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based multiple resonance emitters for highefficiency narrowband green OLEDs with CIE y > 0.7at high doping concentrations†

10-Dibenzothiophenyl-9,9-diphenylacridane-

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Multiple resonance emitters are attractive for high-color-purity organic light-emitting diodes (OLEDs) because of their unique narrowband emissions; however, they are typically used at low doping concentrations (≤15 wt%) due to aggregation-caused quenching and spectral broadening induced by planar molecular skeletons. Here, we report two multiple resonance emitters (BThPAc-1 and BThPAc-2) consisting of a 10-dibenzothiophenyl-9,9-diphenylacridane segment for efficient narrowband green emission at high doping concentrations. The dibenzothiophenyl-9,9-diphenylacridane segment contains two carbon-bridged phenyl rings as steric groups to inhibit intermolecular aggregation and a dibenzothiophene unit to extend conjugation and red-shift the emission to the green region. The resultant emitters exhibit narrowband emissions that peaked at 509−510 nm with a full width at half-maximum (FWHM) of 32 nm in 1 wt% doping films, which are maintained at less than 35 nm even in neat films. Remarkably, OLEDs employing the emitters reveal pure-green electroluminescence with a maximum external quantum efficiency of 20.3% and CIE coordinates of (0.18, 0.72) at 30 wt% doping concentration, which represents the best color coordinates for green multiple resonance OLEDs at high doping concentrations.

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1 Introduction

Thermally activated delayed fluorescence (TADF) emitters are attractive for organic light-emitting diodes because of their capability to utilize triplet excitons by the reverse intersystem crossing (RISC) process without the use of noble metals. To realize the rapid RISC process, a small energy gap ($\Delta E_{\rm ST}$) between singlet (S₁) and triplet (T₁) states is required for TADF emitters.¹⁻⁴ In this regard, TADF emitters are generally designed based on twisted donor–acceptor (D–A) architecture to minimize the overlap between the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) to reduce $\Delta E_{\rm ST}$.⁵⁻⁷ However, such D–A type TADF

emitters are accompanied by strong intramolecular charge transfer (ICT) character with large structural relaxation in excited states, leading to a large Stokes shift and a broad emission spectrum with full width at half-maximum (FWHM) larger than 70 nm, unfavourable for application in high-resolution displays requiring pure red/green/blue (R/G/B) emissions.⁸⁻¹⁰

Recently, multiple resonance thermally activated delayed fluorescence (MR-TADF) emitters have attracted much attention of researchers because of their narrowband emissions that can overcome the drawbacks of broad emission bands for D-A type TADF emitters. 11-15 In general, MR-TADF emitters are composed of polycyclic aromatic hydrocarbon (PAH) skeletons embedded with electron-rich atoms (nitrogen, oxygen, sulfur, and selenium) and electron-deficient atoms/groups (boron/carbonyl) with opposite electronic effects. 16-27 This arrangement of heteroatoms induces the atomic separation of the HOMO and LUMO with non-bonding orbitals, resulting in reduced vibration relaxation and narrowband emissions with FWHM less than 40 nm.28-33 By rational design of molecular structures, fullcolor MR-TADF emitters with emission covering the whole visible light range and EQE > 30% have been developed.34-39 Despite this progress, MR emitters still suffer from severe aggregation-caused quenching (ACQ) and spectral broadening

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at high doping concentrations due to their planar structure and strong intermolecular aggregation, making them always used at relatively low-concentration doped films (≤15 wt%).40-51 Although much attention has been paid to the introduction of bulky groups into MR-TADF for providing steric hindrance to reduce intermolecular interaction,52-63 it is still challenging to develop MR emitters with green emission (CIE y value > 0.70) that are insensitive to doping concentrations. 64-69 For instance, 5,11-diphenyl-5,11-dihydroindolo[3,2-b]carbazole (ICz)-based emitters (BN-ICz-1 and BN-ICz-2)50 have been reported to exhibit reduced vibration and a small FWHM of 21 nm, giving green emission at 522 nm with CIE coordinates of (0.24, 0.73) in OLED devices. Nevertheless, these emitters worked at low doping concentrations of 1-5 wt%. Similarly, MR emitters with the nitrogen atom fused at different positions⁴⁹ (BN-TP-N4) have been reported to show high-color-purity green emission with a CIE y value of 0.70 at 3 wt% doping concentration but deteriorating to CIE y < 0.70 at 5 wt% concentration. Actually, green MR emitters with CIE y > 0.70 at high doping concentrations (>15 wt%) are not reported in the literature so far.

Here, we report two green MR-TADF emitters (BThPAc-1 and BThPAc-2) consisting of a 10-dibenzothiophenyl-9,9-diphenylacridane-based B, N-doped polycyclic skeleton for efficient narrowband OLEDs with high color-purity even at high doping concentration (Fig. 1). This molecular design utilizes two carbon-bridged phenyl rings of a dibenzothiophenyl-9,9diphenylacridane segment as to provide steric hindrance to suppress aggregation of the polycyclic skeleton, and a dibenzothiophene unit to extend conjugation and red-shift the emission to the green region. As a result, the emitters BThPAc-1 and BThPAc-2 exhibit green emissions at 509–510 nm with a narrow FWHM of 32 nm at 1 wt% doping concentration, which are maintained at 515-520 nm with FWHM of 34-35 nm even in neat films. OLED devices based on the emitters exhibit green electroluminescence with an emission peak at 514 nm, FWHM of 33 nm, CIE coordinates of (0.16, 0.71) and a maximum EQE of

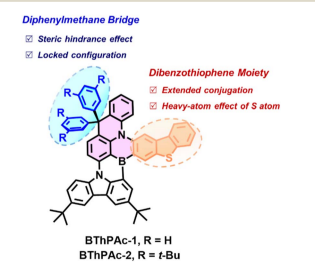


Fig. 1 Molecular design of multiple resonance emitters BThPAc-1 and BThPAc-2.

26.5% at a low doping concentration of 3 wt%. Remarkably, at a high doping concentration of 30 wt%, the OLEDs still maintain the high color purity with CIE coordinates of (0.18, 0.72) for a maximum EQE of 20.3%, representing the best color coordinates for green MR-TADF OLEDs at high doping concentrations (>15 wt%, Table S7†).

2 Results and discussion

The synthesis of BThPAc-1 and BThPAc-2 is outlined in Scheme 1. 1-Bromo-2-chloro-3-fluorobenzene was used as a starting material which underwent nucleophilic substitution with 3,6-bis(tert-butyl)-carbazole and the Buchwald coupling reaction with methyl 2-aminobenzoate to afford the ester intermediate 2. Subsequently, 2 was reacted with phenylmagnesium bromide or 3,5-di-tert-butylphenylmagnesium bromide and then with boron trifluoride etherate to form the key acridan intermediates 3a and 3b, followed by coupling with 2-bromodibenzothiophene to afford the chloride precursors 4a and 4b. The precursors were lithiated and underwent intramolecular electrophilic borylation with boron tribromide and N,N-diisopropylethylamine to give the desired products BThPAc-1 and BThPAc-2 in yields of 40-60%. To investigate the role of the fused thiophene moiety, the MR emitter without the dibenzothiophene moiety (PAc) was also synthesized from 3a through the same synthetic procedure except by using bromobenzene instead of 2-bromodibenzothiophene to form the chloride precursor. The synthesized compounds were identified by ¹H and ¹³C NMR spectra and mass spectra (Fig. S13–S34†). Thermogravimetric analysis shows that both BThPAc-1 and BThPAc-2 exhibit good thermal stability with decomposition temperatures (5% weight) above 400 °C, suitable for the preparation of OLED devices by the vacuum evaporation process. The electrochemical properties of BThPAc-1 and BThPAc-2 were characterized by cyclic voltammetry, which show reversible oxidation waves with onset potentials ($E_{\text{onset}}^{\text{ox}}$) at 0.55 and 0.49 V, and irreversible reduction waves with onset potentials (E_{onset}^{red}) at -1.98 and -1.99 V, respectively. According to the oxidation and reduction onset potentials, HOMO/LUMO levels were determined to be -5.35/-1.98 eV and -5.29/-2.02 eV for **BThPAc-1** and BThPAc-2, respectively.

To investigate the crystallographic structure of the emitters, single crystals are grown by slowly evaporating the solvent of a solution of the samples in a dichloromethane/n-hexane mixture (1/2, v/v) at 20 °C to form needle-like crystals of BThPAc-1 that can be characterized by synchrotron radiation X-ray diffraction. As shown in Fig. 2, the geometry of BThPAc-1 exhibits a quasi-planar configuration for the polycyclic skeleton with typical B-C, N-C and S-C bond lengths of 1.535-1.536 Å, 1.395–1.431 Å and 1.739–1.754 Å, respectively. Remarkably, the two carbon-bridged phenyl groups are almost perpendicular to the polycyclic skeleton plane, providing steric hindrance for the central MR skeleton. In the packing model, a strong steric effect of the perpendicular phenyl groups forces two adjacent BThPAc-1 molecules to stack in a head-to-head and tail-to-tail pattern. Two types of $\pi \cdots \pi$ interactions are observed which belong to two carbazole groups with a distance of 3.554 Å and two

Scheme 1 Synthetic routes for BThPAc-1 and BThPAc-2.

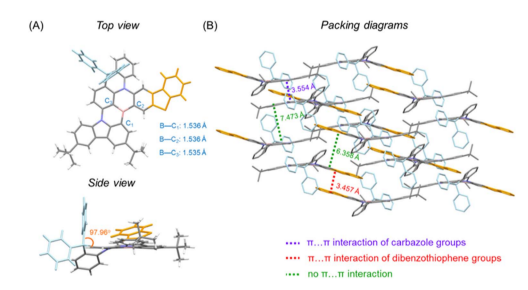


Fig. 2 (A) Single-crystal structure and (B) packing diagrams for BThPAc-1

dibenzothiophene groups with a distance of 3.457 Å, respectively, corresponding to quasi-dimers. However, other π -groups are staggered due to steric hindrance of the carbon-bridged phenyl rings, leading to enlarged distances over 6.0 Å and effectively suppressed intermolecular interactions.

To gain insight into photophysical properties of two emitters, density functional theory (DFT) and time-dependent DFT (TD-DFT) calculations are carried out at the M062x/def2-SVP level to calculate their optimized ground and excited state geometries and frontier molecular orbital (FMO) distributions. To accurately determine excited state energy levels for the emitters, a second-order approximate coupled-cluster with singles and doubles (SCS-CC2) model is also employed using the cc-pVDZ basis set. As depicted in Fig. 3 and S1-S4, HOMOs are mainly localized on nitrogen atoms and their ortho-/para-positions of the surrounding phenyl rings, while

LUMOs are localized on boron atoms and their ortho-/parapositions, corresponding to atomically separated FMO distributions for typical MR emitters. Notably, compared to PAc without the dibenzothiophenyl moiety, BThPAc-1 and BThPAc-2 exhibit extended HOMO and LUMO distributions to peripheral dibenzothiophenyl moieties, indicating larger conjugation of the polycyclic skeleton. Consequently, S1 energy levels of BThPAc-1 and BThPAc-2 (2.91-2.94 eV) are lower than that of **PAc** (3.08 eV). The ΔE_{ST} values for **BThPAc-1** and **BThPAc-2** are 0.11-0.12 eV, which are small enough to promote the RISC process. Meanwhile, the emitters exhibit a small root mean square deviation (RMSD) of 0.09 and a small reorganization energy of 0.14-0.16 eV between So and St states, indicating a small structural change upon excitation. It is noted that BThPAc-1 and BThPAc-2 have considerable SOC matrix elements (SOCME) between singlet states (S_n) and triplet states

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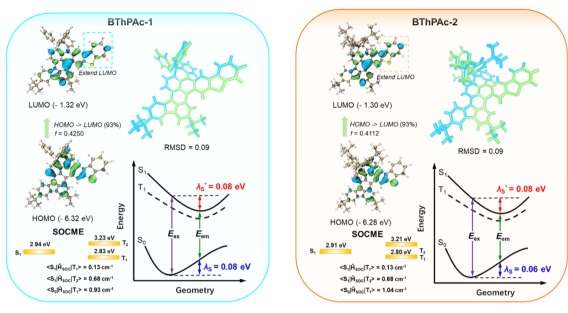


Fig. 3 HOMO/LUMO distributions, SOCME and reorganization energy for BThPAc-1 and BThPAc-2.

 (T_n) (0.13–0.68 for S_1 – T_n (n=1–2) and 0.91–1.04 for S_0 – T_1). Such SOC matrix element values are much larger than those of **PAc** without the dibenzothiophenyl moiety (0.13–0.14 for S_1 – T_n (n=1–2) and 0.73 for S_0 – T_1), indicating that the sulfur atom in the dibenzothiophenyl moiety can play a crucial role in enhancing spin–orbital coupling.

To investigate photophysical properties, UV-vis absorption and photoluminescence spectra of BThPAc-1 and BThPAc-2 in dilute toluene solution (10⁻⁵ M) were measured. As shown in Fig. 4(A) and (B), BThPAc-1 and BThPAc-2 show two absorption bands at 404-405 nm and 485-488 nm in toluene, which are assigned to π - π * transitions of polycyclic skeletons and intramolecular charge-transfer (ICT) transitions, respectively (Table 1). While for PL spectra, BThPAc-1 and BThPAc-2 exhibit intense and sharp emissions at 508 and 509 nm in toluene, respectively, corresponding to a small Stokes shift of 21-23 nm typical for the multi-resonance emitters. Compared to PAc with an emission maximum of 484 nm, the emission peaks for BThPAc-1 and BThPAc-2 are red-shifted by 24-25 nm, consistent with their extended conjugation and reduced S₁ energy level, as predicted by TD-DFT calculations. It is noted that the FWHM values for BThPAc-1 and BThPAc-2 (~30 nm) are the same as that of PAc, indicating that extension of conjugation barely affects the narrowband emission characteristics for the MR skeleton. PL spectra of the emitters in solvents with different polarities show a slight red-shift and broadening as the solvent polarity increases, consistent with their short-range ICT characteristic. According to onsets of fluorescence and phosphorescence spectra at 77 K, the experimental S₁/T₁ state energy levels are 2.57/2.41 eV and 2.55/2.40 eV for BThPAc-1 and **BThPAc-2**, respectively, giving small ΔE_{ST} values of 0.15–0.16 eV. The PLQY for the emitters measured in doped films (5 wt% doping ratios in mCP) are 88-89%, which are higher than that of PAc (PLQY = 83%). To demonstrate the effect of bridged

phenyl rings on aggregation behaviors in the solid state, the dependence of PL spectra of the emitters on the doping concentration is investigated. As shown in Fig. 4(E) and (F), BThPAc-1 exhibits narrowband green emission at 509 nm with an FWHM of 32 nm at a low concentration of 1 wt%. As the doping concentration gradually increases to 100 wt%, the emission band is red-shifted by 11 nm (from 509 to 520 nm), while the FWHM is slightly increased to 35 nm. For BThPAc-2 containing *tert*-butyl groups in two phenyl units, the red shift of emission is decreased by 5 nm (from 510 to 515 nm), and the FWHM value is broadened by only 2 nm (from 32 to 34 nm). These results indicate that intermolecular interaction and aggregation-caused spectral broadening can be efficiently suppressed by incorporating two carbon-bridged phenyl units into the MR skeleton.

Transient PL decay spectra of **BThPAc-1** and **BThPAc-2** are shown in Fig. 4(C) and (D), which reveal both prompt fluorescence with lifetimes of 7.0–7.1 ns and delayed fluorescence with lifetimes in the range of 12.7–13.0 μ s. As the temperature increases from 200 K to 300 K, the delayed components of two emitters increase corresponding to the thermally activated triplet-to-singlet up-conversion process. The ratios of delayed components are 30% and 32% for **BThPAc-1** and **BThPAc-2**, respectively, giving rate constants of prompt fluorescence (k_r), intersystem crossing ($k_{\rm ISC}$) and $k_{\rm RISC}$ of 0.9–1.0 \times 10⁸, 4.2–4.3 \times 10⁷ and 1.1–1.2 \times 10⁵ for **BThPAc-1** and **BThPAc-2**, respectively. Notably, $k_{\rm ISC}$ and $k_{\rm RISC}$ for **BThPAc-1** and **BThPAc-2** are much larger than those of **PAc** ($k_{\rm ISC} = 1.5 \times 10^7$ and $k_{\rm RISC} = 0.6 \times 10^5$), indicating that intersystem crossing between singlet and triplet states can be accelerated by the enhanced SOC.

To investigate electroluminescence properties of the two emitters, OLEDs with the configuration of indium tin oxide (ITO)/(1,1-bis[(di-4-tolylamino)phenyl]cyclohexane) (TAPC) (90 nm)/(tris(4-carbazolyl-9-ylphenyl)amine) (TCTA) (5 nm)/3,3'-

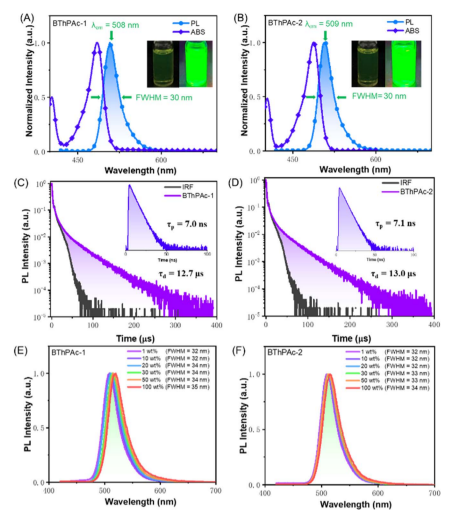


Fig. 4 (A and B) UV-visible absorption and PL spectra of BThPAc-1 and BThPAc-2 in toluene with a concentration of 10^{-5} M. (C and D) Transient PL decay curves for BThPAc-1 and BThPAc-2 doped films in 1 wt% mCP. (E and F) PL spectra of BThPAc-1 and BThPAc-2 doped films in mCP with different dopant concentrations.

Table 1 Summary of photophysical properties of BThPAc-1 and BThPAc-2

	$\lambda_{ m abs}^{a}$ [nm]	$\lambda_{\mathrm{em}}^{}b}$ [nm]	FWHM ^c [nm]	$\Phi_{ ext{PLQY}}{}^d$	$ au_{\mathrm{p}}/ au_{\mathrm{d}}^{}e} \ [\mathrm{ns}\ \mu\mathrm{s}^{-1}]$	$k_{ m r}^f \ [10^7 \ { m S}^{-1}]$	$k_{\rm ISC}^{\ \ f} \ [10^7 \ { m S}^{-1}]$	$k_{ m RISC}^{\ f}$ $\left[10^5 \ { m S}^{-1} ight]$	S_1/T_1^g [eV]	$\Delta E_{\rm ST}^{ h}$ [eV]
BThPAc-1	485	508	30	0.88	7.0/12.7	9.0	4.3	1.1	2.57/2.41	0.16
BThPAc-2	488	509	30	0.89	7.1/13.0	9.0	4.2	1.2	2.55/2.40	0.15

^a Absorption peak measured in toluene solution $(10^{-5} \, \mathrm{M})$. ^b Emission peak measured in toluene solution $(10^{-5} \, \mathrm{M})$. ^c Full width at half maximum of the PL spectrum. ^d Photoluminescence quantum yield measured in doping films in mCP (5 wt%). ^e Prompt and delayed fluorescence lifetimes of doping films. ^f k_r , $k_{\rm ISC}$ and $k_{\rm RISC}$ represent the rate constants of radiative decay of S_1 , intersystem crossing from S_1 to S_1 and reverse intersystem crossing from S_1 to S_1 . ^g Singlet (S_1) and triplet (S_1) state energy levels estimated from fluorescence and phosphorescence spectra. ^h Energy gap between S_1 and S_1 .

bis(N-carbazolyl)-1,1'-biphenyl (mCBP) (5 nm)/2-(5-(4,6-diphenyl-1,3,5-triazin-2-yl)-2-methylphenyl)-1-phenyl-benzo[d] imidazole (BIZ-6Me-TRZ): 1–30 wt% emitter (30 nm)/(5-tri(m-pyrid-3-yl-phenyl)benzene) (TmPyPB) (45 nm)/LiF (1 nm)/Al (150 nm) were fabricated. The chemical structure and energy diagram of materials used for OLED devices are illustrated in Fig. 5(A) and (B), and the device performance is displayed in

Fig. 5(C)–(E). As shown in Fig. 5(D), devices using 1 wt% BThPAc-1 and BThPAc-2 as emitters exhibit emission peaks at 512 and 511 nm, accompanied by FWHM of 34 and 33 nm, respectively, corresponding to Commission International de l'Eclairage (CIE) coordinates of (0.15, 0.70) and (0.14, 0.69) (Table 2). As the emitter concentration increases from 1 to 30 wt%, the emission peak shifts slightly from 511–512 nm to

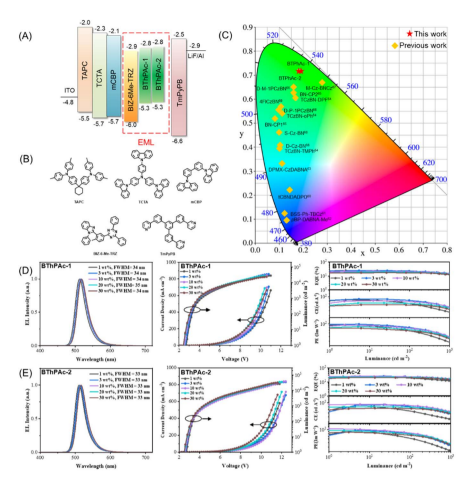


Fig. 5 (A) Device configuration for BThPAc-1 and BThPAc-2. (B) Chemical structures of materials used for the devices. (C) CIE coordinates for BThPAc-1, BThPAc-2 and reported aggregation-resistant MR-TADF emitters at high doping concentrations (>15 wt%) (Table S7†). (D and E) OLED device performance for BThPAc-1 and BThPAc-2.

517–518 nm, and the FWHMs are maintained at 33–34 nm, giving CIE coordinates of (0.19, 0.72) and (0.18, 0.72) for **BThPAc-1** and **BThPAc-2**, respectively, which are the best color coordinates for green MR-TADF OLEDs at high doping concentrations (>15 wt%) (Table S7†).

The current density (J)-voltage (V)-luminance (L) and EQE-L-luminous efficiency (LE) characteristics of the OLEDs are shown in Fig. 5(D) and (E). The devices show low turn-on voltages (at a luminance of 1 cd m $^{-2}$) of 2.4–2.7 V for **BThPAc-1** and **BThPAc-2**. From the EQE-L characteristics of devices, it can be

Table 2 Summary of OLED device performance for BThPAc-1 and BThPAc-2

	Doping concentration	${V_{ m on}}^a \ [{ m V}]$	$L_{ m max}^{\ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ $	$\operatorname{CE}_{\max}^{c}$ [cd A ⁻¹]	$\operatorname{PE_{max}}^d$ $[\operatorname{Im} \operatorname{W}^{-1}]$	$\frac{\mathrm{EQE_{Max/100/1000}}^{e}}{[\%]}$	FWHM ^f [nm]	$\lambda_{\mathrm{EL}}^{g}$ [nm]	CIE^h (x, y)
BThPAc-1	1 wt%	2.67	28 560	84.2	94.5	25.2/20.6/12.4	33.9	512	0.152, 0.702
	3 wt%	2.61	35 443	87.9	98.7	24.8/22.8/15.5	34.3	516	0.171, 0.718
	10 wt%	2.47	30 277	79.6	93.1	22.0/20.9/15.2	34.5	517	0.181, 0.721
	20 wt%	2.41	27 497	69.5	83.6	19.0/18.5/14.0	34.7	518	0.187, 0.721
	30 wt%	2.41	23 844	62.7	75.5	17.0/16.9/12.9	34.4	518	0.191, 0.720
BThPAc-2	1 wt%	2.68	26 918	79.9	86.9	24.7/19.6/12.1	32.7	511	0.143, 0.693
	3 wt%	2.64	35 737	89.9	100.9	26.5/22.0/14.4	32.7	514	0.157, 0.711
	10 wt%	2.60	39 268	91.2	106.8	25.8/24.1/16.4	33.1	516	0.171, 0.721
	20 wt%	2.47	36 141	80.0	95.0	22.4/21.6/15.5	33.0	517	0.174, 0.722
	30 wt%	2.44	32 504	72.6	85.8	20.3/20.0/14.6	32.9	517	0.176, 0.723

 $[^]a$ Turn-on voltage at a luminance of 1 cd m $^{-2}$. b Maximum luminance. c Maximum luminous efficiency. d Maximum power efficiency. e External quantum efficiency at maximum and at luminances of 100/1000 cd m $^{-2}$. f Full-width at half-maximum at 1000 cd m $^{-2}$. g EL maximum at 1000 cd m $^{-2}$.

seen that the maximum EQEs are obtained at doping concentrations of 1 wt% and 3 wt%, which are 25.2% and 26.5% for BThPAc-1 and BThPAc-2, respectively, much higher than that of **PAc** (EQE_{max} = 20.7%) at the same doping concentration. At the practical luminance of 100 cd A⁻¹, EQEs for BThPAc-1 and BThPAc-2 are maintained at 20.6% and 22.0%, respectively. The maximum luminous efficiencies (LEs) are 87.9 and 91.2 cd A^{-1} for BThPAc-1 and BThPAc-2, respectively, corresponding to maximum power efficiencies (PEs) of 98.7 and 106.8 Im W⁻¹, respectively, which are about two-fold that of **PAc** (48.5 Im W⁻¹). Importantly, BThPAc-1 and BThPAc-2 still exhibit high efficiency at a high doping concentration of 30 wt%, with the maximum EQE/LE/PE of 17.0%/62.7 cd A⁻¹/75.5 Im W⁻¹ for **BThPAc-1** and 20.3%/72.6 cd $A^{-1}/85.8$ Im W^{-1} for **BThPAc-2**. These results suggest that the DBDPA-based MR-TADF emitters can be an effective approach to realize high device efficiency for green emission with CIE y > 0.7 even at relatively high doping concentrations (>15 wt%).

3 Conclusions

In summary, we have developed two multiple resonance emitters (BThPAc-1 and BThPAc-2) containing a 10-dibenzothiophenyl-9,9-diphenylacridane-based B, N-doped polycyclic aromatic skeleton for efficient narrowband green emission at high doping concentrations. The two carbon-bridged phenyl rings of the dibenzothiophenyl-9,9-diphenylacridane segment provide steric hindrance to suppress the aggregation of the polycyclic skeleton, while the dibenzothiophene moiety could extend the conjugation to red-shift the emission to the green region. As a result, BThPAc-1 and BThPAc-2 exhibit green emissions of 509-510 nm with a narrow FWHM of 32 nm at 1 wt% doping concentration, which are maintained at 515-520 nm with narrow FWHMs of 34-35 nm as the doping concentration increases to 100 wt%. Meanwhile, the sulfur atom can enhance spin-orbital coupling by the heavy-atom effect, leading to the accelerated RISC process with a rate constant of 1.1-1.2 \times 10⁵ s⁻¹. OLED devices based on the emitters exhibit green emission with an emission peak at 514 nm, an FWHM of 33 nm and a maximum EQE of 26.5% at 3 wt% doping concentration. Importantly, the OLED devices still maintain the small FWHM of 33 nm and CIE coordinates of (0.18, 0.72), together with a high maximum EQE of 20.3% even at a doping concentration of 30 wt%, which is remarkable for green MR-TADF emitters with CIE y > 0.7 at high doping concentrations. These findings provide a facile and effective molecular design strategy for the development of aggregationresistant multi-resonance emitters to realize efficient narrowband green emissions in wide doping concentration ranges.

Data availability

The data supporting this article have been included as part of the ESI. \dagger

Author contributions

Rui Zhong synthesized the emitters and characterized the photophysical property. Mengyu Wang and Shumeng Wang fabricated and characterized the OLED devices. Xingdong Wang and Shiyang Shao provided guidance on synthesis and photophysical characterization. Rui Zhong and Shiyang Shao wrote the manuscript. L. W. supervised and directed this study.

Conflicts of interest

There are no conflicts to declare.

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