Creation of Bifunctional Materials: Improve Electron-Transporting Ability of Light Emitters Based on AIE-Active 2,3,4,5-Tetraphenylsiloles

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2,3,4,5-Tetraphenylsiloles are excellent solid-state light emitters featured aggregation-induced emission (AIE) characteristics, but those that can efficiently function as both light-emitting and electron-transporting layers in one organic light-emitting diode (OLED) are much rare. To address this issue, herein, three tailored n-type light emitters comprised of 2,3,4,5-tetraphenylsilole and dimesitylboryl functional groups are designed and synthesized. The new siloles are fully characterized by standard spectroscopic and crystallographic methods with satisfactory results. Their thermal stabilities, electronic structures, photophysical properties, electrochemical behaviors and applications in OLEDs are investigated. These new siloles exhibit AIE characteristics with high emission efficiencies in solid films, and possess lower LUMO energy levels than their parents, 2,3,4,5-tetraphenylsiloles. The double-layer OLEDs [ITO/NPB (60 nm)/silole (60 nm)/LiF (1 nm)/Al (100 nm)] fabricated by adopting the new siloles as both light emitter and electron transporter afford excellent performances, with high electroluminescence efficiencies up to 13.9 cd A⁻¹, 4.35% and 11.6 lm W⁻¹, which are increased greatly relative to those attained from the triple-layer devices with an additional electron-transporting layer. These results demonstrate effective access to n-type solid-state emissive materials with practical utility.

1. Introduction

Organic luminescent materials have been the subject of intense academic and commercial interest over the past two decades because of their various practical applications such as light emitters for organic light-emitting diodes (OLEDs). Although considerable progresses have been made in the development of organic emitters, major challenges still remain. One stubborn problem is that many “conventional” luminophors that show good emissions in solutions suffer from aggregation-caused quenching (ACQ) in the condensed phase. This becomes one of the thorny obstacles to the evolution of OLEDs, for most light emitters have to be fabricated into solid films when used in devices. Various chemical, physical and engineering approaches have been proposed to mitigate the ACQ effect. But these methods often lead to some side effects, rendering efficient solid-state emitters infrequent. It will be nice, if a system can be developed, in which light emission is enhanced rather than quenched by aggregation. In 2001, Tang’s group found that 1-methyl-1,2,3,4,5-pentaphenylsilole (MPPS) was almost non-fluorescent in solution but it could emit strong light when aggregated in poor solvents or fabricated into solid films, demonstrating an intriguing phenomenon of aggregation-induced emission (AIE). Based on this finding, many AIE-active luminogens are developed, making pure solid films with high emission efficiencies readily achievable. This paves a new avenue to realize efficient non-doped OLEDs, which avoid the complicated and hard-to-control processes of the doping techniques used to alleviate the ACQ effect of light emitters.

Another challenge for next generation of high-performance OLEDs is to simplify device configuration and cut down the cost, without sacrificing device efficiency. In this regard, design and synthesis of new luminescent materials that can simultaneously serve as light emitter and hole-/electron-transporting material in one device is a promising alternative. Compared with organic hole-transporting (p-type) luminogens, electron-transporting (n-type) materials with low
electron injection barrier and fast electron mobility in addition to excellent solid-state emission are scarce.\[8\]

Among numerous AIE luminogens, siloles have attracted considerable attention because of not only bright emissions in solid films but also unique electronic structures.\[9\] The effective $\sigma^*-\pi^*$ conjugation between the $\sigma^*$ orbitals of two exocyclic silicon-carbon bonds and $\pi^*$ orbital of the butadiene moiety imparts the low-lying lowest unoccupied molecular orbital (LUMO) and high electron affinity to siloles.\[10\] So far, efficient solid-state emitters and electron transporter have been developed based on siloles.\[11\] Multiple OLEDs utilizing two different kinds of silole derivatives as light-emitting and electron-transporting layers, respectively, had been tried by Kafafi et al., which afforded high efficiencies up to 4.1\%.\[12\] However, the combination of both merits of siloles to generate efficient n-type light emitters has been barely realized. In our previous studies, we found that OLEDs fabricated by using 2,3,4,5-tetraphenylsiloles (Scheme 1) as light-emitting layers afforded remarkably high electroluminescence (EL) efficiency.\[13\] But these devices need an electron-transporting layer to balance the holes and electrons, implying that the 2,3,4,5-tetraphenylsiloles are excellent light emitters but their electron-transporting ability needs to be further enhanced. In view of this, three tailored silole derivatives consisting of 2,3,4,5-tetraphenylsiloles and bimesitylboryl groups are designed and synthesized in this work (Scheme 1). It is envisioned that the incorporation of inherently electron-deficient group, dimesitylboryl, is beneficial to advances in electron affinity and electron-transporting ability.\[14\] The thermal, photophysical, electronic, electrochemical, and electroluminescent properties of the new siloles are investigated. The results reveal that these new siloles are efficient bifunctional materials for OLEDs.

### 2. Results and Discussion

#### 2.1. Synthesis

The target 2,3,4,5-tetraphenylsiloles carrying dimesitylboryl groups, (MesB)$_2$DMTPS, (MesB)$_2$MPPS and (MesB)$_2$HPS, were facilely prepared according to the synthetic routes outlined in Scheme 1. The detailed procedures and characterization data are given in the Experimental section. Briefly, compounds 2\[15\] and 3\[16\] were prepared according to the methods described in the literature. The treatments of 3 with lithium 1-naphthalenide (LiNaph), and subsequent ZnCl$_2$-TMEDA generated 2,5-meta-lated silole intermediates (4). Without purification, 4 underwent coupling reaction with 2 in the presence of a palladium catalyst, affording final compounds (MesB)$_2$DMTPS, (MesB)$_2$MPPS and (MesB)$_2$HPS in 80, 50 and 40\% yields, respectively. The new siloles were fully characterized by NMR and mass spectros-copies, which confirmed their molecular structures. They are soluble in common organic solvents including THF, dichloromethane, chloroform, toluene, etc., but insoluble in water.

![Scheme 1. Synthetic routes to dimesitylboryl-functionalized 2,3,4,5-tetraphenylsiloles. LiNaph = lithium 1-naphthalenide; TMEDA =N,N,N',N'-tetramethylethylenediamine.](image-url)
The thermal properties of (MesB)$_2$DMTPS, (MesB)$_2$MPPS and (MesB)$_2$HPS were evaluated by thermogravimetric analysis (TGA) and differential scanning calorimetry (DSC). The results show that they possess good thermal stability with decomposition temperatures ($T_d$) of 240–289 °C, according to 5% loss of initial weight. High glass-transition temperatures ($T_g$) of 121–131 °C (Table 1) are recorded from the new siloles, indicating that they are morphologically stable. The good thermal and morphological stabilities enable them to serve as active materials for OLEDs.

2.2. Crystal Structure

Single crystals of (MesB)$_2$DMTPS and (MesB)$_2$HPS were grown from THF/ethanol mixture, and analyzed by X-ray diffraction crystallography. Figure 1 displays the crystal structures of (MesB)$_2$DMTPS and (MesB)$_2$HPS. It can be seen that in the crystalline state, both siloles show twisted conformations and no π–π stacking interactions are found between aromatic rings. Figure 2 illustrates the molecular packing patterns of (MesB)$_2$DMTPS and (MesB)$_2$HPS in the crystalline state. The (MesB)$_2$HPS molecules are arranged more regularly and tightly than (MesB)$_2$DMTPS molecules. The close packing of (MesB)$_2$HPS suggest that it should have better carrier-transporting ability than (MesB)$_2$DMTPS.

2.3. Optical Property

The absorption spectra of the new siloles in dilute THF solutions (10 µM) are shown in Figure 3A. The spectral profiles of (MesB)$_2$DMTPS, (MesB)$_2$MPPS and (MesB)$_2$HPS are similar, with maxima at 392, 395, and 396 nm, respectively, associated with π–π* transitions. The photoluminescence (PL) spectra of three siloles in dilute THF solutions (10 µM) are shown in Figure 3B. Like most silole derivatives, these new siloles show weak emissions in the solution state. Only faint PL signals peaked at 512, 516 and 523 nm are recorded for (MesB)$_2$DMTPS, (MesB)$_2$MPPS and (MesB)$_2$HPS, respectively. The fluorescence quantum yields (Φ) of (MesB)$_2$DMTPS, (MesB)$_2$MPPS, and (MesB)$_2$HPS in dilute THF solutions are as low as 0.87, 1.35, and 1.38%, respectively, estimated using quinine sulfate (Φ = 54.6% in 0.1 N sulfuric acid) as standard, which indicates that they are practically very weak emitters when dissolved in good solvents. These siloles become highly emissive in the solid state. The films of (MesB)$_2$DMTPS, (MesB)$_2$MPPS, and (MesB)$_2$HPS

Table 1. Optical, electronic and thermal properties of (MesB)$_2$DMTPS, (MesB)$_2$MPPS and (MesB)$_2$HPS.

| Compounds      | $\lambda_{ab}^{a)}$ [nm] | $\lambda_{em}^{b)}$ (ΦF $^c))$ [nm][%] | $T_g$/$T_d$ [°C] | HOMO/LUMO [eV] | $E_g$ [eV] |
|----------------|--------------------------|----------------------------------------|------------------|---------------|------------|
| (MesB)$_2$DMTPS| 392                      | 512(0.87)                              | 121/240          | −5.6/−3.0     | 2.6        |
| (MesB)$_2$MPPS | 395                      | 516(1.35)                              | 131/269          | −5.7/−3.1     | 2.6        |
| (MesB)$_2$HPS  | 396                      | 523(1.38)                              | 123/289          | −5.7/−3.1     | 2.6        |

$^a)$ Absorption maximum in dilute THF solution (10 µM); $^b)$ Emission maximum in THF solution; $^c)$ Fluorescence quantum yield in THF solution (soln) determined using quinine sulfate (Φ = 54.6% in 0.1 N sulfuric acid) as standard or in the film state by a calibrated integrating sphere.

Figure 1. ORTEP drawings of (MesB)$_2$DMTPS (CCDC 948629) and (MesB)$_2$HPS (CCDC 948630).
fluoresce intensely with emission maxima at 516, 524, and 526 nm, respectively (Figure S1, Supporting Information). Only slight bathochromic shifts are observed in the PL spectra of the films compared to those of the solutions, which is attributive to the propeller-like molecular conformations as well as the branched dimethylboryl groups that have impeded close π–stacking interactions. The Φ_F values of the films, determined by integrating sphere, are as high as 56, 58, and 62% for (MesB)_2 DMTPS, (MesB)_2 MPPS, and (MesB)_2 HPS, revealing that they are AIE-active and are excellent solid-state emitters.

The AIE attributes of the new siloles are further confirmed by their emission behaviors in THF/water mixtures. Figure 4A illustrates the PL spectra of (MesB)_2 MPPS in THF/water mixtures as an example. It can be seen that when the water content is low, the emission is weak. But when the water content becomes high, the emission intensity enhances swiftly (Figure 4B). Similar emission behaviors are also recorded from (MesB)_2 DMTPS and (MesB)_2 HPS in THF/water mixtures (Figure S2, Supporting Information). Since (MesB)_2 DMTPS, (MesB)_2 MPPS, and (MesB)_2 HPS are insoluble in water, their molecules must have aggregated in aqueous medium. The intramolecular rotation that is active in the solution state is restricted due to steric hindrance in the condensed phase. The nonradiative relaxation channel is thus blocked and radiative decay of the excited state is promoted, rendering molecules highly emissive. These results manifest that the new siloles are AIE-active indeed.

2.4. Theoretical Calculation

To gain a deep insight into the electronic structures of the new siloles, theoretical calculations are performed by density function theory (DFT) method with a B3LYP/6–31G(d) basis set using Gaussian 09 package. The optimized structures and orbital distributions of the HOMOs and LUMOs of (MesB)_2 DMTPS, (MesB)_2 MPPS, and (MesB)_2 HPS are shown in Figure 5. All the molecules adopt twisted conformations, which would hamper the strong intermolecular interactions between the molecules. In LUMOs, significant contributions from the boron centers are observed, owing to the ππ*–ππ* conjugation between empty π-orbital of boron atom and π* orbitals of the phenyl rings. Meanwhile, the electronic cloud of LUMOs can also delocalize to exocyclic Si–C bonds through σ*–π* conjugation. Such distinctive electronic structures result in low LUMO energy levels of −1.97, −2.05, and −2.04 eV for (MesB)_2 DMTPS, (MesB)_2 MPPS, and (MesB)_2 HPS, respectively. The values are much lower than those of DMTPS (−1.59 eV) and other 2,3,4,5-tetraphenylsiloles,[17] demonstrating that the incorporation of dimethylboryl groups has effectively lowered the LUMO energy levels. The electron injection and transport become more favorable in the new siloles than their parents.

2.5. Electrochemical Property

The electrochemical properties of (MesB)_2 DMTPS, (MesB)_2 MPPS, and (MesB)_2 HPS were investigated by cyclic voltammetry (CV) in dichloromethane solution with 0.1 M tetrabutylammonium hexafluorophosphate as the supporting electrolyte at a scan rate of 50 mV s⁻¹ using platinum as the working electrode and saturated calomel electrode (SCE) as the reference electrode. The new siloles exhibit similar CV curves (Figure 6). The oxidation onset potentials (E_onset) of (MesB)_2 DMTPS, (MesB)_2 MPPS, and (MesB)_2 HPS occur at 1.2, 1.3, and 1.3 V, respectively. The energy levels of HOMO (HOMO = −(4.4 + E_onset)) and LUMO (LUMO = −(HOMO + E_g)) were determined by E_onset and optical band gaps. The HOMO energy levels are calculated to be −5.6, −5.7, and −5.7 eV, and the LUMO energy levels are −3.0, −3.1, and −3.1 eV for (MesB)_2 DMTPS, (MesB)_2 MPPS, and (MesB)_2 HPS, respectively. The LUMO energy levels of the present siloles are equal to or even lower than those of widely used electron-transporting materials, for example, tris(8-hydroxyquinoline) aluminum (Alq₃, −3.0 eV) and 1,3,5-tris(N-phenylbenzimidazol-2-yl)benzene.
(TPBi, −2.7 eV), indicating their great potential as electron-transporting materials for OLEDs.

2.6. Electroluminescence

The efficient solid-state PL emissions of (MesB)₂DMTPS, (MesB)₂MPPS, and (MesB)₂HPS inspire us to evaluate their electroluminescence (EL) properties in non-doped OLEDs. Triple-layer OLEDs with a configuration of ITO/NPB (60 nm)/silole (20 nm)/TPBi (40 nm)/LiF (1 nm)/Al (100 nm) (Device I) are fabricated, in which (MesB)₂DMTPS, (MesB)₂MPPS or (MesB)₂HPS serve as light-emitting layer (EML), N,N′-di(1-naphthyl)-N,N′-diphenyl-benzidine (NPB) acts as hole-transporting layer (HTL) and TPBi functions as electron-transporting layer (ETL). The devices I of the new siloles radiate yellow lights, whereas the PL emissions of the films are in the green region. Taking (MesB)₂MPPS as an example, its EL maximum is located at 552 nm, which is red-shifted by 28 nm in comparison with its PL in film (Figure 7A). Such kind of red shifts could be ascribed to space charge accumulation caused by the imbalance of charge transport. The performances of devices I summarized in Table 2 are moderate and the turn-on voltages are relatively high. These results indicate that the standard device configuration with three active layers may not suitable for the present siloles.

Considering the low-lying LUMO energy levels of the new siloles, it is envisioned that they may function as electron transporter in addition to light emitter in device. To confirm this, double-layer OLEDs with a configuration of ITO/NPB (60 nm)/silole (60 nm)/LiF (1 nm)/Al (100 nm) (Device II) are fabricated, in which (MesB)₂DMTPS, (MesB)₂MPPS, or (MesB)₂HPS serve as light-emitting layer (EML) and TPBi functions as electron-transporting layer (ETL). The devices I of the new siloles radiate yellow lights, whereas the PL emissions of the films are in the green region. Taking (MesB)₂MPPS as an example, its EL maximum is located at 552 nm, which is red-shifted by 28 nm in comparison with its PL in film (Figure 7A). Such kind of red shifts could be ascribed to space charge accumulation caused by the imbalance of charge transport. The performances of devices I summarized in Table 2 are moderate and the turn-on voltages are relatively high. These results indicate that the standard device configuration with three active layers may not suitable for the present siloles.

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PL spectrum of film (524 nm) (Figure 7A), indicating that the holes and electrons get more balanced, and the exciton recombination zone has been confined inside the silole layer.

The device II adopting (MesB) 2 MPPS as both EML and ETL is turned on at a much lower voltage (3.9 V) than its device I with TPBi as ETL (7.5 V). It also shows higher current densities, and better luminance at the same voltages than device I (Figure 7B).

The maximum luminance ($L_{\text{max}}$) of device II is 13 900 cd m$^{-2}$, which is higher than that of device I (9610 cd m$^{-2}$). Figure 7C,D depict the current efficiency and external quantum efficiency as a function of luminance for both devices I and II of (MesB) 2 MPPS. It can be seen that both efficiencies of device II are advanced noticeably. Excellent maximum current ($\eta_C$), and external quantum ($\eta_{\text{ext}}$) efficiencies of 13.0 cd A$^{-1}$, and 4.12%, respectively, are attained by device II, which are almost twice as much as those of device I (6.6 cd A$^{-1}$, and 2.13%). The maximum power efficiency ($\eta_P$) of device II reaches 10.5 lm W$^{-1}$, being about four-fold higher than that of device I (2.4 lm W$^{-1}$).

Significantly, device II shows a small roll-off in EL efficiencies when luminance increases, and a high current efficiency of 7.0 cd A$^{-1}$ is reserved at a luminance of 1000 cd m$^{-2}$. This is very important because the lack of a hole-blocking layer often causes a large efficiency roll-off at high luminance range. Similarly, device II of (MesB) 2 HPS also shows better performances than device I. The turn-on voltage of device II is decreased (4.3 vs. 5.4 V), and the peak EL efficiencies are enhanced obviously ($\eta_C$: 13.9 vs 8.4 cd A$^{-1}$; $\eta_{\text{ext}}$: 4.35 vs 2.62%; $\eta_P$: 11.6 vs 4.1 lm W$^{-1}$) (Table 2 and Supporting Information Figure S3). These values are much better than those obtained from double-layer OLEDs of the reported n-type light emitters with close emission colors, such as copolymers of silafluorene and 2,1,3-benzothiadiazole (10.6 cd A$^{-1}$; 3.81%),[19] folded tetraphenylethene derivatives (7.9 cd A$^{-1}$; 3.1%),[31] dimesitylboryl-functionalized tetraphenylethene (7.13 cd A$^{-1}$; 2.7%),[26] cyano-substituted pyridine derivatives (5.4 cd A$^{-1}$),[21] Alq 3 (3.3 cd A$^{-1}$),[22] spiro-bisilole derivatives (1.98 cd A$^{-1}$),[23] and so forth.

Since both (MesB) 2 MPPS and (MesB) 2 HPS have closer LUMO energy levels to that of LiF/Al, the electron-injection is more favorable from LiF/Al to (MesB) 2 MPPS and (MesB) 2 HPS than to TPBi (Figure 8). The TPBi layer is virtually not necessary. It may even cause adverse effect to device performances because of the contact resistance at the interfaces. In addition, the excellent electron-transporting ability stemmed from the synergistic effect of silole ring and boron atoms leads to the shift of the current density-voltage characteristics to lower
voltages. At the same time, electron accumulation and exciton quenching at the cathode are reduced. The balance of holes and electrons is improved, and thus the carrier recombination probability is increased, leading to advance in EL performances.

The current and external quantum efficiencies of device II of (MesB)₂DMTPS are increased slightly relative to those of device I (Table 2, Figure S4, Supporting Information). However, the turn-on voltage is increased and the luminance is decreased. Since electron-transporting ability is not dependent solely on the energy levels of the materials, the arrangement of the molecules in solid film also functions as a crucial factor. Close packing of molecules is conducive to the carrier transport.

Table 2. EL performances of OLEDs based on (MesB)₂DMTPS, (MesB)₂MPPS, and (MesB)₂HPS. Abbreviations: \( \lambda_{\text{EL}} \) = electroluminescence maximum, \( V_{\text{on}} \) = turn-on voltage at 1 cd m\(^{-2} \), \( L_{\text{max}} \) = maximum luminance, \( \eta_c \) = maximum current efficiency, \( \eta_{\text{ext}} \) = maximum external quantum efficiency, \( \eta_p \) = maximum power efficiency, CIE = Commission Internationale de l’Eclairage coordinates.

| Emitter       | Device | \( \lambda_{\text{EL}} \) [nm] | \( V_{\text{on}} \) [V] | \( L_{\text{max}} \) [cd m\(^{-2}\)] | \( \eta_c \) [%] | \( \eta_{\text{ext}} \) [%] | \( \eta_p \) [lm W\(^{-1}\)] | CIE (x, y) |
|---------------|--------|-------------------------------|-------------------------|--------------------------------------|-----------------|-----------------------------|-----------------------------|-------------|
| (MesB)₂DMTPS  | I      | 540                           | 6.9                     | 10500                                 | 7.4             | 2.25                        | 3.2                         | 0.35, 0.55 |
|               | II     | 536                           | 6.9                     | 1810                                  | 7.8             | 2.35                        | 2.3                         | 0.35, 0.57 |
| (MesB)₂MPPS   | I      | 552                           | 7.5                     | 9610                                  | 6.6             | 2.13                        | 2.4                         | 0.40, 0.54 |
|               | II     | 520                           | 3.9                     | 13900                                 | 13.0            | 4.12                        | 10.5                        | 0.30, 0.56 |
| (MesB)₂HPS    | I      | 548                           | 5.4                     | 15200                                 | 8.4             | 2.62                        | 4.1                         | 0.39, 0.55 |
|               | II     | 524                           | 4.3                     | 12200                                 | 13.9            | 4.35                        | 11.6                        | 0.33, 0.56 |

\( ^{a} \) Device configuration: ITO/NPB (60 nm)/silole (20 nm)/TPBi (40 nm)/LiF (1 nm)/Al (100 nm) (Device I); ITO/NPB (60 nm)/silole (60 nm)/LiF (1 nm)/Al (100 nm) (Device II).
Figure 8. Energy level diagrams of multilayer EL devices of (MesB)₂MPPS with or without TPBi as electron-transporting layer.

geometry illustrated in Figure 2, we can see that (MesB)₂HPS molecules adopt a much tighter packing manner than (MesB)₂DMTPS. Thereby, it is highly possible that (MesB)₂HPS molecules could pile up more closely than (MesB)₂DMTPS molecules even in amorphous films. This renders easier electron injection and better electron transport, and hence, superior EL properties of (MesB)₂HPS to (MesB)₂DMTPS. It should also be noted that the devices have not yet been optimized thoroughly. Many factors, for example, cathode materials, carrier injection layers, hole-transporting layers, the layer thicknesses, and substrate temperature for film deposition, have subtle impacts on the balance of carrier injection, carrier mobility, carrier recombination zone and efficiency, and so on. Therefore, given the excellent optoelectronic property of the new siloles, more efficient OLEDs are expected via device engineering.

3. Conclusions

In summary, three thermally stable 2,3,4,5-tetraphenylsiloles modified with dimesitylboryl groups are synthesized, and their molecular structures and arrangement in the crystalline state are investigated. The new siloles are weakly fluorescent in solutions but become highly emissive in the aggregate state, presenting AIE characteristics. Thanks to the synergistic effect between silole ring and dimesitylboryl groups, the LUMO energy levels of the new siloles are lowered and the electron-transporting ability is improved. These merits enable them to serve simultaneously as bifunctional materials of light emitter and electron transporter in OLEDs. Efficient double-layer devices are achieved based on them, exhibiting greatly enhanced performances relative to triple-layer devices with these siloles as light emitters and TPBi as electron transporter. To the best of our knowledge, (MesB)₂MPPS and (MesB)₂HPS are among the best n-type luminescent materials used in the non-doped OLEDs. These results demonstrate an effective molecular design approach to bifunctional candidates (n-type light emitters) for OLEDs, for the sake of simplifying device configuration and reducing manufacturing cost.

4. Experimental Section

Materials and Instruments: THF was distilled from sodium benzenophenone ketyl under dry nitrogen immediately prior to use. All other chemicals and reagents were purchased from Aldrich and J&K Scientific Ltd., and used as received without further purification. 1H and 13C NMR spectra were measured on a Bruker AV 400 or AV 300 spectrometer in deuterated dichloromethane or chloroform using tetramethylsilane (TMS; δ = 0) as internal reference. UV spectra were measured on a Milton Roy Spectronic 3000 Array spectrophotometer. PL spectra were recorded on a Perkin-Elmer LS 55 spectrophotometer. High resolution mass spectra (HRMS) were recorded on a QCT premier CAB048 mass spectrometer operating in MALDI-TOF mode. Single crystal X-ray diffraction intensity data were collected on a Bruker Smart Apex CCD diffractometer with graphite monochromated Mo Kα radiation. Processing of the intensity data was carried out using the SAINT and SADABS routines, and the structure and refinement were conducted using the SHELXL suite of X-ray programs (version 6.10). TGA analysis was carried on a TA TGA Q5000 under dry nitrogen at a heating rate of 10 °C min⁻¹. The ground-state geometries were optimized using the density functional with B3LYP hybrid functional at the basis set level of 6–31G(d). All the calculations were performed using Gaussian 09 package.

Devices Fabrication: The devices were fabricated on 80 nm-ITO coated glass with a sheet resistance of 25 Ω cm⁻². Prior to loading into the pretreatment chamber, the ITO-coated glass was soaked in ultrasonic cleaning agent for 30 min, followed by spraying with de-ionized water for 10 min, soaking in ultrasonic de-ionized water for 30 min, and oven-baking for 1 h. The cleaned samples were treated by perfluoromethane plasma with a power of 100 W, gas flow of 50 sccm, and pressure of 0.2 Torr for 10 s in the pretreatment chamber. The samples were transferred to the organic chamber with a base pressure of 7 × 10⁻⁷ Torr for the deposition of NPB, emitter (and TPBi), which served as hole-transport, light-emitting (and electron-transport) layers, respectively. The samples were then transferred to the metal chamber for cathode deposition which comprised of lithium fluoride (LiF) capped with aluminum (Al). The light-emitting area was 4 mm². The current density-voltage characteristics of the devices were measured by a HP4145B semiconductor parameter analyzer. The forward direction photons emitted from the devices were detected by a calibrated UDT PIN-25D silicon photodiode. The luminance and external quantum efficiencies of the devices were inferred from the photocurrent of the photodiode. The electroluminescence spectra were obtained by a PR650 spectrophotometer. All measurements were carried out under air at room temperature without device encapsulation.

Preparation of Nanoaggregates: Stock THF solutions of the siloles with a concentration of 10⁻⁴ M were prepared. Aliquots of the stock solution were transferred to 10 mL volumetric flasks. After appropriate amounts of THF were added, water was added dropwise under vigorous stirring to furnish 10⁻⁵ M solutions with different water contents (0–90 vol%). The PL measurements of the resultant solutions were then performed immediately.

Synthesis of [2,5-Bis(4-(dimesitylboryl)phenoxy)-1,1-dimethyl-3,4-diundecylcarbene]((MesB)₂DMTPS): A solution of lithium 1-naphthalenide (LiNaph) was prepared by stirring a mixture of lithium naphthalenide (2.56 g, 20 mmol) and lithium granular (0.14 g, 20 mmol) in dry THF (30 mL) for 4 h at room temperature under nitrogen. A solution of 3 (1.3 g, 5 mmol) in THF (20 mL) was then added dropwise into the solution of LiNaph, and the resultant mixture was stirred for 30 min at room temperature. The solution was cooled to ~10 °C, into which ZnCl₂·TMEDA (6.3 g, 25 mmol) and 20 mL of THF were added. After the fine suspension was stirred for 1h, Pd(PPh₃)₄Cl₂ (100 mg) and a solution of 2 (4.04 g, 10 mmol) in 10 mL THF were added. After reflux for 12 h, the reaction was cooled to room temperature and terminated by addition of 2 M hydrochloric acid. The mixture was poured into water and extracted with dichloromethane. The organic layer was washed successively with aqueous sodium chloride solution and water and then dried over magnesium sulfate. After filtration, the solvent was evaporated under reduced pressure and the residue was purified by silica-gel column chromatography using hexane as eluent. Recrystallization gave a yellow solid product in ~80% yield. 1H NMR (400 MHz, CDCl₃, δ (ppm): 7.20 (d, J = 8.0 Hz, 4H), 7.04–6.95 (m, 6H), 6.89 (d, J = 8.0 Hz, 4H), 6.81–6.79 (m, 12H), 2.27 (s, 12H), 1.95 (s, 24H), 0.47 (s, 6H). 13C

NMR (75 MHz, CDCl3), δ (ppm): 155.0, 144.0, 143.0, 142.6, 141.8, 140.8, 138.5, 138.3, 136.3, 129.9, 128.3, 128.0, 127.3, 126.4, 23.3, 21.2, −3.9. HRMS (MALDI-TOF): m/z 910.5288 [M⁺, calcd 910.5276].

Synthesis of 2,5-Bis(4-(dimesitylboryl)phenyl)-1,1,3,4-tetraphenylsilole ((MesB)2HPS). The procedure was analogous to that described for (MesB)2MPPS. Yellow solid; yield 50%. 1H NMR (400 MHz, CDCl3), δ (ppm): 7.63 (d, J = 8.0 Hz, 2H), 7.38–7.30 (m, 3H), 7.10 (d, J = 8.0 Hz, 4H), 7.06–7.08 (m, 6H), 6.87 (d, J = 8.0 Hz, 4H), 6.82 (d, J = 8.0 Hz, 4H), 6.76 (s, 7H), 2.24 (s, 12H), 1.90 (s, 24H), 0.80 (s, 3H). 13C NMR (100 MHz, CDCl3), δ (ppm): 157.6, 143.8, 143.3, 141.8, 141.3, 140.7, 138.6, 138.3, 136.2, 134.6, 133.2, 129.9, 128.5, 128.2, 128.0, 128.0, 127.4, 126.6, 23.3, 21.2, −6.3. HRMS (MALDI-TOF): m/z 972.5442 [M⁺, calcd 972.5433].

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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