Synthesis, Photophysical and Electronic Properties of New Red-to-NIR Emitting Donor–Acceptor Pyrene Derivatives

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Abstract: We synthesized new pyrene derivatives with strong bis(para-methoxyphenyl)amine donors at the 2,7-positions and π-azacene acceptors at the K-region of pyrene. The compounds possess a strong intramolecular charge transfer, leading to unusual properties such as emission in the red to NIR region (700 nm), which has not been reported before for monomeric pyrenes. Detailed photophysical studies reveal very long intrinsic lifetimes of > 100 ns for the new compounds, which is typical for 2,7-substituted pyrenes but not for K-region substituted pyrenes. The incorporation of strong donors and acceptors leads to very low reduction and oxidation potentials, and spectroelectrochemical studies show that the compounds are on the borderline between localized Robin-Day class-II and delocalized Robin-Day class-III species.

Introduction

The polycyclic aromatic hydrocarbon (PAH) pyrene is among the most widely studied chromophores and possesses some unique properties such as intense blue emission and an exceptionally long-lived excited singlet state together with excimer and exciplex formation.[3] In addition to its photophysical properties, it has high chemical stability and charge-carrier mobility. Therefore, pyrene derivatives have been used in a broad range of applications in diverse scientific fields such as organic light-emitting diodes (OLEDs), organic field-effect transistors (OFETs) and organic photovoltaic cells (OPVs).[4] Furthermore, they have been used for sensing of temperature,[5] pressure,[5] or pH,[5] or to detect guest molecules such as O₂ or NH₃,[6] organic molecules,[7] or metals[8] and to construct covalent organic frameworks.[9] Its monomer and excimer fluorescence have also been used to study the properties such as dynamics or interactions of macromolecules[10] and lipids.[11] It remarkably long fluorescence lifetime of 354 ns, compared to other PAHs, makes pyrene exceptionally well-suited for further applications such as the determination of cellular oxygen concentrations or reactive oxygen species (ROS) in biological systems.[12] Furthermore, 2-phenylethynylpyrenes have also been used as fluorescent labels for DNA.[13]

To adjust the properties for a specific application, electron donors and/or acceptors are often introduced onto a PAH core as they strongly modulate the frontier orbitals. Substituting PAHs with donors and acceptors gives derivatives with properties such as a permanent dipole moment, charge transfer (CT) excited states, strong solvatochromism, environmentally influenced photophysics, the possibility for energy or electron transfer, and narrowed energy gaps.[14, 15] Furthermore, such chromophores can absorb and emit in the near-infrared (NIR) region, which is in demand for applications such as bioimaging and cell recognition, as NIR light penetrates deeper into biological tissues, is less damaging than visible or UV light, and gives minimum interference from background autofluorescence by biomolecules.[16] In general, materials with high HOMO energies, such as the compound N,N'-diphenyl-N,N'-bis(3-methylphenyl)-(1,1'-biphenyl)-4,4'-diamine (TPD), are especially useful for hole transport.[17] Common π-donors that have been used in dyes to boost HOMO energies include amines, with a lone pair on the nitrogen, such as diislylamo, diethylamino, dimethylamino or carbazolyl moieties.[18] Diarylamine are among the strongest π-electron donors and have been employed in diverse applications[19] due to their outstanding physical, photochemical and electrochemical properties, and they are easy to synthesize and handle. [20] A methoxy
group at the position *para* to the nitrogen not only increases the electron donating strength of diarylamines, but enables reversible oxidations. Nevertheless, a significant drawback can be energy loss due to the rotational motion of the phenyl rings which can result in a reduction in luminescence efficiency. The compound 1,1,7,7-tetramethyl-julolidine is known to be an even stronger \( \pi \)-donor than diarylamines. The julolidine moiety has been thoroughly studied since its discovery in 1892 by Pinkus, and is used in a wide range of dyes. Its nitrogen lone pair is conformationally restricted to remain parallel to the aromatic system, in our case, the pyrene moiety, in both the ground and excited states, and our previous studies revealed a significantly enhanced electron donating effect on the pyrene core compared to diarylamines.

Pyrene exhibits ten peripheral reactive positions, which can be classified into three sets of chemically inequivalent sites (Figure 1a). Thus, the position of substitution is very important and, typically, the 1-, 3-, 6- and 8-positions are functionalized by electrophilic substitution reactions as the HOMO has its largest coefficients at these positions (Figure 1b). Therefore, \( \pi \)-orbitals of substituents at the 1-, 3-, 6-, and 8-positions mix very efficiently with pyrene’s HOMO/LUMO orbitals. However, unsymmetrical substitution at these positions is rather challenging due to the numerous possible isomers (Scheme 1).

Substituents at the 2,7-positions do not interact with the HOMO/LUMO orbitals of pyrene as they lie on the nodal plane (Figure 1b). However, they can interact strongly with the HOMO−1 and LUMO+1 of pyrene that have nonzero contributions at these positions. We previously reported that the photophysical properties of pyrenes with substituents at the 2-position significantly differ from those with substituents at the 1-position. Moreover, previous studies demonstrated that strong \( \pi \)-donors/acceptors are able to switch the energetic ordering of its HOMO/HOMO−1 and LUMO/LUMO+1, respectively, which results in greatly influenced photophysical and redox properties of these derivatives. Unsymmetrical substitution at these positions is straightforward via iridium-catalyzed C–H borylation, and the introduction of julolidine-like donors at the 2,7-positions (I, Scheme 2) even resulted in an unusual green luminescence.

The K-region (positions 4,5,9,10, Figure 1) has large HOMO/LUMO as well as HOMO−1/LUMO+1 contributions. Ottonelli and co-workers showed that the contribution to the HOMO/LUMO increases in the order: 2,7 < 4,5,9,10 < 1,3,6,8, while contributions to the HOMO−1/LUMO+1 increase in the order...
4,5,9,10 < 1,3,6,8 < 2.7. The K-region possesses alkene-like rather than aromatic character. Hence, reagents such as osmium tetroxide, which are known to react with alkene double bonds, selectively react with pyrene at the 4,5,9,10-positions. Müller and co-workers reported a protocol for the asymmetric substitution at these sites and presented a few examples with emission into the orange region of the spectrum (613 nm in THF) (Scheme 1).[14, 38]

There are also reports on pyrenes that have donor and acceptor moieties on alternative positions, besides the typical three categories (vide supra). Hu and co-workers developed unsymmetric pyrenes with donors at the 1,3-positions and acceptors at the 5,9-positions that emit in the blue-green region (Scheme 1).[29] Sutherland and co-workers combined the 2,7- and 1,8-positions with the K-region (V and VI, Scheme 2), respectively, and obtained an S1→S0 absorbance up to 900 nm, as it was their objective to obtain strongly redshifted absorptions with high molar absorptivities. However, their derivatives are not emissive and thus studies are missing on the influence on the excited state properties and how additional K-region substituents influence 2,7-substituted pyrenes.[40,41]

In our previous reports we showed that the influence of a second amine donor at the 2,7-positions on the occupied orbitals is larger than that of a cyano or even Bmes, acceptor on the unoccupied orbitals and thus emission from the D–acceptor compound is further redshifted than the D–A derivative (II, Scheme 2). Therefore, our initial objective was to utilize two julolidine-like donors at the 2,7-positions and add additional acceptors to the pyrene core to obtain a new class of D–A pyrene derivative with unparalleled photophysical properties such as red to NIR emission and interesting redox behavior. Moreover, we were interested to see how much more the S1 state can be influenced by the addition of acceptors to the K-region.

For the acceptor unit, we chose n-azaacenes fused to the K-region of pyrene that were reported by Mateo-Alonso and co-workers and which possess strong π-accepting properties (VII and VIII, Scheme 2).[43] N-azaacenes have attracted much attention for their outstanding electronic properties and application in organic electronic devices and as anion radicals. In particular, pyrene-fused azaacenes are even more stable than their N-azaacene analogues and emit strongly in the blue-to-green region of the electromagnetic spectrum.[43,44]

Results and Discussion

Synthesis and structural characterization

The procedures used to synthesize the three key compounds are summarized in Schemes 3 and 4. At the beginning of this project, we aimed for derivative 7 (Scheme 3) and its analogue with ketone moieties at all four K-region positions (Scheme S1). The selective RuO4-catalyzed oxidation of pyrene at its 4- and 5-positions is the starting point for the synthesis route and gives the dione 1. This one-step reaction was reported by Harris and co-workers in 2005 as an alternative to the toxic OsO4-catalyzed oxidation of pyrene.[47,48] In this regard, RuO4 is generated in situ from RuCl3·3H2O and NaIO4, and the oxida-
tion takes place in a solvent mixture of MeCN, CH₂Cl₂ and H₂O (1:1:1.25). In 1981, Sharpless and co-workers reported that addition of MeCN to the RuO₄-catalyzed oxidation greatly improves the effectiveness of the reaction. However, this procedure suffers from several disadvantages. Thus, the reaction can only be performed on a small scale and, furthermore, there are various side products such as dialdehydes or acids, which Nowicka et al. identified in detail, that complicate the workup and reduce the yield.

Bodwell and co-workers introduced an improved RuO₄-catalyzed oxidation procedure to obtain 1, which is scalable and eases the workup. This method includes the additive N-methylimidazole, and the solvent MeCN is replaced by THF. In order to perform the Ir-catalyzed C–H borylation, the ketone groups had to be protected as a diketal to prevent unwanted reactions with the Ir-catalyst. The lack of reactivity of oxidized pyrene in other metal-catalyzed reactions has been reported before. Compound 2 was easily obtained by refluxing 1 in toluene with an excess of ethylene glycol for 20 h in the presence of p-toluene-sulfonic acid. The C–H borylation was performed according to our reported procedure and takes place at the 2,7-positions selectively providing 3.
The Bpin moieties were further converted to azide moieties using a procedure reported by Chang and co-workers.\textsuperscript{[32]} They converted arylibronic acids to arylazides in MeOH by using 1.50 equivalents of Na\textsubscript{2}CO\textsubscript{3} as the azide source and 5 mol\% of Cu(OAc)\textsubscript{2} as the catalyst. Subsequent reduction of the azido species 4 with Pd/C and H\textsubscript{2} gave the corresponding primary amine 5. The reaction was stirred at room temperature until monitoring by IR spectroscopy confirmed the complete disappearance of azide moieties (in the region 2000–2270 cm\textsuperscript{-1}) and the appearance of primary amine bands at 3450, 3363 and 1620 cm\textsuperscript{-1} (Figure S26). The amine moieties were alkylated in MeCN with 1-chloro-3-methylbutene in the presence of the base K\textsubscript{2}CO\textsubscript{3} according to the approach used in our previously reported synthesis of 1.\textsuperscript{[33]} The alkylation reaction was complete after 48 h and gave the desired product 6 as a bright yellow solid (42%). Unfortunately, the final step to obtain the desired derivative 7 was unsuccessful. Our aim was to achieve ring closure and simultaneously deprotect the acetal groups in order to regenerate the two ketone moieties at the K-region. However, using a diverse range of acids, the acetal protecting groups were removed first, which most likely generated a pyrene core that was too deactivated to permit the electrophilic ring closure to take place. Therefore, we decided to change our julolidine-like donor moiety to a diarylamine. The new synthesis route is depicted in Scheme 4. Compound 3 was transformed into the corresponding dibromo species 4 by a bromodeboronation reaction.\textsuperscript{[34]} Thus, 3 and CuBr\textsubscript{2} were suspended in a mixture of THF/MeOH/H\textsubscript{2}O (1:1:1) and the reaction mixture was stirred at 90 °C for 4 d. The Buchwald–Hartwig amination, which was performed to obtain 5 from 4 using Pd\textsubscript{2}(dba)\textsubscript{3}·CHCl\textsubscript{3} as the catalyst precursor and XPhos as the ligand, was achieved in a yield of 58%. The deprotection of 5 in a trifluoroacetic acid and water mixture (6:1) is straightforward, and compound 6 was obtained in 81% yield. The cyclocondensation reactions between dione 6 and 2,3-diaminomaleonitrile to give 7, or benzene-1,2-diamine to give 8, were performed in an ethanol/acetic acid mixture (1:1) at 80 °C for 15 h according to the procedure of Mateo-Alonso and co-workers.\textsuperscript{[35,36]}

The solid-state structures of compounds 2, 5, 6, and 7 were determined via single-crystal X-ray diffraction (Figure 2). In compound 7, the biphenyl unit of the pyrene moiety exhibits typical aromatic C=C bond lengths ranging from 1.391(4) to 1.418(3) Å (bonds a, b, f, g, i, and j and their symmetrical equivalents, Figure 2 and Table S2), as is also observed in pyrene. The c and e bonds on the unsubstituted site of the pyrene moiety are slightly longer (1.444(3) and 1.431(4) Å), and the d' bond (1.355(3) Å) is typical of a C=C double bond. This means that it can be viewed as a biphenyl unit constrained to be planar by a -CH=CH- group. It has also recently been observed for many other 2\textsubscript{a} and 2,7-substituted pyrenes by Marder and co-workers.\textsuperscript{[15,29,30]} Similar bond lengths have also been reported for both sides of the pyrene moiety of 2,7-bis(dianislylamino)pyrene,\textsuperscript{[32]} the azacene-free analogue of our compound 7. However, in 7 the c, d, and e bonds on the 4,5-azacene-substituted side of the pyrene moiety are all longer than the equivalent bonds on the unsubstituted side, the d bond (1.426(3) Å) being shorter than the c and e bonds (1.451(3) and 1.453(3) Å) (Table S2). Hence, the incorporation of the azacene moiety at the 4,5-positions has a bond-lengthening effect, and the d bond can no longer be compared with a C=C double bond but rather has aromatic character. Interestingly, the azacene-moiety also effects the central bond which links the two phenyl rings, and is slightly longer (1.435(3) Å) than in the azacene-free analogue compound (1.416(2) Å).\textsuperscript{[32]}

A similar, even more pronounced effect is observed in the unsymmetric compounds with acetics substituted at the 4,5-positions of pyrene, that is, compounds 2, 5', and 6'. While the unsubstituted side of the pyrene core still shows bond distances similar to those of pyrene or compound 7, the 4,5-substituted side shows long bond distances for the c, d, and e bonds (1.515(2)–1.555(3) Å), which are typical of C=C single bonds (Table S2). The central h bond is also further elongated (1.440(3)–1.447(3) Å) with respect to 7.

As is usually observed for aromatic amines, in compound 7 the N–C(pyrene) bond lengths are significantly shorter (1.406(3) and 1.415(3) Å) than the other N–C bonds of the amine moieties (1.422(3)–1.435(2) Å). Similar distances were also reported for the azacene-free pyrene with amines substituted at the 2- and 7- positions.\textsuperscript{[12]} The nitrogen atoms have a
nearly ideal trigonal planar configuration with the sum of the C-N-C angles being between 359.0(6) and 360.0(3)°. The interplanar angles between the NC₃ and pyrene planes (37.45(10)° for N1 and 32.3(2)°–33.2(4)° for the disordered groups bonded to N2) are in a similar range as those between NC₃ and the terminal phenyl rings (35.2(4)°–47.5(2)°) (Table S2). Again, this is in agreement with the NC₃–pyrene angle (31°) which was reported for the analogous azacene-free 2,7-substituted pyrene compound.[32] The methoxyphenyl groups R3 and R4 of 7 are strongly disordered and show a higher degree of rotational freedom than the groups R1 and R2. Indeed, the interplanar angles between the NC₃ planes and the terminal rings vary between 38.2(4)° and 47.5(2)° for the disordered parts of R3 and between 35.2(4)° and 39.0(6)° for those of R4. In addition, important intermolecular interactions involving the methoxyphenyl groups are only present for the R1 and R2 groups but are not observed for the R3 and R4 groups (Table S4).

In the crystal structure of 7, the molecules form π-stacked dimers related by inversion symmetry. Hence, a π-stacking interaction with an interplanar separation of around 3.41–3.47 Å is present between the pyrene core and the azacene group (Table S4). Dimers are arranged edge-to-face in a sandwich-herringbone packing, which is typically observed for pyrene itself and its derivatives (Figure 3).[33] Parallel and inverted molecules that are offset along the a axis exhibit C–C intermolecular interactions between their CN end groups. The molecular packing is further determined by the large steric demand of the amine moieties and the presence of tetrahydrofuran solvent molecules in the crystal lattice. Intermolecular C–H···N and H–H interactions are present between the R1 and R2 methoxyphenyl groups and the pyrene-4,5-azaacene core, while C–H–O interactions exist with the tetrahydrofuran molecule (Table S4). A Hirshfeld surface analysis was performed in order to quantify the nature and type of intermolecular interactions in 7.[35] The disorder of the methoxyphenyl groups was not taken into account, but only the parts of highest occupancies were considered in the analysis. Fingerprint analysis and its breakdown to the individual relative contributions,[36] shows a major contribution from H–H interactions (42%), followed by a significant amount from C–H (22%), N–H (15%), and O–H (10%) interactions (Figures S44 and S45); 5% C–C interactions and around 3.6% N–C interactions are observed as well.

**Photophysical and redox properties**

The absorption spectra of derivatives 6, 7 and 8 are depicted in Figure 4 and are generally similar to that of pyrene, in that the S₁→S₀ absorption is comparably weak with extinction coefficients of ε = 2700–4000 M⁻¹ cm⁻¹. However, for 2,7-substituted pyrenes, these are the largest extinction coefficients reported so far. Thus, the acceptor moieties at the K-region increase how allowed this transition is. Furthermore, the S₁→S₀ absorption has a strong bathochromic shift in the order 8 < 7 < 6, and remarkably broad, covering a range of 6000 cm⁻¹, with no vi-
brational progression, providing an indication of strong charge transfer (CT) character. In particular, dione 6 has a very pronounced bathochromic shift ($\lambda_{\text{max}}(\text{abs}) = 658 \text{ nm}$), which is significantly stronger compared to the analogous dione VI reported by Sutherland and co-workers (Scheme 2).

However, the donor moieties in their derivative are further separated from the pyrene core by alkyne and phenyl spacers. Hence, the donating ability through the 2,7-positions is reduced and, as a result, a weaker CT character is obtained. Thus, derivative VI has a smaller bathochromic shift with $\lambda_{\text{max}}(\text{abs}) = 575 \text{ nm}$ compared to derivative 6. It is interesting to observe that the substituents at the 2,7-positions in these derivatives have such a significant effect on the $S_1 - S_0$ absorption, which is unusual. The CT character of these derivatives is even more evident when comparing them to their analogues without the donor moieties at the 2,7-positions. Mateo-Alonso and co-workers reported that $\lambda_{\text{max}}(\text{em}) = 455 \text{ nm}$ for VII, the analogue of our derivative 7, which demonstrates that the resulting CT character introduced via the additional donors causes a bathochromic shift of 3500 cm$^{-1}$ from the $S_1 - S_0$ absorption in our systems. Compound VIII, the analogue of our derivative 8, has $\lambda_{\text{max}}(\text{abs}) = 435 \text{ nm}$ according to Sahoo et al. Hence, the CT character introduced through our additional donors at the 2,7-positions shifts the $S_1 - S_0$ absorption by 3700 cm$^{-1}$.

Compounds 6 and 7 are not emissive; however, 8 emits in the red to NIR region of the electromagnetic spectrum with $\lambda_{\text{max}}(\text{em}) = 652 \text{ nm}$ in toluene (Table 1), which is intriguing for pyrenes as they usually emit in the blue region and, to the best of our knowledge, such a red-shifted emission has not been reported before for monomeric pyrenes. Even pyrene-fused azaacenes reported previously do not have such a red-shifted emission. Furthermore, compound 8 exhibits significant solvatochroism, as the emission shifts bathochromically with increasing solvent polarity from toluene to CH$_2$Cl$_2$ by 1051 cm$^{-1}$ (48 nm), confirming the CT nature of the lowest energy excited state and a significant change in dipole moment between ground state and excited state. The large CT nature of the excited state becomes more evident when comparing the emission of 8 with previously reported analogues that only possess a donor or acceptor moiety. Derivative VIII emits in the blue region with $\lambda_{\text{max}}(\text{em}) = 471 \text{ nm}$ and compound IV in the green region with $\lambda_{\text{max}}(\text{em}) = 520 \text{ nm}$ (both in CH$_2$Cl$_2$) while we observe a $\lambda_{\text{max}}(\text{em}) = 700 \text{ nm}$ for 8 in CH$_2$Cl$_2$.

Nevertheless, the non-radiative decay rates are significantly increased in CH$_2$Cl$_2$ ($k_{nr} = 11 \times 10^7 \text{ s}^{-1}$), which is fully in line with the energy gap law as the reorganization energy is enhanced in polar solvents. Therefore, the quantum yield is strongly decreased in the polar solvent. The apparent Stokes shifts are very large for pyrenes with values ranging from 3900 to 4900 cm$^{-1}$ and are much larger compared to pyrenes that have D/A groups at the 2,7- or 1,3,6,8-positions only. However, the pyrene derivatives reported by Müllen and co-workers that have D/A moieties at the K-region exhibit even larger Stokes shifts ranging from 4500 to 5400 cm$^{-1}$. Interestingly, the radiative decay rates are rather slow ($k_r = 0.3 - 1.0 \times 10^7 \text{ s}^{-1}$), which is a result of the strong CT character. However, such slow radiative rate constants are also typical of forbidden transitions (Strickler–Berg relation) and, therefore, the lifetimes remain quite long ($\tau = 104 - 293 \text{ ns}$) in this derivative, which is a typical property of 2,7-substituted pyrenes.

There are not many reports on lifetimes of K-region substituted pyrenes, but they are typically in the range of ($\tau_s = 13 - 16 \text{ ns}$), which we recently reported. Müllen and co-workers do not give lifetimes for their K-region D/A derivatives; however, the $S_1 - S_0$ absorptions are significantly more allowed with $\epsilon > 7000 \text{ M}^{-1} \text{ cm}^{-1}$ and, therefore, shorter lifetimes than for 2,7-substituted pyrenes can be assumed. Compared to compound 8, derivative 7 has a more pronounced CT nature, hence its energy gap is even smaller and, thus, it is possible that the non-radiative decay rates are largely increased, so that fluorescence becomes too weak to detect.

### Electrochemistry

In order to determine the impact of the acceptor groups on the K-region in combination with the donors at the 2,7-positions, cyclic voltammetry was performed. The cyclic voltammograms are shown in Figure 5 and the respective reduction and oxidation potentials are given in Table 2. All derivatives exhibit one reversible reduction, whereas compound 6 has the lowest reduction potential with $E_{1/2} = -1.13 \text{ V}$ and 8 the highest with $E_{1/2} = -1.84 \text{ V}$ vs. Fc/Fc$^+$. Hence, the accepting strength of our derivatives is quite strong and decreases in the order 6 $>$ 8. Furthermore, all three derivatives can be reversibly oxidized twice; derivative 8 is the easiest to be oxidized with $E_{1/2} = 0.17$ and 0.39 V, and derivatives 6 and 7 have the same first ($E_{1/2} = 0.28$) and second (0.45 V) oxidation potentials versus Fc/Fc$^+$. Hence, our derivatives possess quite strong donors and the donating strength in 8 is only minimally influenced by the addi-

### Table 1.

| Cpd | Medium | $\lambda_{\text{max}}(\text{nm})$ | $\lambda_{\text{em}}$ | Apparent Stokes shift | $\tau$ | $\Phi$ | $\tau_s$ | $k_r$ | $k_{nr}$ |
|-----|--------|-----------------|------------------|-------------------|-------|-----|-------|------|--------|
| 6   | toluene| 658 (2.7), 408 (7.0), 344 (41), 299 (31) | -                | -                | -     | -   | -     | -    | -      |
| 7   | toluene| 542 (3.1), 427 (8.5), 348 (48), 300 (42) | -                | -                | -     | -   | -     | -    | -      |
| 8   | toluene| 519 (4.0), 417 (21), 399 (17), 354 (42), 333 (76), 317 (78) | 652   | 3930            | 32.3  | 0.31| 104   | 1.0  | 2.1    |
| 8   | CH$_2$Cl$_2$| 521, 416, 397, 352, 332, 316 | 700  | 4908            | 8.8   | 0.03| 293   | 0.3  | 11.0   |
| 8   | solid  | -               | 679  | -               | <0.01 | -   | -     | -    | -      |

(a) Excited at the respective $\lambda_{\text{max}}(\text{abs})$ of $S_1 - S_0$. 

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Upon oxidation to the respective monocations $7^+$ and $8^+$, a band rises in the NIR that is very broad, covering a range of around 7000 cm$^{-1}$. The observed plateau may be caused by two overlapping peaks of a vibronic progression. Thus, the peak maximum depends on the relative intensities of the first and second overtones. For $7^+$, the lowest energy band has a maximum at $\nu_{\text{max}}^{\text{IV-CT}} = 6400$ cm$^{-1}$ (1563 nm, $\epsilon = 15160$ m$^2$ cm$^{-1}$), but for $8^+$, the maximum is $\nu_{\text{max}}^{\text{IV-CT}} = 5350$ cm$^{-1}$ (1869 nm, $\epsilon = 28830$ m$^{-1}$ cm$^{-1}$) with some asymmetry in its shape (Figure S1). The absorption spectra of the monocations $7^+$ and $8^+$ are very similar to that of compound III*, which was reported by Ito and co-workers, for which the lowest energy band has a maximum at 5260 cm$^{-1}$ (1900 nm) and is also slightly asymmetric.\(^{32}\) In general, delocalized (Robin-Day class-III) derivatives possess an asymmetric and narrow lowest energy band, while in localized (Robin-Day class-II) derivatives the lowest energy band results from an intervalence charge transfer (IV-CT) with a well-defined symmetric Gaussian shape.\(^{64}\) For this reason, Ito and co-workers concluded that compound III* is a fully delocalized Robin-Day class-III derivative and assumed an electronic coupling $V$ of $\frac{\hbar}{\tau_{\text{ab}}} = 2608$ cm$^{-1}$.\(^{60}\) However, our TD-DFT computations on the monocations $7^+$ and $8^+$, with a specially adjusted functional (BLYP with 35% exact HF exchange\(^{66}\)), show that the lowest energy excitation is a result of the promotion of an electron from the $\beta$-HOMO to the $\beta$-LUMO orbital and these orbitals show the expected phase behavior for localized Robin-Day class-II compounds (Figure 7). Further analysis of this lowest energy band enables the calculation of the electronic coupling $V$ between the localized mixed valence states according to the Mulliken–Hush theory\(^{66}\) (Equation (1)) where $\mu_{\text{ab}}$ is the transition moment (evaluated by integration of the IV-CT band), and the diabatic dipole moment difference $\Delta\mu_{12}$ was evaluated using the DFT calculated adiabatic dipole moment difference $\Delta\mu_{\text{DFT}}$.\(^{67}\) According to these calculations, both derivatives possess a large dipole moment change between their ground and excited states of $\Delta\mu_{12} = 42$ D ($7^+$) and 43 D ($8^+$), which is typical for Robin-Day class-II compounds.

$$V = \frac{\mu_{\text{ab}}^2}{\Delta\mu_{12}} \quad \text{with} \quad \Delta\mu_{12} = \sqrt{\Delta\mu_{\text{DFT}}^2 + 4\mu_{\text{ab}}^2 ZS} > 1$$

Hence, the electronic coupling $V$ between the localized mixed valence states yields $V = 1279$ cm$^{-1}$ for $7^+$ and

### Spectroelectrochemistry

To gain further information about the electronic structure of the corresponding pyrene $7^+$, $7^{1+}$ and $8^+\ , 8^{1+}$ derivatives, we performed UV/Vis/NIR spectroelectrochemical measurements in CH$_2$Cl$_2$/0.1 m [nBu$_4$N][PF$_6$] (Figure 6).

### Table 2. Cyclic voltammetry results for 6, 7, and 8 in CH$_2$Cl$_2$/0.1 m [nBu$_4$N][PF$_6$] relative to the Fc/Fc$^+$ couple.

| Cpd | $E_{1/2}$ [V] red | $E_{1/2}$ [V] ox | $E_{1/2}$ [V] [ox] |
|-----|------------------|-----------------|-----------------|
| 6   | $-1.13$          | $0.28$          | $0.47$          |
| 7   | $-1.42$          | $0.28$          | $0.45$          |
| 8   | $-1.84$          | $0.17$          | $0.39$          |

### Table 3. Selected optical and electrochemical properties of the pyrene derivatives I to VIII.

| $\lambda_{\text{abs}}$ [nm] | $\lambda_{\text{em}}$ [nm] | $E_{1/2}$ [V] red | $E_{1/2}$ [V] ox | $E_{1/2}$ [V] [ox] |
|-----------------------------|-----------------------------|------------------|-----------------|------------------|
| I$^0$ | 475 (hexane) | 511 (hexane) | – | $-0.18$ (CH$_2$Cl$_2$) | $0.26$ (CH$_2$Cl$_2$) |
| II$^0$ | 481 (THF) | 525 (THF) | – | $-2.51$ (THF) | $+0.29$ (THF) |
| III$^0$ | 450 (hexane) | 464 (hexane) | – | – | – |
| IV$^0$ | 462 (THF) | 497 (THF) | – | – | – |
| V$^0$ | 453 (toluene) | 482 (toluene) | – | – | – |
| VI$^0$ | 630 (CH$_2$Cl$_2$) | – | – | – | – |
| VII$^0$ | 575 (CH$_2$Cl$_2$) | – | – | – | – |
| VII$^0$ | 455 (CH$_2$Cl$_2$) | 538 (CH$_2$Cl$_2$) | – | $-1.43$ (THF) | – |
| VII$^0$ | 435 (CH$_2$Cl$_2$) | 471 (CH$_2$Cl$_2$) | – | – | – |

[a] Redox potentials were measured with the addition of 0.1 m [nBu$_4$N][PF$_6$] and are referenced vs. the Fc/Fc$^+$ couple.

### Figure 5. Cyclic voltammograms of 6, 7, and 8 in CH$_2$Cl$_2$/0.1 m [nBu$_4$N][PF$_6$] with a scan rate of 250 mV s$^{-1}$. 

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1278 cm\(^{-1}\) for \(8^+\). Thus, the electronic coupling \(V\) of both derivatives is smaller than \(1 \text{,}220 \text{,}000 \text{ cm}^{-1}\), which shows that these compounds are class-II compounds, but lie close to the borderline between localized class-II and delocalized class-III compounds, as \(V\) is quite large. This result is also in line with the observed broad and asymmetric band shape. Upon further oxidation to the respective dications, the IV-CT band decreases and the band at 790–792 nm (12 660–12 630 cm\(^{-1}\)) rises significantly (\(7^2^+\): \(\varepsilon_{\text{max}} = 57 000 \text{ m}^{-1} \text{ cm}^{-1}\), \(8^2^+\): \(\varepsilon_{\text{max}} = 54 000 \text{ m}^{-1} \text{ cm}^{-1}\)). Furthermore, both spectra show bands rising at 727 nm (13 800 cm\(^{-1}\)) and 415–421 nm (24 000 cm\(^{-1}\)) while the bands at 320–350 nm (29 000–31 000 cm\(^{-1}\)) decrease, which is also very similar to the observations made for the oxidation to the monocation of compound \(\text{III}\).\[^{[32]}\]

Consequently, the acceptor moieties at the K-region of the pyrene bridge do not have a large influence on the electronic coupling between the two amine donor moieties and thus we prefer to classify compounds \(7^+\) and \(8^+\) as being at the borderline between class-II and class-III.

**DFT and TD-DFT calculations**

To rationalize the observed trends and properties we performed DFT and TD-DFT calculations. The ground state structures were first optimized in the gas-phase at the B3LYP/6-31 + G* level of theory. Previous studies\[^{[28]}\] have shown that range-separated hybrid functionals are necessary to obtain a reliable picture of the nature and relative energetic ordering of the excited states in pyrenes. We have thus used the CAM-B3LYP functional for the subsequent TD-DFT calculations.

The nitrogen 2p\(_z\) orbitals in both pyrene derivatives \(7\) and \(8\) mix very efficiently with the HOMO-1 of the pyrene core, which consequently switches the order of the HOMO and HOMO-1 orbitals. The pyrene-like HOMO (black, Figure 8), on the other hand, mixes with the acceptor unit at the K-region. Hence, the pyrene bridge and the acceptor unit are fully delocalized and thus, this orbital (now HOMO-1) is stabilized by around 0.54 eV in \(7 \) and 0.08 eV in \(8\). A new orbital (blue, Figure 8) of non-bonding character with large coefficients at the nitrogens of the dianisylamine donors is energetically positioned between the new HOMO and HOMO-1 orbitals, which was also observed for derivative IX. The LUMOs of both \(7\) and \(8\) are greatly delocalized over the acceptor units and the pyrene bridge. Hence, these orbitals are considerably stabilized, with the LUMO of \(7\) stabilized by 1.08 eV, and of \(8\) by 0.52 eV compared to the one in pyrene. This pronounced stabilization of the LUMOs in \(7\) and \(8\) is reflected in our cyclic voltammetry studies by their remarkably low reduction potentials of 1.51 eV in \(7\) and 1.82 eV in \(8\).
-1.42 V and -1.84 V vs. Fc/Fc⁺, respectively. The pyrene LUMO and LUMO + 1-like orbitals (grey, Figure 8) are not significantly affected in 7 (now LUMO + 2 and LUMO + 3, respectively) and 8 (now LUMO + 1 and LUMO + 2, respectively). The pyrene LUMO-like orbital is delocalized over the acceptor unit and the pyrene core, while the pyrene LUMO + 1-like one is delocalized over the diisylamine donors and the pyrene core.

The TD-DFT calculations show that the nature of the S₁→S₀ transition changes in such a way that it is no longer a nearly 50:50 weighted configuration interaction of HOMO→-1→LUMO and HOMO→LUMO + 1 as in pyrene (Tables 4 and 5). In 7 and 8, the S₁→S₀ transitions are nearly pure HOMO→LUMO transitions, which have significant CT nature (Figure 8). They are

\[ E_{\text{HOMO}} = -3.20 \text{eV}, \quad E_{\text{LUMO}} = -3.50 \text{eV}, \quad E_{\text{HOMO}+1} = -3.60 \text{eV}, \quad E_{\text{LUMO}+1} = -3.80 \text{eV} \]

\[ \text{S₁} \rightarrow \text{S₀} \]

Table 4. TD-DFT results of the compounds 7⁺, 8⁺, and III⁺ using UBLYP with 35% exact exchange admixture, an SVP basis set and a polarizable continuum model accounting for solvent effects (CH₂Cl₂).

| Configuration (major contribution > 10%) |
|------------------------------------------|
| Configuration (major contribution > 10%) |

Table 5. TD-DFT results (CAM-B3LYP/6-31 + G*) for the five first vertical transitions of pyrene, 7 and 8.

-1.42 V and -1.84 V vs. Fc/Fc⁺, respectively. The pyrene LUMO and LUMO + 1-like orbitals (grey, Figure 8) are not significantly affected in 7 (now LUMO + 2 and LUMO + 3, respectively) and 8 (now LUMO + 1 and LUMO + 2, respectively). The pyrene LUMO-like orbital is delocalized over the acceptor unit and the pyrene core, while the pyrene LUMO + 1-like one is delocalized over the diisylamine donors and the pyrene core.

The TD-DFT calculations show that the nature of the S₁→S₀ transition changes in such a way that it is no longer a nearly 50:50 weighted configuration interaction of HOMO→-1→LUMO and HOMO→LUMO + 1 as in pyrene (Tables 4 and 5). In 7 and 8, the S₁→S₀ transitions are nearly pure HOMO→LUMO transitions, which have significant CT nature (Figure 8). They are
strongly bathochromically shifted in the order 7 > 8 with low oscillator strengths, which matches well with the absorption maxima and extinction coefficients that were measured for this band (7: \( \lambda_{\text{max}} \) (abs) = 542 nm, \( \varepsilon = 3100 \text{ M}^{-1} \text{cm}^{-1} \) and 8: \( \lambda_{\text{max}} \) (abs) = 519 nm, \( \varepsilon = 4000 \text{ M}^{-1} \text{cm}^{-1} \)).

Conclusions

We have reported the synthesis and structural characterization of new pyrene derivatives with donor moieties at the 2,7-positions and an acceptor group at the K-region. In general, unsymmetrically substituted pyrene derivatives remain rare due to their challenging synthesis. The influence of the donors at the 2,7-positions and the acceptors at the K-region on the photophysical and electrochemical properties of the compounds is remarkable. The new derivatives possess very broad absorptions with maxima at 519–658 nm, and tails up to 800 nm, and compound 8 emits in the red to NIR region, which has not previously been reported for monomeric pyrenes to that extent. The intrinsic lifetimes remain rather long (\( \tau_R = 104–293 \text{ ns} \)), which is a typical property of 2,7-substituted pyrenes, whereas the excited states of most K-region substituted pyrenes have lifetimes \( \tau_R \) of only around 13–16 ns. Cyclic voltammetry studies reveal two reversible one-electron oxidations and one reversible reduction for all three derivatives 6, 7 and 8, with very low potentials showing the unique donating and accepting properties of our derivatives. Spectroelectrochemical measurements suggest a strong electronic coupling between the substituents at the 2,7-positions for 7 and 8. Our DFT and TD-DFT calculations indicate that these properties are the result of the very strong donors and acceptors which couple very well with the pyrene orbitals, resulting in a reordering of the occupied orbitals. Consequently, the \( S_2 \) state of these derivatives has strong CT nature giving rise to unparallelled properties.

Experimental Section

General considerations

The catalyst precursors \([\text{Ir(COD)(OMe)}]_{2(6)}, [\text{Pd}_{2}(\text{dba})_{3} \text{CHCl}]_{2(9)}\) were prepared according to literature procedures, \( \text{B}_{2} \text{pin} \), was a gift from AllyChem Co. Ltd. while other starting materials were purchased from commercial sources and used as received. Solvents used for synthesis were HPLC grade, further treated to remove trace water using a commercial solvent purification system from Innovative Technology Inc. and deoxygenated using the freeze-pump-thaw method.

The \( ^1H, ^13C(H), ^1B(H), ^1N \) HMBC NMR spectra were recorded at room temperature in CDCl3 or CD3Cl, solution either on a Bruker Avance I HD 500 (\( ^1H, 500 \text{ MHz}; ^1C, 125 \text{ MHz}; ^1B, 160 \text{ MHz} \)), a Bruker Avance III HD 300 (\( ^1H, 300 \text{ MHz}; ^1C, 75 \text{ MHz}; ^1B, 96 \text{ MHz} \)) or a Bruker Avance 200 (\( ^1H, 200 \text{ MHz} \)). \( ^1H \) NMR and \( ^13C \) spectra are referenced to the residual protonated solvent CDCl3 (\( \delta = 7.26, \text{CDCl}_3 \)) and CHDCl3 (\( \delta = 5.32, \text{CDCl}_3 \)). \( ^1B^1(H) \) NMR signals were referenced to external BF3·OEt2, and \( ^1N \) HMBC signals were referenced to MeNO2 + 10%CDCl3, Chemical shifts are listed in parts per million (ppm) and coupling constants in Hertz (Hz).

HRMS were recorded using a Thermo Scientific Exactive Plus Orbitrap MS system with either an Atmospheric Sample Analysis Probe (ASAP) or by Electrospray Ionization (ESI). Elemental analyses were performed on an Elementar vario MICRO cube elemental analyzer. Cyclic voltammetry experiments were performed using a Gamry Instruments Reference 600 potentiostat. A standard three-electrode cell configuration was employed using a platinum disk working electrode, a platinum wire counter electrode, and a silver wire, separated by a Vycor tip, serving as the reference electrode. Formal redox potentials are referenced to the ferrocene/ferroenium ([CP2Fe]0/+2) redox couple by using decamethylferrocene ([CP2Fe]; \( E_{1/2} = 0.532 \text{ V} \) in CH3Cl) as an internal standard. Tetra-n-butylammonium hexafluorophosphate ([nBu4N][PF6]) or [nBu4N] [Al(O(CF3)2)]3 were employed as supporting electrolytes. Compensation for resistive losses (\( iR \) drop) was employed for all measurements.

Crystals suitable for single-crystal X-ray diffraction were selected, coated in perfluoropolyether oil, and mounted on MiTeGen sample holders. Diffraction data were collected on a Bruker X8 Apex II 4-circle diffractometer with a CCD area detector using Mo-Kα radiation monochromated by graphite (2, 6, 11, 12) or multi-layer focusing mirrors (5, 10). Diffraction data of 7 were collected on a Bruker D8 Quest 4-circle diffractometer with a CMOS area detector (Photon II) and multi-layer mirror monochromated Mo-Kα radiation. The crystals were cooled using Oxford Cryostream or Bruker KryoFlex II low-temperature devices. Data were collected at 100 K. The images were processed and corrected for Lorentz-polarization effects and absorption as implemented in the Bruker software packages. The structures were solved using the intrinsic phasing method (SHELXT) and Fourier expansion technique. All non-hydrogen atoms were refined in anisotropic approximation, with hydrogen atoms bonded to nitrogen atoms “riding” in idealized positions, by full-matrix least squares against \( F^2 \) for all data, using SHELXL software. In compound 11, the coordinates of the hydrogen atom of the water molecule, which lies on a two-fold rotation axis, were refined freely, but restraints were applied to the O-H and H-H distances. In compound 5 the coordinates of the hydrogen atoms bonded to nitrogen were refined freely. The methoxyphenyl groups as well as the tetrahydrofurane solvent molecule are strongly disordered in compound 7. Hence, several restraints had to be applied in the refinement. Diamond3 software was used for graphical representation. Hirstfeld surfaces were calculated and analyzed using the Crystal Explorer program. Other structural information was extracted using Mercury and OLEX2 software. Crystal data and experimental details are listed in Table S1. CCDC 1917153 (2), 1917154 (5), 1917155 (6), 1917156 (7), 1917157 (10), 1917158 (11), and 1917159 (12) contain the supplementary crystallographic data for this paper. These data are provided free of charge by The Cambridge Crystallographic Data Centre.

General photophysical measurements

All photophysical measurements were carried out under an argon atmosphere. All solution state measurements were performed in standard quartz cuvettes (1 cm x 1 cm cross section). UV/Vis absorption spectra were recorded using an Agilent 1100 diode array UV/Vis spectrophotometer. Excitation, emission, lifetime and quantum yield measurements were recorded using an Edinburgh Instruments FLS920 spectrophotometer equipped with a 450 W Xenon arc lamp, double monochromators for the excitation and emission pathways, and a red-sensitive photomultiplier (PMT-R928P) and a near-IR PMT as detectors. The measurements were made in right-angle geometry mode and all spectra were fully corrected for the
spectral response of the instrument. All solutions used in photo-
physical measurements had a concentration lower than $10^{-5}$ mol L$^{-1}$.

Fluorescence quantum yield measurements

Fluorescence quantum yields of the samples were measured using a calibrated integrating sphere (150 mm inner diameter) from Edin-
burgh Instruments combined with the FLS920 spectrophotometer
described above. For solution-state measurements, the longest
wavelength absorption maximum of the compound in the respec-
tive solvent was chosen for the excitation. In order to exclude self-
absorption, the emission spectra were measured with dilute sam-
ple solutions (ca. 0.1 OD at the excitation wavelength).

Fluorescence lifetime measurements

Lifetime measurements were conducted using the time-correlated
single-photon counting method (TCSPC) on the FLS920 spectro-
photometer equipped with a high-speed photomultiplier tube
positioned after a single emission monochromator. Measurements
were made in right-angle geometry mode, and the emission was
collected through a polarizer set to the magic angle. Solutions
were excited with either a 315 (pulse width 932.5 ps), 376 (pulse
width 72.6 ps) or a 472 nm (pulse width 90.6 ps) pulsed diode laser
at repetition rates of 1–5 MHz and were recorded at emission
maxima. Decays were recorded to 10,000 counts in the peak chan-
nel with a record length of at least 4,000 channels. The band-pass
of the monochromator was adjusted to give a signal count rate of
<20 kHz. Iterative reconvolution of the IRF with one decay func-
tion and nonlinear least-squares analysis were used to analyze the
data. The quality of all decay fits was judged to be satisfactory,
based on the calculated values of the reduced $\chi^2$ and Durbin-
Watson parameters and visual inspection of the weighted and au-
tocorrelated residuals.

Spectroelectrochemical measurements

Spectroelectrochemical experiments in reflection mode were per-
formed using an Agilent Cary 5000 Spectrophotometer in combi-
nation with a designed sample compartment consisting of a cylin-
drical PTFE cell with an Infrasil wedge window with an angle of
0.5° and an adjustable three-in-one electrode (6 mm platinum disc
working electrode, 1 mm platinum counter electrode and pseudo
reference electrode). The potentials were adjusted with a Gamry
600 potentiostat and all experiments were performed at room
temperature under an argon atmosphere.

Theoretical studies

All calculations (DFT and TD-DFT) were carried out with the Gaussi-
an 09 (Rev. E01) program package[20] and performed on a
parallel cluster system. GaussView 5.0.9 was used to visualize the
results, to measure calculated structural parameters, and to plot or-
bital surfaces (isovalue: ±0.02 a.u.).[15] The ground-state geomet-
ries were optimized using the B3LYP functional[21] in combination
with the 6-31G(d) basis set.[22] The optimized geometries were con-
firmed to be local minima by performing frequency calculations
and obtaining only positive (real) frequencies. Based on these optim-
ized structures, the lowest-energy gas-phase vertical transitions
were calculated (singlets, 10 states) by TD-DFT, using the Coulomb-
attenuated functional CAM-B3LYP[23] in combination with the 6-
31G + (d,p) basis set. DFT calculations on the monomers of 7 and
8 were carried out, using UBLYP with 35% exact-exchange admix-
ture, the SVP basis set and a polarizable continuum model ac-
counting for solvent effects. The time dependent (TD-DFT) cal-
culations were performed at the same level of theory.[24,25]

Synthesis of the pyrene derivatives

Pyrene-4,5-dione (1): In a round bottom flask pyrene (10.0 g,
49.4 mmol, 1.00 equiv) and RuCl$_3$·xH$_2$O (987 mg, 4.74 mmol,
0.096 equiv) were dissolved in 200 mL of MeCN. NaO$_2$ (423 g,
198 mmol, 4.00 equiv) was dissolved in 250 mL of hot water and
carefully added to the pyrene solution. Another 200 mL of CH$_2$Cl$_2$
were added and the reaction mixture was vigorously stirred for
18 h at room temperature. After filtration of the suspension through
Celite, the filtrate was extracted with CH$_2$Cl$_2$ (3×100 mL) and the
combined organic phases were washed with saturated aqueous Na$_2$SO$_4$
and H$_2$O and then dried over Na$_2$SO$_4$. After re-
moving the solvent under reduced pressure the product was ob-
tained as a dark orange solid (7.56 g, 66%). The spectroscopic data
match those reported previously.[47,50,80] $^{1}$H NMR (200 MHz, CDCl$_3$):
$\delta$ = 8.51 (dd, $J$ = 7 Hz, $J$ = 1 Hz, 2H), 8.19 (dd, $J$ = 8 Hz, $J$ = 1 Hz, 2H),
7.87 (s, 2H), 7.77 (t, $J$ = 8 Hz, 2H) ppm.

Pyrene-4,5-di(ethyleneglycol)ketal (2): Pyrene-4,5-dione (1)
(4.92 g, 21.2 mmol, 1.00 equiv) was suspended in 200 mL of tolu-
ene and ethylene glycol (148 mL, 2.65 mol, 125 equiv) and p-tolue-
nesulfonyl acid (1.81 g, 9.53 mmol, 4.54 equiv) were added. The re-
action mixture was refluxed at 125 °C for 20 h. Toluene was re-
moved under reduced pressure and 400 mL of water. The crude
product was isolated by filtration, washed with water and purified
by flash chromatography (silica, cyclohexane/ethyl acetate 9:1).
The product was isolated as a white solid (3.46 g, 10.8 mmol, 51 %).
The spectroscopic data match those reported previously.[47,50,80] $^{1}$H NMR
(500 MHz, CD$_2$Cl$_2$): $\delta$ = 7.98 (dd, $J$ = 7 Hz, $J$ = 1 Hz, 2H), 7.97 (dd, $J$ =
7 Hz, $J$ = 1 Hz, 2H), 7.82 (s, 2H), 7.73 (dd, $J$ = 8 Hz, $J$ = 7 Hz, 2H),
4.29 (br, 4H), 3.75 (br, 4H) ppm. $^{13}$C NMR (125 MHz, CD$_2$Cl$_2$): $\delta$
= 133.5, 131.8, 129.2, 127.6, 127.1, 126.9, 124.6, 93.9, 62.3. HRMS
(AŠAP): m/z calculated for [C$_9$H$_8$O$_4$ + H$^+$]: 321.1121; found: [M+H]$^+$ 321.1115 ($\Delta m$ = 1.87 ppm). Elemental analysis calcd (%) for C$_9$H$_8$O$_4$: C 74.99, H 5.03; found: C 74.90, H 5.08.

2,7-Bis(Bpin)-4,5-di(ethyleneglycol)ketal-pyrene (3): In a argon-
filled glovebox, a Young’s tube was filled with pyrene-4,5-di(ethylene-
eglycol)ketal (2) (2.91 g, 9.08 mmol, 1.00 equiv), B$_2$pin$_2$ (4.61 g, 18.2 mmol, 2.00 equiv), Ir(COD)(OMe)$_2$
(260 mg, 0.51 mmol, 0.05 equiv), 4,4’-di-tert-butyl-2,2’-dipyridine
(dtbbp) (260 mg, 0.91 mmol, 0.10 equiv) and 50 mL of MTBE. The Young’s tube was sealed, and the reaction mixture was stirred at 80 °C for 20 h. After cooling to room temperature, the crude product was passed through a pad of silica using toluene as the eluent. The solvent was removed under reduced pressure and the crude product was purified via flash chromatography (silica, cyclohexane/ethyl acetate
95:5). The product was obtained as a white solid (4.58 g, 8.00 mmol, 88 %). $^{1}$H NMR (300 MHz, CD$_2$Cl$_2$): $\delta$ = 8.40 (d, $J$ = 1 Hz, 2H), 8.28 (d, $J$ = 1 Hz, 2H), 7.85 (s, 2H), 4.33–4.28 (m, 4H), 3.76–3.71 (m, 4H), 1.41 (s, 24 H) ppm. $^{13}$C NMR (75 MHz, CD$_2$Cl$_2$): $\delta$
= 136.6, 133.0, 131.6, 129.7, 128.6, 127.4, 126.7, 94.0, 84.6, 86.2, 25.2 ppm (one C not observed, likely that attached to B). $^{19}$F NMR (96 MHz, CD$_2$Cl$_2$): $\delta$ = 31.5 ppm. HRMS (AŠAP): m/z calculated for [C$_{29}$H$_{28}$F$_2$O$_6$ + H$^+$]: 753.2826; found: [M+H]$^+$ 753.2799 ($\Delta m$ = 4.71 ppm). Elemental analysis calcd (%) for C$_{29}$H$_{28}$F$_2$O$_6$: C 76.17, H 6.69; found: C 76.69, H 6.97.

2,7-Bis(bromo)-4,5-di(ethyleneglycol)ketal-pyrene (4): The compo-
und 2,7-bis(Bpin)-4,5-di(ethylene glycol)ketal-pyrene (3) (4.58 g,
7.99 mmol, 1.00 equiv) was diluted in 100 mL of THF and heated to
50 °C. Afterwards, 80 mL of MeOH was added and the heating bath
temperature was elevated to 90 °C. A solution of CuBr$_2$ (8.90 g,
40.0 mmol, 6.00 equiv) in 80 mL of H2O was slowly added to this reaction mixture via a dropping funnel. The reaction mixture was stirred for 90 min and an orange solid precipitated. The precipitate was isolated by filtration and washed with water and EDTA (0.1 M) solution. The crude product was purified via flash chromatography (silica, hexane/ethyl acetate 4:1) to give a white solid (1.02 g, 2.13 mmol, 58%). 1H NMR (300 MHz, CDCl3): δ = 7.49 (d, J = 2 Hz, 2H), 7.31 (s, 2H), 7.3, 2H, 7.24 (d, J = 2–7 Hz, 2H), 7.74–7.09 (m, 8H), 6.90–6.86 (m, 8H), 4.11 (br, 4H), 3.80 (s, 12H), 3.65 (br, 8H) ppm. 13Ct(H) NMR (75 MHz, CDCl3): δ = 156.6, 147.6, 141.3, 133.3, 132.0, 127.0, 126.8, 121.3, 119.2, 118.0, 115.1, 93.9, 62.1, 55.9 ppm. 11N (from 1N H HMBC) NMR (300 MHz, CDCl3): –285.8 ppm. HRMS (APCI+): m/z calcd for [C10H6NO3+H]+ 275.0314; found: [M+H]+ 275.0300 ([Δ] = 1.80 ppm). Elemental analysis calcd (%) for C10H6NO3: C 74.40, H 4.74, N 15.86.

2.7-Bis(DPA)-4,5-diene (6): In a round bottom flask, 2,7-bis(DPA)-4,5-diene (5) (500 mg, 645 μmol) was suspended in 10 mL of water. Then, 60 mL of TFA was added dropwise and the reaction mixture was stirred at room temperature for 15 h. The solution was then neutralized with an aqueous solution of NaHCO3 and extracted with ethyl acetate. The organic phases were dried over Na2SO4, and the crude product was washed with methanol and the solvents were removed under reduced pressure. The product 2,7-bis(DPA)-4,5-diene (6) was obtained as a dark green solid (360 mg, 534 μmol, 81%). 1H NMR (300 MHz, CDCl3): δ = 7.88 (d, J = 3 Hz, 2H), 7.43 (d, J = 3 Hz, 2H), 7.39 (s, 2H), 7.15–7.10 (m, 8H), 6.93–6.88 (m, 8H), 3.82 (s, 12H) ppm. 13Ct(H) NMR (75 MHz, CDCl3): δ = 181.1, 157.2, 148.1, 140.2, 132.9, 130.3, 127.5, 127.2, 123.5, 123.1, 122.7, 115.4, 55.9 ppm. 11N (from 11H H HMBC) NMR (300 MHz, CDCl3): –284.9 ppm. HRMS (APCI+): m/z calcd for [C10H6NO3+H]+ 275.0314; found: [M+H]+ 275.0300 ([Δ] = 1.80 ppm). Elemental analysis calcd (%) for C10H6NO3: C 74.40, H 4.74, N 15.86.

2.7-Bis(DPA)-4,5-azacene-CN (7): In a round bottom flask, 2,7-bis(DPA)-4,5-diene (6) (200 mg, 291 μmol), 2,3-diaminomaleonitrile (188 mg, 1.74 mmol), ethanol (35 mL) and acetic acid (35 mL) were stirred at 80°C for 15 h. The solution was then neutralized with an aqueous solution of NaHCO3. The precipitate was collected via filtration and washed with water and ethanol. Product 7 was obtained as an orange solid (360 mg, 534 μmol, 81%).

Syntheses from the initially planned route

2.7-Bis(azido)-4,5-diene(DPA)ketal-pyrene (4’): Compound 3 (6.62 g, 11.6 mmol, 1.00 equiv), NaH (2.26 g, 34.8 mmol) and CuOAc (232 mg, 1.16 mmol) were suspended in 400 mL of MeOH and stirred in a preheated oil bath at 55°C for 48 h. The green precipitate was collected and washed with EDTA solution, water and MeOH. The crude product was purified via column chromatography (silica, hexane/ethyl acetate 4:1) giving the product 4’ as a pale yellow solid (2.58 g, 55%). 1H NMR (200 MHz, CDCl3): δ = 7.75 (s, 2H), 7.64 (d, J = 2 Hz, 2H), 7.56 (d, J = 2 Hz, 2H), 4.32–4.24 (m, 4H), 3.76–3.68 (m, 4H) ppm. 13Ct(H) NMR (125 MHz, CDCl3): δ = 139.7, 155.2, 132.7, 127.5, 124.0, 118.0, 116.8, 93.5, 62.3 ppm. 11N (from 14N H HMBC) NMR (500 MHz, CDCl3): δ = –288.1 ppm. HRMS (ASAP+): m/z calcd for [C10H6NO3+H]+ 403.1149; found: [M+H]+ 403.1145 ([Δ] = 0.09 ppm).

2.7-Bis(amin)-4,5-diene(DPA)ketal-pyrene (5’): To a Schlenk flask, 4’ (1.03 g, 2.56 mmol) and 10% Pd/C (37.6 mg) were added under an inert atmosphere. After the addition of 200 mL of MeOH, the reaction mixture was frozen in liquid N2. Successively, the flask was evacuated and refilled with an excess of hydrogen and evacuated again. The procedure was repeated two more times and, after the last hydrogen flush was performed, the reaction mixture was thawed and stirred at rt. for 16 h. CH2Cl2 was added to the crude product and the suspension was then filtered. Afterwards, the solvent was removed under reduced pressure and the crude product was purified via column chromatography (silica, hexane/ethyl acetate 3:7). The final product 5’ was obtained as a pale yellow solid (440 mg, 49%). 1H NMR (300 MHz, CDCl3): δ = 7.47 (s, 2H), 7.25 (d, J = 2 Hz, 2H), 7.00 (d, J = 2 Hz, 2H), 4.41 (s, 4H), 4.20–4.15 (m, 4H), 3.79 (br, 4H) ppm. 13Ct(H) NMR (75 MHz, CDCl3): δ = 147.0, 134.0, 132.3, 126.8, 120.2, 114.8, 111.2, 94.6, 62.6 ppm. 11N (from 14N H HMBC) NMR (300 MHz, CDCl3): δ = –324.1 ppm. HRMS (ASAP+): m/z calcd for [C10H6NO3+H]+ 351.1339; found: [M+H]+ 351.1324 ([Δ] = 0.47 ppm).

2.7-Bis(DPA)-4,5-azacene-CN-tetras(3-methylbutyl-2-en-1-yl)-4,5-diene(DPA)ketal-pyrene (6’): To a Young's tube, compound 5’ (686 mg, 1.96 mmol), K2CO3 (1.35 g, 9.80 mmol), molecular sieves 4 Å (0.50 g) and 1-chloro-3-methylbut-2-ene (927 μL, 8.23 mmol) were suspended in 60 mL MeCN. The tube was sealed, and the reaction mixture was stirred at 80°C in an oil bath for 48 h. After cooling to rt., the precipitate was collected by filtration and washed with ethyl acetate. The solvent was removed from the filtr-
trate under reduced pressure and the remaining brown oil was pu-

rified by column chromatography (silica, hexane/ethyl acetate 96:4), which yielded product 6 as a yellow solid (518 mg, 42%). 1H NMR 500 MHz, CDCl3: δ = 7.49 (s, 2H), 7.38 (s, 2H), 7.00 (s, 2H), 5.30 (m, 4H), 4.42 (br, 4H), 4.03 (d, J = 6 Hz, 8H), 3.74 (br, 4H), 1.79 (s, 12H), 1.76 (s, 12H) ppm. 13C{1H} NMR (125 MHz, CDCl3): δ = 147.6, 134.8, 132.1, 131.7, 126.8, 122.1, 118.6, 112.9, 110.1, 94.5, 62.3, 49.1, 25.9, 18.1 ppm. HRMS (ASAP®): m/z calc for [C8H12N3O4]+ = 623.3843; found [M+H]+ = 623.3839 ([A] = 0.64 ppm).

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**Conflict of interest**

The authors declare no conflict of interest.

**Keywords:** borylation • K-region • luminescence • polycyclic aromatic hydrocarbons • redox

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does not have a pyrene core; however, for simplification reasons we kept the numbering as they are in pyrene derivatives.

The Stokes shift is defined as the energy difference between the 0-0 transition of the absorption and the emission. However, as the first transition is broad, the band maxima of the absorption and emission have been used to determine the Stokes shift throughout the paper. Consequently, the term "apparent Stokes shift" is used.

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