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Luminescent molecular switches based on dicationic P-doped polycyclic aromatic hydrocarbons†

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We report the efficient synthesis of electron-acceptors based on cationic P-doped polycyclic aromatic hydrocarbons. In particular, exo-skeletal modifications of the π -core were performed and their effect on the optical and redox properties was studied and compared to the effect of endo-skeletal modifications. Combined experimental (spectroelectrochemical and electron paramagnetic resonance) and theoretical studies performed on this entire family provide in-depth characterization of the reduced species and specifically of the stable radical cations formed and afford a guideline for the property tuning of such derivatives. Finally, electrochemical modulation of fluorescence performed on two compounds highlights the potential of these intrinsically switchable electroactive fluorophores.

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Polycyclic aromatic hydrocarbons (PAHs, also called nanographene when their size exceeds 1 nm) are bidimensional organic molecules composed of sp^2 C atoms.¹ This particular chemical composition endows them with unique properties such as strong light absorption and emission, charge transport in the solid state, *etc.* Consequently they are excellent candidates for building semi-conductors or active materials in a wide range of applications linked to molecular electronics. The modification of their carbon framework (ring size, edge configuration *etc.*) endows them with chiroptical or magnetic properties which even enlarge the scope of their potential applications in materials sciences.² Another strategy to tune both the structure and properties is to replace C-atoms by heteroatoms.³ During the last decade, various synthetic approaches allowed introducing single heteroatoms (B, N, O, P, S *etc.*) into the C-framework, therefore endowing it with unique reactivity and electronic properties.⁴ Introducing a second identical heteroatom into the framework has been recently attracting significant interest since this allows the investigation of the electronic communication between the main group elements through a π -conjugated

PAH backbone. As a representative example, Würthner *et al.* described electron-acceptors based on PAHs featuring two σ^3, λ^3 B atoms such as A (Fig. 1).⁵ These authors studied the complex structure–electronic property relationships (based on the position of the B atoms and the size of the PAH) and could also use one of their compounds as an electron acceptor in bulk heterojunction solar cells showing promising power conversion efficiencies of up to 3%. PAHs embedded with two σ^3, λ^4 N⁺ (B in Fig. 1) have also been described by Feng *et al.*⁶ They display viologen-like behavior (*e.g.*, two easily accessible reduction waves including stable radical cations). Among the heteroatoms, P is also of particular interest due to its high variety of valence and coordinance. In our effort to prepare P-containing PAHs,⁷ we recently developed a synthetic approach toward 5 dicationic P-containing PAHs.^{8,9} Various π -backbones could be used and their viologen-like behavior was exemplified on one prototype derivative (C, Fig. 1). This particular behavior was attributed to the presence of two σ^4, λ^4 -P⁺ connected through an even number of sp^2 C-atoms, by analogy with the Hünig strategy.¹⁰ In the present article, we describe the molecular engineering of the C-type PAH platform by using exoskeletal modifications (modification of R₁ and R₂, Fig. 1). The effect on the optical and redox properties is studied and compared to the effect of endoskeletal modifications (π , Fig. 1) based on a joint experimental and theoretical approach. Spectro-electrochemical and EPR studies were carried out on all compounds to investigate in the entire series the “viologen-like” behaviour previously observed for C (Fig. 1). In addition, by taking advantage of the luminescence associated with the redox properties, we demonstrate the electrofluorochromic properties of two

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Fig. 1 Reported examples of PAHs featuring two identical heteroatoms A–C (top) and illustration of our molecular engineering strategy (bottom) (counterions of B–C were omitted for clarity).

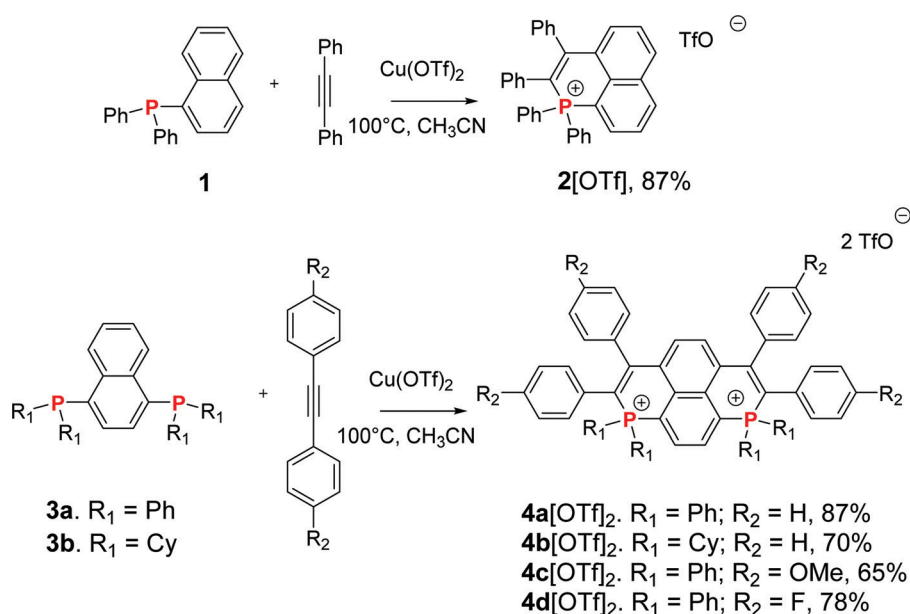
selected compounds. The development of intrinsically switchable electroactive fluorophores remains a challenge for the scientific community.¹¹

Synthesis

To prepare the targeted dicationic P-containing PAHs, we relied on a copper mediated radical approach recently developed by Wang *et al.*¹² In order to have in hand a reference compound, we first reproduced the synthesis of **2**[OTf] (Scheme 1) (see Fig. S16, ESI,[†] for X-ray structure).^{12a} Then, the phosphacyclization of bisphosphane **3a** with phenylacetylene afforded 1,8-bisphosphapyrenium **4a**[OTf]₂ (87% yield, Scheme 1). The synthetic approach is highly versatile as it is compatible with a wide scope of phosphines and alkynes. In order to further tune the optical and redox properties of the bisphosphapyrenium, we

prepared in good yields **4b**[OTf]₂ featuring electron rich exocyclic P-cyclohexyl groups. This compound was characterized by X-ray diffraction and the overall structure displays no important modifications compared to **4a**[OTf]₂ (mostly a planar π -framework and classical P–C bond lengths for phosphonium, Fig. S17, ESI[†]). We then modified the exocyclic C-core by introducing electron-donating *p*-methoxyphenyl groups (**4c**[OTf]₂) and electron poor fluorophenyl (**4d**[OTf]₂) by modification of the alkyne engaged in the phosphacyclization. Introduction of more electron rich (such as aminophenyl) or poor (cyanophenyl) substituents could not be achieved. However, these structural modifications highlight that the bisphosphapyrenium can be chemically tuned through straightforward exoskeletal modifications of various chemical functions (P atoms, C-frameworks).

In short, we prepared three new P-containing PAHs with variable exocyclic P-substituents and C-substituents and a



Scheme 1 Synthesis of **2**[OTf]^{12a} and **4a, 4b–d**[OTf]₂.⁸



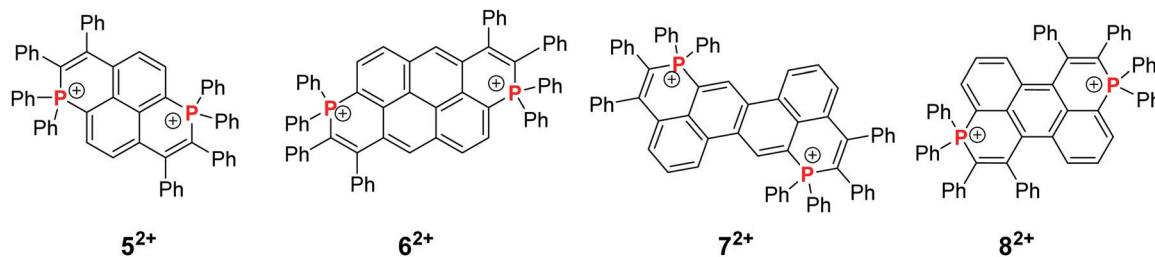


Fig. 2 Molecular structures of previously reported **5–8**²⁺.

reference compound (**2**[OTf]). We thus have in hand an entire family of eight dicationic featuring variable π -cores (**5–8**[OTf]₂, Fig. 2)⁸ or exocyclic substituents (**4a–d**[OTf]₂, Scheme 1).

Optical properties

The spectroscopic properties of **2**[OTf] and **4b–d**[OTf]₂ are investigated in diluted CH₂Cl₂ solutions ($c = 5 \times 10^{-6}$ mol L⁻¹, Fig. 3 and Table 1) and compared to the previously reported **4a**[OTf]₂.^{8,13} **4a**[OTf]₂ displays large absorption bands in the visible region with λ_{abs} at 420 nm.⁸ This transition can be assigned to a delocalized π - π^* transition according to TD-DFT calculations (*vide infra* and Fig. 7). As expected, this absorption

is considerably red-shifted and hyperchromically shifted compared to **2**[OTf] that displays only one P-heterocycle (Fig. 3).

We then focused our attention on the effect of exoskeletal modifications with the study of new derivatives **4b–d**[OTf]₂. Modification of the exocyclic P-substituent (cyclohexyl instead of phenyl) leads to a moderate blue-shift of the absorption ($\lambda_{\text{abs}}(\mathbf{4b}[\text{OTf}]_2) = 409$ nm). While the introduction of a (weak) electron-withdrawing group on the C-framework leaves the absorption spectrum almost unaffected ($\lambda_{\text{abs}}(\mathbf{4d}[\text{OTf}]_2) = 429$ nm), insertion of electron-donating groups strongly affects the absorption with a tail spanning to $\lambda = 518$ nm and a smaller extinction coefficient. In this compound intramolecular charge transfer from the rich methoxy groups toward the electron-deficient polyaromatic platform is envisaged as the cause of these modifications (*vide infra* for TD-DFT rationalization). All derivatives also display fluorescence in solution and a similar trend regarding the effect of donors/acceptors on the emission wavelength can be observed with a gradual red-shift in the series **2–4b–4ad–4c**[OTf]₂. Surprisingly, **4b**[OTf]₂ displays high photoluminescence quantum yield in solution (75%) compared to the other compounds. All derivatives display a large Stokes shift (Table 1), despite their rigid polyaromatic backbone. TD-DFT nevertheless reveals a significant change in geometry between the ground and the excited states (see Fig. S35, ESI[†]). This feature seems general for π -conjugated phosphonium and bisphosphonium and is clearly evidenced when compared with pyrene ($\sigma = 2800$ cm⁻¹) which displays a similar polyaromatic platform to **4a–d**[OTf]₂ and **5**[OTf]₂.^{12b,14,15} All dyes also show solid-state photoluminescence (in powders, Fig. S19, ESI[†]). No clear trend can be drawn on the impact of aggregation on the luminescence wavelength. Except for **4b**[OTf]₂, the luminescence in powder form is higher than in diluted solution. This increase nicely correlates with an emission enhancement induced by the restriction of the rotations of the 8 phenyl rotors and by the global bulkiness of the compounds that prevents π -stacking.

In conclusion, **4a–d**[OTf]₂ display strong absorption in the visible range as well as fluorescence both in solution and in the solid state. We show that exoskeletal modifications of the platforms also allow the tuning of the absorption/emission properties in the visible range.

Redox properties

The electrochemical properties of **2**[OTf] and **4b–d**[OTf]₂ have been investigated by cyclic voltammetry (CV) in dichloromethane



Fig. 3 UV-Vis absorption (top) and normalized emission spectra (bottom) of **2**[OTf] and **4ad**[OTf]₂ in CH₂Cl₂ ($c = 5 \times 10^{-6}$ M).



Table 1 Photophysical and redox data

	λ_{abs}^a [nm]	ϵ^a [M ⁻¹ cm ⁻¹]	λ_{em}^a [nm]	Φ^b (%)	σ (cm ⁻¹)	$\lambda_{\text{em solid}}^c$ [nm]	Φ^{bc} (%)	$E_1^{\text{red d}}$ [V]	$E_2^{\text{red d}}$ [V]
2[OTf]	359	15 000	539	14	9300	507	48	-1.61 ^e	—
4a[OTf] ₂	420	16 000	606	19	7300	552	39	-0.84 ^e	-1.23 ^e
4b[OTf] ₂	409	13 500	538	75	5900	542	52	-0.93 ^e	-1.40 ^e
4c[OTf] ₂	470	8500	625	2	5700	640	10	-0.89 ^f	-1.29 ^f
4d[OTf] ₂	429	11 500	607	13	6800	588	37	-0.82 ^e	-1.18 ^e
5[OTf] ₂	417	23 000	608	14	7600	554	37 ^e	-0.81 ^e	-1.24 ^e
6[OTf] ₂	430	55 000	575	13	6200	555	11 ^e	-0.98 ^e	-1.36 ^e
7[OTf] ₂	432	19 000	560	6	5600	566	19	-1.10 ^e	-1.44 ^f
8[OTf] ₂	522	19 000	672	4	4700	670	16	-0.64 ^e	-1.01 ^e

^a In CH₂Cl₂ (10⁻⁵ M). ^b Measured in a calibrated integrated sphere. ^c Measured in powder. ^d In DCM with Bu₄N⁺PF₆⁻ (0.2 M) at a scan rate of 200 mV s⁻¹. Potentials vs. Fc^{+/0}/Fc. ^e Reversible process ($\Delta E_p < 60$ mV). ^f Quasi-reversible process.

solution (see Fig. 4 and Fig. S19, ESI[†]) and compared to the previously reported **4a** (5–8[OTf]₂ can be found elsewhere).⁸ No oxidation processes are observed with any of the compounds under these experimental conditions. **4a**[OTf]₂ possesses two separated reversible reduction waves at low potential ($E_{\text{red1}}(\mathbf{4a}[\text{OTf}]_2) = -0.84$ V vs. Fc^{+/0}/Fc; $E_{\text{red2}}(\mathbf{4a}[\text{OTf}]_2) = -1.23$ V vs. Fc^{+/0}/Fc, Fig. 4 and Table 1). Comparison with **2**[OTf] (a single reduction wave with $E_{\text{red}} = -1.61$ V vs. Fc^{+/0}/Fc) clearly highlights the crucial role of the two cationic σ^4, λ^4 -P⁺ connected through an even number of sp² C-atoms in the viologen-like behavior of **4a**[OTf]₂.¹⁶ We underline that the first reduction potential of **4a** matches with that of methyl-viologen triflate recorded under the same conditions. Then the properties of the novel compounds **4b–d**[OTf]₂ were examined. As expected, replacing the P–Ph of **4a**[OTf]₂ by a more electron-rich P–Cy leads to a cathodic shift of 0.1 V ($E_{\text{red1}}(\mathbf{4b}[\text{OTf}]_2) = -0.93$ V vs. Fc^{+/0}/Fc). Such a phenomenon is also observed with the insertion of methoxy groups on the C-platform ($E_{\text{red1}}(\mathbf{4c}[\text{OTf}]_2) = -0.89$ V vs. Fc^{+/0}/Fc). For this compound, the two reduction processes are quasi-reversible. The presence of fluorophenyl leaves the reduction almost unaffected ($E_{\text{red1}}(\mathbf{4d}[\text{OTf}]_2) = -0.82$ V vs. Fc^{+/0}/Fc). These observations are consistent with the fact that reinforcing the electronic density on the platform makes the reduction more difficult and has an

impact on the reversibility. However, the fluorophenyl group seems to be a too “weak” acceptor to induce significant changes, in agreement with the optical data. The effect of the exoskeletal modification is thus rather easy to rationalize. As we previously reported, such rationalization is more challenging with the endoskeletal modifications: no clear trend can be drawn regarding the effect of π -extension.

In short, the combined optical and redox study reveals that **4a–d**[OTf]₂ possess all the characteristics of intrinsically switchable electroactive fluorophores.¹¹

Spectroelectrochemistry and EPR

In order to investigate the optical properties of the reduced forms of the dicationic P-containing PAHs, spectroelectrochemical and EPR analyses were performed on all derivatives. We previously reported the properties of **4a**[OTf]₂ for which the 3 redox states including a stable organic radical could be optically characterized.⁸ By studying the 7 other compounds, we show here that the entire family follows the same general trend (Fig. 5 and Fig. S19–S30, ESI[†]). In spectroelectrochemistry, the first reduced state is characterized by the appearance of new bands in the visible-NIR domains, which are characteristic of π -delocalized radicals. Upon further reduction toward the neutral state, these transitions disappear.¹⁷ The persistence in some cases of weak NIR absorption bands in the doubly reduced state probably comes from the presence of traces of the radical cations. The introduction of electron-rich substituents on the platform strongly impacts the stability of the reduced form as no reduced species of **4c**²⁺ could be clearly identified and the redox-triggered modifications appeared irreversible (Fig. S21 and S22, ESI[†]). Regarding the impact of the endoskeletal modifications, it appears that the spectral shapes and maximal wavelength of the absorption of the radical cations both depend on the nature of the platform, with absorption maxima from 662 nm for **7**^{•+} to 1218 nm in the case of **6**^{•+}. The return to the initial state is globally satisfying after the first reduction (see the ESI[†]). The return to the initial state is observed after the second reduction for **4a**²⁺, **4b**²⁺, **4d**²⁺, **6**²⁺ and **8**²⁺ (see Fig. S19–S30, ESI[†]).

Electron paramagnetic resonance of all 8 chemically formed radicals was then investigated. In all cases but **4c**^{•+} and **7**^{•+},

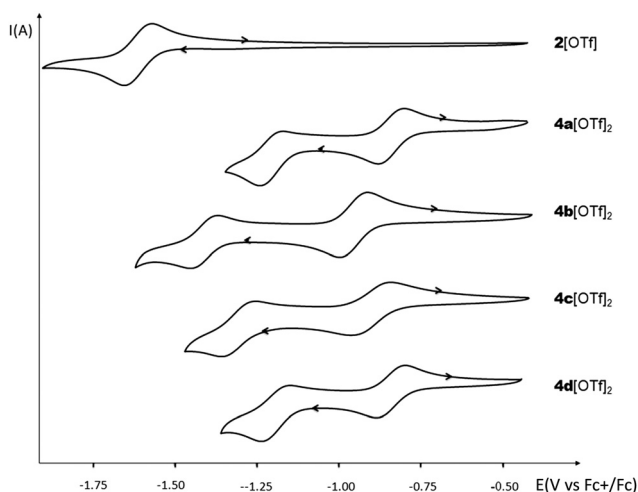


Fig. 4 Cyclic voltammograms of **2**[OTf] and **4a–d**[OTf]₂ ($c = 10^{-3}$ M) in DCM (Bu₄NPF₆ (0.2 M), 200 mV s⁻¹, potentials vs. Fc^{+/0}/Fc).





Fig. 5 Absorption spectra measured upon electrochemical reduction of (a) **4a**[OTf]₂, (b) **4a**[OTf]₂ (c) **6**[OTf]₂, and (d) **8**[OTf]₂ ($c = 5 \times 10^{-4}$ M) in DCM (0.2 M Bu₄NPF₆).

which appear too unstable to be recorded under these conditions, the organic nature of these radicals is confirmed by *g*-factor values in the 2.0032–2.0034 range (Fig. 6 and Fig. S32 and S33, ESI†). The hyperfine structure of all compounds is also

fully consistent with the presence of a radical fully delocalized on the entire planar π -skeleton, without contribution of the exocyclic moieties. Hence, the hyperfine couplings are well reproduced by considering the coupling of the unpaired

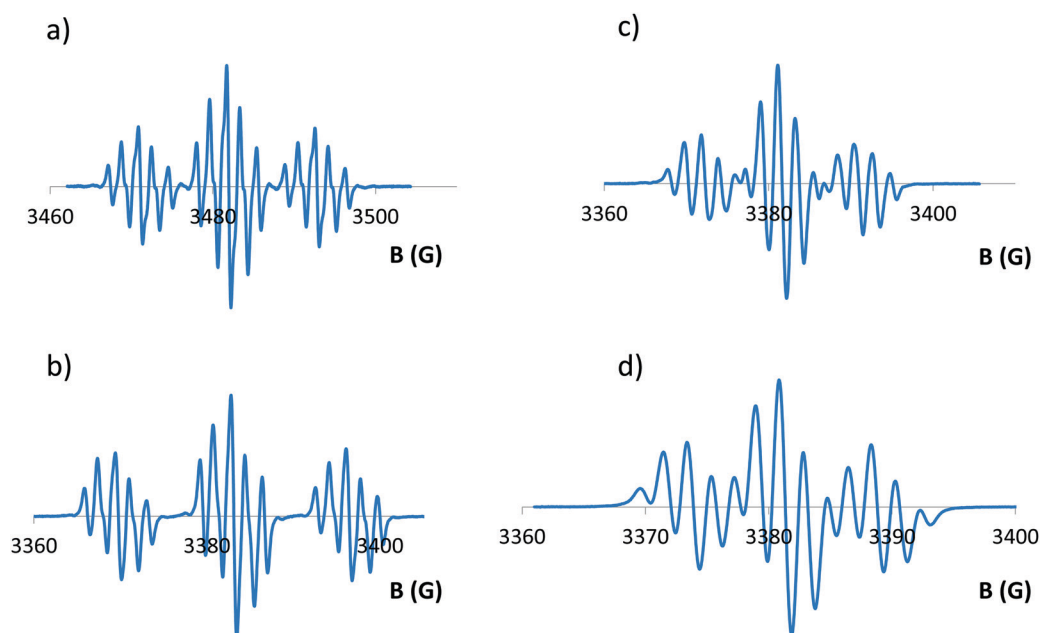


Fig. 6 EPR spectra of the chemically generated **4a**^{••} (a), **4b**^{••} (b), **6**^{••} (c) and **8**^{••} (d) in DCM.



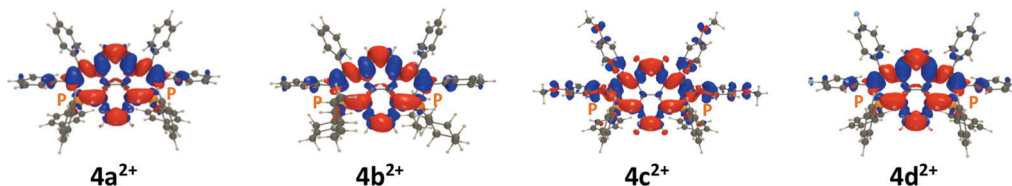


Fig. 7 Electron density difference determined for $4a-d^{2+}$. The blue (red) lobes correspond to a decrease (increase) of density upon photon absorption, respectively. A contour threshold of 8×10^{-4} a.u. was used.

electron with the two P and the different sets of H atoms. Consequently, the modifications of exocyclic substituents in $4b^{+\bullet}$ and $4d^{+\bullet}$ do not affect the hyperfine structure compared to $4a^{+\bullet}$ (Fig. S32, ESI†). The computed spin densities of all radical cations also support these findings (*vide infra*, Fig. 8).

In conclusion, this spectroelectrochemical and EPR study allows generalizing to the entire family of compounds the “viologen-like” behaviour previously observed on $4a[OTf]_2$.⁸

Theoretical calculations

To gain more insights into the properties of the investigated systems, we have carried out DFT and TD-DFT calculations using a protocol described in the ESI.† The TD-DFT calculations return bright S_0-S_1 transitions with a clear $\pi-\pi^*$ character centered on the core of the dyes. The Electron Density Difference (EDD) plots are displayed for all compounds in Fig. 7. One notices that the patterns are very similar for all compounds differing by their exoskeletal substituents that play not a major role in the transition, with the exception of $4c^{2+}$ in which the donor methoxy group loses density upon transition, which is consistent with a slight CT character from the periphery to the core and hence a redshift of the absorption (see Fig. S36, ESI,† for the representation of the frontier orbitals of the four compounds). Consistent with the experimental trend (Fig. 3), the oscillator strength determined by TD-DFT is also significantly smaller in $4c^{2+}$ (0.40) than in $4a,b,d^{2+}$ (0.60–0.66). Regarding the various endoskeletal structures, one notices similar delocalization along all the π -core (Fig. S34, ESI†).⁸

The vertical transition energies determined with TD-DFT using a refined solvation model correspond to absorption wavelengths of 394, 389, 425, 392, 374, 376, 370, and 487 nm, for $4a^{2+}$, $4b^{2+}$, $4c^{2+}$, $4d^{2+}$, 5^{2+} , 6^{2+} , 7^{2+} , and 8^{2+} , respectively. In the exoskeletal series, one clearly notices the significant redshift of $4c^{2+}$, again consistent with the measurements, whereas in the endoskeletal series, the strong bathochromic shift of 8^{2+} is noticeable. We have also determined the vertical emission wavelengths: 599, 513, 722, 610, 622, 450, 462, and 763 nm, for $4a^{2+}$, $4b^{2+}$, $4c^{2+}$, $4d^{2+}$, 5^{2+} , 6^{2+} , 7^{2+} , and 8^{2+} , respectively. Although TD-DFT exaggerates the difference between the systems, the experimental ranking of the various compounds is reasonably reproduced.

The spin density plots are displayed in Fig. 8. As can be seen, the radical is highly delocalized on the π -conjugated core, in all cases, with no significant contributions from the exoskeletal substituents. Therefore, in agreement with the experimental

results, $4a^{+\bullet}$ and $4b^{+\bullet}$ possess similar spin densities. Interestingly, the two central carbon atoms in $4a^{+\bullet}$ show the opposite spin excess, whereas the radical is only delocalized on the rim of the dye in $6^{+\bullet}$, a pattern not found in $8^{+\bullet}$, confirming the specificities of this particular core.

Electrofluorochromism

P-Doped PAHs $4-8^{2+}$ display “viologen-like” redox behavior as illustrated by CV, spectroelectrochemistry and EPR. However, in contrast to most of the viologen derivatives, which fluoresce under very specific conditions only or upon complex molecular engineering,¹⁸ they are intrinsically fluorescent. In order to take advantage of this specific property, the electrochemical modulation of the luminescence of $4a^{2+}$ and 8^{2+} has been investigated. These two derivatives have been chosen because they provide satisfactory electrochemical reversibility and they emit in different regions of the visible spectrum. Compounds with exoskeletal modifications were not tested as we showed that the effect of electron-withdrawing/electron-rich groups is either too weak to induce significant changes or detrimental for the stability of the reduced species. Emission spectroelectrochemistry and simultaneous variations of fluorescence intensity and redox current have been recorded under potential control for

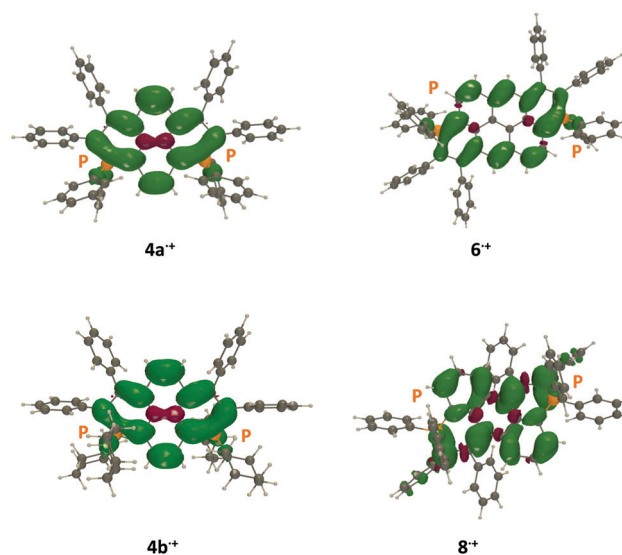


Fig. 8 Representation of the spin density difference in the radical cation of $4a^{+\bullet}$, $4b^{+\bullet}$, $6^{+\bullet}$ and $8^{+\bullet}$ (at their minimal radical cation geometry). The contour threshold is 1×10^{-3} a.u.



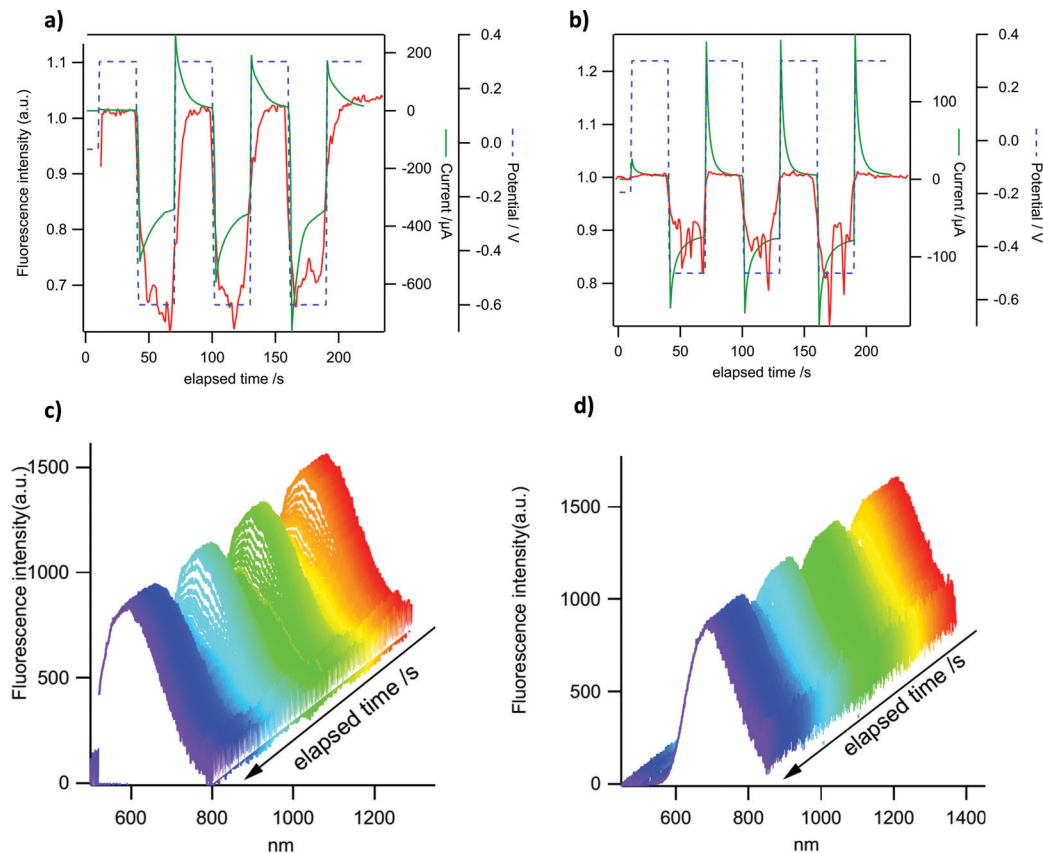


Fig. 9 Top (a and b): modulation of fluorescence intensity (red trace) and electrochemical current (green trace) under potential steps (dashed blue trace) for $4a^{2+}$ (a) and 8^{2+} (b). Bottom (c and d): corresponding variations of emission spectra of $4a^{2+}$ (c) and 8^{2+} (d) with time for the same potential signal as above. Excitation wavelength: 470 nm. The emission is integrated from 520 to 800 nm to give the intensity recorded in (a) and from 600 to 820 nm for the intensity in (b). Concentration of $4a^{2+}$ and 8^{2+} : 1 mM.

$4a^{2+}$ and 8^{2+} in DCM solutions. The ITO substrate is polarized successively at +0.3 V (dicationic state) and -0.6 V for $4a^{2+}$ or -0.5 V for 8^{2+} (radical cation state in each case). The results are shown in Fig. 9a and b. A clear modulation of fluorescence intensity can be seen for both compounds that are synchronous with the electrochemical current variations. One can observe a fluorescence intensity drop upon reduction and a full restoration upon reoxidation to the initial state. These results are consistent with the absorption spectroelectrochemical results in Fig. 5 where a clear increase of absorption can be seen in the wavelength range of emission upon reduction. In the case where the reduced form would be emissive, this impedes observing any emission in this spectral range. The fluorescence intensity modulation remains perfectly stable in three successive cycles for both compounds, although a slight drift can be observed for the fluorescence intensity of $4a^{2+}$. The relative fluorescence intensity drop is larger in $4a^{2+}$ (ca. 30%) than in 8^{2+} (ca. 15%), as a result of the higher electrochemical current for the former compared to the latter. In the potential range explored, the electrochemical conversion is more complete for $4a^{2+}$ leading to a higher emission contrast. This phenomenon can also be seen in the variations of emission spectra with time for the same potential modulation (Fig. 9c and d). This allows us to check that the intensity recorded actually corresponds to

the emission of both compounds (see Fig. 3 for comparison of the emission spectra in the absence of an electrochemical signal). The fluorescence intensity collected in Fig. 9a and b corresponds to the integration of the whole emission spectrum in Fig. 9c and d.

To check that the fluorescence intensity drop can be fully controlled by the applied potential, another potential signal was applied on 8^{2+} (Fig. 10). To enhance the sensitivity, the fluorescence is measured in a microvolume delimited by the substrate and a microelectrode tip (25 μ m) positioned 60 μ m above the ITO substrate. The latter is polarized at more and more negative potentials at each cycle. The results indicate that the fluorescence starts to drop at -0.3 V for the reduction of 8^{2+} (Fig. 10a) exactly when the electrochemical current starts to be significant. The modulation rapidly increases as the potential limit is pushed toward more negative values (from -0.3 to -0.5 V). From -0.5 V to -0.6 V the modulation remains nearly constant because at these potentials the electrochemical conversion remains fairly constant as well¹⁹ and this is confirmed by the amplitude of the electrochemical current. At -0.7 V, the current amplitude starts to increase again due to the start of the second reduction process but the fluorescence intensity does not change much. This gradual increase of the fluorescence intensity modulation can also be seen in Fig. 10b where



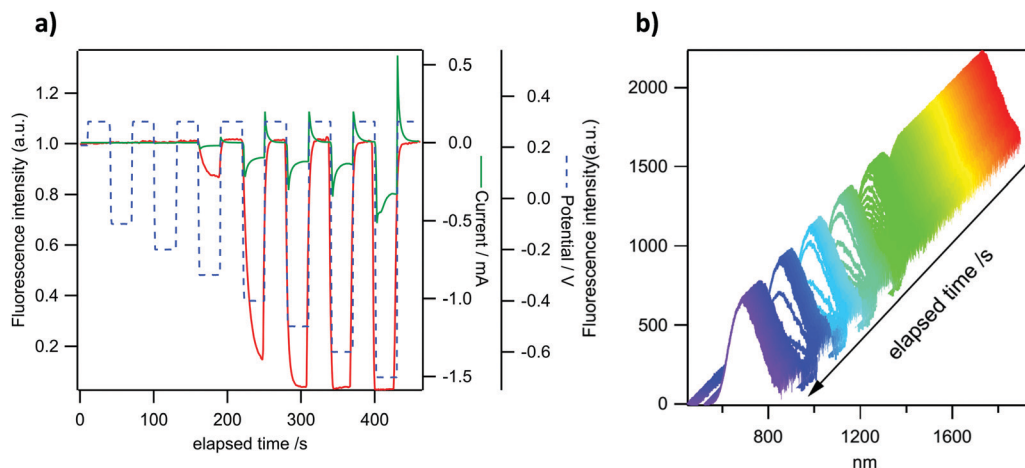


Fig. 10 Modulation of fluorescence intensity (red trace) and electrochemical current (green trace) of compounds 8^{2+} (a) for potential steps of increasing amplitude (dashed blue trace). (b) Corresponding variation of emission spectra with time of compound 8^{2+} for the same potential signal as in (a).

deeper and deeper variations of the intensity can be seen but without modification of the shape and location of the emission spectra. In conclusion, a nearly full contrast between the ‘on’ and ‘off’ states can be obtained at a very moderately negative potential, which endows 8^{2+} with very good electrofluorochromic properties if compared with already well known compounds in this field like *e.g.* tetrazines.²⁰ When considering their emission in the red part of the spectrum, and their initial charged state, they can also be advantageously compared to ruthenium complexes²¹ by offering similar properties but with a reductive quenching. These experiments illustrate that the luminescence of these P-doped PAHs can be electrochemically modulated in the entire visible range, extending the scope of molecules exhibiting efficient luminochromism.²²

In conclusion, synthetic access to three novel dicationic P-doped PAHs and a reference compound is described allowing us to have in hand an entire family of 8 derivatives. Exo-skeletal modifications allow the fine-tuning of their optical and redox properties. 4-8[OTf]₂ are electron acceptors with “viologen-like” redox behaviour. However, in contrast to most of the viologen derivatives, 4-8[OTf]₂ are fluorescent both in solution and in the solid-state. Combined experimental (spectroelectrochemical and EPR) and theoretical studies allowed the more in depth investigation of the reduced species and in particular the stable radical cations formed. This extensive study thus allows generalizing the behaviour previously observed on 4a[OTf]₂ and affords a guideline for the property tuning of such derivatives. Finally, the fluorescence properties of 4[OTf]₂ and 8[OTf]₂ were electrochemically modulated. These first results of electrofluorochromism illustrate the potential of these derivatives to be used as novel luminescent switches or in other relevant applications for multi-stage redox systems such as photoredox catalysis,²³ artificial photosynthesis,²⁴ spintronics²⁵ or energy storage.²⁶

Conflicts of interest

There are no conflicts to declare.

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