Access to tetracoordinate boron-doped polycyclic aromatic hydrocarbons with delayed fluorescence and aggregation-induced emission under mild conditions†

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Boron-doped polycyclic aromatic hydrocarbons (PAHs) have attracted ongoing attention in the field of optoelectronic materials due to their unique optical and redox properties. To investigate the effect of tetracoordinate boron in PAHs bearing N-heterocycles (indole and carbazole), a facile approach to four-coordinate boron-doped PAHs was developed, which does not require elevated temperature and pre-synthesized functionalized boron reagents. Five tetracoordinate boron-doped PAHs (NBNN-1–NBNN-5) were synthesized with different functional groups. Two of them (NBNN-1 and NBNN-2) could further undergo oxidative coupling reactions to form fused off-plane tetracoordinate boron-doped PAHs NBNN-1f and NBNN-2f. The investigation of photophysical properties showed that the UV/vis absorption and fluorescence emission are significantly red-shifted compared to those of the three-coordinate boron-doped counterparts. In addition, the emission of NBNN-1–NBNN-3 consisted of prompt fluorescence and delayed fluorescence. The compounds NBNN-1f and NBNN-2f showed aggregation-induced emission.

Introduction

Chemically doping heteroatoms into aromatic compounds has proven to be one of the most effective methods to tune their optoelectronic properties and intermolecular interactions.1–12 Due to the isoelectronic relationship between boron and carbon cations and between the boron–nitrogen (BN) unit and two carbon atom (CC) motif, boron becomes an important element that can induce novel and complementary properties.13–21 BN-doped aromatic compounds have a long history of evolution since the first example of BN-arene was reported by Alfred Stock in 1926.22 Much the same to many other topics of chemistry, not much attention was paid until the new millennium. Nowadays, the synthesis of BN-doped arenes has become one of the most popular topics in organic chemistry and materials chemistry, and numerous new BN-doped aromatic compounds have been synthesized and demonstrated to have very distinct properties in optoelectronic materials.23–27 Representative applications of B-doped polycyclic aromatic hydrocarbons (PAHs) have been demonstrated in organic field-effect transistors (OFETs)21,28–30 and organic light emitting diodes (OLEDs).25,29,31–37 Particularly, since PAHs decorated with para-positioned boron and nitrogen atoms have been reported to show thermally activated delayed fluorescence (TADF) with narrow full-width at half-maximum (FWHM) of emission based on efficient HOMO–LUMO separation by the multiple resonance effect in 2016,31 BN-doped PAHs have been extensively studied for the next-generation OLEDs.

With the further development of BN-PAH chemistry by advances in the synthetic methodology and by the application potential of BN-doped π-conjugated systems, attention to the structure–property relationship has arisen.38,39 The location of the BN unit was found to be important for the optoelectronic properties because whether the BN unit could fully conjugate with PAHs is highly dependent on its locations.39–44 Although both the number of BN units45–50 and different orientations of BN units51–54 are believed to exhibit non-negligible impact on the optoelectronic properties and intermolecular interactions, only a few of examples have been documented due to the limited synthetic methodology.

In addition, the majority of previous attention has been paid to the tricoordinate boron units in PAHs.55–66 Tetracoordinate boron has been viewed more as a functional group instead of a dopant.61–66 In fact, the coordination number of the boron atom shows noteworthy impact on the photophysical
Results and discussion

For the synthesis of tetracoordinate boron-doped PAHs NBNN-1–NBNN-5, we started from the commercially available 2,6-dibromopyridine, which could undergo the Suzuki coupling reaction with readily available 1-bipin-carbazole and 7-bpin-indole to produce the precursor compounds 1a and 1b in 90% and 46% yields, respectively (Scheme 2).

Our first attempt to introduce the boron atom using BBr3 as the boron source with 1Pr2EtN in o-dichlorobenzene (o-DCB) at 180 °C was successful to produce phenyl group functionalized BN-PAH NBNN-1 when phenylmagnesium bromide was used as the nucleophilic reagent. The existence of o-DCB makes the purification difficult. We tried to use several other solvents with lower boiling points at lower reaction temperatures, all of which generated the target compound NBNN-1 with similar yields around 70%. Usually, the presence of the additive 1Pr2EtN is proven to be essential for the electrophilic borylation, but the absence of 1Pr2EtN did not show any influence on the borylation yield. To our delight, the reaction temperature was finally lowered down to room temperature without 1Pr2EtN in 78% yield (Scheme 2). In addition, the kind of nucleophile was also a problem, and lithium reagents like phenyl lithium could produce compound NBNN-1 in slightly lower yield. While we were preparing the manuscript, the synthesis of NBNN-1 was reported at high temperature with PhBCl2 and the additive 1Pr2EtN. Our mild conditions are probably ascribed to the existence the pyridyl group which could act as a directing group coordinating with BBr3 and make the following borylation of N–H bonds less challenging. This process does not require to break the C–H bonds either. In addition, compared to the pre-functionalized boron reagents RBX2 (X = Cl, Br), the commercially available BBr3 has higher Lewis acidity and reactivity which also play a non-negligible role in the borylation reaction. The mild conditions without using high boiling point solvent make the purification of the product less disturbing. With the optimal conditions in hand, ethynylmagnesium bromide and mesitylmagnesium bromide were used as nucleophiles to generate compounds NBNN-2 and NBNN-3 in 58% and 69% yields, respectively. Compound NBNN-4 was obtained when the reaction was quenched by water without adding any nucleophiles in single-figure yield, however, the yield of NBNN-4 was significantly improved to 46% when 1Pr2EtN was used to quench

![Scheme 1](image1.png)

Representative examples of tetracoordinate B-doped PAHs (a), our previous work (b) and chemical structures of this work (c).
the reaction. We also tried other hydroxy group containing reagents like phenol, which did not give the corresponding product at all. Compound NBNN-5 based on two indole fragments was also synthesized under the optimal conditions. Other nucleophiles like isopropylmagnesium bromide, vinylmagnesium bromide, and methyl lithium could also yield the corresponding tetracoordinate boron-doped PAHs, however, these compounds were slowly decomposing on the silica gel or alumina. Bulky nucleophiles like butyllithium did not react, probably due to the steric hindrance. Compounds NBNN-1 to NBNN-3 were fully characterized by NMR and HRMS spectroscopy. The molecular structures of NBNN-1 and NBNN-3 were further determined by single-crystal X-ray diffraction analysis and are shown in Fig. 1.

Similar to our previous work on synthesized tricoordinate boron-doped PAHs, the proximity and the steric repulsion between the hydrogen atoms of flanking units in NBNN-1 to NBNN-4 make them undergo the oxidative coupling reaction to form the fused forms. By combining 2,3-dichloro-5,6-dicyano-1,4-benzoquinone (DDQ) as the oxidant and TFOH as the acid, the fully fused compounds NBNN-1f, NBNN-2f and NBNN-4f were successfully obtained. Compounds NBNN-1f and NBNN-2f were obtained in isolated yields of 78% and 58%, respectively (Scheme 3). Silica gel or alumina column chromatography was not compatible with NBNN-4f, which decomposed on the silica gel or alumina and was only confirmed by HRMS. NBNN-3 did not undergo the oxidative coupling reaction probably due to the longer distance (3.356 Å, Fig. 1) between two carbon atoms on the carbazole segments. Unlike the tricoordinate boron-doped PAH analogue (E in Scheme 1), the crystal structure of NBNN-1f showed that the four-coordinated boron atom adopts a tetrahedral geometry, which makes the molecular structure off-plane (Fig. 2). The B1–N1, B1–N2, and B1–N3 bond lengths are 1.640 Å, 1.523 Å, and 1.502 Å, respectively. Similar to tricoordinate boron-doped PAH E, the π-stacking interactions of NBNN-1f are limited to two neighbouring molecules with a longer separation distance than E.

The absorption, emission and electrochemical data of these seven tetracoordinate boron-doped PAHs are shown in Fig. 3 and Table 1. The absorption peaks are in agreement with the colors of their THF solutions (Fig. 3b). The general trend of the experimental absorption bands agrees with the time-dependent density functional theory (TD-DFT) calculations (Fig. 4, and see the ESI† for details). The broad absorption bands of NBNN-1 to NBNN-5 between 400 nm and 520 nm are ascribable to S0 → S1 transitions of NBNN-1f and NBNN-4f, with the major transition configuration of HOMO → LUMO and S0 → S2 with the major transition configuration of HOMO → 1 → LUMO. Due to the closer energy gap between the HOMO and HOMO → 1, NBNN-1, NBNN-2 and NBNN-5 only exhibited one peak at 469 nm, 465 nm and 446 nm, respectively. The smaller HOMO–LUMO gap results in red-shifted absorption peaks of NBNN-3, while the larger HOMO–LUMO gap brings about the blue-shifted absorption peaks of NBNN-5. The strong absorption peaks at 310 nm for NBNN-1 to NBNN-4 are likely attributed to the larger conjugation of the carbazole unit than the indole unit in NBNN-5. NBNN-1f and NBNN-2f showed five absorption bands with a maximum at λmax = 531 nm and 528 nm which could be attributed to S0 → S1 with the transition configuration of HOMO → LUMO, respectively. Different from NBNN-1 to NBNN-5, the S0 → S2 transitions of NBNN-1f at 457 nm and NBNN-2f at 439 nm consisted of HOMO → LUMO+1 with charge transfer character. Except NBNN-5, molecules NBNN-1 to NBNN-4 showed near-identical fluorescence spectra with emission peaks between 511 nm and 564 nm, which suggests that the substituents on the boron atom of tetracoordinate boron-doped PAHs have a negligible contribution to fluorescence. The fluorescent peaks in the poly(methyl methacrylate) (PMMA) film of NBNN-1 to NBNN-4 are very similar to those in THF solution, while the λem of NBNN-5 in solid was red-shifted to 540 nm from bright green color to greenish yellow color as shown in Fig. 3b, which is probably because the less crowded structure of NBNN-5 results in the π–π packing in the solid. The quantum yields both in solution and in the PMMA film of the open form of compounds bearing carbazole units are ~25%, except NBNN-1,

![Scheme 3](Image 14x290 to 26x354)

**Scheme 3** Formation of fused tetracoordinate boron-doped PAHs via the oxidative coupling.
which has much higher quantum yields in solution ($\Phi_{pl} = 62.3\%$) and PMMA film ($\Phi_{pl} = 55.7\%$) (Table 1). Compound \textbf{NBNN-5} bearing indole units only has less than 10% quantum yield both in solution and in solid probably due to vibrational relaxation caused by the less crowded structure. Compared to \textbf{NBNN-1} and \textbf{NBNN-2}, the emission peaks of the fused molecules \textbf{NBNN-1f} and \textbf{NBNN-2f} show remarkable red-shift up to 100 nm and exhibit indistinguishable fluorescent spectra ($\lambda_{em} = 660$ nm, $\Phi < 1\%$), while their absorption spectra are also similar with different absorption extinction coefficients. The fluorescence peak of \textbf{NBNN-1f} in the PMMA film was blue-shifted up to 39 nm compared to that in solution (see the ESI† for details), while \textbf{NBNN-2f} in the PMMA film exhibited two emission peaks at 624 nm and 659 nm. The blue-shifted emission peaks of \textbf{NBNN-1f} and \textbf{NBNN-2f} are probably ascribed to the structure constraint, which could also result in aggregation-induced emission (AIE). Both \textbf{NBNN-1f} and \textbf{NBNN-2f} showed a significant increase in quantum yields in solid up to 4%. The low quantum yields of \textbf{NBNN-1f} and \textbf{NBNN-2f} are likely due to the internal conversion between small $S_0\rightarrow S_1$ gaps. The fluorescence peaks in solution and solid are consistent with the emission colors of their THF solutions and PMMA films (Fig. 3b). Ligand 1a has two absorption bands at 294 nm and 364 nm, while Ligand 1b has a broad band at 330 nm and a less intensive band at 364 nm (Fig. S5.16†). These two molecules 1a and 1b showed weak fluorescence with emission maxima at 636 nm and 656 nm, respectively. The quantum yields both in solution and in the PMMA film of the two molecules were less than 0.1%. Compared to the ligands, the boron-doped PAHs exhibited substantially red-shifted absorption spectra with enhanced emission quantum yields. As shown in Table 1 and Fig. S5.5–S5.11† the phosphorescence decays at 77 K were recorded in the range of 200–400 ms for \textbf{NBNN-1–NBNN-5}, while the fused compounds \textbf{NBNN-1f} and \textbf{NBNN-2f} showed much shorter phosphorescence decays (0.012 ms and 5.62 ms, respectively).

The time-resolved photoluminescence spectra were recorded in THF to investigate the photoluminescence decay behaviors of these tetracoordinate boron-doped PAHs. As shown in Table 1 and Fig. 5, compounds \textbf{NBNN-1}, \textbf{NBNN-2} and \textbf{NBNN-3} consisted of prompt fluorescence with $\tau_{pl} = 1.26$ ns, 1.89 ns and 1.60 ns and delayed emission with $\tau_{df} = 38.06$ μs, 8.15 μs and 24.8 μs, respectively. The fully fused compounds \textbf{NBNN-1f} and \textbf{NBNN-2f}, the carbazole-based compound with a hydroxy group on boron \textbf{NBNN-4}, and indole-based compound \textbf{NBNN-5} only exhibited the prompt fluorescence component ($\tau_{pl} = 1.07$ ns, 1.20 ns, 4.69 ns, 1.79 ns, respectively), likely due to the relatively large singlet-triplet band gap.

To further investigate the delayed fluorescence behaviors of \textbf{NBNN-1–NBNN-3}, $\Phi_{pl}$ and the lifetimes of the prompt/delayed components ($\tau_{pl}/\tau_{df}$) were exploited to estimate the photo-physical rate constant of radiative decay ($k_r$), intersystem crossing ($k_{ISC}$) and reverse intersystem crossing of ($k_{RISC}$).44 As shown in Table 2, the $k_r$ value of \textbf{NBNN-2} (8.62 × 10$^{-6}$ s$^{-1}$) is larger than those of \textbf{NBNN-1} (2.38 × 10$^{-7}$ s$^{-1}$) and \textbf{NBNN-3} (3.94 × 10$^{-7}$ s$^{-1}$). $k_{ISC}$ (7.70 × 10$^{-6}$ s$^{-1}$) and $k_{RISC}$ (5.35 × 10$^{-7}$ s$^{-1}$) of tetracoordinate boron-doped PAH \textbf{NBNN-1} with a phenyl group on the boron atom are the largest among those of \textbf{NBNN-1 to NBNN-3}. The $k_r$ value of \textbf{NBNN-2} (8.62 × 10$^{-6}$ s$^{-1}$) is larger than those of \textbf{NBNN-1} (2.38 × 10$^{-7}$ s$^{-1}$) and \textbf{NBNN-3} (3.94 × 10$^{-7}$ s$^{-1}$). It is not clear why the different functional groups on the boron atom of the same skeleton could dramatically change $k_{RISC}$.

Molecules \textbf{NBNN-1} to \textbf{NBNN-5} displayed pseudo-reversible reduction peaks at $\sim$2.0 V (vs. Fe/Fc') and multiple reversible oxidation peaks with the first oxidation peaks at $\sim$0.3 V (vs. Fe/Fc') (Table 1 and ESI†). The fused forms of boron-doped PAHs \textbf{NBNN-1f} and \textbf{NBNN-2f} show almost identical cyclic voltammograms as shown in Fig. 3c. The first oxidation peaks of \textbf{NBNN-1f} and \textbf{NBNN-2f} are lower than those of \textbf{NBNN-1} and \textbf{NBNN-2}, which can be attributed to the high-lying HOMO level (Fig. 4 bottom). The experimental data are in good agreement...
with the calculated HOMO and LUMO. Our previous study on tricoordinate boron-PAHs did not show such rich electrochemical properties. After we measured the CV curves of the ligand 1a and 1b (Fig. 3c, S2.8 and S2.9), we found that the ligand 1a had similar multiple reversible oxidation peaks to the tetracoordinate boron-doped PAHs NBNN-1, NBNN-1f and NBNN-2f, as shown in Fig. 3c, which indicates that the ligand is likely ascribed to the multiple oxidation of NBNN-1, NBNN-1f and NBNN-2f rather than the introduction of the tetracoordinate boron atom. Compared to the reduction of ligands, the tetracoordinate boron-doped PAHs NBNN-1, NBNN-1f and NBNN-2f showed irreversible reduction at ~−2.00 V (vs. Fc/Fc⁺), which indicates that the reduction peaks might originate from the boron atom.

Unlike the tricoordinate boron-doped PAH analogues in which the boron atom has a significant contribution to the LUMO and the HOMO/LUMO spread over the entire molecule, the four-coordinate boron atom in NBNN-1 to NBNN-5, NBNN-1f and NBNN-2f showed irreversible reduction at ~−2.00 V (vs. Fc/Fc⁺), which indicates that the reduction peaks might originate from the boron atom.

Table 1 Photophysical and electrochemical data of NBNN-1 to NBNN-5, NBNN-1f and NBNN-2f

|        | UV/visa | Fluorescence | Phosphorescenceb | Electrochemical datac |
|--------|---------|--------------|------------------|-----------------------|
|        | λabs (nm) | λem a (nm)/λem b (nm) | Φpl a,d (%) | τpl a (ns)/τpl b (µs) | λem (nm) | τph (ms) | Ered (V) | Eox (V) |
| NBNN-1 | 310, 469 | 557/556 | 62.3/55.7 | 1.26/38.06 | 569 | 302 | −2.02 | 0.30 |
| NBNN-1f | 341, 376, 417, 457, 531 | 660/621 | 0.8/4.0 | 1.07/— | 643 | 0.012 | −1.93 | 0.13 |
| NBNN-2 | 308, 465 | 558/562 | 23.4/22.6 | 1.89/8.15 | 567 | 270 | −2.16 | 0.26 |
| NBNN-2f | 332, 359, 404, 439, 528 | 659/624, 659 | 0.8/4.0 | 1.20/— | 638 | 5.62 | −1.96 | 0.19 |
| NBNN-3 | 309, 445, 481 | 564/561 | 24.5/26.6 | 1.60/24.8 | 583 | 213 | −2.09 | 0.27 |
| NBNN-4 | 304, 416, 461 | 551/556 | 34.8/25.3 | 4.69/— | 582 | 377 | −2.06 | 0.28 |
| NBNN-5 | 298, 370, 446 | 523/540 | 8.3/9.5 | 1.79/— | 569, 621 | 355 | — | — |

a 10⁻⁵ M in THF at 298 K. b PMMA film (10 wt%). c vs. Fc/Fc⁺ (ferrocene/ferrocenium) glassy carbon as the working electrode, a Pt wire as reference and counter electrodes, ferrocene (Fc/Fc⁺ = ferrocene/ferrocenium) was added to the solution as the internal reference, scan rate = 100 mV s⁻¹. d Quantum yields were determined using a calibrated integrating sphere. e Not available.

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accompanied by the blue-shift of emission (Fig. 6c). The photoluminescence decays only slightly increased when the water fraction increased. In addition, we found that the AIE behavior of NBNN-1f was accompanied by the blue-shift of emission (Fig. 6c). The quantum yields for prompt ($\Phi_{pf}$) and delayed ($\Phi_{df}$) fluorescence, $\Phi_{pf} + \Phi_{df} = \Phi_{ps}$. The rate constant of fluorescence radiative decay ($k_f = \Phi_{pf}/\tau_{pf}$). Rate constant of ISC ($S_1 \rightarrow T_1$): $k_{ISC} = (1 - \Phi_{pf})/\tau_{pf}$. Rate constant of RISC ($T_1/S_1$): $k_{RISC} = \Phi_{df}/k_{r}T_{1}\tau_{pf}\Phi_{ps}$.

| Cmpd  | $\tau_{pf}$/($\phi_{df}$) (µs)/$\phi_{df}$ (%) | $k_f$ ($10^6$ s$^{-1}$) | $k_{ISC}$ ($10^8$ s$^{-1}$) | $k_{RISC}$ ($10^8$ s$^{-1}$) |
|-------|-----------------------------------------------|--------------------------|----------------------------|-----------------------------|
| NBNN-1 | 1.26/38.06/3.0/59.3  | 2.38                      | 7.70                       | 5.35                        |
| NBNN-2 | 1.89/8.15/16.3/7.1  | 8.62                      | 4.43                       | 0.64                        |
| NBNN-3 | 1.60/24.8/6.3/18.2  | 3.94                      | 5.86                       | 1.24                        |

Table 2 The delayed fluorescence properties of NBNN-1 to NBNN-3 in THF solution

Measured to be 10$^{-3}$ M in THF for NBNN-1 to NBNN-2 at 298 K. $^b$ Emission lifetime for prompt ($\tau_{pf}$) and delayed ($\tau_{df}$) fluorescence. $^c$ Quantum yields for prompt ($\Phi_{pf}$) and delayed ($\Phi_{df}$) fluorescence, $\Phi_{pf} + \Phi_{df} = \Phi_{ps}$. $^d$ Rate constant of fluorescence radiative decay ($S_1 \rightarrow \Phi_0$); $k_f = \Phi_{pf}/\tau_{pf}$.

1f and NBNN-2f have no contribution to the LUMO (Fig. 4), and their LUMO is mainly located on the pyridine and phenyl rings of carbazole and indole connected to pyridine. The calculated HOMO and LUMO prove that the substituents on the four-coordinate boron atom have trivial adjustment on the frontier molecular orbitals. Compared to NBNN-1 and NBNN-2, NBNN-1f and NBNN-2f have smaller HOMO–LUMO gaps due to the notably elevated HOMO and slightly lowered LUMO, which could indicate the reason why the fused product NBNN-4f of NBNN-4 was less stable and slowly decomposed on silica gel or alumina (Scheme 3). We note that the electrostatic potential maps of NBNN-1–NBNN-5, NBNN-1f and NBNN-2f display the development of negative characteristics on the tetracoordinate boron area and positive characteristics on the pyridine unit (Fig. 4 top).

To illustrate the aromaticity of these tetracoordinate boron-doped PAHs, the nucleus-independent chemical shift (NICS(1)) of two C3NBN rings and the 7-memebered C4NBN ring (if exists) around boron atoms was calculated (Fig. 6a, b and ESI†). The two C3NBN rings in NBNN-1–NBNN-5 have positive NICS(1) values (see the ESI†), supporting their antiaromaticity. The two C3NBN rings in NBNN-1f and NBNN-2f have much more positive NICS(1) values than those of NBNN-1–NBNN-5, indicating that they are more antiaromatic. Similar to the tricoordinate boron-doped PAH, the 7-membered C4NBN has a larger NICS(1) value of 13.8 ppm (Fig. 6b). The anisotropy of the induced current density (ACID) calculations (Fig. 6a and b) that are based on the optimized structures further showed that there is no electron density around the boron atom and the central 13-membered C4NC3NC3N ring of NBNN-1f is antiaromatic with an anti-clockwise diatropic ring current circuit.

The fused molecules NBNN-1f and NBNN-2f exhibited AIE behavior, while compounds NBNN-1–NBNN-5 only have normal photoluminescence properties. AIE behaviors of NBNN-1f and NBNN-2f are probably due to their structure constraint. We used NBNN-1f as an example to investigate the AIE phenomenon in detail (Fig. 6e–f). The red emission became stronger after the water content reached 50% in volume in THF, and the emission intensity grew faster when water was further added, as shown in Fig. 6e and e. The THF solution with 90% water displayed bright red emission. The quantum yields gradually increased when the water fraction increased. In addition, we found that the AIE behavior of NBNN-1f was accompanied by the blue-shift of emission (Fig. 6c). The photoluminescence decays only slightly increased when the water fraction increased.

Fig. 6 (a) and (b) ACID plots of NBNN-1 and NBNN-1f, and numbers in red representing NICS(1) in ppm. (c) Emission spectrum change of NBNN-1f in THF with different H2O fractions (vol%). (d) Transient fluorescence curves in THF with different H2O fractions (vol%). (e) Photographs showing the emission intensity increase with increasing H2O fraction. (f) Photographs showing one drop of NBNN-1f solution on a thin-layer chromatography plate with different evaporation timescales.
fractions reached 70% and 90%, as shown in Fig. 6d. The AIE behavior of **NBNN-1f** was further demonstrated through change of emission intensity after adding one drop of **NBNN-1f** THF solution on a thin-layer chromatography plate (Fig. 6f). We observed a notable increase of the red emission under 365 nm UV lamp irradiation.

**Conclusions**

We developed an efficient strategy to construct a series of tetracoordinate boron-doped PAHs with various substituents on boron atoms at room temperature. Some of them were successfully converted to fully fused four-coordinate boron-doped NBN-PAHs via oxidative coupling. The tetracoordinate boron-doped PAHs exhibited delayed fluorescence. All the rings with tetracoordinate boron atoms are aromatic. The fused NBN-PAHs displayed blue-shifted AIE behavior.

**Data availability**

The data sets supporting this article have been uploaded as part of the ESI.

**Author contributions**

L. J., B. Z. and D.-T. Yang conceptualized the work. L. J. developed the methodology and designed experiments. Y. W. recorded the time-resolved emission spectra. D. T. synthesized **NBNN-3**. X. C. and T. M. performed DFT calculations. D.-T. Y. wrote the original draft and created the original visuals. L. J., B. Z. and D.-T. Y. edited the draft and visuals.

**Conflicts of interest**

There are no conflicts to declare.

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