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BN-Substitution in Dithienylpyrenes Prevents Excimer Formation in Solution and in the Solid State

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interactions such as π - π -stacking, dipolar interactions, or the formation of dimers in the excited state. Pyrene is the most prominent example showing the latter as it exhibits a broadened and strongly bathochromically shifted emission band at high concentrations in solution compared to the respective monomers. In the solid state, the impact of intermolecular interactions is even



higher as it determines the crystal packing crucially. In this work, a thiophene-flanked BN-pyrene (BNP) was synthesized and compared with its all-carbon analogue (CCP) in solution and in the solid state by means of crystallography, NMR spectroscopy, UV-vis spectroscopy, and photoluminescence (PL) spectroscopy. In solution, PL spectroscopy revealed the solvent-dependent presence of excimers of CCP at high concentrations. In contrast, no excimers were found in BNP. Clear differences were also observed in the single-crystal packing motifs. While CCP revealed overlapped pyrene planes with centroid distances in the range of classical π -stacking interactions, the BNP scaffolds were displaced and significantly more spatially separated.

INTRODUCTION

Boron-nitrogen-substituted aromatic systems have attracted great interest over the last decade as this formal replacement offers a method to significantly change the physical, chemical, and optoelectronic properties of the molecules. However, the geometrical structures remain similar, at least in the gas phase or in a non-aggregated form in solution.^{1–}

Consequently, the research on the potential applications of BN-aromatics has intensified, and the main emphasis of the current research has especially been placed on the fields of materials science^{5,6} and organic electronics.⁷⁻¹⁰ Other areas have also started to benefit from these developments in the main group chemistry. For instance, BN-substituted pharmaceuticals,¹¹ the BN-analogues of polystyrene and poly-(vinylnaphthalene) with small aromatic BN-units as side groups,^{12–15} and BN-aromatics as ligands for the main group elements and transition metals were investigated.¹⁶⁻¹⁸ The increased interest has prompted the development of numerous synthetic approaches giving facilitated access to substituted BNaromatics.^{19–24}

A BN-substitution in polycyclic aromatic hydrocarbons (PAHs) may not only alter the electronic structure of the single molecules but can also induce the changes in the morphology and aggregation behavior. This in itself results in strongly divergent optical properties²⁵ and provides a powerful tool for synthesizing materials with a precisely attuned photophysical behavior.

Recently, the first organic photovoltaics,¹⁰ organic field-effect transistors,^{7,26} and organic light-emitting diodes (OLEDs)^{8,9} containing BN-substituted compounds have been reported. Compared to the carbonaceous analogues, the highest occupied molecular orbital (HOMO)-lowest unoccupied molecular orbital (LUMO)-gaps of monocyclic azaborinines are usually decreased,²⁷ while fused-ring BN-systems do not allow such a generalized statement. On the one hand, the majority of reports showed that a BN-unit induces a slight decrease of the optical band gaps in comparison with all-carbon analogues.^{25,28–30} On the other hand, several BN-acenes with particularly stabilized HOMO levels have been presented, which are consequently less

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Figure 1. Four series of boron-nitrogen-substituted pyrenes as presented so far.

prone to oxidation.^{31,32} Furthermore, a number of studies showed largely increased quantum yields of BN-PAHs due to decreased non-radiative photochemistry pathways³³ and excellent thermal stabilities up to 400 °C, which are comparable with their carbonaceous congeners, illustrating the high application potential of these structures.^{29,32,34–38} The position of the BN-motif inside the molecule proved to be a crucial determinant of the above-mentioned photophysical properties.^{39–41}

One of the best investigated PAHs is pyrene, which undergoes photoluminescence (PL) in the visible spectrum ($\lambda_{\rm em\ monomer}$ = 384 nm in 0.01 mg mL⁻¹ cyclohexane solution, full band halfwidth: 1900 cm⁻¹) with moderate quantum yields of its unsubstituted form ($\varphi = 0.32$).^{42,43} With the aim of tuning its optical properties, enormous effort has been made to develop reactions to selectively substitute pyrene asymmetrically and symmetrically.^{44–47} In fact, substituted pyrenes may display bathochromically shifted emission bands and largely elevated quantum yields.^{48,49} Therefore, they have become prominent target structures for a variety of applications in materials science.^{50–54}

Pyrene is the textbook example of a PAH that forms excited dimers (excimers) at increased concentrations, resulting from the close interaction between one pyrene molecule in the electronically excited state and the other one in the ground state in coplanar orientation. The unusually high monomer lifetime (up to $\tau_{f_r \text{ monomer}} = 680 \text{ ns}$ in degassed solutions),⁵⁵ caused by a forbidden $S_1 \rightarrow S_0$ transition, favors diffusion-controlled excimer formation.⁵⁶ The other key factors are the planar and rigid geometry of the molecule⁵⁷ as well as the strongly stabilizing short- and long-range interactions of the excited and non-excited pyrene in the optimum stacking orientation.⁵⁸ Consequently, excimer formation occurs before the excited monomer would decay into the electronic ground state. Overall, these determinants induce a low excimer formation energy of pyrene (0.34 eV).⁵⁵

Pyrene excimers show characteristic red-shifted ($\Delta \lambda_{\rm em} = 105$ nm), broadened, and featureless PL emission bands ($\lambda_{\rm em}$ excimer = 489 mm in 1.0 mg mL⁻¹ cyclohexane solution, ⁴³ full band half-width: 3800 cm⁻¹) and separate lifetime components ($\tau_{\rm excimer} = 90$ ns).^{55,58} Numerous works investigated the aggregation-dependent emission and showed that pyrenes may serve as fluorescence probes in biomolecules,⁵⁹ for the detection of environmental changes such as the pH value and temperature^{60,61} or the detection of guest molecules such as heavy metal ions.^{62,63}

However, for many optoelectronic applications, excimer formation can be disadvantageous. Above all, it promotes fluorescence quenching⁶⁴ and limits exciton transfer processes in organic semiconductors.⁶⁵ Furthermore, the broadened shape of the emission spectrum can be undesirable if an OLED with a distinctly specified emission wavelength is desired⁴⁵ because this feature prevents a precise tuning of the emission color and may decrease the fluorescence efficiency in the solid or film state.⁴⁵ Therefore, the number of patents featuring pyrene and its derivatives is lower⁶⁶ compared to other molecules commonly applied in optoelectronics, despite its structural simplicity. In contrast, in pyrenes and other PAHs with suitably large and/or sterically demanding substituents, excimer formation may be suppressed as a close approximation of the π -planes is impeded for steric reasons and the overlap of the π -planes is incomplete.49,58,67-69

To date, four types of BN-substitution patterns in pyrenes have been reported, with one or two BN-moieties in peripheral or internal positions (Figure 1).⁷⁰⁻⁷⁴ The different isomers impressively illustrated the strong influence of the position of the BN-substitutions on the electronic structure and the optical properties as mirrored by very different absorption maxima. According to the discussed unique aggregation behavior of pyrene, the question arises how a BN-pyrene can be characterized in terms of excimer formation capability in solution and the morphologies in the solid state compared to its all-carbon analogue. One of the studies⁷¹ described that sterically demanding mesityl-substituents at the boron atom as in molecule 3 impeded the formation of excimers. Another study reveals that a dimer of 4 did not form excimers in solution, as well.⁷⁵ However, in both cases, it was not evaluated in what manner the BN-unit contributed to this effect compared to a CC-pyrene with an equal substitution pattern.

In the pioneering synthesis of the internally BN-substituted pyrenes by Piers and co-workers,⁷⁰ unsubstituted phenyl- and alkyl-substituted BN-pyrenes (1a-e) were presented (Figure 1).

While the crystal structure of the unsubstituted BN-pyrene **4a** shows a head-tail packing due to the compensation of internal dipoles, an unsubstituted CC-pyrene exhibits a sandwich herringbone pattern.⁷⁶ Alkyl-substituted BN-pyrene **4e** exhibits parallel sheets with a negligible overlap and therefore a lack of π -stacking. This is considered an unfavorable feature with respect to optoelectronic applications as it lowers the solid-state charge carrier mobilities.⁷⁷

Scheme 1. Synthesis of BNP Involving the Concluding Reaction of Ethynylpyridine 7 and Boracycle 8



Scheme 2. Synthesis of CCP Based on a Stille Cross-Coupling Reaction and Subsequent Electrophilic Ring Closure of Biphenyl 15



Based on these works, we hypothesized that the introduction of more suitable substituents should overcome these limitations and implement additional benefits. On the one hand, the potential side groups should not add a large steric hindrance to the system as this might lead to a reduced stacking behavior. On the other hand, the substituents should significantly improve the solubility in organic solvents since PAHs of this size often suffer from low solubility. Furthermore, our goal was to extend the conjugated system and assess how this alteration would affect the absorption and emission maxima. To retain the possibility of post-functionalizations, the substituent should also be easily modifiable. Based on these design principles, we selected thienyl substituents as they fulfill these requirements best and exhibit well-established reactivity.

In this work, we describe the synthesis of internally BNsubstituted 5,9-dithienyl-10b-aza-10c-borapyrene **BNP** and its respective CC-analogue **CCP**. Furthermore, we present an indepth investigation of their properties and aggregation dynamics in solution and in the solid state by means of crystallography, NMR and diffusion-ordered NMR spectroscopy (DOSY), stationary and steady-state absorption and PL spectroscopy, and time-correlated single-photon counting (TCSPC). Our mechanistic hypotheses were further supported by density functional theory (DFT) calculations.

RESULTS AND DISCUSSION

Syntheses. The synthetic route toward **BNP** is based on a report by Piers and co-workers (Scheme 1).⁷⁰ The syntheses were initiated by a Sonogashira cross-coupling reaction of dibromopyridine **5** and ethynylthiophene **6**, affording bis-(ethynylthienyl)pyridine 7 in a yield of 53%. Boracycle **8**, which was required for the final formation of **BNP**, was synthesized in four steps following the literature procedures.

For that purpose, diethynylpentane 9^{78} and di(*n*-butyl)stannane (10)⁷⁹ were reacted to form the stannacycle 11.⁸⁰ A transmetalation of the latter with boron trichloride afforded boracycle 8 (for experimental details, see the Supporting Information, Section 2).⁷⁸ After the reaction of pyridine 7 and the boracycle 8 including a platinum-catalyzed electrophilic cyclization at 115 °C for 15 h in a microwave reactor, the desired BNP was isolated as a yellow crystalline solid in a yield of 60%.

The synthesis of **CCP** is based on a procedure presented by Schreiner and co-workers (Scheme 2).⁸¹ The first key intermediate to be synthesized was 2,6-dibromo-1,1'-biphenyl (13). A Suzuki–Miyaura cross-coupling of 1,3-dibromo-2iodobenzene (12)⁸² and phenylboronic acid afforded biphenyl 13 in a yield of 59%. For the synthesis of the complementary nucleophile 14, a trimethylstannyl group was attached to ethynylthiophene 6 via lithiation, followed by transmetalation with trimethyltin chloride, yielding the stannane 14 in 45% yield. The tin functionality was chosen to access the sterically hindered positions at biphenyl **13** and to ensure a complete conversion into the desired product **15**. Via a Stille crosscoupling reaction between biphenyl **13** and thiophene **14**, bis(thienylethynyl)biphenyl **15** was synthesized in an excellent yield. The concluding electrocyclic ring closure was then performed by heating the reactants with platinum(II) chloride in toluene for 2 d at 120 °C, affording **CCP** as a colorless solid in 36% yield.

While **CCP** was stable under ambient conditions for an indefinite period, **BNP** underwent slow decomposition, which was accelerated in nucleophilic solvents such as water or methanol.

Crystal Structure Analysis. Needle-shaped crystals of **BNP** and **CCP** were obtained by slow evaporation of toluene. The crystal structure of **BNP** (CCDC deposition number: 2040669) was determined via a conventional single-crystal X-ray diffraction experiment at 100 K using a Mo- K_{α} in-house microfocus source.

To obtain the structure of **CCP** (CCDC deposition number: 2040670), a synchrotron X-ray diffraction experiment was conducted at 20 K using the beamline BL02B1 of SPring-8, Japan. For further crystallographic and measurement details, see the Supporting Information (Section 4).

The pyrene scaffolds of BNP and CCP are planar, and the orientations of the peripheral thiophene rings with respect to the pyrene scaffolds are similar ($\varphi = 45^{\circ}$ for **BNP** and $\varphi = 48^{\circ}$ for CCP, Figure 2a,d). However, the dihedral angles between the two planes of the thienyl units differ significantly ($\varphi = 81.8^{\circ}$ for **BNP** and $\varphi = 28.5^{\circ}$ for **CCP**). This has an enormous impact on the crystal packing motifs. Although both BNP and CCP form herringbone patterns such as the reported alkyl-substituted BNpyrene 4e and the unsubstituted pyrene, the moleculemolecule stacking behavior differs significantly. Figure 2b,e reveals that the pyrene units are displaced with respect to each other in BNP, while they overlap in CCP. In contrast, the unsubstituted BN-pyrene 4a forms head-to-tail π -stacks with opposed boron and nitrogen atoms. Different from that, the distorted thienyl units in BNP obviously prevent a close approximation of the molecules, at which the attracting electrostatic forces between boron and nitrogen would dominate the crystal packing.

To further analyze this, a Hirshfeld surface analysis⁸³ was performed. The displaced crystal packing in **BNP** leads to some distinct close atom—atom contacts (red spots on the Hirshfeld surface) in **BNP**, in particular a B…C contact of 3.4 Å distance (Figure 2c).

In CCP, there is no such specific close atom—atom contact, visualized by the absence of any red marker on the Hirshfeld surface. The pyrene centroid—centroid distance in CCP is 4.0 Å, whereas it is 4.8 Å in BNP. The plane-to-plane distances are equal (3.5 Å for both BNP and CCP), a value which was reported for unsubstituted pyrene and BN-pyrene as well.^{70,84} Consequently, in BNP and CCP, a charge carrier transport between the pyrene planes should be possible.^{68,70,85,86} While the overlap area of two stacked molecules of BNP is 35%, it is largely elevated in CCP (58%), which is comparable with unsubstituted pyrene (60%) and a number of substituted pyrenes that throughout exhibit excimer formation in the solid state.⁶⁸ We will discuss the consequences of this overlap angle on the optical properties later in the article.

The fingerprint plots shown in the Supporting Information (Section 4, Figures S9 and S10) confirm the different nature of



Figure 2. Solid-state molecular structures of **BNP** (a) and **CCP** (d) as well as the crystal packing of **BNP** (b) and **CCP** (e). Also presented are the Hirshfeld surfaces with the d_{norm} color coded onto them on a scale from -0.14 (red), 0 (white), to 1.26 (blue) [**BNP**: (c), **CCP**: (f)]. Only the major disorder component is shown for both **BNP** and **CCP**. The atomic displacement parameters are at the 50% probability level.

interactions in both crystal structures with close C···C contacts (indicative of $\pi - \pi$ interactions) accounting for 19% of all contacts in **CCP**, while they are only half as frequent in **BNP** (9.5%, including contacts involving B and N atoms).

The nature of the forces behind the crystal packing contacts can be elucidated with a model energy analysis, here based on calculations at the B3LYP^{87,88} /6-31G(d,p)⁸⁹ level of theory. The analysis shows the total interaction energy of all molecular pairs in the first coordination sphere in relation to the asymmetric unit and the corresponding electrostatic, dispersion, polarization, and Pauli repulsion terms (see the Supporting Information, Section 4, Tables S2 and S3 for a summary of these energy terms).⁹⁰ **BNP** is more stabilized than **CCP** by 14 kJ mol⁻¹ in its crystal packing among the first coordination sphere. This stabilization derives from the significantly higher electrostatic component, which is a direct consequence of the BNsubstitution. In turn, the dispersion term is slightly higher in



Figure 3. Comparison of the proton resonances in the respective ¹H NMR spectra of BNP (top) and CCP (bottom). For better perceptibility, the signals of the solvent (C_6D_6) were omitted.

Table 1. Photophysical P	operties of BNP and	l CCP in DCM Solution	and in the Solid State
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Compound	$\lambda_{\mathrm{abs}} \left(\mathrm{DCM} ight) \ \left[\mathrm{nm} ight]^{a,b}$	$\varepsilon (CHCl_3) [mol^{-1} L^{-1} cm^{-1}]$	$\lambda_{\rm em} ({\rm DCM}) \\ [\rm nm]^{a,b}$	Stokes shift (DCM) $[cm^{-1}]$	Φ_{em} (DCM)	FLT (DCM) [ns]	$\lambda_{ m em} ({ m solid}) \ [{ m nm}]^a$	$\Phi_{ m em}$ (solid)	FLT (solid) [ns]	
BNP	270, 335, 452	15 486	531	3300	0.03	11.8 ^d	541	< 0.01	2.4 ^g	
ССР	283 , 351	23 052	403 , 494 [°]	3700	0.05	3.7, ^e 7.3 ^f	432 , ~500	0.03	$0.6,^{h} 6.4^{i}$	
^{<i>a</i>} Values in bold represent the intensity maxima. ^{<i>b</i>} $c \sim 1 \times 10^{-3}$ mg mL ⁻¹ . ^{<i>c</i>} The local maximum of CCP at λ_{em} = 494 nm was only found at c = 10 mg										
mL ⁻¹ . ${}^{d}\lambda_{em} = 530$ nm. ${}^{e}\lambda_{em} = 420$ nm. ${}^{f}\lambda_{em} = 570$ nm. ${}^{g}\lambda_{em} = 540$ nm. ${}^{h}\lambda_{em} = 432$ nm. ${}^{i}\lambda_{em} = 486$ nm.										

CCP than in BNP. The interaction (plus its symmetry-related inverse) depicted in Figure 2c,f, where the Hirshfeld surface encloses the asymmetric unit, is the most energetically dominant one in both compounds. In both cases, dispersion forces are the main stabilizing factor, but they are 25 kJ mol⁻¹ higher in CCP than in BNP, while in BNP, the electrostatic term is 9 kJ mol⁻¹ higher than in CCP. This indicates that the mechanism of charge transfer in the solid state is different for the two compounds, with considerable consequences for the spectroscopic properties as discussed below.

Summarizing the discussed features, **BNP** and **CCP** are structurally very similar as single molecules, and both form herringbone patterns with a plane distance of 3.5 Å. Although a dominance of electrostatic interactions of boron and nitrogen atoms in the neighboring molecules of **BNP** as a structuredetermining motif is prevented, significant differences in the centroid–centroid distances, overlap areas, and dihedral angles involving the thienyl units are found, which must be a consequence of the increased electrostatic term in **BNP**.

NMR Spectroscopy. All signals in the ¹H and ¹³C{¹H} NMR spectra of **CCP** and **BNP** were located in the aromatic region. As a main trend in the ¹H NMR spectrum of **BNP** (solvent: C_6D_6), the chemical shifts of the outer ring protons differed significantly both in comparison with each other and with their counterparts in **CCP**. While the proton resonances in the pyridine ring (Figure 3, H-a and H-b) are shifted upfield, these are shifted downfield in the borabenzene ring (Figure 3, Hd and H-e). It is moreover significant that the H-h of **BNP** is considerably shifted downfield due to the sterical interaction with the borabenzene ring.

This indicates the high electronic influence of electropositive boron and electronegative nitrogen in close proximity as it determines the degree of aromaticity (see the section on calculations). Interestingly, the shifts of the outer ring protons of **BNP** were far smaller compared to those of **CCP**, when chloroform was chosen as the solvent (see the Supporting Information, Section 3.14 and Figure S1). This could arise from increased charge compensation, as chloroform is a moderate H-bond donor, and is an indication of a different solvent-dependent behavior of both molecules.

We then performed diffusion-ordered NMR spectroscopy (DOSY) experiments of **BNP** in CDCl₃ at concentrations of 9.33 and 0.15 mg mL⁻¹, which was close to the lower detection limit of the NMR spectrometer. The solvodynamic radius was determined to be $9.69 \pm 6.7 \times 10^{-3}$ Å and $10.05 \pm 6.0 \times 10^{-2}$ Å (see the Supporting Information, Section 3.15, Figures S2 and S3, and Table S2). Therefore, it can be assumed that there is no concentration effect on aggregation in the studied range, and there is no evidence of the presence of different ground-state aggregates. As the nature of aggregations was unequivocally proven for CCP (see next chapters), these measurements were not performed for this analogue.

Optical Spectroscopy in Solution. The most significant data from the optical measurements are denoted in Table 1.⁹¹

The stationary absorption and PL spectra were recorded from DCM solutions at $c \sim 1 \times 10^{-3}$ mg mL⁻¹. Regarding the absorption of **BNP** and **CCP**, the intensity maximum of **BNP** was red-shifted by $\Delta \lambda_{abs} = 52$ nm to $\lambda_{abs} = 335$ nm compared to the highest intensity band of **CCP** at $\lambda_{abs} = 283$ nm (Figure 4).

The bathochromic shift in both the absorption and emission has been reported for BN-substituted pyrenes⁷⁰ and is a result of the different electronic structures of CCP and BNP (for a detailed discussion of this effect, see the section on calculations). The lowest energy local maximum of CCP was found at $\lambda_{abs} = 351$ nm.

The observed band shapes differed significantly. While **BNP** displayed one major absorption band with a high intensity ($\varepsilon =$



Figure 4. Normalized absorption (dashed lines) and emission (continuous lines) spectra of **BNP** (red) and **CCP** (blue) measured in DCM at $c \sim 1 \times 10^{-3}$ mg mL⁻¹. The excitation wavelengths were λ_{ex} = 300 nm (**CCP**) and λ_{ex} = 330 nm (**BNP**).

15 486 mol⁻¹ L cm⁻¹ at $\lambda_{abs} = 336$ nm, see the Supporting Information, Section 5.1 and Figure S14, the differences in absorption maximum are due to the solvent used, which was chloroform), the absorption curve of **CCP** was broader, and a protruding maximum intensity band was not present ($\varepsilon = 23052$ mol⁻¹ L cm⁻¹ at $\lambda_{abs} = 283$ nm, see the Supporting Information, Section 5.1 and Figure S15).

The PL spectrum of **CCP** showed a broad, featureless emission band with an intensity maximum at $\lambda_{em} = 403$ nm, in contrast to the vibronic fine structuring of unsubstituted pyrenes with five exposed bands⁵⁶ and the four exposed bands of BN-

pyrene.⁷⁰ For several mono-, di-, and tetrathienyl substitutes of pyrene, this merging effect of the emission bands of the pyrene unit and the thienyl substituents has been reported^{92,93} as a result of the decreasing energies of non-radiative deactivations. This observation is caused by a uniform delocalization of the frontier molecular orbitals among the pyrenyl and thienyl units.^{94,95}

For **BNP**, the intensity maximum was found at $\lambda_{em} = 531$ nm, and in contrast to **CCP**, two shoulder signals were observable ($\lambda_{em} \sim 500$ and 575 nm). This slight vibronic resolution is a first indication that the emissive species is monomeric. With regard to the presented diphenyl-substituted BN-pyrene ($\lambda_{em} = 489$ and 522 nm),⁷⁰ only a slight red shift of $\Delta \lambda_{em} = 9$ nm was determined. In addition, the emission band of **BNP** was narrowed compared to **CCP**, as evidenced by a full width at halfmaximum of 1950 cm⁻¹ for **BNP** compared to 3080 cm⁻¹ for **CCP**.

In comparison with BN-pyrene ($\lambda_{abs} = 321 \text{ nm and } \lambda_{em} = 514 \text{ nm in DCM}$)⁷⁰ and CC-pyrene ($\lambda_{abs} = 335 \text{ nm and } \lambda_{em} = 384 \text{ nm}$ in cyclohexane),⁴³ the peripheral thienyl units induce a bathochromic shift of both absorption and emission of about $\Delta \lambda = 15-20 \text{ nm in all cases}$, reflecting an effective mixing of the respective frontier orbitals of pyrene and thiophene that are involved in the transitions in solution.

The quantum yields of both **BNP** ($\Phi = 0.03$) and **CCP** ($\Phi = 0.05$) were decreased compared to their unsubstituted congeners ($\Phi = 0.15$ for BN-pyrene and $\Phi = 0.32$ for CC-pyrene), which is a consequence of both the increased atomic mass and the formation of additional weak hydrogen bonds of the sulfur atom of the thienyl unit (the S–H interactions account for 12.6% in **BNP** and 14.7% in **CCP**, see the



Figure 5. Normalized concentration-dependent steady-state absorption (a,c) and emission (b,d) spectra for **BNP** (a,b) and **CCP** (c,d) solutions in DCM. The excitation wavelength was $\lambda_{ex} = 330$ nm.

Supporting Information, Section 4 and Figures S10 and S11).^{96,97}

In order to analyze the different species in the ground and excited state, steady-state absorption and PL measurements were performed as a function of concentration (Figure 5, a,b: BNP; c,d: CCP).

Both **BNP** and **CCP** solutions in DCM showed no significant alteration in the absorption curves upon concentration changes. Hence, the aggregation behavior of both compounds in the ground state does not depend on the concentrations in the studied region.

An examination of the PL of **CCP** revealed an emerging broadened band at $\lambda_{em} = 494$ nm at increased concentrations. Such behavior is characteristic for excimer formation in all carbon pyrenes.⁵⁵ The PL spectrum of **BNP** remained nearly unchanged at different concentrations, hinting at the absence of excimers at increased concentrations.

To analyze a potential solvatochromism and possible aggregation behavior, concentration-dependent PL measurements were performed in solvents of different polarities and dielectric constants (cyclohexane, DCM, ethyl acetate, and methanol, see the Supporting Information, Section 5.2 and Figure S16).

For **BNP**, a variation of the solvent did not have a notable impact on the emission maxima and only a minor impact on the intensity of the shoulder signal at $\lambda_{em} \sim 500$ nm (Figure 6, DCM and methanol as examples).



Figure 6. Normalized PL spectra of **BNP** and **CCP** in DCM and methanol at $c = 10 \text{ mg mL}^{-1}$.

As excitation spectra were unchanged up to concentrations as low as $c = 4.3 \times 10^{-6}$ mg mL⁻¹ besides that (see the Supporting Information, Section 5.5 and Figure S24), the presence of a hypsochromically shifted ground-state aggregate in solution is unlikely. Contrary to Piers' work,⁷⁰ no bathochromically shifted band was identified; therefore, excimer formation is not plausible.

In contrast, the intensity of the characteristic excimer emission band of **CCP** was significantly solvent dependent, as it was absent in methanol and most intense in DCM. This could be a solubility issue, as all tested solvents except DCM did not allow an entire dissolution at $c \ge 1$ mg mL⁻¹. Furthermore, methanol may cause more intense interactions of the thienyl units, also leading to decreased pyrene π - π -interactions.⁹⁸ In all solvents at different concentrations, the monomer emission band of CCP remained nearly unchanged.

The PL lifetimes of **CCP** and **BNP** were determined at different concentrations by TCSPC experiments using nondegassed DCM as the solvent (Figure 7). In TCSPC measurements, the emitted single photons are counted as a function of time, resulting in a photon distribution over time.



Figure 7. TCSPC measurements of **BNP** (red, $\lambda_{ex} = 355$ nm and $\lambda_{em} = 530$ nm) and **CCP** (blue and violet tones, $\lambda_{ex} = 371$ nm and $\lambda_{em} = 420$ and 570 nm) in DCM solutions and regarding lifetime fittings (black/white lines). The dynamic range of the measurements was 10³ (**CCP**) and 10⁴ (**BNP**), which explains the different onset values of the respective curves.

The lifetimes (FLT) of **BNP** ($\tau = 11.8$ ns) and **CCP** ($\tau = 3.7$ ns), determined by fitting the decay curves at c = 0.1 mg mL⁻¹, differed remarkably (Figure 7). However, both of them followed single exponential decays in broad sections of the curves.

The relative brevity of the lifetimes is an immediate consequence of the thienyl units in 4,10-positions, which contribute to most of the natural transition orbitals (NTOs) that are involved in the fluorescence process (see the section on calculations and the Supporting Information, Section 6.2 and Figure S29) and enhance the symmetry-forbidden $S_1 \rightarrow S_0$ transitions. In contrast, unsubstituted CC-pyrene (τ up to 680 ns)⁵⁸ and unsubstituted BN-pyrene ($\tau = 70$ ns)⁷⁰ exhibit largely increased lifetimes, which can doubtlessly be attributed to forbidden transitions at least for CC-pyrene.⁶⁹

As oxygen is a well-known quencher of fluorescence, 55,99 we also measured the PL lifetimes in degassed DCM (see the Supporting Information, Section 5.3 and Figures S17 and S18). As expected, the lifetimes were increased, however only in the range of 4–14%, indicating that oxygen did not lead to major quenching.

Recording the lifetime curves of **CCP** at different concentrations (c = 0.1 and 1.0 mg mL⁻¹) allowed us to analyze the decays of both lifetime components ($\lambda_{em} = 420$ nm for the excited monomer and $\lambda_{em} = 570$ nm for the excimer), deriving from the existence of excimers only at higher concentrations. While only monomer decay was observed at c = 0.1 mg mL⁻¹, the excimer decay became apparent at c = 1.0 mg mL⁻¹ ($\tau = 7.3$ ns) when setting the emission wavelength to $\lambda_{em} = 570$ nm. In contrast, only one lifetime of **BNP** was observed when screening low to medium concentrations (Figure 7). This suggests the presence of only one emissive species and contradicts the presence of excimers. When the concentration became very high, a notable decrease of the lifetimes was observed due to

enhanced collision-induced quenching. All lifetimes were independent of the fluence, which was tested by setting it to values between 25 and 500 μ W (see the Supporting Information, Section 5.3 and Figure S19).

To obtain an explicit representation of the differences in excimer formation, we measured the sample emission curves at different time delays using an intensified charge-coupled device (iCCD) setup. The excitation light pulse at $\lambda_{ex} = 355$ nm results from the double frequency of $\lambda_{ex} = 710$ nm, generated by a non-collinear optical parametric amplifier. We then plotted separate emission curves for a range of lifetime periods as well as separate lifetime curves for different wavelength sections. The excimer emission could be visualized by iCCD images.

In highly diluted solutions of **CCP** ($c = 0.05 \text{ mg mL}^{-1}$), no excimer formation could be detected as the decays were comparable for four investigated emission wavelengths, and the emission curve remained almost unchanged for all lifetime sections (Figure 8).



Figure 8. Segmented emission spectra (a) and lifetime curves (b) as well as iCCD images (c) of **CCP** at c = 0.05 mg mL⁻¹ and an excitation wavelength of $\lambda_{ex} = 355$ nm. The solvent was non-degassed DCM.

In contrast, when the concentration was increased to $c = 5 \text{ mg} \text{ mL}^{-1}$, the emission decays at different wavelengths diverged (Figure 9).

While the component of the emission spectrum at $\lambda_{\rm em} = 390-400$ nm decayed with a lifetime of $\tau \sim 3-4$ ns following a single exponential decay, the components at $\lambda_{\rm em} = 500-510$ nm and $\lambda_{\rm em} = 580-590$ nm decayed much more slowly (Figure 9b). Furthermore, the lower energy band at $\lambda_{\rm em} = 507$ nm became the most intense band when displaying the emission components that are associated with long lifetimes (Figure 9a). The results strongly support excimer formation of **CCP** at high concentrations, coexisting with monomers.

In contrast, no secondary emission band was present for **BNP** for both investigated concentrations (c = 0.2 and 5.0 mg mL⁻¹). Moreover, no significant differences were found in the lifetimes of the different sections of emission wavelengths (Figures 10 and 11).



Pump-Probe delay [ns]

Figure 9. Segmented emission spectra (a) and lifetime curves (b) as well as iCCD images (c) of **CCP** at $c = 5.0 \text{ mg mL}^{-1}$ and an excitation wavelength of $\lambda_{ex} = 355 \text{ nm}$. The solvent was non-degassed DCM.



Figure 10. Segmented emission spectra (a) and lifetime curves (b) as well as iCCD images (c) of **BNP** at $c = 0.2 \text{ mg mL}^{-1}$ and an excitation wavelength of $\lambda_{ex} = 355 \text{ nm}$. The solvent was non-degassed DCM.

Optical Spectroscopy in the Solid State. In general, the absorption bands in solution were well reflected by the results from solid-state measurements (see the Supporting Information, Section 5.4 and Figure S22), with negligible deviations of the intensity maxima. However, in contrast to CCP, the bands of BNP were significantly broadened.

Similar PL bands as in solution were also found in the solid state. However, a significantly slower decrease of the PL intensity at the red edge of the spectrum was noted for both derivatives. While a solution of **BNP** in DCM revealed its emission maximum at $\lambda_{em} = 531$ nm, the latter was only slightly shifted to $\lambda_{em} = 541$ nm in the solid state (Figure 12). Moreover,



Figure 11. Segmented emission spectra (a) and lifetime curves (b) as well as iCCD images (c) of **BNP** at $c = 5.0 \text{ mg mL}^{-1}$ and an excitation wavelength of $\lambda_{ex} = 355 \text{ nm}$. The solvent was non-degassed DCM.

the lifetime was significantly reduced in the solid state ($\tau = 2.41$ ns, see the Supporting Information, Section 5.3 and Figure S20).



Figure 12. Normalized PL spectra of **BNP** (red) and **CCP** (blue) in DCM solution (continuous, $c = 10 \text{ mg mL}^{-1}$) and the solid state (dashed).

As no bathochromically shifted bands were present, no evidence is given for the presence of solid-state excimers of **BNP**. This can be explained by the small interplanar overlap area of 35% as can be seen from the crystal structure.

The solid-state emission spectrum of CCP revealed the presence of both the monomeric emissive species ($\lambda_{em} = 432$ nm) and the excimer ($\lambda_{em} \sim 500$ nm), similar to the results from PL spectroscopy in solution at the highest investigated concentration ($c = 10 \text{ mg mL}^{-1}$). Apparently, the large molecule-to-molecule overlap area of 58% allows effective excimer formation in this case. The high intensity of the excimer band in the solid state can be explained by the already closely adjoined molecules (plane-to-plane distance: 3.5 Å in a single crystal), which readily allow excimer formation in contrast to a diffusion-controlled process in solution. Two lifetime components were noticed ($\tau = 0.64 \text{ ns}/6.40 \text{ ns}$, see the Supporting Information, Section 5.3 and Figure S21), which support the presence of excimers.

The excitation spectra of **BNP** and **CCP** are shown in the Supporting Information (Section 5.4 and Figure S23). The quantum yields in the solid state were low ($\Phi < 0.01$ for **BNP** and $\Phi = 0.03$ for **CCP**), which indicates a predominating aggregation-caused quenching (ACQ).^{100,101}

Calculations. We calculated the optimized geometries in S₀ and S₁, NTOs,¹⁰² as well as the absorption and emission spectra of **CCP** and **BNP** with ORCA 4.2.0¹⁰³ using time-dependent DFT (TD-DFT)^{104–106} at the B3LYP^{107–109} /cc-pVDZ¹¹⁰ level of theory (for details, see the Supporting Information, Section 6). The absorption spectra were well reproduced as all local maxima were mapped by calculations with deviations of only $\Delta \lambda_{\rm abs} \pm 15$ nm (Figure 13).

The emission maxima of the calculated bands match with the experimental results in good approximation and even the vibronic bands that suffer from broadening in the experimental spectra were reflected correctly (Figure 14).

Furthermore, a decreased full width at half-maximum of **BNP** compared to **CCP** was obvious. Both molecules clearly fluoresce out of a π - π * state as evidenced by NTO calculations (see the Supporting Information, Section 6.2 and Figure S29). The results indicate that the decisive factor for the bathochromic shifts in the emission of **BNP** compared to its carbonaceous counterpart is indeed the different electronic structure, while aggregation effects may play an additional but subordinate role in the solution PL spectra at low concentrations.

 $NICS(0)^{111}$ (nucleus-independent chemical shift) values were calculated at the MP2¹¹² /cc-pVDZ level of theory. The



Figure 13. Measured normalized absorption bands and calculated absorption sticks of BNP (a) and CCP (b).





Figure 14. Measured (continuous, solvent: DCM) and calculated (dashed) normalized PL spectra of BNP (red) and CCP (blue).

results confirmed the trend that the rings at the long end of the pyrene core were more aromatic (NICS(0) ~ -12.5 ppm for CCP and -11.9 and -13.7 ppm for BNP) than the rings at the short end (NICS(0) = -4.0 ppm for CCP and -6.8 ppm for BNP); moreover, the latter results pointed out an overall increased aromaticity in BNP.

CONCLUSIONS

A BN-dithienylpyrene BNP and its internally CC-substituted counterpart CCP were synthesized successfully by employing cross-coupling reactions, transmetalations, and concluding electrophilic cyclizations. While BNP was obtained in a total yield of 23%, the overall yield of CCP was 21%. The crystal packing of CCP is comparable with the unsubstituted and arylsubstituted pyrenes, showing classical $\pi - \pi$ interactions and a pyrene-pyrene overlap area of 58%. In contrast, the BN-unit of BNP induces a higher electrostatic term to the attraction forces. As the molecular structures are very similar besides that, the dipolar interactions cause a more displaced crystal packing with a significantly reduced overlap area (35%). However, the role of the distorted thienyl units should not be underestimated, as they may prevent a head-to-tail aggregation of the opposed boron and nitrogen atoms. The differences in the spectro-optical behavior of both compounds are immense. Due to the divergent electronic structures, a bathochromic shift of $\Delta \lambda_{\rm em}$ = 128 nm for the emission maximum of BNP was observed compared to CCP, corresponding to the results of TD-DFT calculations. Compared to the unsubstituted derivatives, both emission bands are featureless due to the electronic impact of the thienyl substituents, which moreover cause a bathochromic shift of λ_{abs} and $\lambda_{\rm em} \sim 15-20$ nm. As can be seen from the steady-state experiments, the monomer emission bands of BNP and CCP are barely solvent and concentration dependent. Compared to Piers' BN-pyrene, no evidence was given that BNP forms excimers in solution. In contrast, CCP showed excimer formation at high concentrations ($\lambda_{em \text{ monomer}} = 403 \text{ nm}$ and $\lambda_{em \text{ excimer}} = 494 \text{ nm}$ at c = 10.0 mg mL⁻¹), which is a classical feature of pyrenes. The presence of excimers was also proven by TCSPC lifetime measurements, showing a secondary emission decay. Unlike BNP, CCP shows excimer formation in the solid state as well. The differences in excimer formation can clearly be attributed to the significant distinction in the π - π -overlap areas. Therefore, despite having the same plane-to-plane distance and a larger energy gain upon stacking in BNP, it is a different centroid-centroid length, sideway shift of each molecule, and thus a different transition dipole moment that is likely to be the

reason for an entirely different excited-state photochemistry. We conclude that a BN-substitution in combination with the implementation of flanking thienyl groups can be a useful tool to suppress excimer formation in pyrene solutions and in the solid state. As BNP can be post-functionalized selectively at the thiophene sites and should be electrochemically stable such as the parent BN-pyrene,⁷⁰ the generated knowledge could be fundamental for the design of novel materials for the application in organic electronic devices. While the issue of ACQ limits the potential as an electroluminescent compound,¹¹³ the investigation of the charge carrier generation and transporting capability of BN-PAHs is of high interest. Especially, an analysis of the competing redox processes at the pyrene scaffold and the thienyl units should be performed in the future. Consequently, the implementation of BNP and related BN-pyrenes into fieldeffect transistors as well as electropolymerization should be envisaged.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcc.1c08812.

Experimental procedures, crystal data, NMR spectra, additional optical measurements, and calculation details (PDF)

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Notes

The authors declare no competing financial interest.

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