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Boosting quantum yields and circularly polarized luminescence of penta- and hexahelicenes by doping with two BN-groups†

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The incorporation of boron–nitrogen (BN) units into polycyclic aromatic hydrocarbons (PAHs) as an isoelectronic replacement of two carbon atoms can significantly improve their optical properties, while the geometries are mostly retained. We report the first non- π -extended penta- and hexahelicenes comprising two aromatic 1,2-azaborinine rings. Comparing them with their all-carbon analogs regarding structural, spectral and (chir)optical properties allowed us to quantify the impact of the heteroatoms. In particular, BN-hexahelicene BN[6] exhibited a crystal structure congruent with its analog CC[6], but displayed a fivefold higher fluorescence quantum yield ($\varphi_{\rm fl}=0.17$) and an outstanding luminescence dissymmetry factor ($|g_{\rm lum}|=1.33\times10^{-2}$). Such an unusual magnification of both properties at the same time makes BN-helicenes suitable candidates as circularly polarized luminescence emitters for applications in materials science.

Introduction

Circularly polarized luminescence (CPL) is the differential emission of left- (I_L) and right-handed (I_R) circularly polarized light by chiral luminophores. Recently, CPL-active materials have demonstrated their immense application potential for optoelectronics, $^{2-4}$ 3D displays, 5,6 switches for data storage, 7,8 spintronics 9,10 or chiral sensors. 11,12

Small organic molecules are particularly suitable as CPL emitters due to their structural modifiability.¹³ It allows the

tuning of crucial optical properties like molar absorptivities (ε), fluorescence quantum yields ($\varphi_{\rm fl}$) or emission dissymmetry factors ($g_{\rm lum} = 2 \times (I_{\rm L} - I_{\rm R})/(I_{\rm L} + I_{\rm R})$), quantifying the degree of polarization in the emission. Moreover, structure-related parameters like configurational¹⁴ and thermal¹⁵ stabilities towards decomposition are tunable, which is of high relevance for the processing into functional materials.

Among the various classes of chiral, π -conjugated molecules, helicenes are well-known for their strong optical rotation and electronic circular dichroism (ECD). Carbohelicenes are chiral, screw-shaped polycyclic aromatic hydrocarbons (PAHs) with $n \geq 5$ ortho-fused benzenoid rings. Research studies have especially focused on derivatives of pentahelicene and hexahelicene because they allow chiral resolution at ambient conditions and gram-scale preparation by various synthetic approaches. In

For assessing the suitability of helicenes as CPL-emitting molecules, key prerequisites are high $|g_{\text{lum}}|$ values, but also elevated ε and φ_{fl} . To combine these parameters into one, the CPL brightness factor $B_{\text{CPL}}=0.5\times\varepsilon\times\varphi_{\text{fl}}\times|g_{\text{lum}}|$ has been introduced,²⁰ allowing to compare the suitability of such luminophores for potential applications. Planar PAHs typically display high φ_{fl} values due to electronically allowed π - π * transitions, but the distorted π -planes of helicenes induce a significant decrease ($\varphi_{\text{fl}}=0.04$ for both penta- and hexahelicene). This is a result of accelerated intersystem crossing, leading to non-emissive triplet states.²¹ Furthermore, the dissymmetry factors of carbohelicenes are almost exclusively below 10^{-2} ($|g_{\text{lum}}|=2.7\times10^{-3}$ for penta- and 9×10^{-4} for

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hexahelicene). 22,23 Consequently, pristine carbohelicenes (B_{CPL} < 2 M^{-1} cm $^{-1}$) are barely useful as CPL emitters. 24

Incorporating main group elements like nitrogen, ^{25–27} oxygen, ^{28,29} silicon ^{30,31} or combinations of these (*e.g.* B–O)³² in PAHs in general can modulate the optical properties significantly. However, this also perturbs the molecular architecture and the estimation of structural and electronic influences isolated from each other is complicated. ³³

Therefore, the doping with neighboring boron-nitrogen (BN) units has emerged as a popular strategy especially for planar PAHs, ³⁵⁻³⁸ as it provides isoelectronic and isostructural BN-PAHs. In such compounds, the disparities in properties are almost exclusively caused by altered electronegativities of the heteroatoms, ³⁹ facilitating the comparison with the parent, all-carbon CC-PAHs. ⁴⁰⁻⁴²

Unfortunately, reports of helical BN-derivatives have remained scarce, $^{43-47}$ which especially applies to helicenes with B=N groups in the aromatic backbone. Moreover, experimental studies that compare BN-helicenes with their all-carbon counterparts are rare. To our knowledge, the work from Nowak-Król and Ingleson describes the only non- π -extended B=N-helicenes so far (Fig. 1, top right). Besides a small structural impact, the most distinctive effect was an amplification of the radiative decay ($\varphi_{\rm fl}=0.30$ for the (mono)BN-pentahelicene, $\varphi_{\rm fl}=0.21$ for the BN-hexahelicene), compared to the all-carbon congeners, which were calculated but not synthesized. Nevertheless, this suggests the great potential of this compound class.

Herein, we present the first non- π -extended penta- and hexahelicenes BN[5] and BN[6], respectively, comprising two aromatic boron-nitrogen-containing rings (Fig. 1, bottom). A

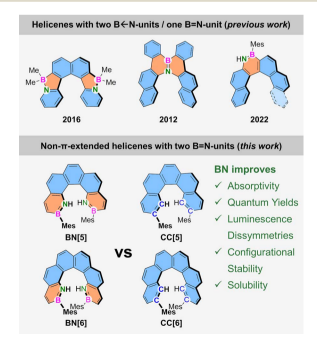


Fig. 1 Top: previously reported BN-penta- and hexahelicenes. Bottom: structures studied in this work and the demonstrated benefits of such a twofold BN-substitution.³⁴

multi-dimensional comparison with their all-carbon analogs CC[5] and CC[6] allowed us to gauge the exact influence of the BN-substitution on structural and (chir)optical properties. Most strikingly, the BN-substitution led to an outstanding CPL brightness of BN[6] ($B_{CPL}=59~M^{-1}~cm^{-1}$) while leaving the geometrical structure almost unaffected.

Results and discussion

Syntheses

BN-helicenes **BN**[5] and **BN**[6] were synthesized *via* transition-metal-catalyzed electrophilic cyclizations of (bis)ethynylarene precursors. These mild conditions^{51–53} were chosen to ensure the integrity of the moderately acidic, *N*-deprotected 1,2-aza-borinine building blocks.

Initially, electrophilic benzene- 1 and naphthalene precursors 2, equipped with two (trimethylsilyl)ethynyl substituents, were synthesized (Scheme 1).⁵² Subsequent Suzuki-Miyaura cross-coupling reactions with nucleophilic mesitylazaborinine 3 (ref. 54) gave TMS-alkynes 4 and 5 in yields of 72% and 75%, respectively. Under basic reaction conditions, deprotected alkynes 6 and 7 were obtained quantitatively.

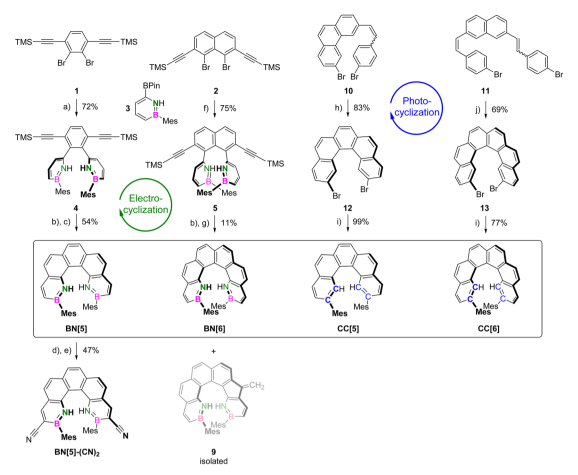
Our previous study of planar BN-PAHs⁵⁵ showed that six-membered rings are formed almost exclusively when reacting the deprotected arylalkynes with $PtCl_2$ at 100 °C. Transferring these conditions to BN-helicenes, however, significantly favored the competing 5-*exo-dig* pathway, leading to dibenzofulvene-like side-products in conversions of more than 40%. Therefore, we investigated different catalytic systems based on Pt(II), Au(II), Au(III) and Ru(II) (see ESI, Tables S1 and S2†).

In the case of **BN**[5], the reaction of phenylalkyne **6** with AuCl₃ at 100 °C for 24 h provided the highest conversion with less than 5% *endo/exo*-side product **8** (see ESI, Table S1†). Derivatization of the latter with piperidine⁵⁶ and chromatographic separation of the resulting Michael-type adduct eventually allowed the isolation of **BN**[5] in 54% yield.

On the other hand, the cyclization of naphthylalkyne 7 to BN[6] was achieved using $[(p\text{-cymene})RuCl_2]_2$ and $AgSbF_6$. Among all catalytic systems tested, this was the only one that favored the formation of two six-membered rings (*ca.* 80% of *endo/endo*) (see ESI, Table S2†). We assume that the reaction with Ru(II) preferably proceeds *via* the η^1 -vinylidene intermediate, assisting the nucleophilic attack at the terminal alkyne carbon. After heating 7 and the catalysts to 170 °C for 1 h in mesitylene and purification as for BN[5], BN[6] was obtained in a yield of 11% (Scheme 1). In contrast to the Ru(II) catalyst, the use of $AuBrPPh_3$ and $AgSbF_6$ allowed to selectively synthesize and isolate *endo/exo*-derivative 9 (see ESI, Section 2.28†).

For the syntheses of CC[5] and CC[6] (phenylethenyl)phenanthrene **10** and bis(phenylethenyl)naphthalene **11** were prepared in total yields of 52% and 18%, respectively. Mallory photocyclization reactions using UV irradiation at $\lambda = 365$ nm in the presence of iodine and propylene oxide gave dibromopenta- **12** and dibromohexahelicene **13**. Subsequently, mesityl substituents were installed at positions 2,13 and 2,15, respectively, *via* Kumada cross-coupling reactions with mesitylmagnesium bromide. Eventually, racemic mixtures of CC[5]

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Scheme 1 Syntheses of BN- and CC-helicenes. Reagents and conditions: (a) **3** (4.0 equiv.), potassium phosphate (3.0 equiv.), $[Pd(dppf)Cl_2]$ (5 mol%), methyl tert-butyl ether (MTBE), water, 90 °C, 2 d; (b) KOH (2.0 equiv.), MeOH, El_2O , 25 °C, 3 h; (c) 1. $AuCl_3$ (30 mol%), mesitylene, 100 °C, 24 h; 2. Piperidine, acetonitrile, 25 °C, 1 h; (d) Bromine (2.2 equiv.), DCM, 0 °C, 2 h; (e) CuCN (10 equiv.), DMF, 170 °C, 19 h; (f) **3** (4.0 equiv.), potassium phosphate (3.0 equiv.), $[Pd_2(dba)_3]$ (5 mol%), SPhos (10 mol%), THF, water, 60 °C, 17 h; g) 1. $[(p-cymene)RuCl_2]_2$ (30 mol%), $AgSbE_6$ (30 mol%), mesitylene, 170 °C, 1 h; 2. Piperidine, acetonitrile, 25 °C, 1 h; (h) Iodine (1.1 equiv.), propylene oxide (500 equiv.), toluene, 365 nm LED, 25 °C, 30 min; (i) MesMgBr (3.0 equiv.), $[Pd(dppf)Cl_2]$ (5 mol%), 1,4-dioxane, 95 °C, 3 d; (j) Iodine (2.2 equiv.), propylene oxide (500 equiv.), toluene, 365 nm LED, 25 °C, 3 h.

and CC[6] were obtained in yields of 82% and 53% over the last two steps (Scheme 1).

Introducing functional groups onto a pre-existing helicene scaffold by electrophilic aromatic substitution is highly desirable but often demanding. In a 1,2-azaborinine, the uneven charge distribution renders all carbon atoms distinct in reactivity.58 In particular, the selective electrophilic bromination at the α -carbon adjacent to boron has been shown to be a viable starting point for post-functionalizations. 59-61 This methodology allowed us to selectively synthesize 3,12-dibrominated BN[5]-Br₂ in 75% yield by the reaction of BN[5] with 2.2 equiv. of bromine. Subsequently, the conversion with an excess of copper(1) cyanide under Rosenmund-von Braun conditions gave BN[5]-(CN)₂ in a yield of 62% (Scheme 1). In contrast, the reaction of CC[5] with bromine gave the C5,C6-dibrominated adduct.62 Upon heating it to 130 °C, rearomatization occurred and a mixture of reactant and the C5-brominated derivative was obtained. This highlights the superior selectivity of BN-PAHs in such a reaction (see ESI, Section 2.29†).

Structural analysis

Single crystals of the racemic target compounds suitable for X-ray diffraction analysis were obtained by vapor diffusion of acetonitrile, methanol, cyclohexane or *n*-hexane into DCM solutions of the helicenes.

Comparing the bond lengths within the terminal rings (B–N: 1.426 ± 0.003 Å, C1–C2: 1.385 ± 0.003 Å, $\Delta d\approx3\%$) showed the expected, relatively small geometric effect of the BN-substitution. Moreover, the C–C bond lengths within outer (1.34–1.38 Å) and inner helices (1.43–1.46 Å) remained almost identical, confirming the high two-dimensional isostructuralism of carbo- and BN-helicenes. ⁶³ On the other hand, the effect of BN on the spatial configurations of the pentahelicenes was remarkable (Fig. 2). Compared to CC[5] ($\phi=52.4^\circ$), the torsion angle (ϕ) between the terminal rings (A) was only $\phi=42.8^\circ$ in BN[5], pointing at attracting interactions involving the NH units in close proximity (A0 (NH···N) = 2.35 Å).

This also has implications on the molecular arrangements. While the packing structures of both compounds consist of

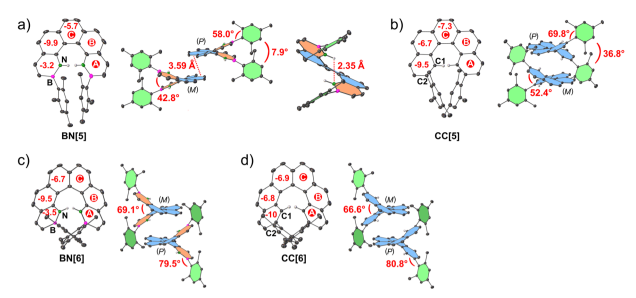


Fig. 2 Solid-state structures as obtained from X-ray diffraction analysis. Thermal ellipsoids are shown at a 50% probability level. Hydrogen atoms (except for N-H and C1-H) are omitted for clarity. NICS(0) values for the individual rings were calculated at the MP2 (ref. 64)/cc-pVDZ⁶⁵ level of theory, as implemented in the Q-Chem 6.0 program package. 66 C_2 symmetry was assumed so that angles and distances are given as average values of both sides of the molecules.

opposite pairs of (*P*)- and (*M*)-enantiomers, there is a significant overlap of the helicene scaffolds (d=3.59 Å between rings C) in a BN[5] crystal, hinting at the presence of π - π -interactions. Furthermore, the mesityl groups of each molecule are oriented almost parallelly ($\varphi=7.9^{\circ}$), which leads to a highly ordered packing arrangement. In CC[5], rings A and B of opposite molecules are arranged co-planarly but displaced. Also, the interplanar angle between the mesityl groups is larger ($\varphi=36.8^{\circ}$). The structural parameters of BN[5]-(CN)₂ were mostly comparable with BN[5] (see ESI, Section 3.3†).

Both hexahelicenes exhibit much more similar metrics ($\varphi = 69.1^{\circ}$ for BN[6], $\varphi = 66.6^{\circ}$ for CC[6] between rings A), identical packing modes (see ESI, Fig. S10 and S14†) and very similar unit-cell parameters. The absence of additional dipolar interactions in BN[6] indicates that for hexahelicenes, the distant BN-units do not play a structure-determining role.

Besides the high planarity of the 1,2-azaborinine rings (averaged $\varphi=2.6^{\circ}$ within rings A), aromaticity was clearly confirmed by nucleus-independent chemical shift (NICS) calculations (NICS(0) < -3 ppm, Fig. 2).⁶⁷

Optical spectroscopy

Absorption and fluorescence spectra as well as $\varphi_{\rm fl}$ and fluorescence lifetimes ($\tau_{\rm fl}$) of the investigated helicenes were acquired in DCM solutions (Table 1).

BN[5], CC[5], BN[6] and CC[6] revealed comparable absorption maxima ($\lambda_{\rm abs} = 266$ –283 nm) with decent molar extinction coefficients ($\varepsilon = 3.4$ –5.2 \times 10⁴ M⁻¹ cm⁻¹ see ESI, Fig. S18–S22†). The absorption bands of both hexahelicenes closely resembled each other with small bathochromic shifts of BN[6] ($\Delta\lambda_{\rm abs} \approx 10$ –15 nm, Fig. 3b). This suggests similar energy orderings of the differently polarized dipole transition moments. 68 Most remarkably, the BN-moieties induced a substantial

intensification of the least energetic 1L_b band, which is associated with fluorescence⁶⁹ (BN[6]: $\varepsilon = 3300~\text{M}^{-1}~\text{cm}^{-1}$ at 424 nm, CC[6]: $\varepsilon = 200~\text{M}^{-1}~\text{cm}^{-1}$ at 414 nm).^{70,71} With respect to the pentahelicenes, the absorption spectra were more different in shape but both revealed shoulder bands at $\lambda_{\rm abs} \approx 400~\text{nm}$. The ε value of the lowest-energy transition of BN[5] was intensified by about an order of magnitude as well (Fig. 3a). This is a typical effect of BN-doping, rendering the symmetry-forbidden $S_0 \to S_1$ transition of all-carbon aromatics partially allowed.⁷²

The fluorescence spectra of all five compounds consisted of at least three distinct vibronic bands, and the Stokes shifts, if determinable, were small (\approx 600 cm⁻¹, Fig. 3). Compared to the very similar emission bands of the pentahelicenes ($\lambda_{\rm fl} \approx$ 410 nm, $\Delta \lambda_{\rm fl} = 2$ nm for the 0–0 bands), the difference between the hexahelicenes was slightly more pronounced ($\lambda_{\rm fl} = 434$ nm for BN[6], $\lambda_{\rm fl} = 423$ nm for CC[6]).

Moreover, the bathochromic effect of a twofold BNsubstitution was identified by comparison with the reported (mono)BN-hexahelicene (Fig. 1, top right, $\lambda_{abs} = 397$ nm, $\lambda_{fl} =$ 419 nm).50 In terms of fluorescence quantum yields, the BNhelicenes were substantially brighter ($\varphi_{\rm fl} = 0.10$ for BN[5], $\varphi_{\rm fl}$ = 0.17 for **BN[6]**) than the sparsely emissive carbohelicenes ($\varphi_{\rm fl}$ = 0.04 for CC[5], $\varphi_{\rm fl}$ = 0.03 for CC[6]). Furthermore, the intensity averaged $\tau_{\rm fl}$ of the heterohelicenes were comparably smaller than those of the studied carbohelicenes. While the difference was just 20% for the hexahelicenes, the lifetime of **BN**[5] ($\tau_{\rm fl} =$ 4.7 ns) was halved compared to CC[5] ($\tau_{\rm fl} = 9.7$ ns). The increase of φ with a concomitant decrease of $\tau_{\rm fl}$ for BN-helicenes can be rationalized with a larger emissive rate of these fluorophores compared to the all-carbon analogs. The influence of the mesityl-substitution on the optical features of the parent carbohelicenes (cf. hexahelicene: $\lambda_{abs} = 410$ nm, $\lambda_{fl} = 420$ nm, $\varphi_{fl} =$ 0.03, $\tau_{\rm fl} = 8.4 \text{ ns})^{76}$ was negligible, emphasizing that the

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Table 1 Experimental, photophysical properties of the investigated helicenes and calculated oscillator strengths f for the lowest-energy transitions

Compound	$\lambda_{\mathrm{abs}}{}^{a,b}$ [nm] (10 ⁻³ × ε [M ⁻¹ cm ⁻¹])	$\lambda_{\mathrm{fl}}^{a,b,c}\left[\mathrm{nm}\right]$	${arphi_{ m fl}}^a$	Stokes shift [cm ⁻¹]	$\tau_{\rm fl}^{\ a} [{\rm ns}]$	$f^f(\lambda [nm])$
BN[5] CC[5] BN[6] CC[6] BN[5]-(CN) ₂	283 (48.2), 363 (14.7), 383 (12.6) 275 (34.2), 308 (27.0), 336 (14.8), 375 (1.0) 280 (52.2), 342 (26.3), 401 (3.1), 424 (3.3) 266 (44.7), 318 (19.1), 390 (0.3), 414 (0.2) 313 (28.0), 390 (14.9), 413 (16.6)	407, 428 409, 425 434, 459 423, 444 424, 446	0.10 0.04 0.17 0.03 0.25	d d 540 510 630	4.7 9.7^e 7.1^e 8.7^e 3.5	0.1599 (370) 0.0010 (372) 0.0418 (400) 0.0027 (387) 0.0053 (415)

 $[^]a$ Measurements performed in DCM solutions at $c=2.4\times10^{-5}$ to 4.0×10^{-5} M. b Bold values represent intensity maxima. c LED, $\lambda_{\rm ex}=370$ nm (BN-helicenes), 3 00 nm (carbohelicenes). d Not determinable because the 0–0 transitions were present as shoulder bands ($\lambda_{\rm abs}\approx400$ nm). e Intensity averaged values because fluorescence decay consisted of more than one species (see ESI, Section 5.3 for more details). f TD-DFT at the B3LYP^{73–75}/cc-pVDZ⁶⁵ level of theory.

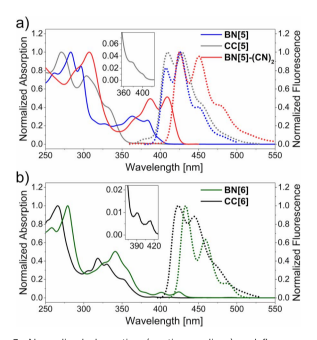


Fig. 3 Normalized absorption (continuous lines) and fluorescence (dotted lines) spectra of pentahelicenes (a) and hexahelicenes (b) in DCM ($c=2.4\times10^{-5}$ to 4.0×10^{-5} M). Insets show the lowest-energy absorption (shoulder) bands of the carbohelicenes.

enhancement of the photophysical properties by the BN-unit should not be limited to this particular substitution pattern.

We further investigated the structural and electronic properties of the helicenes by time-dependent density functional theory (TD-DFT) calculations at the B3LYP^{73–75}/cc-pVDZ⁶⁵ level of theory, as implemented in the Q-Chem 6.0 (ref. 66) and Gaussian 09, Rev. B.01 (ref. 77) program packages. For all helicenes, the emissive S_1 states⁷⁸ mostly consisted of highest occupied (HONTO) and lowest unoccupied (LUNTO) natural transition orbitals (NTOs). Moreover, the carbohelicenes revealed increased contributions of HONTO–1 and LUNTO+1 (see ESI, Section 7.4, Fig. S31†). All NTOs were delocalized over the PAH scaffolds with little to no involvement of the mesitylgroups due to their perpendicular orientations. Most noticeably, BN-substitution induced a symmetry break of the NTOs of BN[5] (C_8 for HONTO, C_2 for LUNTO, both are C_2 symmetric for

CC[5]). This feature is the cause of the higher $\varphi_{\rm fl}$ of the BN-helicenes, because the associated transition is rendered symmetry-allowed. Therefore, the related oscillator strength of BN[5] (f=0.1599) was increased by more than two orders of magnitude compared to CC[5] (f=0.0010). Despite the almost identically shaped NTOs of BN[6] and CC[6], the BN-helicene still revealed a tenfold increase of f. Overall, these results rationalize the experimental increase of ϵ in the low-energy regions.

The change of the optical properties of BN[5] upon installation of cyano-substituents was remarkable and higher than for similarly functionalized carbohelicenes: ⁷⁹ BN[5]-(CN)₂ not only revealed an intensification and a bathochromic shift of the 0–0 absorption band ($\Delta\lambda_{\rm abs}=30$ nm), but was also the most emissive compound ($\varphi_{\rm fl}=0.25$). In combination with the short lifetime ($\tau_{\rm fl}=3.5$ ns), this indicates a particularly high fluorescence emission rate.

Chiral resolution & kinetics of racemization

The targeted helicenes were resolved into their enantiomers by high-performance liquid chromatography (HPLC), employing a chiral stationary phase (CSP). From a practical perspective, the pentahelicenes revealed a better resolution than the hexahelicenes with baseline-separated peaks (see ESI, Fig. S15 for chromatograms†). While CC[6] could not be resolved as a result of its low polarity concomitant with a poor solubility, BN[5] and BN[6] (enantiomeric excess ee = 99% for both) as well as BN[5]- $CN)_2$ (ee = 94%) and CC[5] (ee = 70%) were collected in their highly enantiomerically enriched forms.

This enabled the determination of the half-lives $(t_{1/2\text{rac}})$ and activation parameters of enantiomerization. For that purpose, each enantioenriched pentahelicene was subjected to time-course ECD measurements at 50, 60 and 70 °C. **BN**[5] showed a significantly lower racemization rate $(t_{1/2\text{rac}} \approx 80 \text{ min at } 60 \text{ °C})$ than CC[5] $(t_{1/2\text{rac}} \approx 20 \text{ min})$. This translated into a Gibbs activation energy ΔG^{\ddagger} (25 °C) of 25.7 kcal mol⁻¹ for **BN**[5], which was higher by more than 1 kcal mol⁻¹ compared to CC[5] (24.3 kcal mol⁻¹, Table 2, for Eyring plots and calculations see ESI, Section 4.2†). Despite the bulkiness of the mesityl groups, the 2,13 disubstitution had a comparably small effect on the configurational stability $(\Delta G^{\ddagger}$ (25 °C) = 24.1 kcal mol⁻¹ for

 Fable 2
 Kinetics of racemization and chiroptical properties obtained from ECD and CPL measurements

	Kinetics ^b		ECD^a			$\mathrm{Cb}\Gamma_a$		
Compound	$\Delta G^{\ddagger}\left(25\ \circ \text{C}\right)$ Compound $t_{1/2\text{rac}}\left[\text{min}\right]$ [keal mol ⁻¹]	$\Delta G^{\ddagger}\left(25\ ^{\circ}\mathrm{C}\right)$ [kcal mol $^{-1}$]	$\lambda_{ m max}^{ef} [m nm]$	$\Delta \varepsilon [\mathrm{M}^{-1} \mathrm{cm}^{-1}] (\lambda_{\mathrm{max}} [\mathrm{nm}]) $	$10^3 \times g_{\rm abs} ^g$	$\lambda_{ ext{max}}^{e,h} [ext{nm}]$	$10^3 \times g_{\rm tum} ^i$	$B_{ m CPL}^{j} [{ m M}^{-1} { m cm}^{-1}]$
BN[5]	77.6	25.7	304 (-), 373 (+), 393 (+)	276 (304), 111 (393)	6.6	423, 444	4.2	10.1
CC[5]	21.1	24.3	276 (-), 319 (+), 337 (+), 399 (-)	152 (319), 82 (337)	0.9	408 , 429	4.4	3.0
BN[6] CC[6]	с д		285 (-), 344 (+), 409 (+), 431 (+)	151 (285), 28 (431)	10.8	432 , 459	13.3	59.0
$BN[5]-(CN)_2$	50.3	25.2	291 (+), 313 (–), 389 (+), 412 (+)	187 (313), 127 (412)	8.0	429	5.1	17.8
^a Chiroptical decomposition maxima. ^f Sign	measurements pe n before reaching ns for (P)-enantior	rformed in DCM; a temperature siners. $(-)$ = negati	^a Chiroptical measurements performed in DCM solutions at $c = 1.3 \times 10^{-5}$ to 4.0×10^{-5} M. ^b By time-course ECD measurements in <i>n</i> -heptane. $t_{1/2\text{rag}}$ at 60 °C. ^c Not measurable due to decomposition before reaching a temperature sufficient for racemization. ^a Not determined because chiral resolution by (CSP)-HPLC was unsuccessful. ^c Bold values represent intensity maxima. ^f Signs for (P)-enantiomers. (-) = negative CE, (+) = positive CE. ^g $t_{\text{sbs}} = \Delta \epsilon / \epsilon$ for the lowest energetic CE. ^h $t_{\text{cs}} = 370$ nm (BN-helicenes), ^c $t_{\text{cm}} = 2 \times 100$	10^{-5} M. b By time-course ECD rmined because chiral resoluti for the lowest energetic CE. h l	measurements i on by (CSP)-HPL $_{2x}=370~\mathrm{nm}$ (BN-	in <i>n</i> -heptane. $t_{1/2r}$. C was unsuccessic welicenes), $\lambda_{ m ex}=3$	ac at 60°C. ^c Not ful. ^e Bold values 300 nm (carboheli	measurable due to represent intensity cenes). i $g_{\text{lum}} = 2 \times$

pentahelicene).⁸⁰ In contrast to the pentahelicenes, heating an enantiopure sample of hexahelicene BN[6] to 200 °C for 14 h did not induce any racemization as analyzed by (CSP)-HPLC. Instead, higher temperatures caused a rapid decomposition.

In order to understand the influence of BN- and mesityl-substitution on the racemization, we calculated the respective transition states and activation barriers (see ESI, Section 7.2†). Although the absolute theoretical values of $\Delta G^{\ddagger}(T)$ were higher than the experimental ones, a gain in configurational stability upon BN-substitution was reproduced. We attribute this to the dipolar repulsions of the opposite NH hydrogen atoms, which are in significantly closer proximity in the transition state ($d \approx 1.5 \,\text{Å}$ for BN[5]) than in the C_2 -symmetric ground state ($d = 2.2 \,\text{Å}$ in a crystal). In contrast, the parallelly aligned but distant mesityl groups in the transition states do not appear to play a major role.

Chiroptical spectroscopy

Electronic Circular Dichroism (ECD) measurements of the highly enriched enantiomers throughout resulted in mirrorimaged spectra (Fig. 4 and Table 2).

In all cases, the correlation with the computed rotatory strengths $R_{\rm vel}$ allowed us to assign that the absolute configuration of the first fraction from (CSP)-HPLC was (P), which corresponds to parent penta- and hexahelicenes (Fig. 4).

The (*P*)-enantiomers of BN[5] ($\lambda=393$ nm, $\Delta \varepsilon=+111$ M $^{-1}$ cm $^{-1}$) and BN[5]-(CN) $_2$ ($\lambda=412$ nm, $\Delta \varepsilon=+127$ M $^{-1}$ cm $^{-1}$) revealed intense, positive Cotton effects (CE) in the low-energy region. Comparably sharp, negative CE maxima were located at $\lambda\approx300$ nm. In contrast, the band that represents the 1L_a transition of (*P*)-CC[5] ($\lambda=399$ nm, $\Delta \varepsilon=-1.8$ M $^{-1}$ cm $^{-1}$) was barely visible (Fig. 4b), which closely resembles the parent pentahelicene. Where 22 Moreover, the band for the $^{1}B_b$ transition was mirrored as a positive CE at $\lambda=319$ nm ($\Delta \varepsilon=+152$ M $^{-1}$ cm $^{-1}$).

The ECD spectra of (*P*)-BN[6] and (mono)BN-hexahelicene (Fig. 1, top right)⁵⁰ were similarly shaped, with one strongly negative CE ($\lambda=285$ nm, $\Delta\varepsilon=-151$ M $^{-1}$ cm $^{-1}$ for BN[6], $\lambda=245$ nm, $\Delta\varepsilon=-290$ M $^{-1}$ cm $^{-1}$ for the reported compound) and several less intense CE with opposite sign at higher wavelengths. As for the absorption and emission maxima, a twofold BN-substitution induced a significant bathochromic shift of the whole ECD spectrum ($\Delta\lambda\approx40$ nm).

The experimental absorption dissymmetry factors were typical of helicenes ($|g_{abs}|=6$ –11 \times 10⁻³, cf. 7.6 \times 10⁻³ for pentahelicene²² and 9.2 \times 10⁻³ for hexahelicene,²³ in DCM solutions at 25 °C, see ESI, Section 5.4†). By means of TD-DFT calculations, we analyzed the contributions of electric ($|\mu|$) and magnetic (|m|) transition dipole moments and the angle between both (θ) on absorption dissymmetries. For helicenes, the simplified expression $g_{abs} \approx 4 \times |m| \times \cos(\theta)/|\mu|$ applies.⁸¹ Due to the good reproduction of the experimental transition energies, we made use of the same level of theory as applied previously, although several recent studies reveal the accurate prediction of $|g_{abs}|$ and $|g_{lum}|$ values by using long-range corrected functionals (see ESI, Section 7.7†).⁸²⁻⁸⁴

 $-I_{
m R})/(I_{
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m CPL}=0.5 imes arepsilon imes ert g_{
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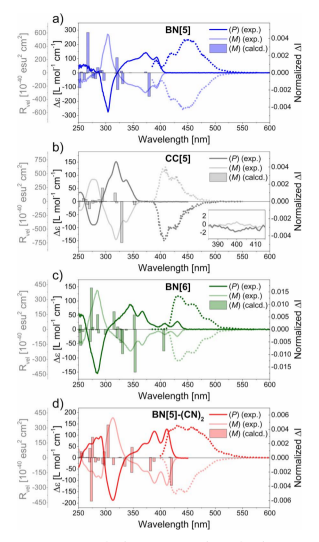


Fig. 4 Experimental ECD (continuous lines) and CPL (dotted lines) spectra in DCM ($c=1.3\times10^{-5}$ to 4.0×10^{-5} M) as well as computed rotatory strengths (bars outlined in grey). ΔI was normalized to the respective global maxima of $|g_{\rm lum}|$.

For BN[5] and BN[5]-(CN)₂, strongly enhanced transition dipole moments (\approx thirtyfold increases of |m| and \approx tenfold increases of |m|), compared to CC[5]), were calculated for the lowest-energy transitions (see ESI, Tables S55 and S56†). However, the favorable, parallel orientations of the transition dipole moments of CC[5] ($\theta=0^\circ$, cf. 114° for BN[5]) balanced out the BN-induced increase of |m|, so that $|g_{abs}|\approx 1.0$ –1.3 \times 10^{-2} for all pentahelicenes. With respect to the $S_0 \rightarrow S_1$ transitions of the hexahelicenes, both | μ | and |m| of BN[6] were enhanced only by a factor of four, compared to CC[6]. Here, the more favorably aligned dipole moments result in a considerably enhanced theoretical absorption dissymmetry.

TD-DFT calculations of the geometry-optimized structures in the S_1 states also allowed us to obtain the theoretical luminescence dissymmetry factors $|g_{\text{lum}}|$ (see ESI, Table S56†). Overall, the absolute values of $|\mu|$, |m| and θ of the $S_1 \rightarrow S_0$ transition were comparable with the $S_0 \rightarrow S_1$ transition, so that $|g_{\text{lum}}| = 0.74-1.02 \times |g_{\text{abs}}|$. Although the calculated bond lengths were

very similar in both electronic states, the disparities in plane-toplane angles were more pronounced, leading to deviating geometries (see ESI, Tables S49–53†).

Despite the previously discussed small Stokes shifts (Table 1), this could contribute to the experimental and theoretical differences between $|g_{abs}|$ and $|g_{lum}|$.

Experimental CPL spectra were obtained and normalized to the respective maxima of g_{lum} (Fig. 4 and ESI, Section 5.5†). In all cases, the sign of the CPL was identical with the sign of the lowest-energy transition in ECD. All pentahelicenes revealed comparable luminescence dissymmetry factors ($|g_{\text{lum}}| = 4-5 \times 10^{-3}$), which is in good agreement with the empirically observed correlation in helicenes ($g_{\text{lum}} \approx 0.61 \times g_{\text{abs}}$) in the case of CC[5] and BN[5]-(CN)₂.⁸⁵ Due to the enhanced fluorescence quantum yields of both BN-pentahelicenes, their CPL brightness factors were considerably elevated ($B_{\text{CPL}} > 10 \text{ M}^{-1} \text{ cm}^{-1}$) compared to CC[5] ($B_{\text{CPL}} = 3.0 \text{ M}^{-1} \text{ cm}^{-1}$).

BN[6] exhibited an outstanding experimental value of $|g_{lum}| = 1.33 \times 10^{-2}$, which is considerably higher than of unsubstituted hexahelicene ($|g_{lum}| = 9 \times 10^{-4}$).²³ In accordance with that, the theoretical values ($|g_{lum}| = 2.3 \times 10^{-2}$ for **BN[6]**, $|g_{lum}| = 0.6 \times 10^{-2}$ for **CC[6]**) indicate a stronger impact of BN-substitution on a hexahelicene than on a pentahelicene. In particular, the much more favorable orientation of the transition dipole moments in **BN[6]** (135°, *cf.* 102° for **CC[6]**) is the most significant contributor to the dramatically improved theoretical $|g_{lum}|$ value.

Compared to several reported helicenes^{84,86} with coordinative B \leftarrow N-bonds ($|g_{\text{lum}}| = 0.25 - 3.5 \times 10^{-3}$ and $B_{\text{CPL}} < 10 \text{ M}^{-1} \text{ cm}^{-1}$), the combination of increased φ_{fl} and $|g_{\text{lum}}|$ values led to a CPL brightness of $B_{\text{CPL}} = 59 \text{ M}^{-1} \text{ cm}^{-1}$ for BN[6]. This value is exceptionally high for a non- π -extended, monomeric helicene, and proves that BN-doping of helicenes is a concept that could foster the development of potent CPL emitting materials.

Conclusions

In summary, we have prepared and investigated the first examples of fully conjugated and non- π -extended (bis)BN-substituted penta- and hexahelicenes. Comparing them with their all-carbon congeners allowed the precise estimation of the influence of such BN-doping.

As seen from the solid-state structures, the helix torsion of pentahelicene BN[5] was significantly smaller than of CC[5]. Owing to a larger spatial separation, this was not the case for BN[6], resulting in congruent structures of both hexahelicenes. The impact of BN-substitution on $\lambda_{\rm abs}$ and $\lambda_{\rm fl}$ was rather small ($\Delta\lambda\approx10$ –15 nm). However, both the intensities of the lowenergy absorptions as well as the $\varphi_{\rm fl}$ values were significantly elevated, peaking at 0.17 (BN[6]) and 0.25 (BN[5]-(CN)₂). TD-DFT calculations allowed us to reproduce the causal, largely increased oscillator strengths of the BN-helicenes, being attributable to the reduction of symmetry. With respect to the configurational stabilities, BN[5] ($t_{1/2\rm rac}\approx80$ min at 60 °C) racemized four times slower than CC[5], which is possibly due to NH-repulsions in the transition state.

Mirror-imaged ECD spectra were obtained and, according to the strengthened low-energy absorptions, the $\Delta\varepsilon$ was considerably intensified in this region for the BN-pentahelicenes, as well. Regarding CPL, the $|g_{\text{lum}}|$ value of BN[6] (1.33 \times 10⁻², cf. hexahelicene: 9 \times 10⁻⁴) was unusually high, as elevated dissymmetry factors are usually associated with a loss of fluorescence efficiency.¹³

Taking into account that a selective dibromination of BN[5] was possible and the boron atoms may be modified with nucleophilic groups at an earlier stage of the reaction sequence,⁸⁷ BN-helicenes with tailored push–pull substitution patterns could be synthesized in the future in order to explore their potential as efficient CPL emitters.

Data availability

Experimental and theoretical details supporting the statements of this article are included in the ESI.†

Author contributions

Y. A. and A. S. conceived the project. Y. A. and N. W. performed the syntheses and routine analytics. Y. A. and S. M. L. performed the chiral resolutions, kinetics and chiroptical measurements, which were supervised by A. G. C. and M. J. P. P. performed the X-ray diffraction analyses. Y. A., S. M. L. and S. K. performed the DFT calculations, which were supervised by T. N. M. M. collected the quantum yields. D. M. performed the lifetime and excitation experiments. The manuscript was written by Y. A and S. M. L. All authors participated in reviewing the manuscript.

Conflicts of interest

There are no conflicts to declare.

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