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A spiro-silafluorene-phenazasiline donor-based efficient blue thermally activated delayed fluorescence emitter and its host-dependent device characteristics†

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We report a blue thermally activated delayed fluorescence compound based on a spiro-silicon-connected silafluorene-phenazasiline donor. By incorporating triphenyltriazine as an acceptor, the compound exhibited well-separated highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) levels and charge transfer-type emission, confirmed by density functional theory (DFT) calculations and its solvatochromic characteristics. The compound exhibited blue emission with a photoluminescence (PL) peak at 469 nm, a high photoluminescence quantum yield (PLQY) of 68%, and a single-triplet energy gap of 0.20 eV in a DPEPO host matrix. Due to its co-planar silafluorene and triphenyltriazine moieties, the compound exhibited a high horizontal emission dipole ratio of 79%. Various electroluminescent devices of single host, mixed host, and interface exciplex structures were fabricated using the compound as the emitter. The external quantum efficiency (EQE), maximum brightness, and Commission Internationale de L'Eclairage (CIE) y-values ranged from 0.63 to 20.6%, 210 to 6930 nit, and 0.120 to 0.206, respectively, depending on the host materials and device structures. Among the fabricated EL devices, the DPEPO host device exhibited the highest EQE of 20.6% with CIE color coordinates of (0.150, 0.184).

1. Introduction

Blue thermally activated delayed fluorescence (TADF) emitters that could harvest triplet excitons *via* reverse intersystem crossing (RISC) have attracted a great deal of attention in the field of organic light-emitting diodes (OLEDs) due to their potential for replacing the current, relatively inefficient blue fluorescence emitters in display applications. ¹⁻⁴ TADF emitters typically incorporate donor–acceptor structures to minimize the singlet–triplet gap by separating the highest occupied molecular orbital (HOMO) and the lowest unoccupied molecular orbital (LUMO). ⁵ The emission color of TADF emitters can be modulated by selecting donors and acceptors with appropriate HOMO and LUMO levels, respectively. Therefore, blue TADF emitters generally employ

Silicon-based compounds are widely used as both the host and light-emitting material in blue OLEDs. High bandgap and triplet energy hosts based on tetraarylsilanes,^{31–33} and blue light emitting materials including oligomers and polymers based on silafluorene or spiro-silafluorene scaffolds^{34–39} are examples. In addition, the introduction of silicon atoms into azaheterocycles has resulted in blue OLEDs with many desirable characteristics. For example, phenazasiline, which is a silicon-bridged diphenylamine, has been successfully applied as the host and emitter material in blue OLEDs due to its high triplet level,

a relatively weak donor or acceptor moiety. Adachi *et al.* reported the first pure blue TADF emitter based on a diphenyl-sulfone acceptor and a dimethylcarbazole donor exhibiting a high external quantum efficiency of 9.9%. Since then, various acceptor moieties, such as triazine, pyrimidine, hosphine oxide, resulting cyanobenzene, hence hence hence hence hence here here acceptor moieties, such as triazine, phosphine oxide, resulting cyanobenzene, hence here hence hence

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Fig. 1 Molecular structure, ground state geometry, highest occupied molecular orbital-lowest unoccupied molecular orbital (HOMO-LUMO) distribution, and excited state energies of SAzTrz.

high bandgap, deep HOMO, and rigid molecular structure. $^{40-42}$ However, only a few TADF emitters containing silicon atoms have been reported to date. 16,43 Moreover, to the best of our knowledge, a spiro-silicon scaffold has not been used for a TADF emitter. This is partly due to synthetic difficulties, including the siliconheterocyclization of a donor (or an acceptor) followed by the cyclization of silafluorene. 41,42

We report herein the first TADF emitter based on a spirosilicon scaffold (Fig. 1). We connected silafluorene and phenazasiline moieties through a spiro-silicon as a donor moiety. The resulting donor, spiro[phenazasiline-10(5H),9'-[9H-9]silafluorene], was combined with a triphenyltriazine acceptor into an efficient blue TADF molecule. The newly designed TADF emitter, SAzTrz, exhibited a blue emission with a peak wavelength of 469 nm, a high photoluminescence quantum yield (PLQY) of 68%, and a high horizontal emission dipole ratio of 79% in a DPEPO host. OLEDs based on SAzTrz as the emitter and DPEPO as the host showed a high external quantum efficiency of 20.6% with blue Commission Internationale de L'Eclairage (CIE) color coordinates of (0.150, 0.184). Furthermore, we investigated the photophysical and electroluminescent (EL) properties of SAzTrz in various host materials to provide guidance regarding further applications of spiro-silafluorene-phenazasiline as the donor moiety in blue TADF emitters.

Results and discussion

Synthesis and characterization

The synthetic route of SAzTrz is illustrated in Scheme 1. Intermediate (1), 5'-(4-methoxybenzyl)-5'H-spiro[dibenzo[b,d]silole-5,10'-dibenzo[b,e][1,4]azasiline], was synthesized by in situ nucleophilic substitution of 2-bromo-N-(2-bromophenyl)-N-(4methoxybenzyl)aniline using 2,2'-dibromo-1,1'-biphenyl. Intermediate (2), 5'H-spiro[dibenzo[b,d]silole-5,10'-dibenzo[b,e][1,4]azasiline], was obtained through a de-protection reaction using DDQ. The final product (3), 5'-(4-(4,6-diphenyl-1,3,5-triazin-2yl)-2-methylphenyl)-5'H-spiro[dibenzo[b,d]silole-5,10'-dibenzo-[b,e][1,4]azasiline] (SAzTrz), was synthesized by Buchwald-Hartwig amination. The obtained intermediates and SAzTrz were characterized by proton nuclear magnetic resonance (1H-NMR), carbon-13 NMR (13C-NMR), and electron ionization (EI) mass spectrometry. Detailed synthesis and characterization procedures are described in the ESI.† The thermal stability of SAzTrz was analyzed by thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC) (Fig. S1, ESI†). SAzTrz was thermally stable up to 360 °C with a glass transition temperature of 245 °C. The HOMO level of SAzTrz was determined as −5.64 eV from the cyclovoltametric measurement and the LUMO level was determined as -2.69 eV by adding the optical bandgap (2.95 eV) (Fig. S2, ESI†).

2.2 Photophysical properties

Density functional theory (DFT) and time-dependent DFT (TD-DFT) calculations were used to predict the geometries, HOMO-LUMO distribution, natural transition orbitals (NTOs), and transition energies of SAzTrz. Calculations of the ground state geometry and excited state geometries were performed with Gaussian09, using the B3LYP functional for the ground state and the CAM-B3LYP functional for excited states. Single point calculations for transition energies and NTOs were performed with Jaguar software (Schrödinger, USA), using the LC-wPBE functional with modified parameter ω = 0.1. The 6-31G(d,p) basis set was used for both geometry optimization

Scheme 1 Synthesis of SAzTrz.

and single point calculations. The dihedral angles of silafluorene/ phenazasiline and phenazasiline/triphenyltriazine were $\sim 90^{\circ}$ in both the ground and excited states. Due to an orthogonal donor and acceptor conformation, the HOMO and LUMO were localized at the donor and acceptor, respectively. NTOs of the lowest singlet vertical emission were almost identical to the HOMO-LUMO distribution, indicating absorption and emission from a charge transfer state (Fig. S3, ESI†). The calculated S₁ vertical emission energy of 2.925 eV matched well with the onset energy of experimental SAzTrz spectra, but the T₁ energy of 2.912 eV was an overestimate compared to the experimental value of 2.79 eV. This was due to overestimation of the locally excited triplet state energy by a long-range-corrected functional, leading to a calculation result indicating that the charge transfer state is the lowest triplet state. The experimentally measured T₁ energy of SAzTrz and the related delayed fluorescence will be discussed below.

The photophysical properties of SAzTrz in various media are shown in Fig. 2, 3 and Fig. S4 (ESI†) and are summarized

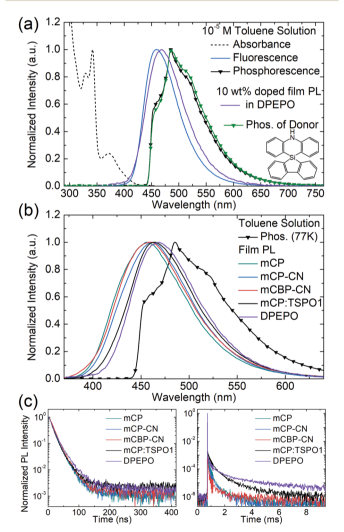


Fig. 2 Photophysical properties of SAzTrz. (a) Absorbance and PL spectra of a toluene solution. Fluorescence: PL at 300 K. Phosphorescence: PL of a frozen toluene solution at 77 K, 100 ms delay. (b) PL spectra and (c) transient PL decay of 10 wt% SAzTrz doped in various host materials.

Table 1 Photophysical properties of SAzTrz

Solution/ film	$\lambda_{\mathrm{PL}}{}^{a}$ (nm)	$\frac{\Delta E_{\mathrm{ST}}^{}b}}{\mathrm{(eV)}}$	PLQY ^c (%)	$\Phi_{ m p}^{d} \ (\%)$	$\Phi_{ m d}^{e} \ (\%)$	$ \tau_p^f $ (ns)	τ _d ^g (μs)	Θ^h (%)
Toluene	458	0.20	_	_	_	_	_	
mCP	457	0.35	40	23	77	15.1	103	_
mCP-CN	462	0.31	43	24	76	15.1	93	_
mCBP-CN	459	0.33	35	35	65	15.2	72	_
mCP:TSPO1	465	0.25	65	11	89	15.6	173	70
DPEPO	469	0.20	68	10	90	16.8	165	79

 a Photoluminescence (PL) peak. b S1–T1 gap. c PL quantum yield. d Prompt portion of total emission. e Delayed portion of total emission. f Prompt decay lifetime. g Delayed decay lifetime. h Horizontal emitting dipole ratio.

in Table 1. The solution PL of SAzTrz was measured in cyclohexane ($\varepsilon = 2.02$), toluene ($\varepsilon = 2.38$), chlorobenzene ($\varepsilon = 5.62$), tetrahydrofuran ($\varepsilon = 7.58$), and acetonitrile ($\varepsilon = 37.5$) (Fig. S4, ESI†). SAzTrz exhibited strong solvatochromism with a red-shift of its PL peak from 410 nm in cyclohexane to 585 nm in acetonitrile, indicating charge transfer-type emission as predicted from TD-DFT calculations. Fig. 2a depicts the absorbance, fluorescence, and phosphorescence spectra of 10⁻⁵ M SAzTrz in toluene. The intense and structured absorption in the range of 310-350 nm could be assigned to the absorption of spiro-silafluorenephenazasiline, while the relatively weak and broad absorption from 330 nm to 430 nm could be assigned to the intramolecular charge transfer excitation. The phosphorescence of SAzTrz in frozen toluene (cooled with liquid N2, 77 K) was measured by integrating its delayed luminescence between 100 and 500 ms after 325 nm ultraviolet (UV) excitation. The phosphorescence spectrum of SAzTrz was identical to that of spiro-silafluorenephenazasiline and the lowest triplet state of SAzTrz could be assigned to the locally excited triplet state of the donor (³LE_D). The ³CT energy could not be assigned directly from the phosphorescence spectrum since it decays nonradiatively through internal conversion to the lowest triplet (³LE_D). However, the estimated ¹CT-³CT energy gap of SAzTrz obtained from TD-DFT calculations is very small (0.013 eV), due to well-separated $S_1 \rightarrow S_0$ and $T_1 \rightarrow S_0$ transition orbitals (Fig. S3, ESI†). Thus, we may infer that the ³CT state of SAzTrz lies very close to the ¹CT state. Recently, Monkman et al. showed that the RISC of TADF emitters occurs from second-order transitions, including the coupling of ³CT-³LE and ³LE-¹CT, and that the rate of RISC is maximized when the ¹CT-³LE gap is minimized. ⁴⁴ Hence, the RISC efficiency (Φ_{RISC}) of SAzTrz will be strongly affected by the ¹CT energy, which is determined by the polarity of the host matrix.

To evaluate the performance and emission properties of SAzTrz as the emitter for electroluminescence devices, the photophysical properties of doped films were investigated prior to OLED fabrication. For host materials, mCP ($T_1 = 3.02 \text{ eV}$), mCP-CN ($T_1 = 3.0 \text{ eV}$), mCBP-CN ($T_1 = 2.81 \text{ eV}$), ⁴⁵ TSPO1 $(T_1 = 3.36 \text{ eV})^{46}$ and DPEPO $(T_1 = 3.00 \text{ eV})^{47}$ were chosen to confine triplet excitons within the TADF emitter. The emission peak of SAzTrz red-shifted from 457 nm (mCP) to 469 nm (DPEPO) with increasing polarity of the host material (Fig. 2b). We also determined the singlet-triplet energy gap (ΔE_{ST}) of

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SAzTrz in various host materials from the difference in emission onset energy between fluorescence at 300 K in a doped film and phosphorescence at 77 K in a toluene solution. $\Delta E_{\rm ST}$ ranged from 0.35 eV in mCP to 0.20 eV in DPEPO. $\Delta E_{\rm ST}$ is the barrier for reverse intersystem crossing (RISC) from a triplet exciton to a singlet exciton, and the effect of ΔE_{ST} on delayed fluorescence can be observed from the transient PL decay (Fig. 2(c)). The delayed fluorescence of SAzTrz in mCP:TSPO1 and DPEPO persisted for over 10 ms and the delayed portion of the total emission (Φ_d) was as high as 90%. However, in mCP, mCP-CN, and mCBP-CN, the delayed fluorescence of SAzTrz ended at around 4 ms with a delayed portion of $\Phi_d = 65-77\%$, indicating decreased RISC efficiency compared to that of SAzTrz in the mCP:TSPO1 and DPEPO hosts. The longer lifetimes of SAzTrz delayed fluorescence in mCP:TSPO1 (173 µs) and DPEPO (165 µs) compared to that of SAzTrz in other host materials (72-103 µs) can be attributed to an enhanced ISC/ RISC cycle by small $\Delta E_{\rm ST}$ values. The delayed PL lifetime of SAzTrz was especially short in the mCBP-CN host. Here we may infer that the triplet exciton is not perfectly confined to SAzTrz, due to the similar triplet energies of mCBP-CN (2.81 eV) and SAzTrz (2.79 eV).

The emitting dipole orientations of SAzTrz in the mCP: TSPO1 and DPEPO hosts were determined by angle-dependent PL measurements in doped films, and by optical simulations (Fig. S5, ESI†). The proportion of horizontally oriented emitting dipoles (Θ) of SAzTrz was 70% for SAzTrz in mCP:TSPO1 and 79% in DPEPO (note that $\Theta = 66.7\%$ indicates an isotropic orientation). Therefore, SAzTrz exhibited a preferred horizontal orientation in both host materials. This is due to a parallel alignment between the emitting dipole moment and the longaxis of the SAzTrz molecule, and a coplanar arrangement between the silafluorene and triphenyltriazine moieties.

The photophysical properties of SAzTrz and blue TADF emitters based on similar donor moieties were compared and are shown in Fig. 3, Fig. S6, S7 and Table S1 (ESI†). These three blue TADF emitters, SAzTrz, DTPDDA, 43 and SpiroAC-TRZ, 8 all incorporate triphenyltriazine as an acceptor but incorporate spiro-silafluorene-phenazasiline, diphenyl-phenazasiline, and spiro-fluorene-acridine as donors, respectively. Among these TADF emitters, SAzTrz shows the deepest PL emission for both the toluene solution (Fig. 3(b)) and doped film in mCP:TSPO1 (Fig. S6, ESI†). The bond length of Si-C is 1.86 Å, which is longer than the C-C bond (1.53 Å), and it results in a larger C-N-C bond angle (127.1°) of diphenylamine in SAzTrz compared to that of 121.5° for SpiroAC-TRZ (Fig. S7, ESI†). Hence, the conjugation within the donor moiety of SAzTrz is weaker than that of SpiroAC-TRZ, leading to a larger HOMO-LUMO gap and higher S1 energy of SAzTrz. Although it is not as significant as the difference of the PL spectra of SAzTrz and SpiroAC-TRZ, SAzTrz also shows a blue-shifted PL spectrum compared to DTPDDA. Formation of spiro-silicon (linking two phenyls into silafluorene) slightly increases the C-N-C bond angle of the diphenylamine and results in a blueshifted PL spectrum. Formation of spiro-silafluorene also effects the molecular orientation of the emitter doped in host materials.

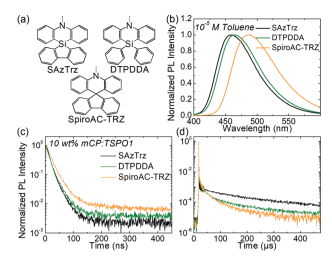


Fig. 3 Comparison of the photophysical properties of SAzTrz and blue TADF emitters based on similar donor moieties. (a) Donor structure of SAzTrz, DTPDDA, and SpiroAC-TRZ. (b) PL spectra in toluene. (c and d) Transient PL decay in mCP:TSPO1.

As the silafluorene and triphenyltriazine of SAzTrz are co-planar, it exhibits a higher Θ of 70% than DTPDDA (66%).

Modification of the donor structure gives rise to a difference in the PL decay curves. The prompt decay rate is the fastest for SAzTrz (Fig. 3(c)) and we may infer from the PLQY and prompt portion of the total emission (Φ_p) (Table S1, ESI†) that the nonradiative decay rate of the singlet state is the largest for SAzTrz. We expected that linking two freely rotating phenyls (DTPDDA) into silafluorene (SAzTrz) would suppress nonradiative relaxation but this was not the result. In the delayed emission region, the decay rate of DTPDDA is faster than the decay rate of SAzTrz until 200 μs and then becomes similar. The larger change in the delayed decay rate of DTPDDA implies that it has a broader distribution of the molecular configuration than SAzTrz. The difference in delayed decay tendency of SAzTrz and DTPDDA might also originate from different RISC mechanisms. As shown in Fig. S6 (ESI†), DTPDDA exhibits broad CT-like phosphorescence whereas the phosphorescence of SAzTrz is from the donor. As the lowest triplet state of DTPDDA is ³CT and the locally excited triplet of the donor and acceptor is much higher than ¹CT and ³CT, ³CT would be upconverted to ¹CT by direct RISC. On the other hand, the lowest triplet of SAzTrz is ³LE, and ³CT lies between ¹CT and ³LE. Therefore, the main RISC mechanism of SAzTrz would be population of 3CT from 3LE by vibronic coupling followed by secondorder coupling of ¹CT and ³CT induced by spin-orbit coupling of ¹CT-³LE and vibronic coupling of ³LE-³CT.⁴⁴

The device performances of TADF OLEDs based on SAzTrz, DTPDDA, and SpiroAC-TRZ are summarized in Table S2 (ESI†). The RISC efficiencies of the emitters estimated by optical simulations based on the PLQY and Θ are compared.

2.3 Electroluminescent properties

Several EL devices were fabricated to investigate the EL properties of SAzTrz and their dependence on the host material and device structure (Fig. 4, 5, and Table 2). The structures of the fabricated OLEDs were as follows: (a) ITO (70 nm)/6 wt% ReO₃:mCP (45 nm)/mCP (10 nm)/10 wt% SAzTrz-mCP:TSPO1 (1:1) (30 nm)/TSPO1 (10 nm)/6 wt% Rb₂CO₃:TSPO1 (45 nm)/LiF (0.7 nm)/Al (100 nm), (b) ITO (70 nm)/MoO₃ (1 nm)/H2 (45 nm)/ mCBP (5 nm)/10,20 wt% SAzTrz:mCBP-CN (15 nm)/CN-T2T (50 nm)/LiF (0.7 nm)/Al (100 nm), and (c) ITO (70 nm)/MoO₃ (1 nm)/H2 (45 nm)/mCP (5 nm)/10 wt% SAzTrz:mCP, mCP-CN, or DPEPO (15 nm)/DPEPO (5 nm)/TPBi (45 nm)/LiF (0.7 nm)/Al (100 nm).

Device (a) employs a mixed host of mCP:TSPO1, a thick (30 nm) emitting layer (EML), a doped charge transport layer to attain charge balance in the EML and a wide recombination zone. Device (b) utilizes an interface exciplex (Fig. S8, ESI†) composed of mCBP-CN:CN-T2T to form excitons on the emitter by energy transfer from the interface exciplex rather than direct charge trapping on the emitter. Device (c) is a single-host structured device incorporating mCP, mCP-CN, and DPEPO as host materials with hole/electron blocking layers.

As expected from the photophysical properties of SAzTrz in various host materials, the DPEPO device exhibited the highest external quantum efficiency (EQE) (20.6%), followed by an EQE of 8.2% with the mCP:TSPO1 device. Theoretically achievable EQEs were calculated as a function of the thickness of the transport layers by optical simulations based on the PLQY and

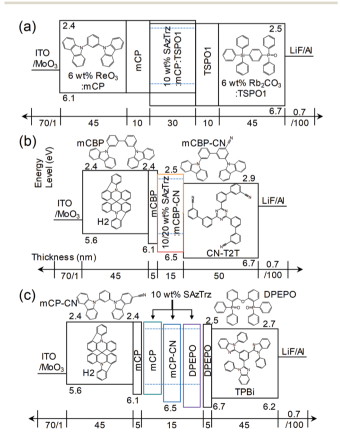


Fig. 4 Device structures of organic light-emitting diodes (OLEDs) based on SAzTrz. (a) mCP:TSPO1 mixed host structure. (b) mCBP-CN:CN-T2T interface exciplex structure. (c) Single host device structures based on mCP, mCP-CN, and DPEPO hosts

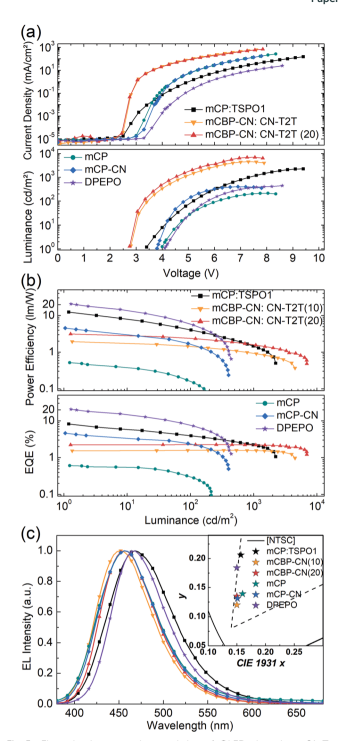


Fig. 5 Electroluminescent characteristics of OLEDs based on SAzTrz. (a) Current density-voltage-luminance curves. (b) Current density-power efficiency/external quantum efficiency (EQE) curves. (c) EL spectrum and CIE color coordinates (inset) at 200 cd m⁻²

 Θ of the emitter, assuming that the RISC efficiency (Φ_{RISC}) is 100% (Fig. S9, ESI†). The EQE of the DPEPO device was close to the calculated maximum achievable EQE of 20.8%, implying that the RISC efficiency was nearly 100%. Conversely, the EQE of the mCP:TSPO1 device was much smaller than the calculated EQE of 20.2%, implying a relatively lower amount of harvested

Electroluminescence properties of OLEDs based on SAzTrz Table 2

Device	EQE_{max}^{a} (%)	EQE_{100}^{b} (%)	${ m PE}_{ m max}^{c} \left({ m lm} \ { m W}^{-1} ight)$	$\lambda_{\mathrm{EL}}^{}d}$ (nm)	CIE^e	Turn on $f(V)$	$L_{\text{max}}^{g} \left(\text{cd m}^{-2} \right)$
mCP:TSPO1	8.2	3.9	12.4	468	(0.157, 0.206)	3.4	2240
mCBP-CN:CN-T2T (10 wt%)	1.6	1.6	1.9	452	(0.150, 0.120)	2.8	4500
mCBP-CN:CN-T2T (20 wt%)	2.3	2.3	3.1	456	(0.150, 0.134)	2.8	6930
mCP	0.6	0.3	0.5	456	(0.160, 0.139)	4.0	210
mCP-CN	4.6	2.3	4.6	456	(0.151, 0.133)	3.8	400
DPEPO	20.6	7.6	20.7	468	(0.150, 0.184)	4.1	440

^a Maximum external quantum efficiency (EQE). ^b EQE at 100 cd m⁻². ^c Maximum power efficiency (PE). ^d Electroluminescence (EL) peak. ^e Commission Internationale de L'Eclairage (CIE) 1931 coordinates at 200 cd m⁻². ^f Turn on voltage. ^g Maximum luminance.

triplets. Other devices, including those based on the interface exciplex structure and single-host structures with mCP and mCP-CN hosts, showed poor efficiency with EQEs under 5%, indicating that the majority of the emission from these devices originated from singlet excitons instead of upconverted triplets.

The maximum luminance of the fabricated devices did not trend with efficiency performances with regard to host materials. Devices employing an interface exciplex based on mCBP-CN:CN-T2T exhibited the highest luminance up to 6930 nit, while the luminance values of mixed- and single-host devices were 2240 nit and 210-440 nit, respectively. The large differences in maximum luminance observed among the devices can be attributed to the triplet lifetime of the emitter. For the case of the interface exciplex device, most of the excitons are formed at the interface between mCBP-CN and CN-T2T, and only singlet exciplexes are transferred to the emitter via Förster energy transfer. In addition, the triplet lifetime of SAzTrz ($T_1 = 2.79$ eV) was shortest in mCBP-CN $(T_1 = 2.81 \text{ eV})$ due to their similar triplet energies. Hence, the population of "dark" triplet states in SAzTrz was smallest in the interfacial exciplex device, resulting in the highest luminance. With a single-host device, the majority of excitons are formed by charge trapping at the emitter. Triplets are confined to the SAzTrz molecule in the mCP and mCP-CN hosts, which leads to a large population of dark triplet states and low brightness. Although the RISC process of SAzTrz is efficient in a DPEPO host, the triplet lifetime is the longest due to a prolonged ISC/RISC cycle and the efficiency decreases rapidly at high current densities, resulting in a low maximum luminance. The exciton dynamics of SAzTrz in mCP:TSPO1 are similar to those of SAzTrz in DPEPO, but the wider exciton recombination zone and thicker EML of the former result in higher luminance.

3. Conclusion

In summary, we designed and synthesized the first TADF emitter containing a spiro-silicon scaffold in its molecular structure. Spiro-silicon-connected silafluorene-phenazasiline and triphenyltriazine were used as the donor and acceptor moieties, respectively, resulting in a D- π -A structured TADF molecule, SAzTrz. The lowest triplet energy of silafluorene-phenazasiline was 2.79 eV, which was high enough to yield a small singlettriplet energy gap ($\Delta E_{\rm ST}$ = 0.20 eV) with efficient triplet upconversion (Φ_{RISC} = 1). The photophysical properties of SAzTrz were investigated in various host materials and OLEDs were fabricated with mixed-host, single-host, and interface exciplex structures. Highly efficient blue OLEDs were realized with DPEPO as a host material, attaining an EQE value of 20.6% with CIE coordinates of (0.150, 184).

4. Experimental section

4.1 Fabrication of organic thin films and OLEDs

Organic thin films for photoluminescence (PL) measurements were fabricated by thermal deposition onto pre-cleaned fused silica substrates at a base pressure of $<5 \times 10^{-7}$ Torr. OLEDs were fabricated by thermal deposition onto 70 nm-thick patterned ITO glass substrates at a base pressure of $<5 \times 10^{-7}$ Torr. Before deposition of the organic layers, the ITO substrates were pre-cleaned with acetone and isopropyl alcohol and then exposed to UV-ozone for 10 minutes. Samples for photoluminescence and electroluminescence measurements were encapsulated in a nitrogen-filled glove box.

4.2 Measurements of absorption, photoluminescence, and electroluminescence

The absorbance spectrum of 10⁻⁵ M SAzTrz dissolved in toluene was measured with a Varian Cary 5000 ultravioletvisible near-infrared (UV-vis-NIR) spectrometer. The PLQY was measured using a monochromator-attached photomultiplier tube (PMT) with a PL sample in an integrating sphere (LabSphere, USA) using a continuous wave 325 nm He/Cd laser (Kimmon Koha, Japan) for excitation. Photoluminescence was measured with a CCD spectrometer (Maya 2000; Ocean Optics, USA) using a He/Cd laser (325 nm) for excitation. Phosphorescence of the toluene solution was measured by freezing the solution in liquid nitrogen at 77 K. For angle-dependent PL measurements, p-polarized light emitted from PL samples was measured by attaching the film substrate to a half-cylinder lens with index-matching oil and changing the angle between the sample and the detector from -90° to 90° using a motorized rotational stage. Transient PL was measured with a streak camera (Hamamatsu Photonics, Japan) using a nitrogen laser (337 nm; Usho Optical Systems, Japan) as the excitation source. The *I-V-L* curves of the OLEDs were measured using a programmable source meter (Keithley 2400; Keithley Instruments, USA) and a spectrophotometer (Spectrascan PR650; Photo Research, USA).

4.3 Thermogravimetric analysis and cyclic voltammetry

Thermal analyses were performed with a TGA-2100 thermogravimetric analyzer in a nitrogen atmosphere at a heating rate of 10 °C min⁻¹. Cyclic voltammetry (CV) was performed using a CH instruments electrochemical analyzer. A three-electrode system composed of a glassy carbon working electrode, a platinum wire counter electrode, and a Ag/AgCl reference electrode was used. A 0.1 M solution of tetrabutylammonium hexafluorophosphate (TBAP) in chloroform was used as the electrolyte. The ferrocenium/ferrocene (4.43 eV) redox couple was used as an internal standard. Cyclic voltammograms were obtained at a scan rate of 100 mV s⁻¹. The HOMO level was obtained from the CV oxidation onset and the LUMO level was obtained by adding the optical bandgap energy to the HOMO level.

4.4 Synthesis

A detailed synthesis procedure and characterization of SAzTrz are described in the ESI.†

Author contributions

Y. K., Y.-H. K., and S.-K. K. designed and synthesized the emitter compound. S.-J. W. designed and carried out the photoluminescence and electroluminescence experiments. S.-J. W. and J.-J. Kim wrote the manuscript. S.-J. W. and Y. K. equally contributed to this work.

Conflicts of interest

There are no conflicts to declare.

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