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Molecular design of DBA-type five-membered heterocyclic rings to achieve 200% exciton utilization for electroluminescence[†]

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Achieving high exciton utilization is a long-cherished goal in the development of organic light-emitting diode materials. Herein, a three-step mechanism is proposed to achieve 200% exciton utilization: (i) hot triplet exciton (T₂) conversion to singlet S₁; (ii) singlet fission from S₁ into two T₁; (iii) and then a Dexter energy transfer to phosphors. The requirement is that S₁ should lie slightly lower than or close to T₂ and twice as high as T₁ in energy. For this, a scenario is put forward to design a series of donor-bridge-acceptor (DBA) type molecules with $2E(T_1) \leq E(S_1) < E(T_2)$, in which the Baird-type aromatic pyrazoline ring is used as a bridge owing to its stabilized T₁ (1.30–1.74 eV) and different kinds of donors and acceptors are linked to the bridge for regulating S₁ (2.35–3.87 eV) and T₂ (2.44–3.96 eV). The ultrafast spectroscopy and sensitization measurements for one compound (TPA-DBPrz) fully confirm the theoretical predictions.

Introduction

There has been a longstanding challenge to beat a 25% spin statistical limit¹ for organic electroluminescence devices.^{2–5} Such a limit is exceeded by organometallic molecules with emission from the triplet state whose exciton utilization could reach 100% with the aid of the strong spin-orbit coupling of heavy metals.^{6,7} The singlet formation ratio for organic

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^e Institute of chemistry, Chinese Academy of Sciences, Beijing, 100049, P. R. China ^f School of Chemical Engineering, Dalian University of Technology, 2 Ling Gong The maximum exciton utilization for conventional electroluminescence, such as electrophosphorescence and TADF, is limited to 100%. Here, we suggest a novel photophysical scenario to achieve 200% exciton utilization, namely, we theoretically design a host-guest molecular material system where the host molecule possesses an excited state structure as $E(T_2) > E(S_1) \ge 2E(T_1)$ so that both reverse intersystem crossing $(T_2 \rightarrow S_1)$ and singlet fission $(S_1 \rightarrow T_1 + T_1)$ can occur and the resulting 200% T_1 can efficiently transfer to the guest molecule, which is an efficient triplet emitter. In this way, for one electrically pumped carrier, we can eventually obtain two photons. We further suggest a plausible molecular structure for the host by virtue of Baird's aromaticity based on a five-membered heterocyclic ring as a bridge (B) to link different donors (D) and acceptors (A) with the aid of computational chemistry to search the peculiar excited structure.

polymers is also demonstrated to exceed 25%⁸ because of the spin-dependent exciton formation rates with $k_{\rm s} > k_{\rm T}$.^{1,9,10} Triplet-triplet annihilation (TTA) which converts electropumped triplets to singlets via $T_1 + T_1 \rightarrow S_1 + S_0$ could realize 62.5% exciton in the theoretical limit.¹¹ Recently, a thermally activated delayed fluorescence (TADF) mechanism is proposed to achieve 100% by upconverting triplets to singlets via reverse intersystem crossing (RISC).^{12,13} Besides, the hot exciton mechanism¹⁴ is made to not only promote the exciton utilization by the RISC between higher-lying triplets and singlets, but also to allow large oscillator strength in the singlet state.^{15,16} Very recently, π -radical compounds with spin-half are proposed to be promising organic light-emitting diode (OLED) emitters to break the 25% limit.¹⁷ In addition, singlet fission (SF) process in the host materials is used to achieve 100.8% exciton utilization in OLEDs,¹⁸ in which the theoretical limit is 125%. Here, we propose a new mechanism by combining the RISC of the hot triplet exciton (HE) T_2 to S_1 and the SF $S_1 + S_0 \rightarrow T_1 + T_1$ in the same host material (HE-SF). Overall, the exciton utilization can reach up to 200% in principle (Fig. 1): the RISC process converts one electro-pumped hot triplet exciton (T_2) to one singlet exciton (S_1) , and then the SF process turns this S_1 into

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Fig. 1 Mechanism toward achieving 200% exciton utilization theoretically in OLEDs (DET: Dexter energy transfer).

two triplet excitons (T₁), and then Dexter energy transfer occurs from the host material to the guest material, and finally the guest material emits light. The efficient occurrence of RISC needs the S₁ to be slightly lower than or close to T₂ in energy $(E(T_2) \ge E(S_1))$, and the T₂ to be far larger than T₁ in energy (large $\Delta E(T_1T_2)$) to suppress the nonradiative rate from T₂ to T₁ and an efficient SF process basically requires the energy of S₁ to be larger than twice that of T₁, $E(S_1) \ge 2E(T_1)$.^{19,20} However, it is a formidable challenge to design molecules with large $\Delta E(T_1T_2)$ or $\Delta E(S_1T_1)$, let alone both. Our work here aims to search for some building blocks in five-membered heterocyclic rings with $E(T_2) \ge E(S_1) \ge 2E(T_1)$ by manipulating the aromaticity for excellent HE-SF materials.

In this work, we first demonstrate that partially conjugated five-membered heterocyclic rings (*i.e.*, pyrazoline) naturally

have low $E(T_1)$ owing to high aromaticity in T_1 obeying Baird's rule,^{21,22} and high $E(S_1)$ and $E(T_2)$ due to the energetically wellseparated frontier orbitals. A series of donor-bridge-acceptor (DBA) systems are then designed based on the pyrazoline ring as a bridge *via* introducing different donor or acceptor moieties (see Scheme 1). For the constructed pyrazoline derivatives, the low $E(T_1)$ is consistently maintained because the transition density of T_1 is still localized on the pyrazoline ring and the high $E(S_1)$ and $E(T_2)$ are generated *via* transition involving the substituted donors or acceptors, which results in $E(T_2) \ge E(S_1) \ge$ $2E(T_1)$. With that, a novel series of HE-SF molecules are predicted through theoretical calculations and proved by the transient absorption spectroscopy spectra of TPA-DBPrz, which, to the best of our knowledge, has not been previously reported.

Results and discussion

Five-membered heterocyclic rings with large triplet-triplet and singlet-triplet gaps

According to Baird's rule, the aromaticity is always changed upon triplet excitation and the aromatic triplet states are more stable with lower energy than antiaromatic ones.^{22–24} It is hence expected to be a good way to find organic systems with large triplet–triplet and singlet–triplet gaps *via* changing their aromaticity. A series of five-membered rings are chosen, including cyclopentene, 3-pyrroline, pyrazoline, pyrazole, imidazole and pyrrole (Scheme 1), which are frequently used as fundamental units in functional molecules. The geometrical structures of



Scheme 1 Chemical structures of five-membered rings and constructed DBA-type molecules in this work (red: donor; blue: acceptor; black: bridge).



Fig. 2 (a) Optimized bond lengths (inside: S_0 -geometry; outside: T_1 -geometry). (b) Natural transition orbitals (NTOs) of T_1 at T_1 -geometry. (c) AICD plots in the T_1 state where the induced current density vectors are denoted by the arrows for six five-membered rings.

these five-numbered rings are optimized *via* SA4-CASSCF/cc-PVTZ with active space (10e, 10o) and the energies and properties of low-lying excited states are calculated *via* MS4-CASPT2/ cc-PVTZ, as given in Fig. 2, 3 and Table 1. Upon excitation, the double bonds are elongated to a greater extent in cyclopentene (0.130 Å), 3-pyrroline (0.158 Å) and pyrazoline (0.184 Å) than those (*ca.* 0.100 Å) in the other three rings (Fig. 2a), which would largely weaken the coupling between the p-orbitals of two atoms, decreasing the degree of molecular conjugation (Fig. 2b). As seen in Table 1, the energy condition of $2E(T_1) \leq E(S_1) < E(T_2)$ is met naturally in the three ring cyclopentene,



Fig. 3 Frontier orbital energies and the involved transitions for the low-lying states T_1 (orange), S_1 (purple) and T_2 (green).

 $\label{eq:Table 1} \mbox{ Table 1 Vertical excitation energies (unit: eV) at their T_1-geometries of six five-numbered rings$

Compound	$E(T_1)$	$E(T_2)$	$E(S_1)$	$2E(T_1)$	$\Delta E(T_1T_2)$
Cyclopentene	1.01	6.04	3.77	2.02	5.03
3-Pyrroline	1.26	4.09	3.56	2.52	2.83
Pyrazoline	1.54	4.68	3.55	3.08	3.14
Pyrazole	2.84	4.43	4.97	5.68	1.59
Imidazole	3.44	5.10	5.44	6.88	1.66
Pyrrole	3.35	4.84	5.34	6.70	1.49

3-pyrroline and pyrazoline moieties, while the other three rings possess high $E(T_1)$ and relatively small $\Delta E(T_1T_2)$. The anisotropy of the induced current density²⁵ (AICD) plotted in Fig. 2c indicates that cyclopentene, 3-pyrroline and pyrazoline rings in T₁ are aromatic with the clockwise ring currents, while the other three rings are antiaromatic in T₁ with anticlockwise ring currents. It is easily understood that pyrazoline (π_3^4) is a [4n] electron system and it is typically aromatic in T₁. Thus, cyclopentene, 3-pyrroline and pyrazoline retain stabilized T_1 states with low energy as seen in Table 1. Furthermore, the frontier orbitals exhibit energetically well-separated highest occupied molecular orbital (HOMO) and HOMO-n ($n \ge 1$) orbitals in cyclopentene, 3-pyrroline and pyrazoline (Fig. 3), which lead to large $\Delta E(T_1T_2)$ and $\Delta E(S_1T_2)$ because the T_1 states are dominated by the transition from the HOMO to lowest unoccupied molecular orbital (LUMO), while the S1 and T2 states stem from the transition from deeper occupied orbitals to the LUMO or higher unoccupied orbitals. All these results indicate that cyclopentene, 3-pyrroline and pyrazoline with aromatic triplets would be good triplet-stabilized candidates to satisfy $2E(T_1) \leq$ $E(S_1) < E(T_2)$ for the HE-FS materials. A similar conclusion can be obtained by the TDA-DFT approach as shown in the ESI.[†] Moreover, the extended pyrazoline derivatives (pyrazoline-a, pyrazoline-b and pyrazoline-c in Scheme 1) also exhibit the same feature of energy gap among the low-lying excited states (Table S3, ESI⁺), because their T₁ states mainly originate from the pyrazoline ring, while T₂ and S₁ are significantly determined by the introduced chemical groups (Fig. S1a, ESI[†]). These further prove that the partially conjugated rings with aromatic T₁ can serve as triplet-stabilized moieties for the HE-FS materials.

Construction of pyrazoline derivatives with large triplet-triplet and singlet-triplet gaps

In principle, cyclopentene, 3-pyrroline and pyrazoline can be used to be the bridge in the DBA compounds. Pyrazoline derivatives have been investigated as hole-transport materials,²⁶ fluorescent sensors²⁷ and medicines²⁸ owing to high stability and wonderful photophysical properties. The pyrazoline ring hence is chosen here as a representative example to be a bridge (B) to construct 25 compounds by introducing different donor (D) and acceptor (A) groups, namely, DBA type compounds (see Scheme 1). Among them, the electron-withdrawing ability of the acceptors becomes stronger from FB to Py to BN owing to the lower LUMO energies and the electron-donating ability of the donors becomes stronger from TP to Cz, PTZ and PZ due to the higher HOMO energies (Table S4, ESI[†]).

Compounds **1–9** with different D or A at the same substituted 1- and 3-positions all possess nearly coplanar conformations with high rigidity. Upon excitation, significant modifications occur on the three bond lengths, including C = N (L₁), B-R₂ (L₂) and B-R₁ (L₃) (Scheme 1), rather than angles and dihedral angles for the low-lying excited states (Table S5, ESI†). From S₀ to T₁, L₁ is considerably elongated by 0.081–0.136 Å, which is similar to the change in a single pyrazoline ring, while L₂ and L₃ are slightly shortened by 0.002–0.070 Å. As expected, the S₁, T₁ and T₂ exhibit quite different transition properties *via* analysing



are divided into three parts (B, A and D) that denote bridge, acceptor and donor moieties, respectively. The percentage of the distribution is reflected by different colours.

their hole-electron distribution heat maps²⁹ (Fig. 4) and natural transition orbitals (NTOs) (Fig. S2, ESI[†]) for these molecules. T₁ is a locally excited (LE) state originating from the pyrazoline ring, resulting in low $E(T_1)$ of *ca.* 1.50 eV in Fig. 5, very close to that of a single pyrazoline ring. T₂ is a LE dominant state stemming from either an acceptor or donor group, leading to high $E(T_2)$. Thus, $E(T_1)$ is much smaller than $E(T_2)$. By contrast, S₁ acquires a delocalized transition with obvious charge-transfer (CT) character. The resultant $E(S_1)$ is smaller than $E(T_2)$ (Fig. 5a). Overall, the nine compounds all meet the energy conditions $2E(T_1) \leq E(S_1) < E(T_2)$. What's more, their optical band gaps vary in a wide visible region of 2.30–3.27 eV (Table S6, ESI[†]). These indicate them to be promising candidates for achieving 200% exciton utilization in OLEDs.

The position dependence of the substituents on the lowlying excited states is investigated by changing the position of BF and B in compound **1** with the largest $\Delta E(T_1T_2)$ (2.19 eV) among the nine molecules, (compounds **10–18** in Scheme 1). It can be seen from Fig. 5 that for compounds **11–18**, $E(T_1)$ remains very low, fluctuating between 1.30 and 1.71 eV, regardless of the positions of BF and B on the pyrazoline ring and the $2E(T_1) \leq E(S_1) < E(T_2)$ is well held because of their similar transition properties with **1–9** except a little bit of violation of **11** and **14** (Fig. 4 and Fig. S2, ESI†).

The effect of the number of substituents on the excited-state property is studied by adding a donor or acceptor at the 4- or 5site of the pyrazoline ring to compound 1 (compounds **19–24** in Scheme 1), respectively. The excitation energies and NTOs are given in Fig. 5 and Table S6 (ESI[†]), Fig. S2 (ESI[†]) indicating that the $E(S_1)$, $E(T_1)$ and $E(T_2)$ of these compounds are all similar to those of compound 1, suggesting that the substituents linked to the sp³ hybridized carbon atom (R₃ and R₄) have no effect on the excited-state properties. Therefore, the introduction of R₃ or R₄ can improve physical properties such as the film-forming property and glass-transition temperature, *etc.*, while maintaining excellent excited-state electronic properties.

Experimental validation of the theoretically designed TPA-DBPrz molecule

Keeping the above design strategy in mind, we build a compound named TPA-DBPrz with R_1 = phenyl, R_2 = phenyl, R_3 = H and R_4 = TPA, with a large enough molecular weight to meet the requirement for the application in OLEDs. The calculated $E(S_1)$ is 3.12 eV at the S_1 -geometry and the $E(T_1)$ and $E(T_2)$ are 1.55 eV and 3.23 eV at the T_1 -goemetry, respectively, meeting $2E(T_1) \le E(S_1) < E(T_2)$ with $\Delta E(T_1T_2)$ of 1.68 eV. Moreover, the spin-orbit coupling between S_1 and T_2 is large enough (0.55 cm⁻¹), beneficial to the RISC from T_2 to S_1 .

To confirm the theoretical predictions, TPA-DBPrz is synthesized and characterized by ¹H NMR, ¹³C NMR, high resolution mass spectrometry and elemental analysis as given in the ESI,† and its photophysical properties are measured in chloromethane



Fig. 5 Excitation energies of the S₁, T₁ and T₂ states of 1-24 for the designed compounds at T₁-geometry at the TDA/LC- ω PBE*/6-31G(d) level.



Fig. 6 (a) The normalized UV-vis and emission spectra; (b) the fs-TA spectra and (c) the corresponding dynamics curves of TPA-DBPrz (10 μ M) in DCM; (d) comparisons between the fs-TA spectrum at 250 ps and fitted curves and ns-TA of Ru(bpy)₃Cl₂ (20 μ M) and TPA-DBPrz (1 μ M) at 2 μ s.

(DCM) solution with the results shown in Fig. 6. The emission maximum peak appears at 453 nm (Fig. 6a). Femtosecond transient absorption spectra (fs-TA) excited at 400 nm exhibit two features: an initial component with an excited state absorption (ESA) band at about 582 nm and a delayed new component with a strong ESA band at 830 nm, as well as two accompanying weak ESA bands at 527 nm and 675 nm (Fig. 6b and d). The former is promptly generated upon excitation and then disappears rapidly, which is attributed to the $S_1 \rightarrow S_n$ transition. The latter dominates the TA spectrum after 250 ps (populated from the S1 state) and lasts for several nanoseconds. To assign this spectral signature, the nanosecond transient absorption spectroscopy (ns-TA) spectrum of TPA-DBPrz sensitized by Ru(bpy)₃Cl₂ is measured under excitation at 532 nm (Fig. S3 and S4, ESI⁺), and it is seen that a mixed solution of Ru(bpy)₃Cl₂&TPA-DBPrz exhibits obvious ESA at 860/640/510 nm, corresponding to T1 of TPA-DBPrz sensitized by $Ru(bpy)_3Cl_2$. It is shown that the TA spectrum at 250 ps of TPA-DBPrz in Fig. 6d closely resembles the sensitized triplet reference spectrum (Fig. 6d). Moreover, the calculated intersystem crossing (ISC) rate from S1 to T1 using our self-developed thermal vibration correlation function (TVCF) rate formalism $^{30\text{--}32}$ is 6.00 \times 10 5 s $^{-1}$, which is far slower than the experimentally measured conversion rate from singlet to triplet state, which is in the order of picoseconds. Thus, we can assign the long-lived component to be a triplet-oriented species, namely the correlated triplet state of (TT). The slight spectral shift probably comes from different configurations between

metastable (TT) and free triplet. As shown in Fig. 6c, there is a close correlation between the decay of 582 nm and the rise of 830 nm, reflecting the conversion from S₁ to (TT) states. From the rise in evolution of the 830 nm curve, the rate constant of S₁ \rightarrow (TT) is estimated to be 103 ps⁻¹. Moreover, the ESA band positions agree well with the calculated excitation energies of the three lowest-lying excited states with significant oscillator strengths (f), which are 1.68 eV (f = 0.0067) of T₁ \rightarrow T₂, 1.83 eV (f = 0.0001) of T₁ \rightarrow T₃ and 2.19 eV (f = 0.0022) of T₁ \rightarrow T₄, respectively, based on the optimized T₁-geometry (Table S7, ESI†). Therefore, the 830 nm curve in fs-TA is pointed to be T₁ \rightarrow T₂ absorption, which fully confirms the theoretically predicted large $\Delta E(T_1T_2)$ in TPA-DBPrz. Unfortunately, the T₁ energy cannot be measured because of the absence of detectable phosphorescence even at 77 K.

Conclusions

A three-step mechanism is proposed to realize 200% exciton utilization in theory for electroluminescence by combining a hot exciton mechanism with singlet fission (HE-SF): the electropumped triplets form hot excitons (T₂) and then go to S₁, then undergo a fission process to generate two T₁, which eventually could be harvested to give phosphorescence. The energy level of the HE-SF molecule is required to be $2E(T_1) \le E(S_1) < E(T_2)$. For this purpose, an excited-state aromaticity strategy is put forward in which Baird-type aromatic five-membered heterocyclic rings are used as a bridge owing to their stabilized T₁ and then kinds of donors and acceptors are linked to the bridge to regulate S1 and T2. The calculated results of the designed 24 DBA-type molecules indicate that T_1 is a LE state concentrated on the pyrazoline ring with $E(T_1)$ of 1.30–1.74 eV, T₂ is a LE state centred upon donor or acceptor moieties with much larger $E(T_2)$, while S_1 is an electronic state with obvious charge-transfer character from donors to acceptors and its energy is close to $E(T_2)$. Consequently, the condition $2E(T_1) \leq$ $E(S_1) < E(T_2)$ is met with large $\Delta E(T_1T_2)$ and $\Delta E(S_1T_1)$. Finally, a designed promising TPA-DBPrz compound is synthesized and the fs-TA spectrum of TPA-DBPrz and the ns-TA spectrum of TPA-DBPrz&Ru(bpy)₃Cl₂ in DCM both fully confirm large $\Delta E(T_1T_2)$ of 1.49 eV. Due to the low $E(T_1)$ level of these pyrazoline derivatives, they are prone to sensitize near-infrared (NIR) phosphorescence materials, which could largely improve the efficiency of NIR OLED devices. In addition, based on the above strategy, a non-dopant phosphorescent emitter would also be designed to directly achieve 200% exciton utilization in principle. This work provides a new avenue that promises to break the limits of exciton utilization in OLEDs and a facile scenario for designing materials with the required energy conditions in practice.

Author contributions

Z. S., Q. P. and Q. S. conceived the project. Q. S. designed the molecules and carried out the calculations. Q. O., D. H., and L. W. helped with the calculation process and provided inspiring suggestions for improvement. Y. C. and Y. W. synthesized the molecules. Y. W., C. G., H. D., J. Z., C. Z. and H. F. conducted the ultrafast spectroscopy and sensitization measurements. All the authors contributed to the data analysis and writing the paper.

Conflicts of interest

There are no conflicts to declare.

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