


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# Weaving host matrices with intermolecular hydrogen bonds for high-efficiency white thermally activated delayed fluorescence†

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A thermally activated delayed fluorescence (TADF) white organic light-emitting diode (WOLED) holds great promise for low-cost, large-scale lighting applications. Nevertheless, manipulating exciton allocation in a white TADF single layer is still a challenge. Herein, we demonstrate that the exciton kinetic process of dually doped white TADF films is strongly dependent on the grid regularity of the host matrix. Intermolecular hydrogen bonds (IHBs) are used to weave the matrices of two host molecules DPEQPO and DPSQPO featuring four phosphine oxide (PO) groups and different IHB orientations. The DPSQPO matrix forms regular grids to uniformly disperse and separate dopants, while DPEQPO exhibits chaotic IHBs, in turn inducing a heterogeneous dopant distribution. As a consequence, in both photoluminescence and electroluminescence processes, in contrast to DPEQPO hosted systems with comparable singlet Förster resonance energy transfer and triplet Dexter energy transfer, DPSQPO provides a FRET-predominant exciton allocation between blue and yellow dopants, which markedly suppresses triplet quenching and improves the white color purity, resulting in a state-of-the-art external quantum efficiency up to 24.2% of its single-emissive-layer pure-white TADF diode, in contrast to 16.0% for DPEQPO based analogs. These results indicate the significance of host engineering for exciton kinetics and suggest the feasibility of host grid design for developing high-performance TADF lighting.

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## 1. Introduction

Thermally activated delayed fluorescence (TADF) holds immense potential for application in white organic light-emitting diodes (WOLEDs) for diverse daily lighting applications,<sup>1–3</sup> owing to its merits of 100% theoretical internal quantum efficiency (IQE,  $\eta_{IQE}$ ), low cost and environmental friendliness.<sup>4–7</sup> One of the intrinsic advantage of TADF materials is that they are pure-organic donor–acceptor (D–A) systems featuring strong intramolecular charge transfer (CT), giving rise to negligible singlet–triplet splitting ( $\Delta E_{ST}$ )<sup>8,9</sup> and feasible reverse inter-system crossing (RISC) for up-conversion from nonradiative triplets to radiative singlets.<sup>10</sup> Therefore, despite showing potentially 100% exciton harvesting and desired sustainability, TADF molecules have high polarity and strong intermolecular interactions, causing serious collision and dipole–interaction

induced exciton quenching and marked efficiency roll-off. Consequently, doped emissive layers (EMLs) featuring TADF emitters dispersed in an inertia host matrix were widely used in high-efficiency monochromatic diodes, partially leading to the rapid improvement of external quantum efficiencies ( $\eta_{EQE}$ ) beyond 25%.<sup>11–18</sup> These advances lead to developing high-performance pure-TADF WOLEDs,<sup>19–25</sup> especially based on single-EML structures<sup>26–33</sup> to maximize the advantages of organic materials and devices in low cost and large-scale production.

However, with CT-featuring first singlet ( $S_1$ ) states (<sup>1</sup>CT), the emission colors of TADF emitters are directly proportional to molecular polarities. Therefore, dipole–dipole interactions<sup>34,35</sup> between blue and other high-polarity emitters worsen the quenching of multicomponent white TADF systems. Furthermore, compared to fluorescence (FL) and phosphorescence (PH) counterparts, the exciton allocation between different TADF emitters is markedly more complicated through not only singlet Förster resonance energy transfer (FRET), but also charge-exchange based triplet Dexter energy transfer (DET) and intermolecular CT.<sup>36,37</sup> These processes should be accurately balanced to rationalize exciton allocation for achieving both high efficiencies and satisfactory white color purity.<sup>38,39</sup> Moreover, the short-range feature of DET makes it often accompanied by intermolecular interaction induced triplet quenching,<sup>40</sup>

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*e.g.* singlet–triplet (STA) and triplet–triplet annihilation (TTA).<sup>41</sup> Therefore, combining the state-of-the-art monochromatic efficiencies into a single-EML TADF WOLED is still a formidable challenge, because of mutual quenching between different emitters and their competition in the exciton allocation process.<sup>42–45</sup>

Considering the different distance dependences of energy transfer,<sup>46,47</sup> CT and quenching effect,<sup>48</sup> the rational dispersion of TADF molecules in host matrices is crucial to overcome this challenge, since intermolecular interactions and contributions of FRET, DET and intermolecular CT to exciton allocation can be accurately modulated and optimized by adjusting dispersive uniformity in the host matrix. Compared to FL and PH counterparts, most blue/white TADF diodes adopted high doping concentrations of blue emitters ( $\geq 10\%$ ) to take use of intermolecular CT channels for RISC enhancement.<sup>49–52</sup> In EMLs of these devices, hosts with too weak intermolecular interactions cannot effectively disperse blue dopants. Conversely, too strong intermolecular interactions of host matrices cause the concurrent heterogeneous distributions of both host and dopant molecules. It means the “ideal” host materials for TADF WOLEDs should have appropriate intermolecular interactions to form regular gridding matrices uniformly filled with dopant molecules. In our previous studies, compared to too strong  $\pi$ – $\pi$  stacking, intermolecular hydrogen bond (IHB) networks in host matrices were more effective to separate dopants and avoid worsening host–host and host–dopant quenching.<sup>53–56</sup> It is also noted that IHB modes directly influenced the electroluminescence (EL) performance of the devices, which actually provides a desired platform to get a deep insight into the accurate relationship between the regularity of host grids and exciton-involved PL and EL processes in single-layer white TADF systems.

In this contribution, we construct two host molecules named **DPEQPO** and **DPSQPO** respectively featuring diphenylether (DPE) and diphenylsulfide (DPS) skeletons substituted with four diphenylphosphine oxide (DPPO) groups (Scheme S1†). The bond angle reduction from ether to sulfide induces the orientations of P=O groups to change from four different directions for **DPEQPO** to more concentrated but sufficient two directions for **DPSQPO**. Consequently, in contrast to the chaotic and complicated IHB network of **DPEQPO**, **DPSQPO** exhibits high regularity with ordered IHB grids, giving rise to the uniform dispersion of both blue and yellow TADF dopants in its dually doped single-layer film. The rationally weaved **DPSQPO** matrix endowed the single-EML pure-white and warm-white diodes with a state-of-the-art  $\eta_{\text{EQE}}$  up to 24.2% and 22.4%, respectively, which are improved by 70% and 40%, compared to **DPEQPO** based analogs. For both PL and EL processes, in addition to dramatically improved quantum efficiencies, the delayed fluorescence (DF) lifetimes ( $\tau_{\text{DF}}$ ) of **DPSQPO** based white TADF systems are far shorter than those of **DPEQPO** based analogs. These results suggest that uniform dopant dispersion and separation in the **DPSQPO** matrix make FRET predominant in exciton allocation, and the effective restraint of DET actually accelerates exciton radiation, and simultaneously suppresses triplet quenching. Thus, host intermolecular interaction

optimization and host-dopant synergism are crucial for developing high-performance single-layer white TADF systems.

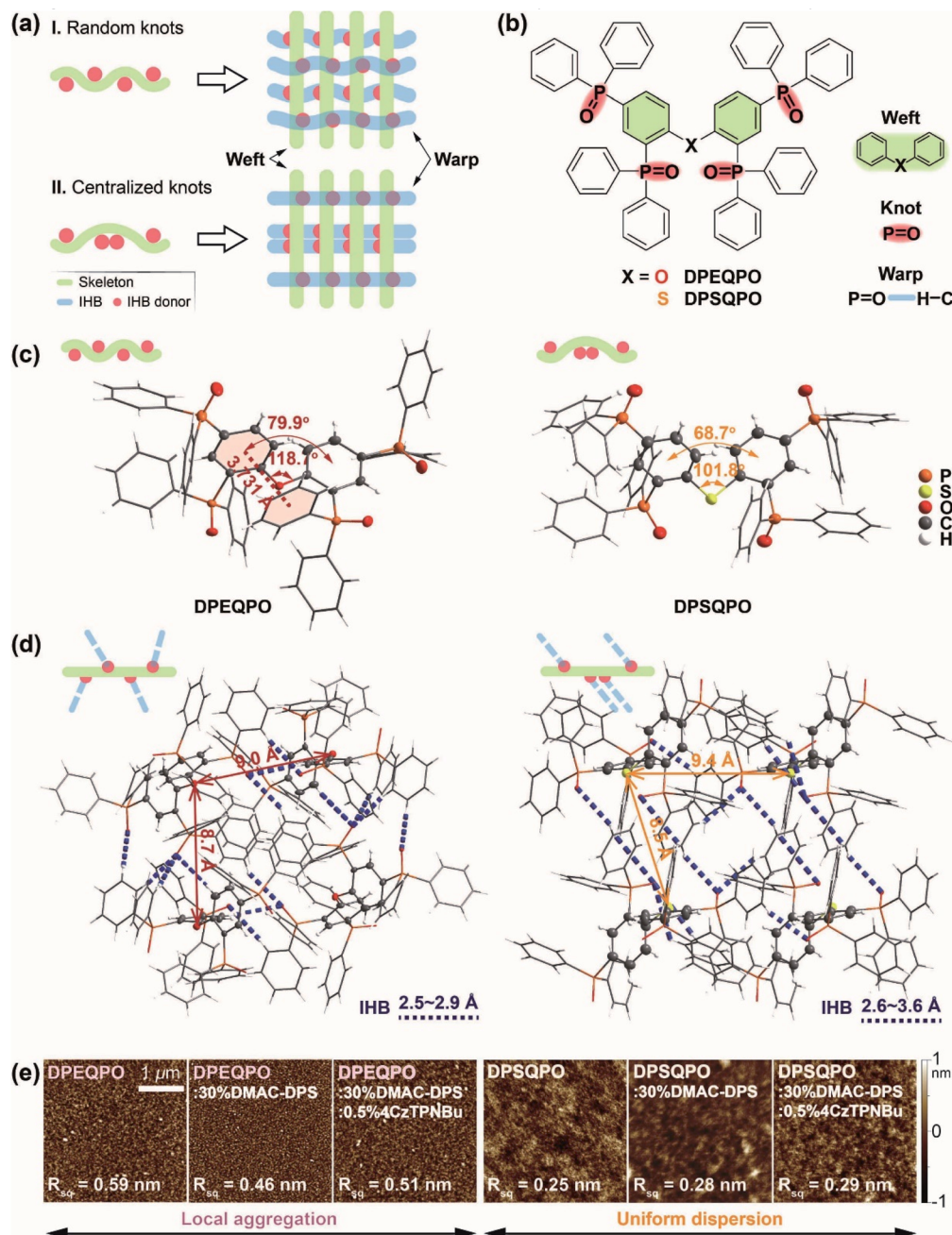
## 2. Results and discussion

### 2.1 Design, structures and morphological properties

To form continuous IHB networks, host molecules should contain more than one hydrogen-bond donor group. In the cases of **DPEQPO** and **DPSQPO**, collectively named **DPXQPO**, their four P=O groups make two-dimensionally structural weaving feasible. The plain weave models show that the grid structure is directly dependent on the knot arrangement, which provides two conventional four-knot models (Fig. 1a): (i) Model I: wefts with four random and scattered knots linked with warps to constitute a complicated but dense cross structure; (ii) Model II: pairwise centralizing knots on wefts improves weft and warp regularity, and relatively enlarges grid gaps. In **DPXQPO** matrices, P=O groups serve as knots on DPE and DPS skeletons (wefts) to form P=O...H IHBs (warps) between adjacent molecules (Fig. 1b). Single-crystal structures show that the angles of the C–O–C bond in **DPEQPO** and the C–S–C bond in **DPSQPO** are respectively 118.7° and 101.8°, respectively, close to the typical values of 120° and 103° (Fig. 1c). The larger C–O–C bond angle corresponds to a dihedral angle of 79.9° for the DPE skeleton, in contrast to 68.7° of the DPS skeleton. Therefore, the larger mutual steric hindrance between two DPPOs at the *ortho* positions of more linear DPE gives rise to the different orientations of one inward and the other outward P=O bonds. Due to a similar steric effect, the P=O orientations of two *para*-DPPOs in **DPEQPO** are also different. As a result, all P=O groups of **DPEQPO** point to different directions. In contrast, a more folded DPS skeleton provides more space to the two *ortho*-DPPO groups of **DPSQPO**, leading to two inward P=O bonds with the same orientation. The identical *meta*-position steric hindrances of these *ortho*-DPPOs further induce the same P=O orientations of two *para*-DPPOs.

Consequently, in accord with Model I, **DPEQPO** has four differently orientated P=O groups alternately on both sides of the DPE skeleton. The P=O arrangement in **DPSQPO** is similar to that in Model II, namely two *ortho*-substituted P=O groups centralized on one side of the DPS skeleton, accompanied by two P=O groups with the same orientation at the *para*-positions on the other side. As predicated by plain weave models, the single-crystal packing diagram of **DPEQPO** reveals that strong IHBs are formed between adjacent molecules with bond lengths in a range of 2.5–2.9 Å and especially diverse bond directions, which further constitute a complicated, disordered and interlaced IHB network (Fig. 1d). In contrast, **DPSQPO** displays grid-like IHB networks with simply two main bond directions and elongated bond lengths of 2.6–3.6 Å. The strong and accumulated IHB interactions markedly shorten the intermolecular distances of the **DPEQPO** matrix. More notably, **DPEQPO** exhibits four different molecular orientations. Each molecule of the **DPEQPO** tetramer contributes one phenyl to form a quasi-*J*-aggregate along the *b* axis with an average distance of 3.5 Å in the inner space (Fig. S1†). Not enough gaps between adjacent molecules actually reflect the remarkable self-





**Fig. 1** Design strategy of host matrices with different weaving modes. (a) Weaving modes of a quadruple system featuring a skeleton substituted with four hydrogen-bond donors. Two plain weave models are borrowed to illustrate the correlation between hydrogen-bond donor arrangements and intermolecular hydrogen-bond (IHB) network formation: (I) four hydrogen-bond donors (red dot, corresponding to knots) randomly disperse on the skeleton, corresponding to the weft, leading to diverse and multidirectional IHB networks; (II) two sets of two hydrogen-bond donors respectively locate on each side of the skeleton, constituting IHB grids but reducing bond directions. (b) Chemical structures of DPXQPO (X = S for DPSQPO and O for DPEQPO), in which skeletons diphenyl sulfide (DPS) and diphenyl ether (DPE) serve as the weft, and P=O acceptors are like knots to form IHBs, corresponding to warps. (c) Single-crystal structures of DPXQPO, in which the dihedral angle and C–X–C angle respectively decrease from 79.9° and 118.7° of DPEQPO to 68.7° and 101.8° of DPSQPO. (d) Packing diagrams of DPXQPO single crystals. Adjacent four molecules are included to show intermolecular interactions. Distances between adjacent molecules are marked with arrows. IHBs are highlighted as blue dashed lines. (e) Atom force microscopy (AFM) images of neat and doped films based on DPXQPO matrices. DPXQPO:30% DMAC-DPS and DPXQPO:30% DMAC-DPS:0.5% 4CzTPNBu are blue and white-emitting films, in which DMAC-DPS and 4CzTPNBu are blue and yellow TADF emitters, respectively.

aggregation tendency of **DPEQPO**. In comparison, owing to the regular intermolecular interactions, only two molecular orientations are observed in the **DPSQPO** packing diagram (Fig. S2†).

Despite the slightly shorter intermolecular distance of **DPSQPO** dimer, the closest centroid–centroid distance between phenyls reaches 4.4 Å, beyond the critical value for  $\pi$ – $\pi$  stacking.



Meanwhile, the dimer–dimer distance in the tetramer is elongated by 0.5 Å. Thus, the **DPSQPO** tetramer constitutes a grid gap unit, which is further connected by a regular IHB network. It is obvious that compared to **DPEQPO** with a dense and intricate packing mode, **DPSQPO** is undoubtedly superior in alleviating local aggregation and improving doping uniformity.

The decomposition temperatures of **DPXQPO** are equally beyond 420 °C, indicating their similar device fabricability by vacuum evaporation (Fig. S3 and Table S1†). But, the melting point ( $T_m$ ) of **DPEQPO** is 46 °C lower than that of **DPSQPO**, since an ordered and uniform molecular arrangement is more effective to improve packing stability than locally enhanced intermolecular interactions. The atom force microscopy (AFM) images of the vacuum evaporated films show that the surface roughness of the neat **DPEQPO** film (0.59 nm) is twice that of the neat **DPSQPO** film (0.25 nm) (Fig. 1e and S4†). The granular patterns further indicate local aggregations of the neat **DPEQPO** film. Singly doped blue-emitting and dually doped white-emitting films were prepared based on a blue TADF emitter bis[4-(9,9-dimethyl-9,10-dihydroacridine)phenyl]sulfone (**DMAC-DPS**) and a yellow TADF emitter 2,3,5,6-tetrakis(3,6-di-*tert*-butyl)carbazol-9-yl)-1,4-dicyanobenzene (**4CzTPNBu**). **DPEQPO**:30% **DMAC-DPS** and **DPEQPO**:30% **DMAC-DPS**:0.5% **4CzTPNBu** films reveal nearly identical roughness and granular patterns, indicating unchanged aggregation. Similarly, **DMAC-DPS** and **4CzTPNBu** doped **DPSQPO** films preserve the good surface morphology, verifying the uniform dopant dispersion in **DPSQPO** grids.

## 2.2 WOLED performance

To figure out the influence of host matrix regularity on EL performance, we fabricated single-EML pure-TADF WOLEDs with a conventional four-layer configuration of ITO|MoO<sub>3</sub> (6 nm)|NPB (50 nm)|**mCP** (5 nm)|**DPXQPO**:*x*% **DMAC-DPS**:*y*% **4CzTPNBu** (20 nm)|**DPXQPO** (5 nm)|BPhen (45 nm)|LiF (1 nm)|Al, in which *N,N'*-bis-(1-naphthalenyl)-*N,N'*-bis-phenyl-(1,1'-biphenyl)-4,4'-diamine (NPB) and 4,7-diphenyl-1,10-phenanthroline (BPhen) were hole and electron transporting layers, and 5-nm thin layers of 1,3-bis(carbazol-9-yl)benzene (**mCP**) and **DPXQPO** were used to block exciton migration, respectively (Fig. 2a). *x* and *y* were respectively tuned in ranges of 20–40 and 0.2–1.5 (Fig. S5–S7†). The singly doped blue OLEDs achieved the best performance at *x* = 30, in which **DPSQPO** based devices realized the maximum  $\eta_{\text{EQE}}$  reaching 18.8%, 1.4 fold that of **DPEQPO** based analogs (Fig. S5 and Table S2†). Simultaneously, at *x* = 30, **DPSQPO** effectively reduced EQE roll-off to 20% at 1000 cd m<sup>-2</sup>, compared to 25% of **DPEQPO** based devices. It is noted that the efficiencies of **DPSQPO**:*x*% **DMAC-DPS** based devices were markedly reduced by increasing *x* from 30 to 40, in contrast to the nearly unchanged efficiencies of **DPEQPO** based analogs. Furthermore, the EL spectra of **DPSQPO** based blue devices were nearly independent on *x*, but the EL emissions of **DPEQPO** based analogs gradually shifted red when increasing *x*. It means that the intermolecular interaction-induced concentration quenching of **DMAC-DPS** can be hardly alleviated in **DPEQPO** based devices, but can be effectively suppressed in the EMLs of **DPSQPO**:*x*% **DMAC-DPS**.

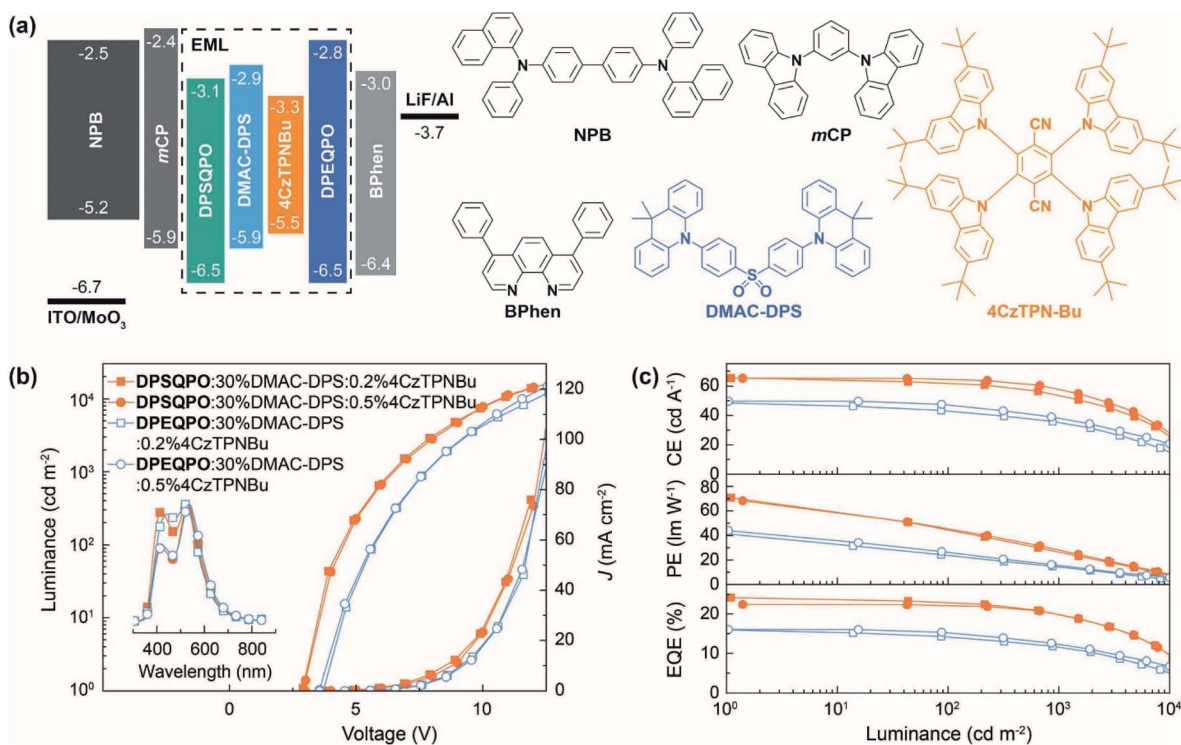


Fig. 2 EL performance of white TADF devices. (a) Structures of single-emissive layer WOLEDs, ITO|MoO<sub>3</sub> (6 nm)|NPB (50 nm)|**mCP** (5 nm)|**DPXQPO**:30 wt% **DMAC-DPS**:*y*% **4CzTPNBu** (20 nm)|**DPXQPO** (5 nm)|BPhen (35 nm)|LiF (1 nm)|Al (*y* = 0.2 for pure-white and 0.5 for warm-white), and chemical structures of the employed materials. (b) Current density (*J*)–voltage–luminance characteristics and EL spectra of the devices. (c) Relationship between the current efficiency (CE), power efficiency (PE) and external quantum efficiency (EQE) and luminance of the WOLEDs.



The EL spectra of all the devices based on dually doped EMLs containing DMAC-DPS and 4CzTPNBu showed blue and yellow components respectively at  $\sim 470$  and  $\sim 550$  nm (Fig. 2b, S6a and S7a†). EL chromaticity was readily tuned by gradually increasing the 4CzTPNBu concentration ( $y$ %), causing emission color change from pure white to warm white, corresponding to Commission Internationale de l'Eclairage (CIE) coordinates and correlated color temperature (CCT) close to standard Illuminants D65 and A (Fig. S6b and S7b†). In the practical brightness range of  $1000$ – $10\,000$   $\text{cd m}^{-2}$ , the EL emissions of these WOLEDs were early unchanged, corresponding to the negligible variations of CIE coordinates ( $x, y$ ) and CCT values were within  $0.02$  for  $x + y$  and  $300$  K, respectively. The excellent chromatic stabilities make these devices competent for diverse lighting applications. It is noteworthy that the relative blue intensities of **DPSQPO** based WOLEDs were higher than those of **DPEQPO** based analogs, reflecting more excitons allocated to 4CzTPNBu in **DPEQPO** hosted EMLs due to enhanced blue-to-yellow energy transfer.

Compared to the corresponding blue diodes, 4CzTPNBu incorporation in **DPEQPO** based EMLs largely increased the brightness of the WOLEDs (Fig. 2b and S6c and Table S2†). The maximum luminance of **DPEQPO** based WOLEDs was directly proportional to  $y$  when  $y \leq 1.0$  and then kept stable at  $y = 1.5$ . Furthermore, codoping 4CzTPNBu hardly influences driving voltages and current densities ( $J$ ). It means that in the EMLs of **DPEQPO**:30% DMAC-DPS: $y$ % 4CzTPNBu, energy transfer from DMAC-DPS to 4CzTPNBu actually alleviated the self-quenching of the former, thereby facilitating exciton radiation. The driving voltages and  $J$  of **DPSQPO** based WOLEDs were also comparable at  $y \leq 0.5$  (Fig. 2b and S7c†). At the critical concentration ( $y = 1.0$ ), direct carrier injection, capture and recombination by 4CzTPNBu made a significant contribution to exciton formation and utilization, leading to markedly reduced driving voltages and increased  $J$  and luminance. So, it implies that 4CzTPNBu molecules in **DPEQPO** hosted films were mostly embedded by DMAC-DPS aggregates; meanwhile, for **DPSQPO**:30% DMAC-DPS: $y$ % 4CzTPNBu, both the dopants are dispersed uniformly and separated from each other in the **DPSQPO** grids.

**DPSQPO**:30% DMAC-DPS:0.2% 4CzTPNBu endowed its diode with the state-of-the-art efficiencies, *e.g.* a maximum current efficiency (CE,  $\eta_{\text{CE}}$ ), power efficiency (PE,  $\eta_{\text{PE}}$ ) and  $\eta_{\text{EQE}}$  of  $65.3$   $\text{cd A}^{-1}$ ,  $70.8$   $\text{lm W}^{-1}$  and  $24.2\%$ , which are among the best results from single-EML pure-TADF WOLEDs (Fig. 2c and S7d and Table S2†). At  $y = 0.5$ , **DPSQPO** still supported the highest efficiencies with the maxima of  $65.2$   $\text{cd A}^{-1}$ ,  $68.2$   $\text{lm W}^{-1}$  and  $22.4\%$  for its WOLED. At  $1000$   $\text{cd m}^{-2}$ , both these two WOLEDs still preserved the favorable efficiencies beyond  $55$   $\text{cd A}^{-1}$ ,  $30$   $\text{lm W}^{-1}$  and  $20\%$ . In contrast, **DPEQPO** based WOLEDs achieved the highest efficiencies at  $y = 1.0$  with the maxima of  $49.7$   $\text{cd A}^{-1}$ ,  $44.0$   $\text{lm W}^{-1}$  and  $16.1\%$ , which rapidly decreased to  $39.0$   $\text{cd A}^{-1}$ ,  $16.2$   $\text{lm W}^{-1}$  and  $12.5\%$  at  $1000$   $\text{cd m}^{-2}$  (Fig. 2c and S6d†). It is noted that compared to the corresponding blue diodes, codoping  $0.2\%$  4CzTPNBu increased the maximum  $\eta_{\text{EQE}}$  by  $29\%$  for the **DPSQPO** hosted WOLED, but had negligible influence on the  $\eta_{\text{EQE}}$  of the **DPEQPO** based device. So, the **DPSQPO** matrix is superior to **DPEQPO**, in terms of improving

the synergism between blue and yellow dopants in exciton utilization.

Furthermore, in accord with the situations of the volt-ampere characteristics for these WOLEDs, the efficiencies of **DPEQPO** hosted WOLEDs were nearly independent on the 4CzTPNBu concentration (Fig. 2c and S6d†). The variation percentage of the maximum  $\eta_{\text{EQE}}$  was within  $11\%$  at  $y = 0.2$ – $1.5$ . It is convincing that codoping 4CzTPNBu in the **DPEQPO** matrix had a positive effect on alleviating triplet concentration quenching on DMAC-DPS by triplet energy transfer, but was roughly counteracted by dopant–dopant quenching. In comparison, the efficiencies of **DPSQPO** hosted WOLEDs at  $y = 0.2$  and  $0.5$  were also nearly identical. However, further increasing  $y$  to  $1.0$  led to a sharp efficiency decrease by more than  $20\%$ . At  $y = 1.5$ , the  $\eta_{\text{EQE}}$  values of both **DPXQPO** hosted WOLEDs were already comparable. Obviously, in the **DPSQPO** matrix, at  $y < 1.0$ , 4CzTPNBu is markedly more effective in quenching alleviation and exciton utilization improvement. But, at  $y \geq 1.0$ , dramatic efficiency reduction revealed that dopant–dopant quenching and competition in exciton allocation became dominant.

The different dependencies of efficiencies on dopant concentrations for **DPXQPO** hosted diodes indicates that **DPSQPO** is much more efficient to prevent the concentration quenching of DMAC-DPS and mutual quenching between DMAC-DPS and 4CzTPNBu; meanwhile, even at a low doping concentration, dopant concentration quenching was already serious in the **DPEQPO** matrix. These results suggest that the device performances of **DPXQPO** are directly correlated with their dispersive abilities. Within a rational range of doping concentration, the **DPSQPO** matrix can uniformly disperse and effectively separate dopants to alleviate mutual interaction induced quenching. In contrast, the **DPEQPO** matrix seemed incapable of preventing intermolecular interactions between dopants. It is consistent with its self-aggregation tendency as shown in the single-crystal packing diagram, which in turn leads to the heterogeneous distributions of dopants and consequently worsened concentration quenching. Furthermore, owing to effective quenching suppression, **DPSQPO** simultaneously endowed its WOLEDs with a doubled device duration time, compared to its DPE based analogs (Fig. S8†).

### 2.3 Theoretical simulation

Because intermolecular interactions markedly influence excited-state properties, it is indispensable to figure out the relationship between the transition characteristics of **DPXQPO** matrices and their different intermolecular interactions. Models of monomers and tetramers for density functional theory (DFT) and time-dependent DFT (TDDFT) simulations are established according to single-crystal data (Fig. 3 and S8†).

The DFT simulation of ground states ( $S_0$ ) shows that in addition to the high electronegativity of ether oxygen atoms, the combined electron-withdrawing effects of the four P=O groups of **DPEQPO** on the DPE skeleton result in the major contribution of DPE to unoccupied molecular orbitals, and the occupied molecular orbitals are mainly localized on the phenyls of



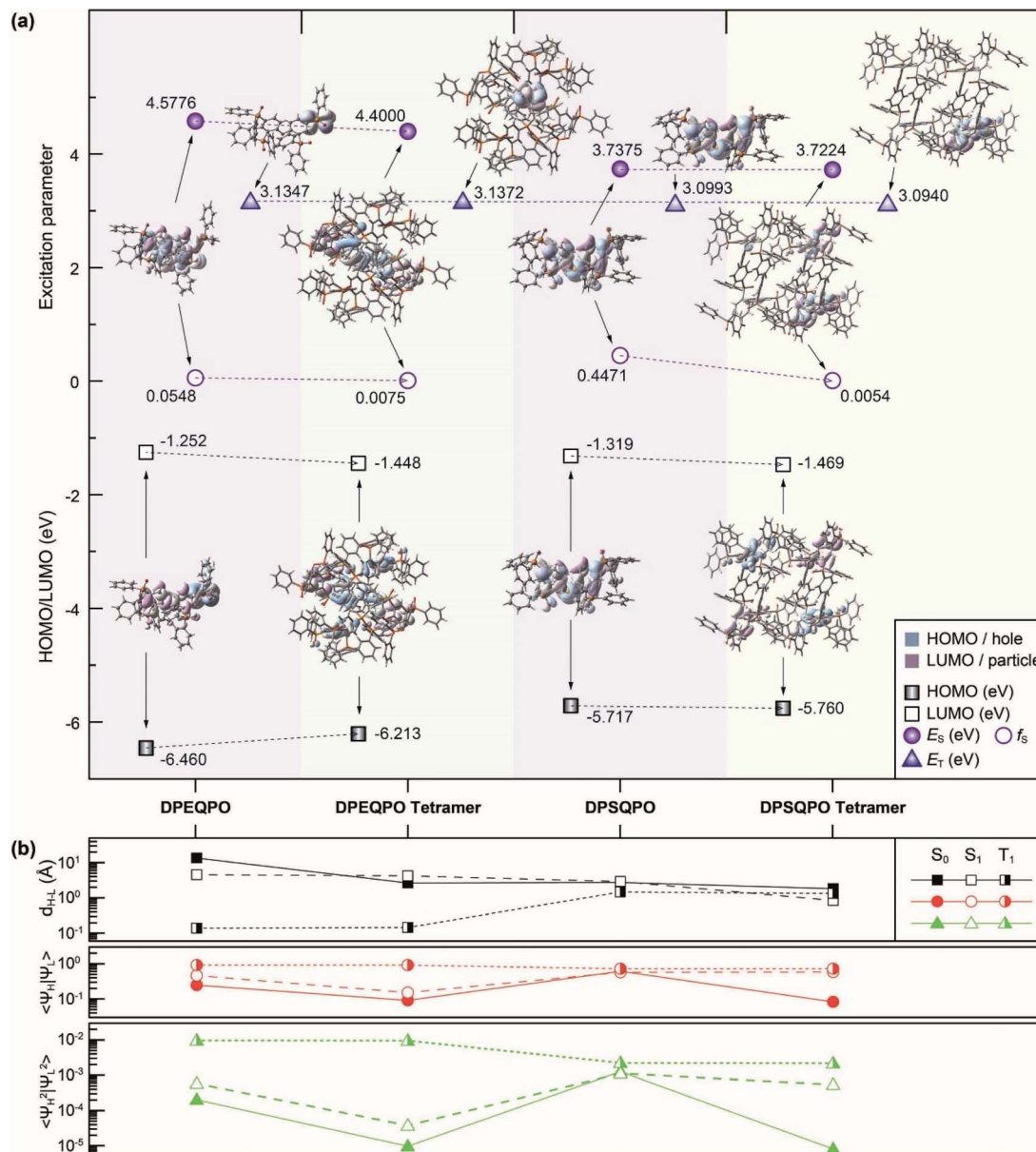


Fig. 3 Variation of ground and excited-state characteristics after host weaving estimated by Gaussian simulations. (a) Contours and energy levels of the Frontier molecular orbitals for ground states (S<sub>0</sub>) and "holes" and "particles" for the first singlet (S<sub>1</sub>) and triplet (T<sub>1</sub>) excited states of DPXQPO single molecules and tetramers. The geometries and packings are generated according to single crystal data. E<sub>S</sub>, E<sub>T</sub> and f<sub>s</sub> refer to the excited energy levels of the S<sub>1</sub> and T<sub>1</sub> states and the singlet oscillator strength. (b) Variation of the centroid distance (d<sub>H-L</sub>), overlap integral (⟨ψ<sub>H</sub>|ψ<sub>L</sub>⟩) of the Frontier orbitals and electronic cloud overlap probabilities (⟨ψ<sub>H</sub><sup>2</sup>|ψ<sub>L</sub><sup>2</sup>⟩) of the Frontier orbitals for the S<sub>0</sub>, S<sub>1</sub> and T<sub>1</sub> states of single molecules and tetramers.

electron-deficient DPPO groups (Fig. 3a and S9<sup>†</sup>). In contrast, the highest occupied (HOMO) and the lowest unoccupied (LUMO) molecular orbitals of **DPSQPO** are mainly contributed by the DPS skeleton, but slightly dispersed to P=O groups, due to the ambipolar characteristics of the sulfur-containing moiety.<sup>57</sup> Cyclic voltammetry (CV) analysis shows that the lowest occupied molecular orbital (LUMO) energy level of **DPSQPO** (−3.11 eV) is 0.28 eV deeper than that of **DPEQPO**, manifesting the stronger electron injecting ability of the former (Fig. S10a<sup>†</sup>). However, the ultraviolet photoelectron spectra (UPS) of **DPXQPO** films show that the ionization potential and electron

affinity of **DPSQPO** are 7.48 and 4.46 eV, which are 0.83 and 0.57 eV lower than those of **DPEQPO** (Fig. S11<sup>†</sup>). Therefore, in solid states, the **DPEQPO** matrix improves electron injection, based on its stronger intermolecular interactions, but its hole injection ability is weakened, due to the embedment of the hole-transporting skeleton. In contrast, the **DPSQPO** matrix provides balanced hole and electron injecting ability (Table S3<sup>†</sup>).

Both **DPXQPO** exhibit locally excited singlet and triplet states. For S<sub>0</sub> → S<sub>1</sub> excitation, the "holes" and "electrons" of **DPXQPO** are localized on their DPE/DPS skeletons. Nonetheless, the calculated first singlet (S<sub>1</sub>) energy level of **DPEQPO**



reaches 4.58 eV, markedly higher than that of **DPSQPO** (3.74 eV). In contrast, the singlet oscillator strength ( $f_s$ ) of **DPSQPO** (0.4471) is one order of magnitude larger than that of **DPEQPO**. Therefore, compared to **DPEQPO**, **DPSQPO** is superior in forming singlet excitons and transferring to the dopant by FRET. In contrast, the first triplet ( $T_1$ ) energy levels of **DPXQPO** ( $\sim 3.1$  eV) are nearly identical. But, the “hole” and “electron” of **DPEQPO** for  $S_0 \rightarrow T_1$  excitation are centralized on one phenyl of its peripheral DPPO group, in contrast to the  $T_1$  location of **DPSQPO** still on the embedded DPS skeleton. The exposed  $T_1$  state of **DPEQPO** is more beneficial to charge-exchange-based triplet DET. So, FRET would be predominant for **DPSQPO**, in contrast to comparable FRET and DET for **DPEQPO**.

The  $S_1$  states of **DPXQPO** tetramers are mainly and equally contributed by two molecules, decreasing  $f_s$  by 1–2 orders of magnitude. Different to the  $S_1$  state of the **DPSQPO** tetramer like the simply duplicated  $S_1$  state of the monomer, the singlet “hole” and “particle” of the **DPEQPO** tetramer are separately distributed on two phenyls and two DPE skeletons, which is similar to the FMO distributions. Thus, the  $S_1$  energy levels of the **DPSQPO** monomer and tetramer are identical, but the **DPEQPO** tetramer reveals an  $S_1$  energy level 0.18 eV lower than that of the monomer. Furthermore, despite similar  $f_s$  values, the high singlet-state similarity of the **DPSQPO** monomer and tetramer makes the mutual transformation between the  $S_1$  states feasible, thereby combining the advantages of high transition probability for exciton formation and uniform dispersion for FRET. Meanwhile, identical to monomers, the  $T_1$  states of **DPXQPO** tetramers are centralized on a single molecule, accompanied by unchanged  $T_1$  energy levels. As a consequence, the respective predominance of **DPXQPO** molecules in FRET and DET is further enhanced in their matrices.

The quantitative analysis of Frontier orbital parameters shows that at the  $S_0$  states, despite the reduced HOMO–LUMO centroid distance ( $d_{H-L}$ ), the FMO wave-function overlap integral ( $\langle \Psi_H | \Psi_L \rangle$ ) and electron-cloud-density overlap integral ( $\langle \Psi_H^2 | \Psi_L^2 \rangle$ ) of the **DPXQPO** tetramer are reduced, in accord with their enhanced FMO separation. However, in contrast to the equal singlet  $d_{H-L}$  of the **DPEQPO** monomer and tetramer, the  $d_{H-L}$  of the **DPSQPO** tetramer is shorter than that of its monomer. Therefore, although the singlet “holes” and “electrons” of **DPXQPO** tetramers are more dispersive than monomers, the  $S_1$  state of the **DPSQPO** tetramer basically preserves the  $\langle \Psi_H | \Psi_L \rangle$  and  $\langle \Psi_H^2 | \Psi_L^2 \rangle$  of its monomer, compared to markedly decreased  $\langle \Psi_H | \Psi_L \rangle$  and  $\langle \Psi_H^2 | \Psi_L^2 \rangle$  of the **DPEQPO** tetramer. Therefore, the **DPSQPO** tetramer can form more stable singlet excitons for facilitating FRET to dopants. In contrast, the triplet-state properties of the monomers and tetramers are almost identical. In this case, the triplet  $d_{H-L}$  of the **DPEQPO** tetramer is shorter than that of the **DPSQPO** tetramer, and the corresponding  $\langle \Psi_H | \Psi_L \rangle$  and  $\langle \Psi_H^2 | \Psi_L^2 \rangle$  of the former are larger than those of the latter. Therefore, DET is dominant in the **DPEQPO** matrix with a more stable  $T_1$  state.

## 2.4 Photophysical investigation

The electronic absorption spectra of **DPXQPO** in dilute  $\text{CH}_2\text{Cl}_2$  solution ( $10^{-6}$  mol  $\text{L}^{-1}$ ) consist of three main bands peaked

around 228, 262 and 310 nm, attributed to the  $\pi \rightarrow \pi^*$  transitions of phenyls and  $n \rightarrow \pi^*$  transitions of P=O and ether/sulfide moieties (Fig. 4a). The absorption tail of **DPSQPO** extends to 410 nm, corresponding to the  $S_1$  energy level of 3.01 eV, which is lower than 3.28 eV of **DPEQPO** (Table S1†). Furthermore, the intensities of  $n \rightarrow \pi^*$  transitions for **DPSQPO** are markedly higher than those of **DPEQPO**. As estimated with Mulliken's relationship,<sup>58</sup> the single-molecular  $f_s$  of **DPSQPO** is 0.2441, 1.5 fold that of **DPEQPO** (0.1594), which is in accord with TDDFT results. Compared to **DPEQPO**, the steady-state photoluminescence (PL) spectrum of the solution of **DPSQPO** peaking at 365 nm is also red shifted by 54 nm. Moreover, the full width at half maximum (FWHM) of FL emission from **DPEQPO** is as small as 33 nm, which is less than half that of **DPSQPO** (68 nm), due to the high molecular rigidity of **DPEQPO** enhanced by intramolecular  $\pi$ – $\pi$  interaction between its *ortho*-DPPO and DPE skeleton, as shown in the single crystal structure (Fig. 1c). The PH spectra of **DPXQPO** in dilute solutions were recorded with time-resolved technology at 77 K (inset of Fig. 4a). As estimated from the first PH peak wavelengths, the  $T_1$  energy levels of **DPXQPO** are beyond 2.9 eV, and are high enough to support positive DET to blue TADF emitters. The  $T_1$  difference between **DPXQPO** is only 0.07 eV, about one fourth of their  $S_1$  difference. Thus, **DPSQPO** simultaneously achieves a comparably high  $T_1$  energy level and improved electrical properties.

It is noteworthy that in addition to the single-molecular main peak at 313 nm, the vacuum-evaporated **DPEQPO** neat film reveals an aggregation-originated shoulder peaked at 370 nm and another long emission tail centered 470 nm, which manifests the heterogeneous distribution of intermolecular interactions and local self-aggregation in the **DPEQPO** matrix (Fig. S12†). In contrast, the **DPSQPO** neat film exhibits a single-peak emission with a smaller redshift of 34 nm, reflecting the homogeneous and weaker intermolecular interactions in the **DPSQPO** matrix. This result suggests the preservation of packing regularity and grid-like uniformity in **DPSQPO** based thin films. Despite aggregation, the  $T_1$  energy levels of **DPXQPO** films are equal to 3.03 eV, and are high enough to support positive energy transfer to DMAC-DPS (Fig. S13†). In addition, despite the small singlet–triplet splitting energy, the time decay curve of the **DPSQPO** film reveals a monoexponential fitting with a lifetime value of  $\sim 2$  ns (Fig. S14†). Therefore, emission from the **DPSQPO** film corresponds to fluorescence rather than TADF.

DMAC-DPS was further doped in the **DPXQPO** matrix by vacuum evaporation. The PL spectra of the resultant **DPXQPO**: $x\%$  DMAC-DPS films ( $x = 10$ – $40$ , with an interval of 10) correspond to typical blue emissions from the dopants (Fig. S15 and S16†). However, it is noted that increasing  $x$  induces the marked emission-peak redshift of **DPEQPO** hosted films by 16 nm from 463 to 479 nm, which is obviously induced by gradually worsened aggregation (Fig. S14a†). Due to aggregation, intermolecular CT and concentration quenching is simultaneously in direct proportion to  $x$ , giving rise to elongated prompt fluorescence (PF) and shortened delayed fluorescence (DF) time decays of **DPEQPO**: $x\%$  DMAC-DPS films (Fig. S15b†). In comparison, the emission redshift of **DPSQPO**: $x\%$  DMAC-



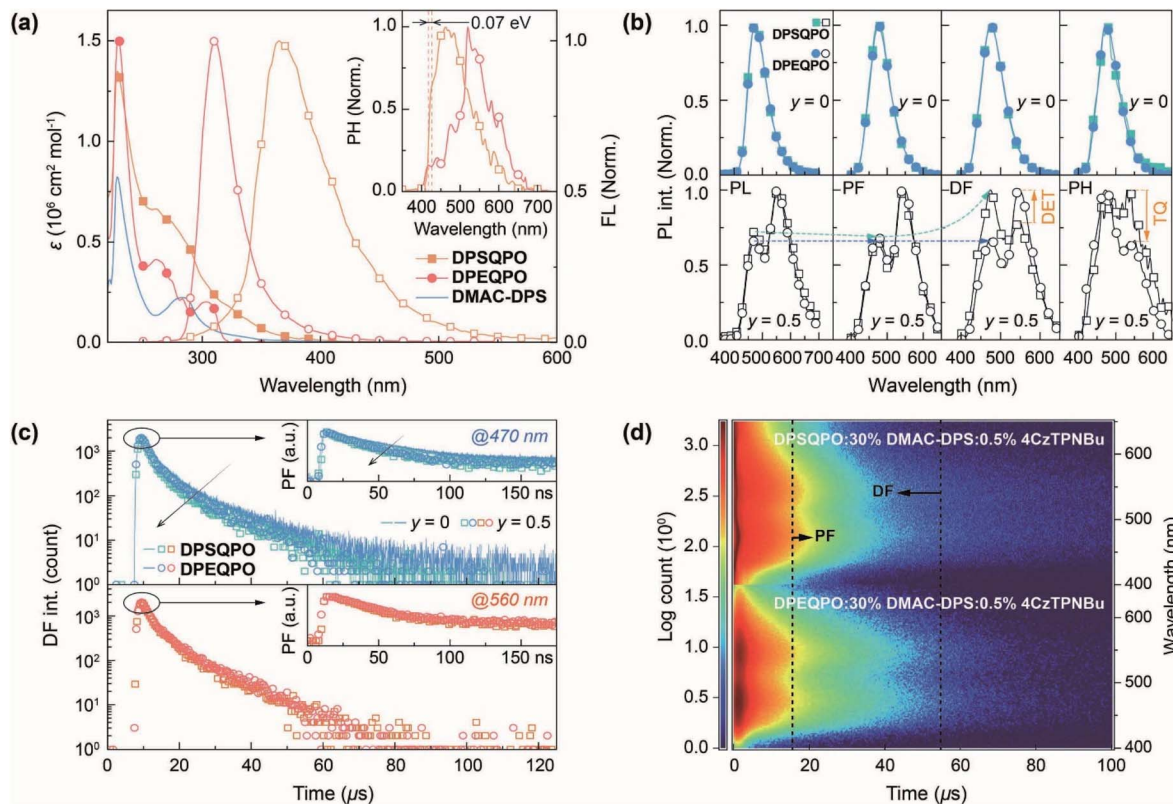


Fig. 4 Photophysical properties of DPXQPO. (a) Electronic absorption and photoluminescence (PL) spectra of DPXQPO and absorption spectrum of DMAC-DPS in dilute  $\text{CH}_2\text{Cl}_2$  solutions ( $10^{-6}$  mol  $\text{L}^{-1}$ ). Fluorescence (FL) spectra were recorded at room temperature. Phosphorescence (PH) spectra were recorded at 77 K with time resolution technology after a delay of 100  $\mu\text{s}$  to exclude delay fluorescence. (b) PL, prompt (PF) ( $\leq 1$   $\mu\text{s}$ ) and delayed fluorescence (DF) (1–100  $\mu\text{s}$ ) and PH ( $> 100$   $\mu\text{s}$ ) spectra of blue and white-emitting films of DPXQPO:30% DMAC-DPS:y% 4CzTPNBu ( $y = 0$  for blue; 0.5 for white). Film thickness is 100 nm. DET and TQ refer to Dexter energy transfer and triplet quenching. (c) Time decay curves of blue and yellow PF and DF components from DPXQPO:30% DMAC-DPS:0.5% 4CzTPNBu films, respectively, recorded at 470 and 560 nm. PF and DF decays of DPXQPO:30% DMAC-DPS are included for comparison. (d) Transient emission contours of DPXQPO:30% DMAC-DPS:0.5% 4CzTPNBu films. PF and DF differences are marked with arrows.

DPS films is limited to 7 nm at  $x = 40$ , accompanied by  $x$ -independent PF and DF time decays (Fig. S16<sup>†</sup>). Therefore, the DPXQPO matrix can indeed effectively alleviate dopant aggregation and concentration quenching. Furthermore, the PF and DF decays of a conventional blue host bis[2-[di(phenyl)phosphino]-phenyl]ether oxide (DPEPO) ( $T_1 > 3.2$  eV) were also measured for comparison (Fig. S17<sup>†</sup>), and are almost identical to those of DPEQPO. This in turn demonstrates the positive energy transfer from DPXQPO to DMAC-DPS.

At  $x = 30$ , the PL, PF, DF and PH spectra of DPXQPO:30% DMAC-DPS films are almost identical; however, the corresponding dually doped DPXQPO:30% DMAC-DPS:y% 4CzTPNBu films ( $y = 0.2$ –1.5) reveal totally different emission spectra (Fig. 4b and S18<sup>†</sup>). At the same  $y$ , the intensity ratios of blue and yellow peaks in the PL spectra of DPXQPO:30% DMAC-DPS:y% 4CzTPNBu films are markedly larger than those for DPEQPO hosted analogs. More importantly, for DPXQPO hosted films, yellow PF components are relatively stronger than yellow DF components; meanwhile, the blue/yellow ratios of PF and DF components for DPEQPO hosted films are nearly the same. A more detailed comparison of the emission spectra of DPXQPO:30% DMAC-DPS:0.5% 4CzTPNBu films indicates that

despite their PL and PF spectra with the same blue/yellow ratios, the DF spectra of the DPXQPO hosted film is remarkably blue-peak dominant, in contrast to the yellow-peak dominant DF emission of the DPEQPO hosted analog (Fig. 4b). In contrast, the relative intensity of the yellow PH component for the DPXQPO hosted film is higher than that for the DPEQPO hosted analog. Compared to DPEQPO:30% DMAC-DPS, codoping 0.5% 4CzTPNBu markedly reduces both the PF and DF lifetimes of DMAC-DPS, due to blue-to-yellow FRET and DET (Fig. 4c). DPXQPO:30% DMAC-DPS:0.5% 4CzTPNBu reveals a similar blue PF lifetime reduction, but its blue DF decay is basically unchanged. Therefore, compared to DPEQPO, the DPXQPO matrix limits triplet DET to 4CzTPNBu, which should shorten the yellow DF lifetime. However, both dually doped white-emitting films display similar yellow PF and DF decay curves. It means that the DPXQPO matrix simultaneously alleviates triplet quenching induced lifetime reduction.

The time-resolved emission spectra (TRES) of DPXQPO:30% DMAC-DPS:0.5% 4CzTPNBu films further indicate that in the DPXQPO matrix, PF emissions are predominant for both blue and yellow components, whose intensities are far beyond DF emissions; meanwhile, the DPEQPO matrix slightly reduces the



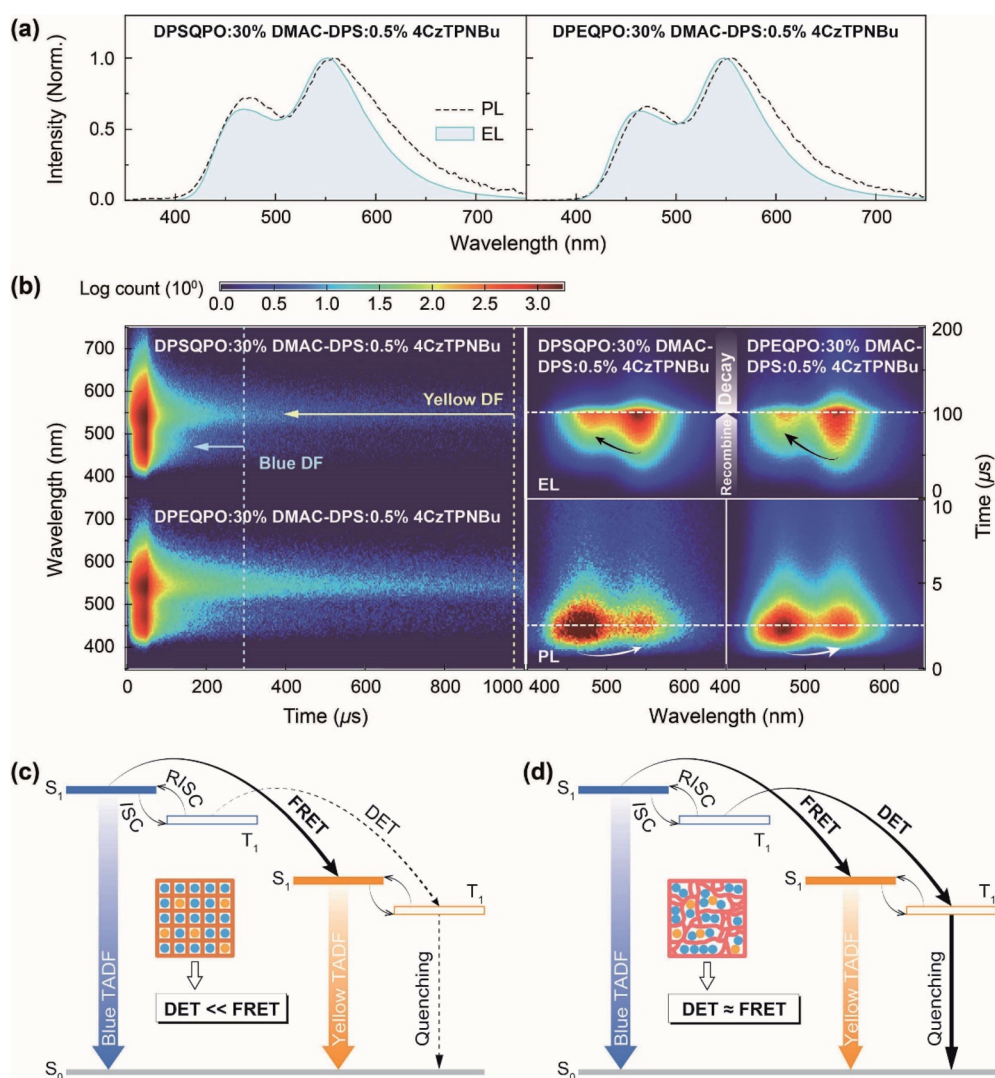
PF proportion, but significantly increases the DF contribution. Considering the short-range feature of the DET mechanism and the collision-inducible character of triplet quenching, the **DPSQPO** matrix effectively separates dopant molecules through grid-like uniform dispersion, thereby simultaneously suppressing intermolecular-interaction based triplet energy transfer and concentration quenching. In contrast, the heterogeneous distribution of DMAC-DPS in the **DPEQPO** matrix increases the probability of doping 4CzTPNBu directly into or close to DMAC-DPS aggregates, thus enhancing DET and worsening triplet quenching.

## 2.5 Exciton kinetics

The steady-state PL and EL spectra of **DPEQPO:30% DMAC-DPS:0.5% 4CzTPNBu** are nearly overlapped, manifesting the absolute predominance of energy transfer in the exciton

allocation of **DPEQPO** hosted devices (Fig. 5a). However, the blue EL component of **DPSQPO:30% DMAC-DPS:0.5% 4CzTPNBu** is weaker than its blue PL component, indicating the considerable incorporation of direct charge capture and recombination by 4CzTPNBu into exciton allocation. It is rational that in the **DPEQPO** matrix, directly interacting DMAC-DPS and 4CzTPNBu both with ambipolar characteristics mutually compete in terms of charge capture. Therefore, the effective separation and uniform dispersion of DMAC-DPS and 4CzTPNBu in the unipolar **DPSQPO** matrix amplify the advantage of 4CzTPNBu with the deepest LUMO and the shallowest HOMO in direct charge capture.

Transient EL emission spectra further show that compared to the **DPEQPO:30% DMAC-DPS:0.5% 4CzTPNBu** based device, the **DPSQPO** matrix simultaneously and dramatically shortened the blue and yellow emission decays of its WOLED by about 100



**Fig. 5** Exciton allocation in DPXQPO based WOLEDs. (a) Comparison of the PL and EL spectra of DPXQPO:30% DMAC-DPS:0.5% 4CzTPNBu systems. (b) EL transient emission contours of DPXQPO:30% DMAC-DPS:0.5% 4CzTPNBu based devices (left) and the comparison of exciton formation and transfer in PL and EL processes. (c and d) Exciton allocation mechanisms of binary complementary white-emitting systems in a uniformly weaved matrix like **DPSQPO** (c) and a locally aggregated matrix like **DPEQPO** (d) regarding energy transfer. FRET and DET refer to Förster resonance energy transfer and Dexter energy transfer. ISC and RISC are intersystem crossing and reverse ISC.



and 700  $\mu\text{s}$ , respectively (Fig. 5b left). Obviously, the **DPSQPO** matrix effectively accelerated exciton formation and radiation, which is beneficial to quenching suppression. Furthermore, compared to the blue PL decay (Fig. 4d), the EL decay of the blue component for **DPSQPO**:30% DMAC-DPS:0.5% 4CzTPNBu was slightly elongated by 10–20  $\mu\text{s}$ , due to delays for carrier injection, transportation and recombination. In contrast, the time decay of its yellow EL DF was largely increased by 200  $\mu\text{s}$ , further reflecting the predominance of charge transfer and capture in triplet allocation to 4CzTPNBu rather than DET. In contrast, compared to PL decays, the blue and yellow EL components of **DPEQPO**:30% DMAC-DPS:0.5% 4CzTPNBu were markedly elongated by 200 and 900  $\mu\text{s}$ , respectively. Obviously, DMAC-DPS aggregation in the **DPEQPO** matrix enhanced intermolecular charge transfer, which significantly extended carrier recombination and blue EL time decay. Furthermore, effective DET from DMAC-DPS markedly enhanced triplet allocation to 4CzTPNBu, further increasing the yellow DF EL proportion and elongating the yellow EL decay.

Different to simultaneous photoexcitation processes in **DPXQPO** hosted films, the EL kinetic process shows that carrier recombination in the **DPEQPO** based device occurred earlier than that in the **DPSQPO** based analog (Fig. 5b right). It is because TADF molecules with low concentrations are uniformly dispersed, carriers mainly transport in inertia **DPSQPO** matrix, which postponed carrier capture and recombination by TADF emitters; meanwhile, in the **DPEQPO** matrix, dopant aggregation actually increased the probability of exciton formation. For both **DPXQPO** hosted EMLs, exciton formation is completely determined by the statistical advantage for photoexcitation at the first, but then carrier capture ability for electro-excitation. Therefore, excitons were firstly generated on DMAC-DPS with a high concentration in the PL process, and then on 4CzTPNBu due to its advantageous FMOs in the EL process. Although exciton formation on DMAC-DPS in the **DPEQPO** based device was also earlier than that in the **DPSQPO** based analog, in accord with the PL process, the blue emission intensity of the former was always markedly lower than that of the latter, manifesting the synchronism of exciton formation and blue-to-yellow energy transfer in the **DPEQPO** based device. Compared to long-distance FRET, short-range DET is more dependent on high exciton concentration; therefore, markedly higher blue exciton concentration in the **DPSQPO** based WOLED suggests limited incorporation of DET in exciton allocation, compared to the **DPEQPO** based device. Therefore, carrier injection in EMLs would still be mainly supported by **DPXQPO**, rather than DMAC-DPS or 4CzTPNBu. Exciton formation through either energy transfer or carrier capture occurs during the carrier transportation process. In this sense, the driving voltages of the devices should be in accord with the FMO energy levels of **DPXQPO**, leading to the lower driving voltages of **DPSQPO** based devices.

So, as illustrated in Fig. 5c and d, the **DPSQPO** matrix provides high grid regularity to uniformly disperse and separate blue and yellow dopants, inducing FRET-predominant exciton allocation. The suppression of dopant–dopant interactions not only alleviates exciton quenching, but also avoids excessive

exciton allocation to yellow dopants by DET. In contrast, the local aggregations of the **DPEQPO** matrix and TADF dopants enhance intermolecular charge transfer and blue–yellow interactions, accelerating DET and worsening mutual quenching. Therefore, rationalizing the intermolecular interaction network to weave host grids is effective for optimizing the exciton kinetic process in high-efficiency single-layer white TADF systems.

### 3. Conclusions

We demonstrate the significance of host grid regularity with two similar host molecules **DPEQPO** and **DPSQPO**, whose IHB networks are differentiated by their DPE and DPS skeleton induced different orientations of P=O hydrogen-bond donors. Compared to **DPEQPO** with a complicated IHB network-induced heterogeneous dispersion, **DPSQPO** featuring highly regular IHB grids supported a state-of-the-art  $\eta_{\text{EQE}}$  up to 24.2% for its single-EML pure-white TADF diode, which is 1.5 fold the maximum  $\eta_{\text{EQE}}$  for **DPEQPO** based analogs. Both steady-state and time-resolved PL and EL spectra indicate different exciton allocation and kinetic processes in **DPXQPO** hosted films and WOLEDs. Compared to the **DPEQPO** matrix, **DPSQPO** uniformly disperses and effectively separates TADF dopants, and therefore preserves more blue DF components for its white TADF systems by limiting blue-to-yellow DET. FRET-predominant exciton allocation in the **DPSQPO** matrix not only shortens emission decays and restrains dopant–dopant interactions for quenching suppression, but also improves white emission purity. These results suggest that host grid regularization is a feasible strategy for optimizing exciton kinetics, establishing a fundamental basis for developing high-performance TADF WOLEDs for daily lighting applications.

### Data availability

All the data are available from the authors.

### Author contributions

H. X. and J. Z. conceived the project. Y. T., H. W., Y. M., N. Z., and Y. L. performed the experiments. H. X., Y. T., H. W. and J. Z. analyzed the data and wrote the paper. All authors commented on the manuscript.

### Conflicts of interest

There are no conflicts to declare.

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## Notes and references

- J.-i. Nishide, H. Nakanotani, Y. Hiraga and C. Adachi, *Appl. Phys. Lett.*, 2014, **104**, 233304.
- B. S. Kim, K. S. Yook and J. Y. Lee, *Sci. Rep.*, 2014, **4**, 6019.
- S. Y. Lee, T. Yasuda, Y. S. Yang, Q. Zhang and C. Adachi, *Angew. Chem., Int. Ed.*, 2014, **53**, 6402–6406.
- Z. Yang, Z. Mao, Z. Xie, Y. Zhang, S. Liu, J. Zhao, J. Xu, Z. Chi and M. P. Aldred, *Chem. Soc. Rev.*, 2017, **46**, 915–1016.
- M. Y. Wong and E. Zysman-Colman, *Adv. Mater.*, 2017, **29**, 160544.
- Y. Tao, K. Yuan, T. Chen, P. Xu, H. Li, R. Chen, C. Zheng, L. Zhang and W. Huang, *Adv. Mater.*, 2014, **26**, 7931–7958.
- H. Uoyama, K. Goushi, K. Shizu, H. Nomura and C. Adachi, *Nature*, 2012, **492**, 234–238.
- S. Huang, Q. Zhang, Y. Shiota, T. Nakagawa, K. Kuwabara, K. Yoshizawa and C. Adachi, *J. Chem. Theory Comput.*, 2013, **9**, 3872–3877.
- Y. Liu, C. Li, Z. Ren, S. Yan and M. R. Bryce, *Nat. Rev. Mater.*, 2018, **3**, 18020.
- T. J. Penfold, F. B. Dias and A. P. Monkman, *Chem. Commun.*, 2018, **54**, 3926–3935.
- T.-A. Lin, T. Chatterjee, W.-L. Tsai, W.-K. Lee, M.-J. Wu, M. Jiao, K.-C. Pan, C.-L. Yi, C.-L. Chung, K.-T. Wong and C.-C. Wu, *Adv. Mater.*, 2016, **28**, 6976–6983.
- T.-L. Wu, M.-J. Huang, C.-C. Lin, P.-Y. Huang, T.-Y. Chou, R.-W. Chen-Cheng, H.-W. Lin, R.-S. Liu and C.-H. Cheng, *Nat. Photonics*, 2018, **12**, 235–240.
- H. Kaji, H. Suzuki, T. Fukushima, K. Shizu, K. Suzuki, S. Kubo, T. Komino, H. Oiwa, F. Suzuki, A. Wakamiya, Y. Murata and C. Adachi, *Nat. Commun.*, 2015, **6**, 8476.
- W. Zeng, H.-Y. Lai, W.-K. Lee, M. Jiao, Y.-J. Shiu, C. Zhong, S. Gong, T. Zhou, G. Xie, M. Sarma, K.-T. Wong, C.-C. Wu and C. Yang, *Adv. Mater.*, 2018, **30**, 1704961.
- Y. Xiang, P. Li, S. Gong, Y.-H. Huang, C.-Y. Wang, C. Zhong, W. Zeng, Z. Chen, W.-K. Lee, X. Yin, C.-C. Wu and C. Yang, *Sci. Adv.*, 2020, **6**, eaba7855.
- X. Tang, L.-S. Cui, H.-C. Li, A. J. Gillett, F. Auras, Y.-K. Qu, C. Zhong, S. T. E. Jones, Z.-Q. Jiang, R. H. Friend and L.-S. Liao, *Nat. Mater.*, 2020, **19**, 1332–1338.
- Y.-L. Zhang, Q. Ran, Q. Wang, Y. Liu, C. Hänisch, S. Reineke, J. Fan and L.-S. Liao, *Adv. Mater.*, 2019, **31**, 1902368.
- B. Zhao, H. Wang, C. Han, P. Ma, Z. Li, P. Chang and H. Xu, *Angew. Chem., Int. Ed.*, 2020, **59**, 19042–19047.
- Y. Li, X.-L. Li, D. Chen, X. Cai, G. Xie, Z. He, Y.-C. Wu, A. Lien, Y. Cao and S.-J. Su, *Adv. Funct. Mater.*, 2016, **26**, 6904–6912.
- J. Li, D. Ding, Y. Tao, Y. Wei, R. Chen, L. Xie, W. Huang and H. Xu, *Adv. Mater.*, 2016, **28**, 3122–3130.
- X. Lv, W. Zhang, D. Ding, C. Han, Z. Huang, S. Xiang, Q. Zhang, H. Xu and L. Wang, *Adv. Opt. Mater.*, 2018, **6**, 1800165.
- Z. Wu, Y. Liu, L. Yu, C. Zhao, D. Yang, X. Qiao, J. Chen, C. Yang, H. Kleemann, K. Leo and D. Ma, *Nat. Commun.*, 2019, **10**, 2380.
- J. Rao, X. Liu, X. Li, L. Yang, L. Zhao, S. Wang, J. Ding and L. Wang, *Angew. Chem., Int. Ed.*, 2020, **59**, 1320–1326.
- J. Zeng, J. Guo, H. Liu, Z. Zhao and B. Z. Tang, *Adv. Funct. Mater.*, 2020, **30**, 2000019.
- C. Zhang, Y. Lu, Z. Liu, Y. Zhang, X. Wang, D. Zhang and L. Duan, *Adv. Mater.*, 2020, **32**, 2004040.
- W. Li, B. Li, X. Cai, L. Gan, Z. Xu, W. Li, K. Liu, D. Chen and S.-J. Su, *Angew. Chem., Int. Ed.*, 2019, **58**, 11301–11305.
- R. Du, C. Duan, Y. Li, J. Zhang, C. Han and H. Xu, *Chem. Eng. J.*, 2019, **374**, 471–478.
- F. Gao, R. Du, Y. Wei and H. Xu, *Opt. Lett.*, 2019, **44**, 5727–5730.
- R. Du, C. Duan, Y. Li, J. Zhang, C. Han and H. Xu, *Chem. Eng. J.*, 2020, **392**, 124870.
- C. Han, W. Yang and H. Xu, *Chem. Eng. J.*, 2020, **401**, 126049.
- J. Sun, J. Zhang, Q. Liang, Y. Wei, C. Duan, C. Han and H. Xu, *Adv. Funct. Mater.*, 2020, **30**, 1908568.
- J. Zhang, C. Han, F. Du, C. Duan, Y. Wei and H. Xu, *Adv. Funct. Mater.*, 2020, **30**, 2005165.
- Y. Li, Z. Li, J. Zhang, C. Han, C. Duan and H. Xu, *Adv. Funct. Mater.*, 2021, 2011169.
- C. Han, Z. Zhang, D. Ding and H. Xu, *Chem*, 2018, **4**, 2154–2167.
- Z. Zhang, D. Ding, Y. Wei, J. Zhang, C. Han and H. Xu, *Chem. Eng. J.*, 2020, **382**, 122485.
- Q. Wang and D. Ma, *Chem. Soc. Rev.*, 2010, **39**, 2387–2398.
- L. Gan, Z. Xu, Z. Wang, B. Li, W. Li, X. Cai, K. Liu, Q. Liang and S.-J. Su, *Adv. Funct. Mater.*, 2019, **29**, 1808088.
- H. Nakanotani, T. Furukawa, K. Morimoto and C. Adachi, *Sci. Adv.*, 2016, **2**, e1501470.
- Y. R. Sun, N. C. Giebink, H. Kanno, B. W. Ma, M. E. Thompson and S. R. Forrest, *Nature*, 2006, **440**, 908–912.
- S. Reineke, K. Walzer and K. Leo, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2007, **75**, 125328.
- K. Masui, H. Nakanotani and C. Adachi, *Org. Electron.*, 2013, **14**, 2721–2726.
- X. Tang, Y. Li, Y.-K. Qu, C.-C. Peng, A. Khan, Z.-Q. Jiang and L.-S. Liao, *Adv. Funct. Mater.*, 2020, **30**, 1910633.
- D. Ding, Z. Wang, C. Li, J. Zhang, C. Duan, Y. Wei and H. Xu, *Adv. Mater.*, 2020, **32**, 1906950.
- P. Wei, D. Zhang and L. Duan, *Adv. Funct. Mater.*, 2019, **29**, 1907083.
- C. Han, R. Du, H. Xu, S. Han, P. Ma, J. Bian, C. Duan, Y. Wei, M. Sun, X. Liu and W. Huang, *Nat. Commun.*, 2021, **12**, 3640.
- T. Förster, *Ann. Phys.*, 1948, **437**, 55–75.
- D. L. Dexter, *J. Chem. Phys.*, 1953, **21**, 836–850.
- J. Kalinowski, W. Stampor, J. Mecedilzdotytk, M. Cocchi, D. Virgili, V. Fattori and P. Di Marco, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2002, **66**, 235321.
- G. X. Yifan Li, S. Gong, K. Wu and C. Yang, *Chem. Sci.*, 2016, **7**, 5441–5447.
- X.-L. Chen, J.-H. Jia, R. Yu, J.-Z. Liao, M.-X. Yang and C.-Z. Lu, *Angew. Chem., Int. Ed.*, 2017, **56**, 15006–15009.
- G. Xie, J. Luo, M. Huang, T. Chen, K. Wu, S. Gong and C. Yang, *Adv. Mater.*, 2017, **29**, 1604223.



- 52 S. Shao, J. Hu, X. Wang, L. Wang, X. Jing and F. Wang, *J. Am. Chem. Soc.*, 2017, **139**, 17739–17742.
- 53 H. Yang, Q. Liang, C. Han, J. Zhang and H. Xu, *Adv. Mater.*, 2017, **29**, 1700553.
- 54 F. Gao, R. Du, F. Jiao, G. Lu, J. Zhang, C. Han and H. Xu, *Adv. Opt. Mater.*, 2020, **8**, 2000052.
- 55 Y. Shi, K. Wang, Y. Tsuchiya, W. Liu, T. Komino, X. Fan, D. Sun, G. Dai, J. Chen, M. Zhang, C. Zheng, S. Xiong, X. Ou, J. Yu, J. Jie, C.-S. Lee, C. Adachi and X. Zhang, *Mater. Horiz.*, 2020, **7**, 2734–2740.
- 56 Y.-Z. Shi, K. Wang, X.-C. Fan, J.-X. Chen, X.-M. Ou, J. Yu, J.-S. Jie, C.-S. Lee and X.-H. Zhang, *Adv. Opt. Mater.*, 2021, **9**, 2100461.
- 57 S.-C. Dong, C.-H. Gao, X.-D. Yuan, L.-S. Cui, Z.-Q. Jiang, S.-T. Lee and L.-S. Liao, *Org. Electron.*, 2013, **14**, 902–908.
- 58 R. S. Mulliken, *J. Chem. Phys.*, 1939, **7**, 14–20.

