists in solution during the CV acquisition.

The UV-Vis absorption spectra of the four emitters in dilute toluene are shown in Fig. 3(b). The spectra show two prominent spectral features, with two intense bands centered at ~390 nm and a hypochromic broadband at ~470 nm, which are assigned to locally excited (LE)  $\pi$ – $\pi^*$  transitions of the donor and acceptor moieties,<sup>30,31</sup> and intramolecular charge transfer (ICT) transitions of the DMAC donor to the acceptor moiety, respectively.<sup>24,30</sup> The ICT band expectedly shifts to lower energies with the introduction of an increasing number of N atoms at positions 4/5 of the acceptor; for instance, the ICT band at 465 nm of **DMACBPN** ( $\epsilon = 2.5 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ ) is red-shifted compared to that of **DMACBP** ( $\lambda_{\text{abs}} = 458 \text{ nm}$ ,  $\epsilon = 1.8 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$ ) while those of **DMACPyBP** and **DMACPyBPN** absorb at essentially the same energy at 474 and 475 nm ( $\epsilon = 3.2 \times 10^3 \text{ M}^{-1} \text{ cm}^{-1}$  for both), which match well with the



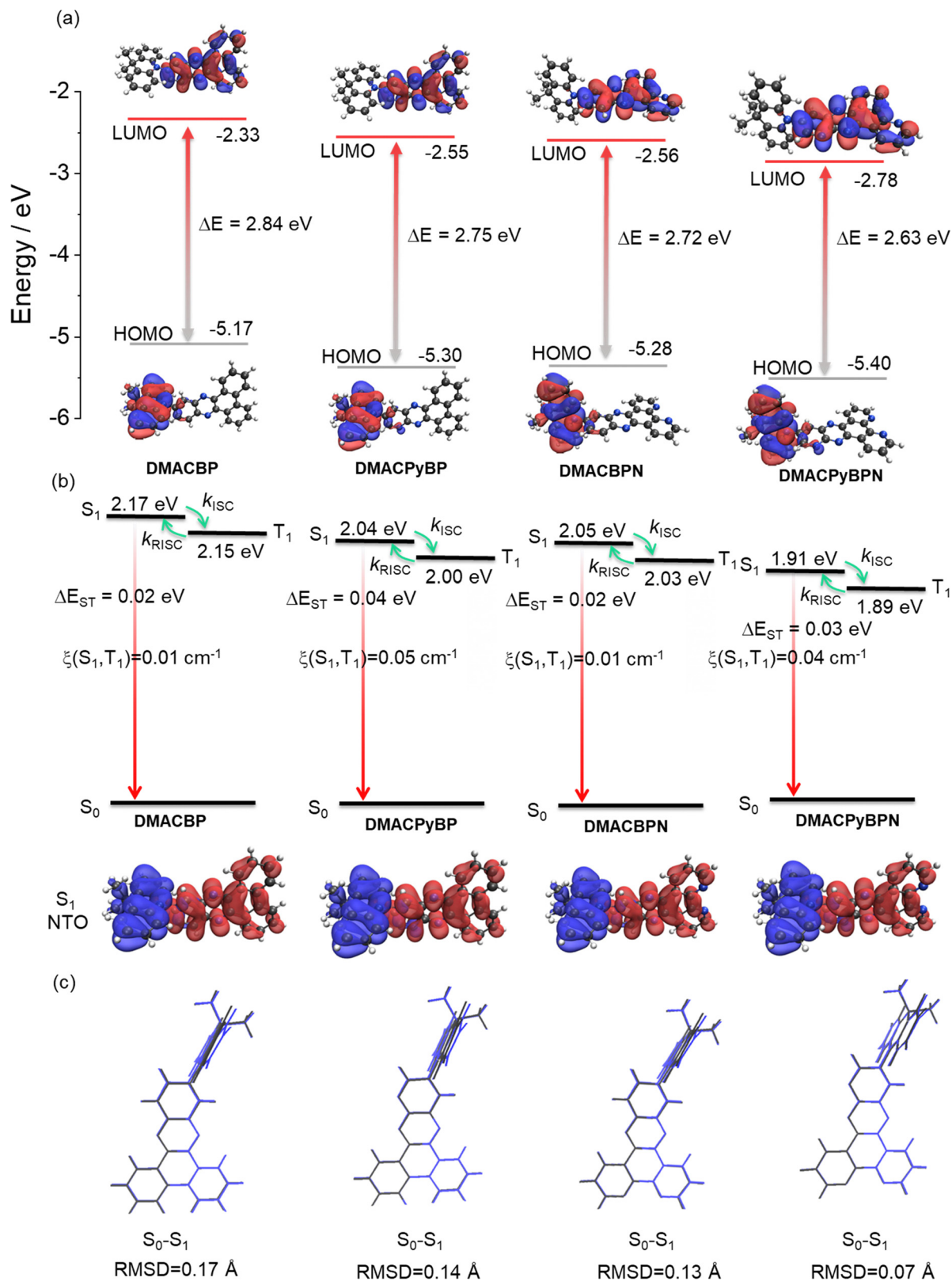
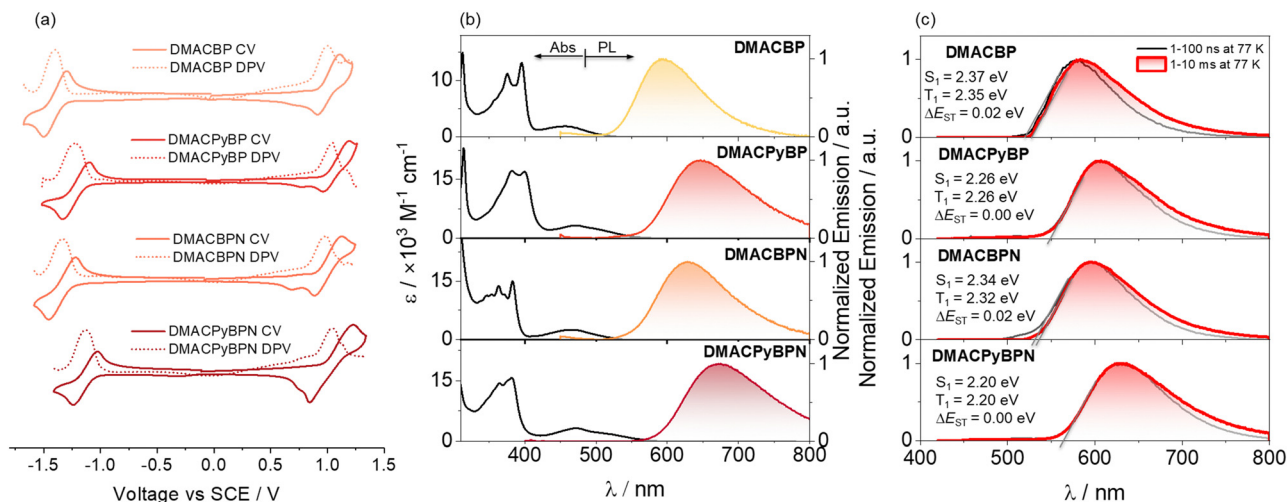


Fig. 2 (a) Frontier molecular orbitals (isovalue: 0.02) and (b) Vertical excitation energy levels, natural transition orbitals (unoccupied (hole, blue) & occupied (electron, red)), (isovalue: 0.02) of  $S_1$  of **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** calculated at the optimized  $S_0$  geometry in the gas phase at the PBE0/6-31G(d,p) level. (c) Comparison of optimized structures of **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** at  $S_0$  (grey) and  $S_1$  (blue) states.





**Fig. 3** (a) Cyclic and differential pulse voltammograms measured in degassed DCM with 0.1 M  $[\text{tBu}_4\text{N}]\text{PF}_6$  as the supporting electrolyte and  $\text{Fc}/\text{Fc}^+$  as the internal reference (0.46 V vs. SCE).<sup>29</sup> Scan rate = 100  $\text{mV s}^{-1}$ . (b) UV-vis absorption and steady-state photoluminescence (SSPL) spectra of **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** recorded in toluene at room temperature ( $\lambda_{\text{exc}} = 375 \text{ nm}$ ). (c) Prompt fluorescence (1–100 ns) and phosphorescence spectra (1–10 ms) in 2-MeTHF at 77 K of **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** ( $\lambda_{\text{exc}} = 375 \text{ nm}$ ).

TD-DFT calculated  $S_1$  energies (Fig. 2(b)). Positive photoluminescence (PL) solvatochromism is observed for all compounds (Fig. S29 and Table S1, ESI†), which is consistent with the ICT nature of the emissive excited state.

The steady-state photoluminescence (SSPL) spectra in toluene gradually red-shift from **DMACBP** ( $\lambda_{\text{PL}} = 595 \text{ nm}$ ) to **DMACBPN** ( $\lambda_{\text{PL}} = 630 \text{ nm}$ ) and **DMACPyBPN** ( $\lambda_{\text{PL}} = 672 \text{ nm}$ ), which coincides with an increasing number of nitrogen atoms in the acceptors (Fig. 3(b)). However, **DMACPyBP** ( $\lambda_{\text{PL}} = 645 \text{ nm}$ ) containing three nitrogen atoms within the acceptor exhibits a slightly red-shifted emission than **DMACBPN** ( $\lambda_{\text{PL}} = 630 \text{ nm}$ ), which contains four nitrogen atoms within the acceptor. This behavior correlates with the calculated  $S_1$  energies (Fig. 2(b)) and demonstrated that acceptor **PyBP** with 10-position N showed stronger electron-withdrawing abilities than **BPN** acceptor with two N in [1,10] phenanthroline, following the trend of acceptor strength as shown in Fig. 1(a). The corresponding optical bandgaps, determined from the intersection point of the normalized absorption and emission spectra,<sup>32–34</sup> for **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** are 2.37 eV, 2.20 eV, 2.31 eV and 2.12 eV, respectively (Fig. S30, ESI†), which is consistent with the

trend in emission energies (Fig. S31, ESI† and Table 1). The photoluminescence quantum yields ( $\Phi_{\text{PL}}$ ) for **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** in degassed toluene solution are 35%, 20%, 38% and 6%, respectively. When exposed to oxygen, these values significantly decrease to 9%, 7%, 8%, and 3% respectively. The energies of the onset of the prompt fluorescence and phosphorescence spectra in 2-MeTHF glass at 77 K were used to determine the  $S_1$  and  $T_1$  energies (Fig. 3(c)). The  $S_1$  energies are 2.37, 2.26, 2.34 and 2.20 eV, while the  $T_1$  energies are 2.35, 2.26, 2.32 and 2.20 eV, for **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN**, respectively. As both the prompt fluorescence and phosphorescence spectra are structureless, it is reasonable to assign both the  $S_1$  and  $T_1$  states as possessing significant CT character. The corresponding  $\Delta E_{\text{ST}}$  values for **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** are then 0.02 eV, 0.00 eV, 0.02 eV and 0.00 eV, respectively, which align with the TDA-DFT calculated results (Fig. 2(b)).

From the difference in transient PL decay behavior under aerated and degassed conditions in toluene, it is clear that the delayed emission is significantly quenched by the presence of oxygen (Fig. 4); interestingly, the quenching of the PL in

**Table 1** Summary of photophysical properties of nitrogen-doped PAH acceptor-based compounds

| Molecule         | $\lambda_{\text{abs}}/(\epsilon/\times 10^3 \text{ M}^{-1} \text{ cm}^{-1})/\text{nm}^a$ | $\lambda_{\text{PL}}^a/\text{nm}$ | $\tau_{\text{p}}/\text{ns}$ | $\tau_{\text{d}}/\mu\text{s}^a$ | $\lambda_{\text{PL}}^b/\text{nm}$ | $\Phi_{\text{PL}}^b/\%$ | $S_1/T_1^c/\text{eV}$ | $\Delta E_{\text{ST}}^c/\text{eV}$ | $\tau_{\text{p}}/\text{ns}^d$ | $\tau_{\text{d}}/\mu\text{s}^d$ | $k_{\text{ISC}}^e/\times 10^7 \text{ s}^{-1}$ | $k_{\text{RISC}}^e/\times 10^4 \text{ s}^{-1}$ | $E_{\text{HOMO}}^f/\text{eV}$ | $E_{\text{LUMO}}^f/\text{eV}$ | $E_{\text{g}}^g/\text{eV}$ |
|------------------|------------------------------------------------------------------------------------------|-----------------------------------|-----------------------------|---------------------------------|-----------------------------------|-------------------------|-----------------------|------------------------------------|-------------------------------|---------------------------------|-----------------------------------------------|------------------------------------------------|-------------------------------|-------------------------------|----------------------------|
| <b>DMACBP</b>    | 395 (13), 458(1.8)                                                                       | 595                               | 18                          | 146                             | 568                               | 75(56)                  | 2.33/2.31             | 0.02                               | 15.3                          | 130.3                           | 1.7                                           | 1.0                                            | −5.34                         | −2.94                         | 2.40                       |
| <b>DMACPyBP</b>  | 400 (17), 474(3.2)                                                                       | 645                               | 20                          | 3.5                             | 601                               | 47(41)                  | 2.32/2.31             | 0.01                               | 21.1                          | 5.9                             | 0.6                                           | 19.4                                           | −5.39                         | −3.12                         | 2.27                       |
| <b>DMACBPN</b>   | 383 (15), 465(2.5)                                                                       | 630                               | 27                          | 19.7                            | 586                               | 71(52)                  | 2.26/2.25             | 0.01                               | 22.4                          | 47.2                            | 1.2                                           | 2.9                                            | −5.33                         | −3.00                         | 2.33                       |
| <b>DMACPyBPN</b> | 382 (16), 475(3.2)                                                                       | 672                               | 10                          | 2.3                             | 606                               | 37(30)                  | 2.32/2.31             | 0.01                               | 23.2                          | 2.9                             | 0.8                                           | 42.6                                           | −5.39                         | −3.20                         | 2.19                       |

<sup>a</sup> In PhMe at 298 K ( $\lambda_{\text{exc}} = 340 \text{ nm}$ ). <sup>b</sup> Spin-coated 2 wt% emitters doped in CBP films and  $\Phi_{\text{PL}}$  values were determined using an integrating sphere ( $\lambda_{\text{exc}} = 340 \text{ nm}$ ). Values quoted are under  $\text{N}_2$ . Values in parentheses are in air. <sup>c</sup>  $S_1$  was obtained from the onset of the prompt emission (time-gated window: 1–100 ns) ms measured in doped film at 77 K and  $T_1$  was obtained from the onset of the phosphorescence spectrum (time-gated window: 1–10 ms) measured in doped film at 77 K. <sup>d</sup> PL lifetime. <sup>e</sup>  $k_{\text{ISC}}$  = intersystem crossing rate constant from  $S_1$  to  $T_1$  states;  $k_{\text{RISC}}$  = reverse intersystem crossing rate constant. <sup>f</sup> In DCM with 0.1 M  $[\text{tBu}_4\text{N}]\text{PF}_6$  as the supporting electrolyte and  $\text{Fc}/\text{Fc}^+$  as the internal reference (0.46 V vs. SCE).<sup>29</sup> The HOMO and LUMO energies were determined using  $E_{\text{HOMO/LUMO}} = -(E_{\text{ox}}/E_{\text{red}} + 4.8) \text{ eV}$ , where  $E_{\text{ox}}$  and  $E_{\text{red}}$  are anodic and cathodic peak potentials, respectively, obtained from the DPV versus  $\text{Fc}/\text{Fc}^+$ .<sup>29</sup> <sup>g</sup>  $E_{\text{g}} = |E_{\text{HOMO}} - E_{\text{LUMO}}|$ .





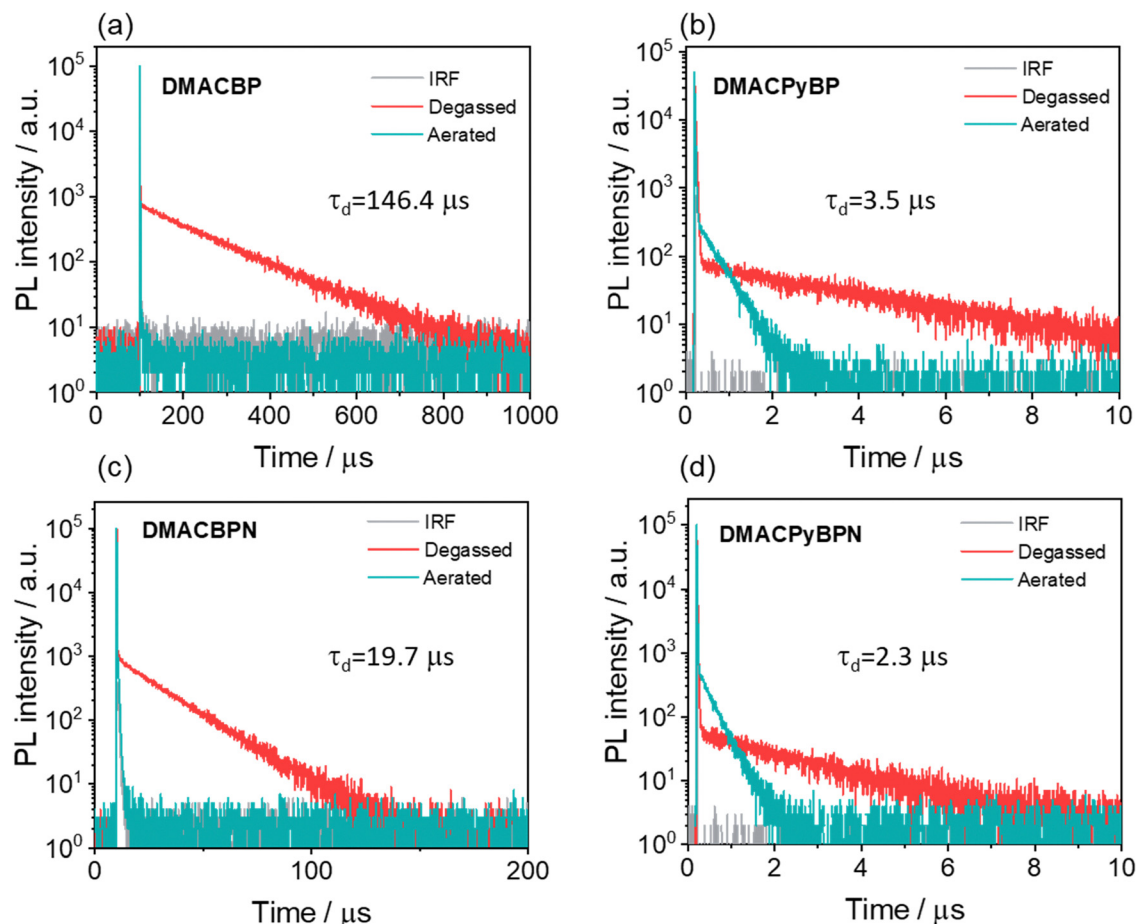


Fig. 4 Time-resolved PL decay of (a) **DMACBP**, (b) **DMACPyBP**, (c) **DMACBPN** and (d) **DMACPyBPN** in degassed and aerated toluene (time window: 1 ns to 200  $\mu$ s or 1 ns to 1 ms measured by MCS, 1 ns to 10  $\mu$ s measured by TCSPC,  $\lambda_{\text{exc}} = 375$  nm).

**DMACBP** and **DMACBPN** is more significant than in **DMACPyBP** and **DMACPyBPN**, the latter two of which contain an additional pyridine ring fused to the pyrazine in the acceptor. The increased quenching of the PL of the former two compounds may be due to their longer delayed lifetimes. The emission intensity is likewise strongly affected by the presence of oxygen, especially for **DMACBP** and **DMACBPN**, while for **DMACPyBP** and **DMACPyBPN**, the enhancement in intensity upon oxygen removal was much smaller (Fig. S32, ESI $^{\dagger}$ ). The PL decays with biexponential kinetics, with prompt fluorescence lifetimes,  $\tau_p$ , of 18 ns, 21 ns, 27 ns and 10 ns (Fig. S33, ESI $^{\dagger}$ ), and delayed fluorescence lifetimes,  $\tau_d$ , of 146.4  $\mu$ s, 3.5  $\mu$ s, 19.7  $\mu$ s and 2.3  $\mu$ s, respectively, for **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN**. The corresponding rate constants of intersystem crossing ( $k_{\text{ISC}}$ ) for **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** are  $4.1 \times 10^7 \text{ s}^{-1}$ ,  $3.3 \times 10^7 \text{ s}^{-1}$ ,  $2.9 \times 10^7 \text{ s}^{-1}$  and  $5.0 \times 10^7 \text{ s}^{-1}$ , respectively, while the rate constants for reverse intersystem crossing ( $k_{\text{RISC}}$ ) for **DMACPyBP** ( $81.6 \times 10^4 \text{ s}^{-1}$ ) and **DMACPyBPN** ( $87.0 \times 10^4 \text{ s}^{-1}$ ) are much faster than those of **DMACBP** ( $2.7 \times 10^4 \text{ s}^{-1}$ ) and **DMACBPN** ( $24.1 \times 10^4 \text{ s}^{-1}$ ), due in part to the smaller  $\Delta E_{\text{ST}}$  in the former two that is correlated to the introduction of a nitrogen atom at the 10-position in **DMACPyBP** and **DMACPyBPN**.

We next investigated the photophysical properties of the four emitters in an OLED-relevant host, 4,4'-N,N'-dicarbazole-biphenyl (CBP), as this host matrix has sufficiently high triplet energy ( $T_1 = 2.56 \text{ eV}$ ) to confine the excitons onto the emitter.<sup>35,36</sup> The PL of the doped films of varying dopant concentrations from 2–10 wt% of emitters was first investigated (Fig. S36, ESI $^{\dagger}$ ) and the corresponding  $\Phi_{\text{PL}}$  values are summarized in Table S2 (ESI $^{\dagger}$ ). As the doping concentration is increased, the emission of all of the compounds is red-shifted from yellow to red (Fig. S34, ESI $^{\dagger}$ ), and the highest  $\Phi_{\text{PL}}$  values are at a doping concentration of 2 wt%. The SSPL spectra of **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** at a doping concentration of 2 wt% exhibit broad and unstructured emission with  $\lambda_{\text{PL}}$  at 568, 601, 586, and 606 nm, respectively (Fig. 5(a)). The trend in  $\lambda_{\text{PL}}$  mirrors those in toluene (Fig. 3(b)). The  $\Phi_{\text{PL}}$  values of these films of **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** are 75, 47, 71 and 37%, respectively, under a  $\text{N}_2$  atmosphere (Fig. 5(b) and Table 1). The decrease in  $\Phi_{\text{PL}}$  for **DMACPyBP/DMACPyBPN** can be ascribed to their smaller energy gaps, thus leading to increased non-radiative decay.<sup>37</sup> However, the difference in the distribution of N atoms in the BP and BPN (two additional nitrogen atoms on the 4- and 5-position) acceptors has only a relatively small impact on the acceptor strength,





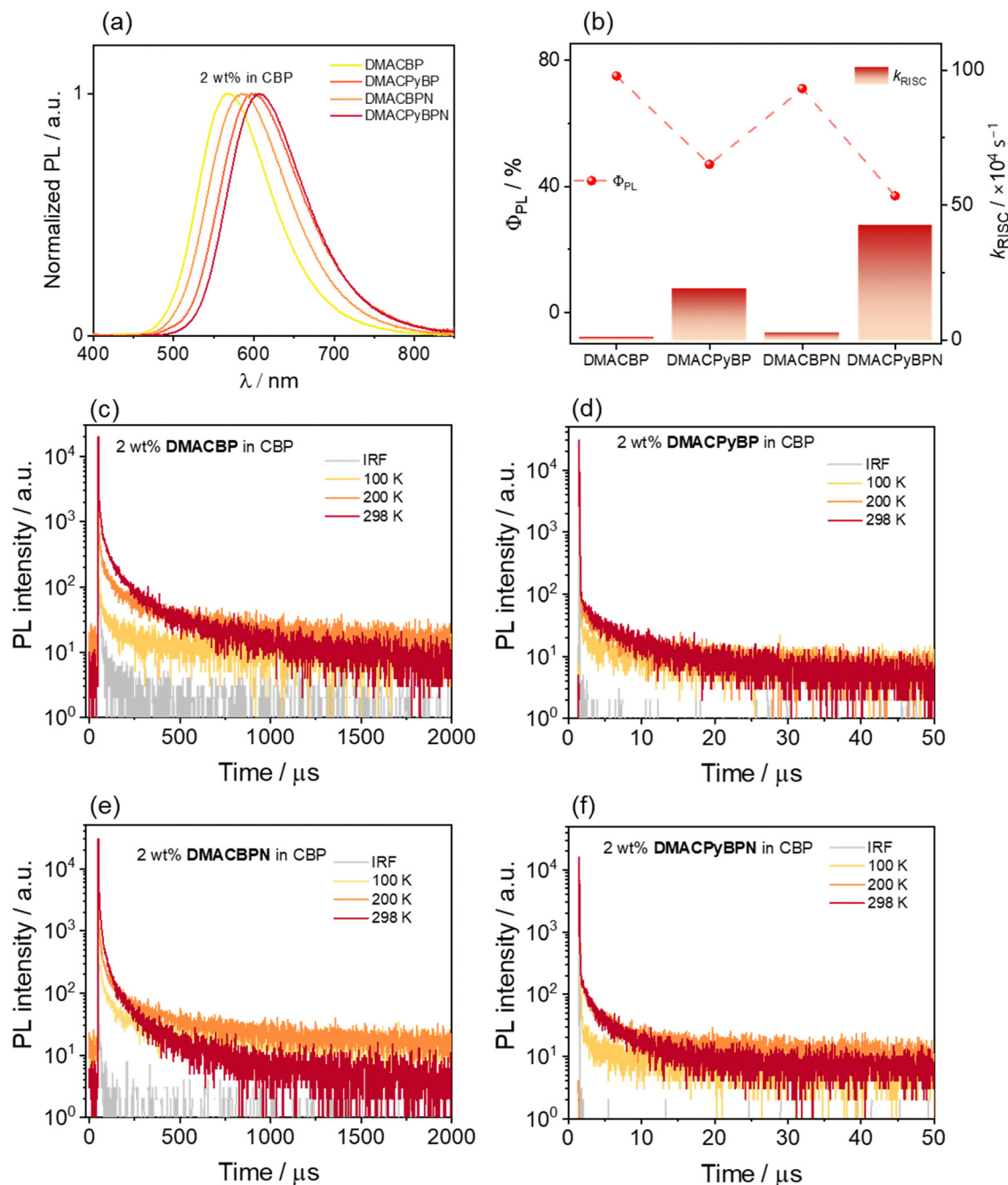


Fig. 5 (a) PL spectra of the DMACBP, DMACPyBP, DMACBPN and DMACPyBPN in 2 wt% doped films in CBP ( $\lambda_{\text{exc}} = 360 \text{ nm}$ ); (b)  $\Phi_{\text{PL}}$  and  $k_{\text{RISC}}$  for each of the four emitters; temperature-dependent TRPL decays of (c) 2 wt% DMACBP, (d) 2 wt% DMACPyBP, (e) 2 wt% DMACBPN and (f) 2 wt% DMACPyBPN doped CBP films ( $\lambda_{\text{exc}} = 375 \text{ nm}$ ).

with a small red shift in the emission from 568 nm for DMACBP to 586 nm for DMACBPN; as well, there is only a negligible decrease in  $\Phi_{\text{PL}}$  between DMACBP and DMACBPN (Table 1 and Fig. 5(b)).<sup>38</sup> The  $\Phi_{\text{PL}}$  values of DMACBP and DMACBPN exhibited a larger decrease in the presence of air at 56 and 52%, respectively, compared to 41% for DMACPyBP and 30% for DMACPyBPN, indicating a greater proportion of triplet excitons in DMACBP and DMACBPN. As shown in Fig. 5, all four compounds showed multiexponential decay kinetics, with average prompt

fluorescence lifetimes, average  $\tau_{\text{p}}$ , of 15.3, 21.1, 22.4, and 23.2 ns (Fig. S35, ESI†) and average delayed emission lifetimes, average  $\tau_{\text{d}}$ , of 130.3, 5.9, 47.2, and 2.9  $\mu\text{s}$  at room temperature for DMACBP, DMACPyBP, DMACBPN and DMACPyBPN, respectively. The corresponding  $k_{\text{ISC}}$  values for DMACBP, DMACPyBP, DMACBPN and DMACPyBPN in the 2 wt% doped films in CBP are  $1.7 \times 10^7 \text{ s}^{-1}$ ,  $0.6 \times 10^7 \text{ s}^{-1}$ ,  $1.2 \times 10^7 \text{ s}^{-1}$  and  $0.8 \times 10^7 \text{ s}^{-1}$ , respectively, while  $k_{\text{RISC}}$  for DMACPyBP and DMACPyBPN reached  $1.94 \times 10^5 \text{ s}^{-1}$  and  $4.26 \times 10^5 \text{ s}^{-1}$ , more than ten times

higher than those of **DMACBP** at  $0.10 \times 10^5 \text{ s}^{-1}$  and **DMACBPN** at  $0.29 \times 10^5 \text{ s}^{-1}$  (Fig. 5(b)), following a similar trend to that observed for toluene. The introduction of a nitrogen atom at the 10-position in **DMACPyBP** and **DMACPyBPN** leads to a much stronger CT and faster  $k_{\text{RISC}}$ . Temperature-dependent time-resolved PL (TRPL) analysis shows that the intensity of the delayed emission increases with increasing temperatures for all 4 compounds, confirming that these compounds emit *via* TADF (Fig. 5(c) and (d)). The energy levels of the  $S_1$  and  $T_1$  states were estimated from the onset of the corresponding prompt fluorescence and phosphorescence spectra of the 2 wt% doped films in CBP at 77 K (Fig. S36, ESI†). As expected, **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** have, respectively, small  $\Delta E_{\text{ST}}$  values of 0.02, 0.01, 0.01 and 0.01 eV, values consistent with those in 2-MeTHF glass.

### Device characterization

To evaluate the electroluminescence (EL) performance of these emitters, OLEDs employing different doping concentration emitters (2, 5, 8 and 11 wt%, the data for 5 and 8 wt% can be found in Fig. S38 and Table S3, ESI†) doped in CBP films as the emissive layer (EML) was fabricated. The OLEDs have the following configuration: ITO (indium tin oxide)/TAPC (4,4'-(cyclohexane-1,1-diyl)bis(*N,N*-di-*p*-tolylaniline)) (40 nm)/TCTA (tris(4-(9*H*-carbazol-9-yl)phenyl)amine) (10 nm)/CBP (10 nm)/CBP:*x* wt% dopants (20 nm)/TmPyPB (3,3'-(5'-(3-(pyridin-3-yl)phenyl)-[1,1':3',1''-terphenyl]-3,3''-diyl)dipyridine) (45 nm)/LiF (1 nm)/Al (Fig. 6(a)). ITO and Al serve as the anode and the cathode, respectively, while TAPC, TCTA and TmPyPB are,

respectively, used as the hole-transporting layer, the electron-blocking layer, and the electron-transporting layer.

As shown in Fig. 6(b), the OLEDs with **DMACBP**, **DMACPyBP**, **DMACBPN** and **DMACPyBPN** at a doping concentration of 2 wt% in the CBP host exhibit yellow-to-red EL with the peak emission,  $\lambda_{\text{EL}}$ , at 576, 600, 588 and 608 nm, respectively, which match the PL in CBP very well (Fig. 5(a)). The corresponding CIE coordinates are (0.48, 0.50), (0.56, 0.44), (0.53, 0.46) and (0.57, 0.42). The devices with **DMACBP**, **DMACPyBP**, **DMACBPN**, and **DMACPyBPN** showed  $\text{EQE}_{\text{max}}$  of 18.5, 13.7, 19.4 and 10.5%, respectively, reflecting the variations in their intrinsic  $\Phi_{\text{PL}}$  (Fig. 6(c) and Table 2). The maximum current efficiency ( $\text{CE}_{\text{max}}$ ) and power efficiency ( $\text{PE}_{\text{max}}$ ) values are 58.7  $\text{cd A}^{-1}$  and 54.2  $\text{lm W}^{-1}$  for the devices with **DMACBP**, 28.9  $\text{cd A}^{-1}$  and 26.7  $\text{lm W}^{-1}$  for the devices with **DMACPyBP**, 48.9  $\text{cd A}^{-1}$  and 46.5  $\text{lm W}^{-1}$  for the devices with **DMACBPN**, and 17.4  $\text{cd A}^{-1}$  and 16.6  $\text{lm W}^{-1}$  for the devices with **DMACPyBPN** (Fig. 6(d)), respectively. The efficiency roll-off behavior is noticeably different between the four devices. At a luminance of 100  $\text{cd m}^{-2}$ , a large efficiency roll-off of 40.5% and 24.7% was observed in the devices with **DMACBP** and **DMACBPN**, respectively, which decreased considerably to 8.0% and 4.7% for the devices with **DMACPyBP** and **DMACPyBPN**, respectively. The EQE decreased to 3.7% at a luminance of 1000  $\text{cd m}^{-2}$  for the device with **DMACBP**; however, for the device with **DMACPyBP**, the  $\text{EQE}_{1000}$  was 7.5%. Similarly, at the same current density of 1  $\text{mA cm}^{-2}$ , the devices with **DMACBP** and **DMACBPN** exhibit much larger efficiency roll-off of 58% and 36%, respectively, than the devices with **DMACPyBP** (16%) and **DMACPyBPN** (14%)

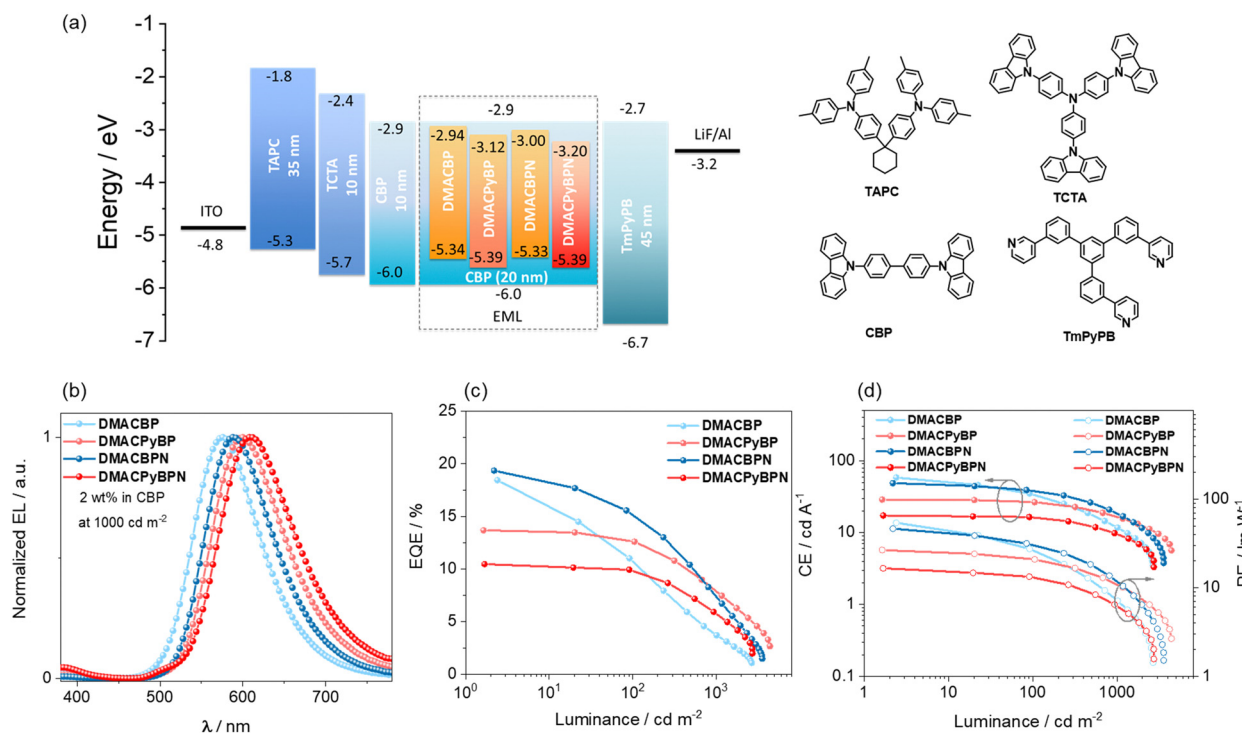


Fig. 6 (a) Energy level diagram and molecular structure of materials employed in the devices; (b) EL spectra of the devices; (c) external quantum efficiency versus luminance curves for the devices; (d) current efficiency and power efficiency versus luminance curves for the devices.

Table 2 Electroluminescence data for the devices

| Emitter          | Concentration/<br>wt% | $V_{on}^a$ /<br>V | $CE_{max}^b$ /<br>A <sup>-1</sup> cd | $PE_{max}^c$ /lm<br>W <sup>-1</sup> | $EQE_{max/100/1000}^d$ /<br>% | Efficiency roll-off<br>100/1000 <sup>e</sup> /<br>% | $L_{max}^f$ /cd<br>m <sup>-2</sup> | $\lambda_{EL}^g$ /<br>nm | CIE <sup>h</sup> (x,<br>y) |
|------------------|-----------------------|-------------------|--------------------------------------|-------------------------------------|-------------------------------|-----------------------------------------------------|------------------------------------|--------------------------|----------------------------|
| <b>DMACBP</b>    | 2.0%                  | 3.4               | 58.7                                 | 54.2                                | 18.5/11.0/3.7                 | 40/80                                               | 2665                               | 576                      | 0.48, 0.50                 |
|                  | 11.0%                 | 3.5               | 38.1                                 | 34.2                                | 14.6/12.0/5.6                 | 18/62                                               | 4332                               | 584                      | 0.52, 0.47                 |
| <b>DMACPyBP</b>  | 2.0%                  | 3.4               | 28.9                                 | 26.7                                | 13.7/12.6/7.5                 | 8/45                                                | 4346                               | 600                      | 0.56, 0.44                 |
|                  | 11.0%                 | 3.4               | 11.6                                 | 10.7                                | 9.0/8.3/4.7                   | 8/47                                                | 3369                               | 620                      | 0.61, 0.39                 |
| <b>DMACBPN</b>   | 2.0%                  | 3.3               | 48.9                                 | 46.5                                | 19.4/14.6/6.7                 | 25/65                                               | 3551                               | 588                      | 0.53, 0.46                 |
|                  | 11.0%                 | 3.1               | 31.9                                 | 32.3                                | 15.9/13.4/7.5                 | 16/53                                               | 4315                               | 600                      | 0.57, 0.43                 |
| <b>DMACPyBPN</b> | 2%                    | 3.3               | 17.4                                 | 16.6                                | 10.5/10.0/5.0                 | 5/52                                                | 2723                               | 608                      | 0.57, 0.42                 |
|                  | 11%                   | 3.2               | 4.1                                  | 4.0                                 | 5.4/5.1/2.0                   | 5/63                                                | 1428                               | 640                      | 0.63, 0.37                 |

<sup>a</sup> Voltage at 1 cd m<sup>-2</sup>. <sup>b</sup> Maximum current efficiency. <sup>c</sup> Maximum power efficiency. <sup>d</sup> Maximum external quantum efficiency at 100 cd m<sup>-2</sup>/at 1000 cd m<sup>-2</sup>. <sup>e</sup> EQE roll off at 100 cd m<sup>-2</sup>/at 1000 cd m<sup>-2</sup>. <sup>f</sup> Maximum luminance. <sup>g</sup> EL emission peak at 1000 cd m<sup>-2</sup>. <sup>h</sup> Commission Internationale de L'Éclairage coordinates.

(as shown in Fig. S37, ESI†). A plausible explanation for this difference in efficiency roll-off behavior is due to the faster  $k_{RISC}$  and shorter  $\tau_d$  in **DMACPyBP** and **DMACPyBPN**, thus leading to smaller triplet exciton populations at high current densities in the devices with these two emitters. The **DMACPyBP**-based OLEDs showed remarkably low-efficiency roll-off, especially compared to other DMAC-containing orange-red TADF OLEDs (Fig. S1, ESI†).<sup>23,24,30</sup> Increasing the doping concentration to 5 wt%, 8 wt% or even to 11 wt% does not lead to a further improvement in performance and the EL spectra red-shifted progressively (Fig. S38 and Table S3, ESI†). At 11 wt% doping, the OLEDs emit in the range of 600–640 nm and showed an EQE<sub>max</sub> of 5.5–15.9% (Table S3, ESI†).

## Conclusion

We have developed a family of four orange-to-red TADF compounds whose structures differ in the number of nitrogen atoms contained within the conjugated acceptor core. The structure–property relationship among the four compounds has been systematically investigated using DFT calculations and a combination of photophysical and electrochemical studies; OLEDs were also fabricated using these materials as emitters. It was found that increasing the nitrogen atom content in the acceptor (BP) of the D–A type compounds results in a more stabilized LUMO, a smaller  $\Delta E_{ST}$  and a faster  $k_{RISC}$ . In particular, the presence of nitrogen at the 10-position of BP notably enhances  $k_{RISC}$ , as demonstrated in **DMACPyBP** and **DMACPyBPN** compared to **DMACBP** and **DMACBPN**, despite the small decrease in  $\Phi_{PL}$ . OLEDs with **DMACBPN** showed a maximum EQE of 19.4% and an electroluminescence maximum of 588 nm. As the N/C ratio increased, the EL spectra of the corresponding devices progressively red-shifted, with the reddest-emitting device ( $\lambda_{EL}$  = 640 nm) employing **DMACPyBPN**. Furthermore, the **DMACPyBP**/**DMACPyBPN**-based OLEDs showed a remarkably low-efficiency roll-off of only 8 and 5%, compared to 40 and 25% at 100 cd m<sup>-2</sup> for the devices with **DMACBP**/**DMACBPN**, respectively.

## Conflicts of interest

There are no conflict to declare.

## Acknowledgements

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