

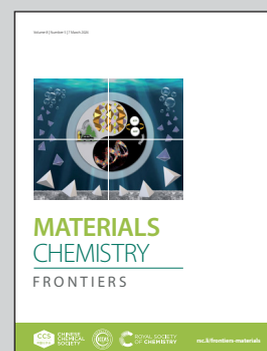


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Influence of the pendant substituent at the C1 position
 of a spirobifluorene scaffold on the electronic properties

The present work reports the synthesis and the study
 of a series of organic semi-conductors constructed on
 an emerging molecular fragment, namely C1-substituted
 Spirobifluorene.

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Influence of the pendant substituent at the C1 position of a spirobifluorene scaffold on the electronic properties†

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1-Substituted fluorenes have emerged in recent years as being among the most efficient functional materials for phosphorescence and thermally activated delayed fluorescence OLEDs. The position C1 of the fluorene is indeed highly interesting to design functional materials with specific properties. This scaffold displays two main characteristics. First, there is a strong π -conjugation disruption between the pendant substituent attached at C1 and the substituted fluorene. Second, the specific geometry of the C1-fluorene allows through-space interactions between the substituent attached at C1 and the cofacial building unit attached at C9. However, to date, this family of organic-semi-conductors has not been intensively studied yet. In this work, we report the synthesis and the structural, electrochemical and photophysical properties of four 1-substituted spirobifluorene derivatives, constructed with widely known extended π -conjugated systems: naphthalene, anthracene, phenanthrene and pyrene at the C1 position of spirobifluorene. Using a structure–property relationship approach, we show how the nature of the substituent grafted at C1 significantly modifies the electronic properties (HOMO/LUMO energy levels, triplet state energy levels, etc.). As 1-substituted spirobifluorenes are undoubtedly very promising organic semi-conductors, such a study provides fundamental knowledge to construct future efficient organic materials for specific applications.

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Introduction

9,9'-Spirobifluorene (SBF) is one of the most important building blocks used in the synthesis of organic semi-conductors (OSCs) for organic electronics (OEs).^{1–5} The most studied position of substitution is the C2 position, known to significantly increase the conjugation length between the fluorene and its substituent.^{1,2,6} However, in recent years, manipulating the substitution pattern of SBF (positions of substitution C1, C3 or C4) has appeared as a very efficient tool to further tune the electronic properties of SBF-based materials.¹ The particularity of the substitution at C1,^{7–9} C3,^{8,10–16} or C4^{8,17–21} is the restriction of the electronic coupling between the fluorene and the attached substituent. This key characteristic

has been advantageously used to design high triplet energy materials ($E_T > 2.7$ eV), which are particularly attractive for hosting phosphors in phosphorescence OLEDs (PhOLEDs).^{22–24} Poly-substituted organic materials with specific properties have also been investigated, taking advantage from the different SBF positions (C1^{25–28} and C3^{29–31}).

In a C4-substituted SBF, constructed on a twisted *ortho* linkage, the π -conjugation restriction is induced by strong steric hindrance between the fluorene and its substituent.^{1,17} In a C3-substituted SBF, the π -conjugation restriction is caused by the electronic decoupling induced by the presence of a *meta* linkage. The C1 position combines these two features, *meta* linkage and strong steric hindrance, between the fluorene and the substituent. Thanks to the strong electronic decoupling induced by these two features, a C1-substituted SBF scaffold has appeared as the most efficient regioisomer to construct high E_T materials, recently providing important advances in the field of PhOLEDs with the most efficient blue and white PhOLEDs ever reported in the literature.²³ The substitution at the C1-position of SBF displays another interesting characteristic that is the intramolecular interaction occurring between the pendant substituent at C1 and the cofacial fluorene.

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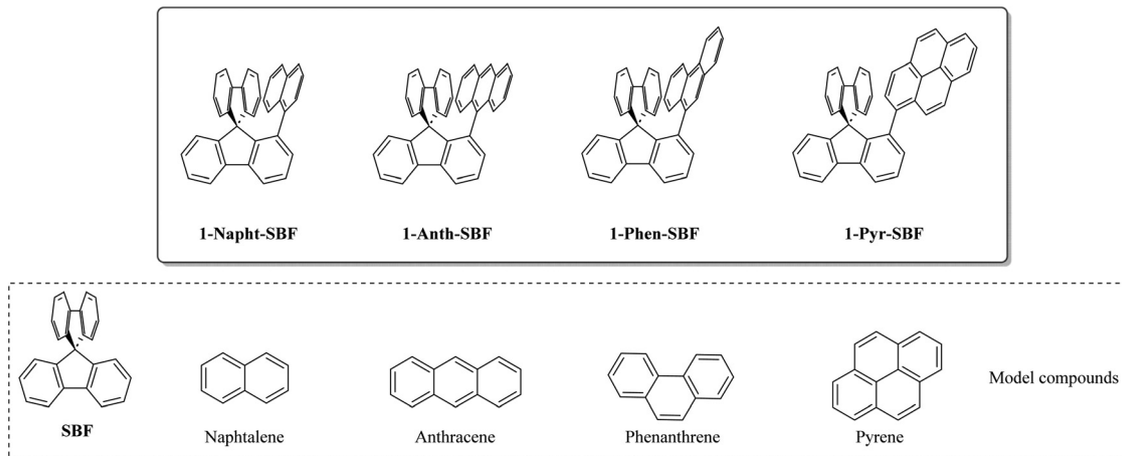


Chart 1 Molecular structures of **1-Napht-SBF**, **1-Phen-SBF**, **1-Anth-SBF** and **1-Pyr-SBF** and constituting building blocks (spirobifluorene, naphthalene, phenanthrene, anthracene and pyrene).

This characteristic has been efficiently used in thermally activated delayed fluorescence (TADF) based OLEDs, providing cofacial donor/acceptor systems.^{32–36}

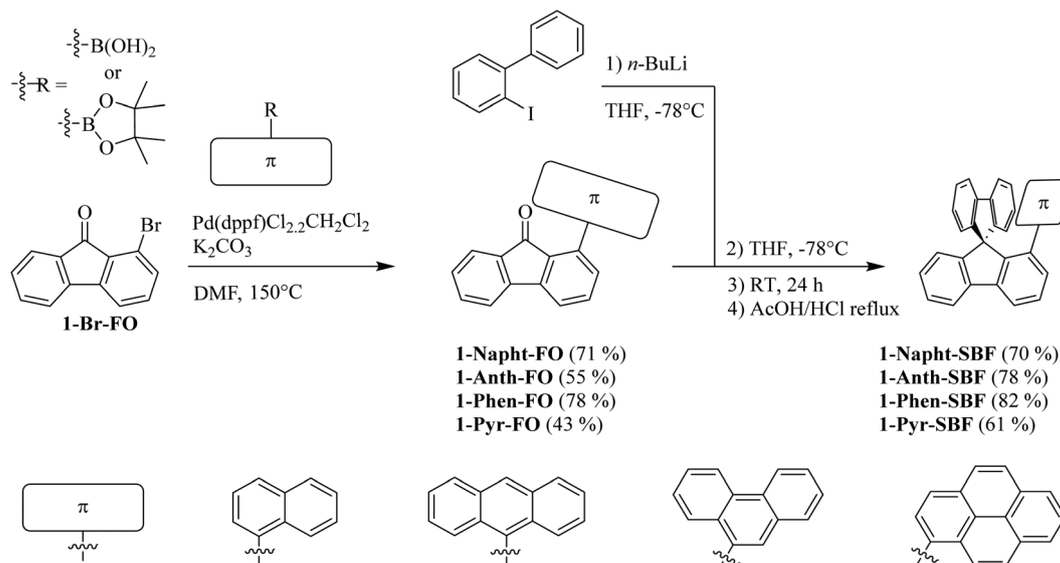
Thus, the substitution at C1 has many advantages, which have allowed designing high efficiency materials for electronics. However, as almost all the C1-based materials reported to date were designed to host phosphors in PhOLEDs, only small molecular units with short π -conjugation pathways were attached such as phenyl,⁷ biphenyl,²³ carbazole,⁷ and *m*-terphenyl.²³ Thus, in order to increase the molecular diversity of the emerging family of 1-substituted SBFs and to well apprehend their electronic properties, we wish to report herein a series of 1-substituted SBFs incorporating various π -conjugated molecular fragments possessing different conjugation lengths, namely naphthyl, phenanthrenyl, anthracenyl or pyrenyl. The present work reports a detailed structure–property relationship study of 1-naphthyl-, 1-phenanthrenyl-, 1-anthracenyl- and

1-pyrenyl-spirobifluorenes (**1-Napht-SBF**, **1-Phen-SBF**, **1-Anth-SBF** and **1-Pyr-SBF**, respectively, Chart 1) with their constituting building blocks (naphthalene, phenanthrene, anthracene and pyrene) and the structurally related analogue spirobifluorene (SBF). We show the importance of the substituent in the modulation of the electronic properties (HOMO/LUMO energy levels, optical transitions, triplet state energy E_T , charge transport, *etc.*). This feature provides an interesting degree of tuning for this emerging family of OSCs.

Results and discussion

Synthesis

Since 1-bromo-fluorenone (**1-Br-FO**) can be obtained at the multi-gram scale,⁷ the synthesis of **1-Napht-SBF**, **1-Phen-SBF**, **1-Anth-SBF** and **1-Pyr-SBF** is straightforward with high yield (Scheme 1).



Scheme 1 Synthesis of **1-Napht-SBF**, **1-Phen-SBF**, **1-Anth-SBF** and **1-Pyr-SBF**.



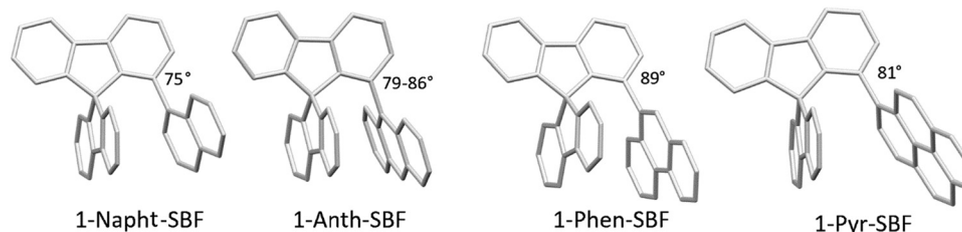


Fig. 1 Molecular structures from X-ray diffraction of **1-Napht-SBF**, **1-Anth-SBF**, **1-Phen-SBF** and **1-Pyr-SBF** and the angles between the substituted phenyl ring and the phenyl of the substituent.

1-Br-FO was obtained as previously reported using a one pot efficient synthesis method reported by Sorensen and coworkers.³⁷ The aryl fragments were then attached *via* Suzuki-Miyaura cross-coupling to give the corresponding 1-aryl-fluorenones with yields varying from 43 to 78%. The synthesis of the targeted 1-aryl-spirobifluorenes was then carried out using a classical two-step procedure: the lithium-iodine exchange of 2-iodobiphenyl with *n*-butyllithium at low temperatures, followed by addition of **1-Br-**

FO affording the corresponding fluoreno (not isolated), which was further involved in an intramolecular cyclisation reaction (AcOH/HCl) to provide the corresponding 1-aryl-spirobifluorenes with overall yields in the range of 61 to 82% from 1-aryl-fluorenones.

Structural properties

The structural arrangement of the four derivatives obtained by X-ray diffraction are depicted in Fig. 1. The most important

	d_{C-C} (Å)	d_1 (Å)	θ_1 (°)	d_2 (Å)	θ_2 (°)
1-Napht-SBF Ph/Ph	3.90	1.61	24.4	1.41	21.21
1-Napht-SBF Cp/Ph	3.68	0.89	14.05	2.18	36.38
1-Phen-SBF Ph/Ph	3.63	1.54	25.23	1.18	18.97
1-Phen-SBF Ph/Ph	3.78	1.03	15.81	1.71	26.97
1-Phen-SBF Cp/Ph	3.61	1.13	18.27	1.78	29.63
1-Anth-SBF (Mol1- d_{C-C} 1) Ph/Ph	3.77	1.51	23.64	1.10	16.92
1-Anth-SBF (Mol1- d_{C-C} 2) Ph/Ph	3.86	0.47	11.08	1.88	29.25
1-Anth-SBF (Mol1- d_{C-C} 3) Cp/Ph	3.78	2.18	35.26	1.00	15.35
1-Anth-SBF (Mol2- d_{C-C} 1) Ph/Ph	3.74	1.65	26.09	0.9	13.92
1-Anth-SBF (Mol2- d_{C-C} 2) Ph/Ph	3.78	0.82	12.16	1.65	25.92
1-Anth-SBF (Mol2- d_{C-C} 3) Cp/Ph	3.70	1.16	18.25	2.09	34.5
1-Anth-SBF (Mol3- d_{C-C} 1) Ph/Ph	3.58	0.747	12.06	1.31	21.46
1-Anth-SBF (Mol3- d_{C-C} 2) Ph/Ph	3.88	0.57	8.44	1.79	27.38
1-Anth-SBF (Mol3- d_{C-C} 3) Cp/Ph	3.70	2.11	34.68	1.11	17.43
1-Anth-SBF (Mol4- d_{C-C} 1) Ph/Ph	3.76	0.56	8.36	1.73	27.36
1-Anth-SBF (Mol4- d_{C-C} 2) Ph/Ph	3.75	0.62	9.55	1.304	20.35
1-Anth-SBF (Mol4- d_{C-C} 3) Cp/Ph	3.74	2.10	34.09	0.88	13.67
1-Pyr-SBF Cp/Ph	3.76	2.18	35.46	0.49	7.59
1-Pyr-SBF Ph/Ph	No $d_{C-C} < 4$ Å				

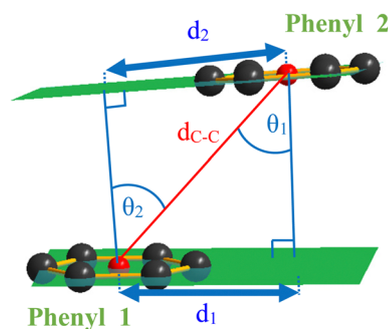


Fig. 2 Left: selected crystallographic data of **1-Napht-SBF**, **1-Anth-SBF**, **1-Phen-SBF** and **1-Pyr-SBF**. Right: reprinted (adapted) with permission from *J. Phys. Chem. C* 2019, 123, 19094–19104. Copyright 2023 American Chemical Society.



structural feature is the relative position of the pendant substituent with respect to the phenyl ring of the substituted

fluorene. For the four molecules, the angle between the substituted phenyl ring and the grafted phenyl of the substituent is

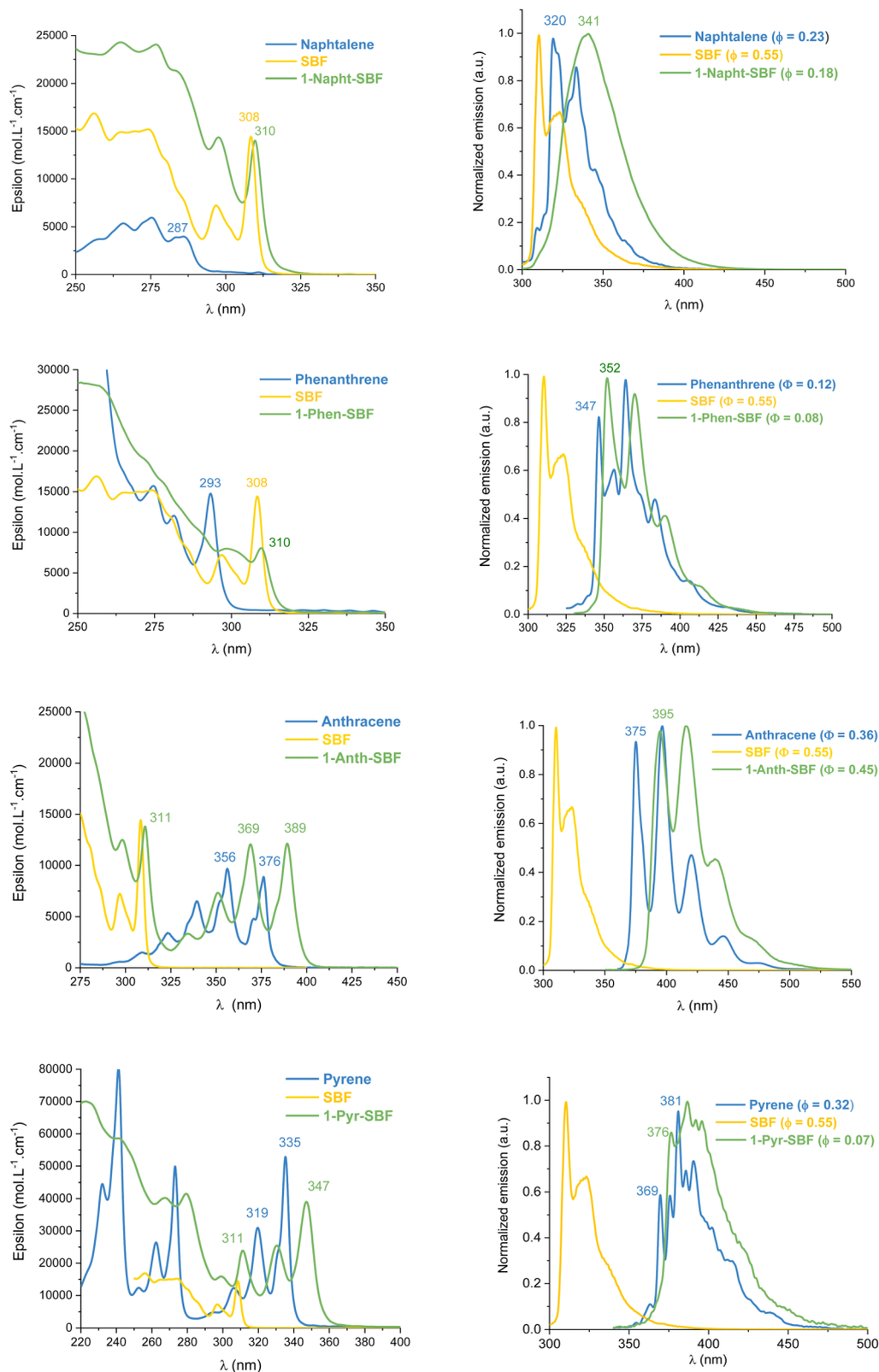


Fig. 3 Absorption (left) and emission (right) at room temperature in cyclohexane of 1-aryl-SBFs (green), SBF (yellow) and the substituents (blue).



Table 1 Selected electronic data for 1-Phen-SBF, 1-Naph-SBF, 1-Pyr-SBF and 1-Anth-SBF and their corresponding building units

	1-Naph-SBF	Naphthalene	1-Phen-SBF	Phenanthrene	1-Pyr-SBF	Pyrene	1-Anth-SBF	Anthracene	SBF ^{a,5}
λ_{abs}^a [nm]	265 (2.4)	257 (0.37)	298 (0.79)	274 (1.5)	221 (7.0)	232 (4.4)	298 (1.24)	309 (0.15)	297
λ_{em}^a [nm]	277 (2.4)	265 (0.53)	310 (0.8)	281 (1.2)	243 (5.8)	241 (8.0)	311 (1.38)	324 (0.34)	(0.72) 308
λ_{abs}^a ($\times 10^3$ L mol ⁻¹ cm ⁻¹)									(1.45)
283 (2.1)	275 (0.59)	275 (0.59)	293 (1.47)	267 (4.0)	252 (1.24)	335 (0.33)	335 (0.33)	339 (0.66)	
298 (1.4)	283 (0.39)	283 (0.39)	279 (4.1)	279 (4.1)	262 (2.62)	351 (0.73)	351 (0.73)	352 (0.35)	
310 (1.4)	286 (0.39)	286 (0.39)	299 (1.5)	299 (1.5)	273 (4.98)	369 (1.2)	369 (1.2)	356 (0.97)	
			311 (2.3)	311 (2.3)	294 (0.47)	389 (1.2)	389 (1.2)	371 (0.48)	
			331 (2.5)	331 (2.5)	306 (1.21)	—	—	376 (0.88)	
			347 (3.9)	347 (3.9)	319 (3.09)	—	—	—	
			335 (5.3)	335 (5.3)	370, 376, 381, 386, 390, 395, 416, 439, 397, 413, 427, 439, 468 467	—	—	375, 396, 420, 446, 477	310, 323
θ^b	0.18	0.23	0.08	0.12	0.07	0.32	0.45	0.36	0.55
τ_s [ns] ^c	4.8	16.4	15	14.3	1.9	17.2	4.8	4.2	4.6
k_f [$\times 10^7$ s ⁻¹]	3.8	14	0.5	0.8	3.7	1.9	9.4	8.6	12
k_{nr} [$\times 10^7$ s ⁻¹]	17	47	6.1	6.1	49	4.0	11	15	10
E_s [eV] ^d	3.94	4.03	3.58	3.63	3.34	3.52	3.23	3.36	4.07
E_T [eV] exp ^e	2.61	2.65	2.64	2.71	2.07	2.29	—	—	2.89
E_T [eV] calc ^f	2.31	2.36	2.38	2.45	1.80	1.83	—	—	—
τ_T [s] ^g	2.7	2.8	3.6	3.4	0.6	—	—	—	5.4
Ox potentials [V vs. SCE] ^g	1.58, 1.77(sh), 1.92	1.75, 1.99 (sh)	1.55, 1.88 (sh), 2.02	1.69, 2.1	1.22, 1.82	1.22(sh), 1.29, 1.44, >2.0	1.12, 1.82, 2.17	1.23, 1.92	1.67, 1.86
HOMO [eV]	-5.90	-5.94	-5.85	-5.89	-5.52	-5.54	-5.43	-5.46	-5.95
LUMO [eV]	-5.91	-6.14	-5.88	-6.08	-5.52	-5.67	-5.40	-5.57	-5.99
E_I^h	-1.97	-1.87	-2.00	-1.95	-2.29	-2.31	-2.36	-2.42	-1.74
E_I^i	-1.36	-1.41	-1.41	-1.42	-1.81	-1.87	-1.95	-2.04	-1.26
ΔE [eV]									
Opt ⁱ	3.94	4.25	3.93	4.16	3.47	3.64	3.10	3.26	3.97
E_I^k	3.93	4.07	3.85	3.94	3.23	3.23	3.07	3.04	4.21
Cal ^f	4.55	4.73	4.47	4.66	3.71	3.80	3.45	3.53	4.73
μ_h (cm ² V ⁻¹ s ⁻¹)	1.1×10^{-6}	—	3.5×10^{-6}	—	0.15×10^{-6}	—	3.2×10^{-6}	—	—

^a In cyclohexane. ^b Quantum yield θ measured with quinine sulphate as the reference. ^c $\lambda_{\text{exc}} = 310$ nm. ^d From the onset of fluorescence spectra in cyclohexane. ^e From the pic maximum in 2-MeTHF at 77 K. ^f From theoretical calculations (TD-DFT, b3lyp, 6-311+g(d,p)). ^g From CV in Bu₄NPF₆ 0.2 M in DCM for oxidation. ^h From CV in Bu₄NPF₆ 0.1 M in DMF. ⁱ From the onset of the absorption spectrum in cyclohexane.

very high: 75° for **1-Napht-SBF**, 79° and 86° for **1-Anth-SBF** (four molecules are indeed present in the asymmetric unit), 81° for **1-Pyr-SBF** and 89° for **1-Phen-SBF**. This feature indicates a significant conjugation breaking between the fluorene and its substituent at C1. The model compound **1-Phenyl-SBF**, previously reported in the literature, displays an angle of 75° .⁸ Thus, from phenyl to naphthalene, the dihedral angle remains identical, showing that the steric hindrance is not increased when an additional phenyl ring is fused. However, the fusion of an additional phenyl ring in anthracene increases the dihedral angle (from 80° to 86°). The highest value is obtained for **1-Phen-SBF**, 89° , in which the fused phenyl rings are in *ortho* and *meta* positions of the 1-SBF linkage.

Due to the particular geometry of the 1-SBF scaffold, the molecular structures of the four compounds display many short intramolecular C–C distances between the two cofacial fragments (see the ESI†). These C–C distances are shorter than the sum of their van der Waals radii (3.4 \AA)³⁸ and translate a sterically hindered environment. In order to evaluate the strength of these interactions between the pendant substituent and the cofacial fluorene, three structural parameters have been evaluated for the cofacial phenyl rings, namely ring-

centroid/ring-centroid distance d_{C-C} , vertical displacements d_1 and d_2 and slippage angles θ_1 and θ_2 (Fig. 2, see the ESI† for methods). As discussed by Janiak and coworkers, these parameters can reflect the strength of the interactions between two phenyl rings.^{39–43} Indeed, ring-centroid/ring-centroid distances below 3.8 \AA , vertical displacements below 1.5 \AA , and ring slippage angles below 25° indicate strong interactions between two interacting phenyl rings. In the present case, the central cyclopentadienyl ring of the fluorene will also be considered.

All molecules display short ring-centroid/ring-centroid distances between two cofacial phenyl or cyclopentadienyl rings (below 3.9 \AA). Very small distances in the series are detected, notably for **1-Phen-SBF** and **1-Anth-SBF** with short d_{C-C} of 3.61 and 3.58 \AA , respectively. Many phenyl/phenyl and cyclopentadienyl/phenyl couples also display very short vertical displacements (below 1.5 \AA) and ring slippage angles (below 25°) satisfying the above-mentioned criteria. Three of the four molecules present in the asymmetric unit of **1-Anth-SBF** fully satisfy the three Janiak's criteria, indicating strong intramolecular interactions between the facing anthracene and fluorene units. Compounds **1-Phen-SBF** and **1-Napht-SBF** also present several phenyl/phenyl and cyclopentadienyl/phenyl

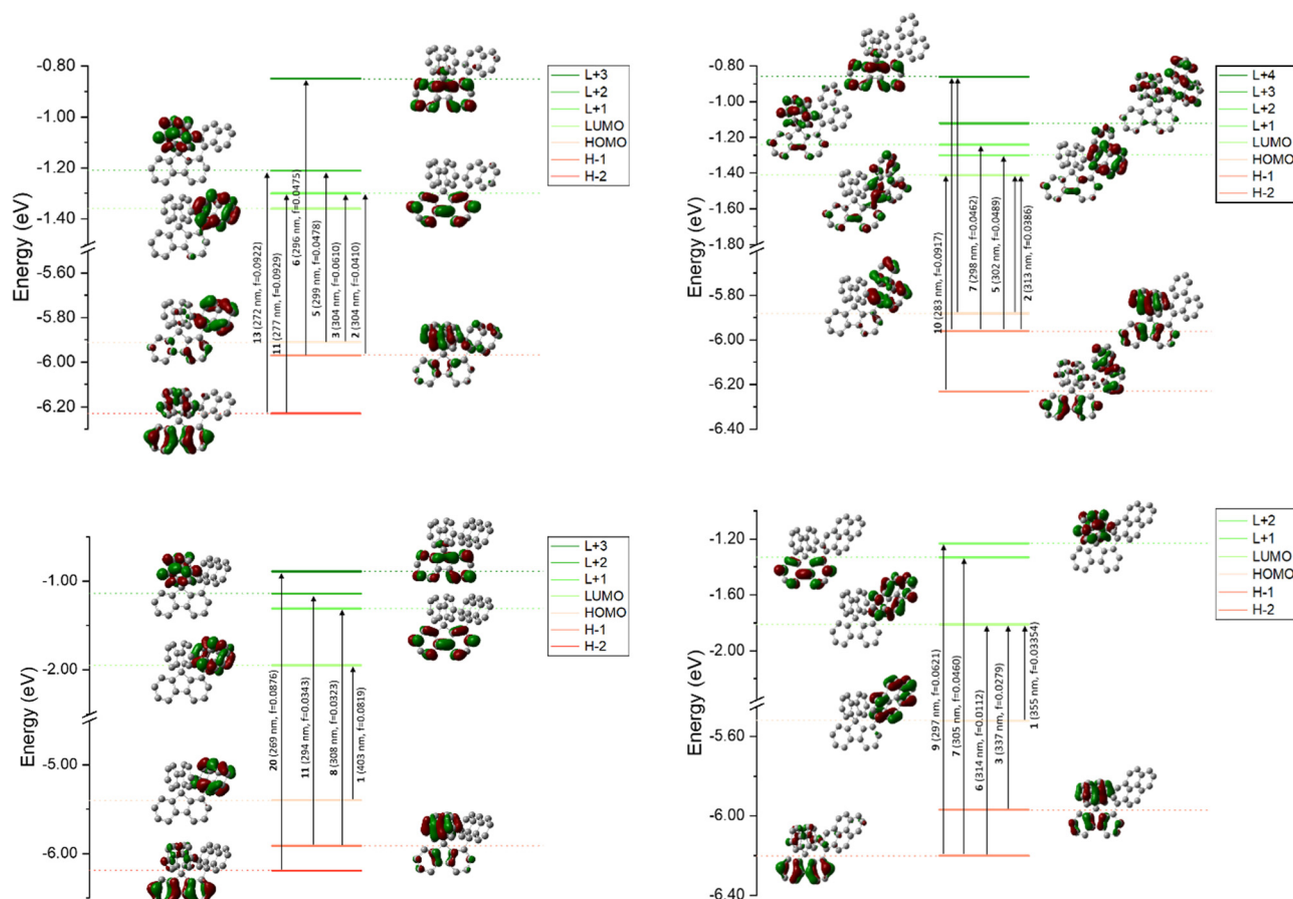


Fig. 4 Representation of the energy levels and the molecular orbitals involved in the main electronic transitions of **1-Napht-SBF** (top-left), **1-Phen-SBF** (top-right), **1-Anth-SBF** (bottom-left) and **1-Pyr-SBF** (bottom-right) as obtained by TD-DFT, B3LYP/6-311 + G(d,p) shown with an isovalue of 0.04 (e Bohr⁻³)^{1/2}.



interactions. These interactions partially satisfy all the above-mentioned criteria and translate therefore moderately to strong π - π interactions between the cofacial systems. In contrast, **1-Pyr-SBF** only displays one short d_{C-C} and the pyrene unit is therefore less influenced by the facing fluorene. This shows that the C1 grafting allows not only modulation of the fluorene-substituent direct conjugation, but also fine tuning of the interactions between the second SBF-included fluorene and the grafted substituent at C1. Especially, these data show that the substitution at C1 provides a sterically hindered environment which seems to govern the π - π interactions between the cofacial systems, and these interactions are modulated by the size of the grafted substituent.

Photophysical properties

UV-Vis absorption and emission spectra of the four compounds, **1-Napht-SBF**, **1-Anth-SBF**, **1-Phen-SBF** and **1-Pyr-SBF**, were recorded in dilute cyclohexane solution at room temperature (Fig. 3) in addition to their corresponding building units, either naphthalene for **1-Napht-SBF**, anthracene for **1-Anth-SBF**, phenanthrene for **1-Phen-SBF** and pyrene for **1-Pyr-SBF** and **SBF**. The photophysical properties are collected in Table 1 (and in Table S1, ESI[†] for their building units).

For the four 1-SBF derivatives, the absorption ranges from 250 to 400 nm (Fig. 3-left). First, it should be noted that unsubstituted **SBF** exhibits two thin absorption bands at *ca.* 297 and 308 nm (π - π^* transitions). Moreover, the aromatic substituents display absorption bands, which can be classified in three categories:⁴⁴ the first bands are weak ($\epsilon = 10^2$ - 10^3 mol L⁻¹ cm⁻¹) and may be hidden by the others (they can be seen above 300 nm for naphthalene), and the second bands are moderately intense ($\epsilon = 10^4$ mol L⁻¹ cm⁻¹) with a vibrational structure displaying peaks spaced by 1400 cm⁻¹ and the third bands are very intense ($\epsilon = 10^5$ mol L⁻¹ cm⁻¹) and unstructured. The bands corresponding to each fragment (both SBF and the substituents) can be found in the 1-substituted SBF. It should be noted that the bands corresponding to anthracene and pyrene are clearly red-shifted in **1-Anth-SBF** and **1-Pyr-SBF** in comparison to the model compounds (by 13 and 12 nm, respectively) while the other bands corresponding to the **SBF** seem relatively unchanged. There is indeed a slight red shift of 2 nm between the **SBF** building block and the corresponding band of the 1-substituted SBFs. This shows that the substituent is not completely isolated from the SBF backbone (through bonds and/or through-space interactions). As the dihedral angle between the C1-substituted fluorene and the substituent attached is very high in these molecules, π -conjugation extension by through-bond interactions should be very weak. However, as shown above, through-space interactions occur and these interactions might be responsible for the red shift observed in the absorption spectra. This hypothesis will be verified below by cyclic voltammetry and discussed throughout this work.

Theoretical calculations were performed using DFT and TD-DFT (Fig. 4). For both **1-Napht-SBF** and **1-Phen-SBF**, HOMO/HOMO-1 and LUMO/LUMO+1 levels are approximately <0.1 eV. The HOMO of **1-Napht-SBF** is spread out on a

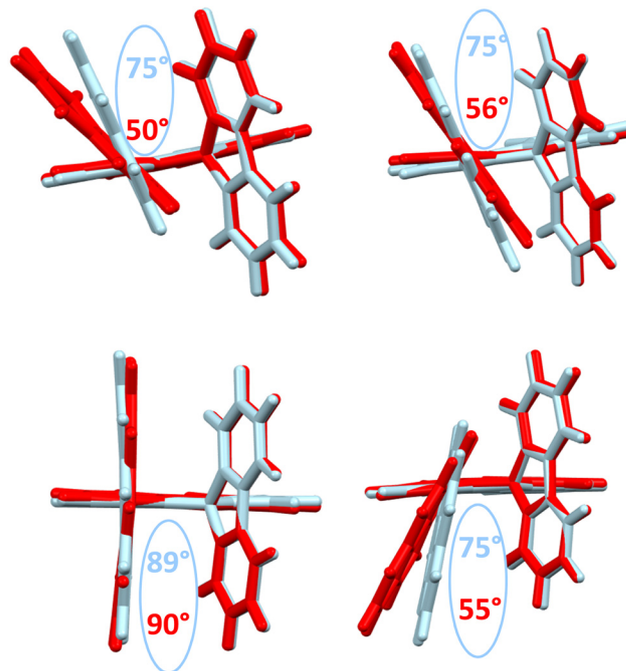


Fig. 5 Superposition of optimized geometries: S_0 (ground state, light blue) and S_1 (first singlet excited state, red). Top: **1-Napht-SBF** (left) and **1-Phen-SBF** (right) and bottom: **1-Anth-SBF** (left) and **1-Pyr-SBF** (right). On each structure, the modeled dihedral angle between the substituted fluorene and the substituent in the S_0 and S_1 states is presented.

naphthyl unit and, to a lesser extent, substituted fluorene. Interestingly, HOMO-1, which is very close in energy to the HOMO, is spread out on both the naphthyl unit and its cofacial fluorene, translating the potential cofacial interaction. For **1-Napht-SBF**, the TD-DFT calculations indicate that the band experimentally found at 310 nm is due to two transitions (corresponding to transitions from the ground to the second and the third excited states). In addition to the HOMO-1 \rightarrow LUMO+1 transition localized on the SBF core (found in **SBF**), the HOMO \rightarrow LUMO+1 transition occurs from the naphthyl unit to the substituted fluorene. By comparison, **1-Phenyl-SBF** displays a very intense HOMO/LUMO transition⁸ with both molecular orbitals centred on the fluorene, showing how the substituent can significantly modify the nature of the transitions.

With the phenanthrene attached, the HOMO/LUMO transition of **1-Phen-SBF** is detected with both the HOMO and the LUMO mainly spread out on the phenanthrene core. However, the LUMO of **1-Phen-SBF** displays more density on the fluorene than the LUMO of **1-Napht-SBF** whereas the opposite is observed for the HOMO. This is due to the different π -conjugation pathway observed in phenanthrene *vs.* naphthalene as phenanthrene can be seen as a naphthalene unit with an additional phenyl core fused in C3 and C4 positions. This results in the modification of different transitions in **1-Phen-SBF**.

Both the HOMO and the LUMO of **1-Anth-SBF** and **1-Pyr-SBF** are exclusively spread out on the pendant substituent with no participation of the fluorenyl units. The HOMO/LUMO transitions in these two compounds fully involve the anthracene or



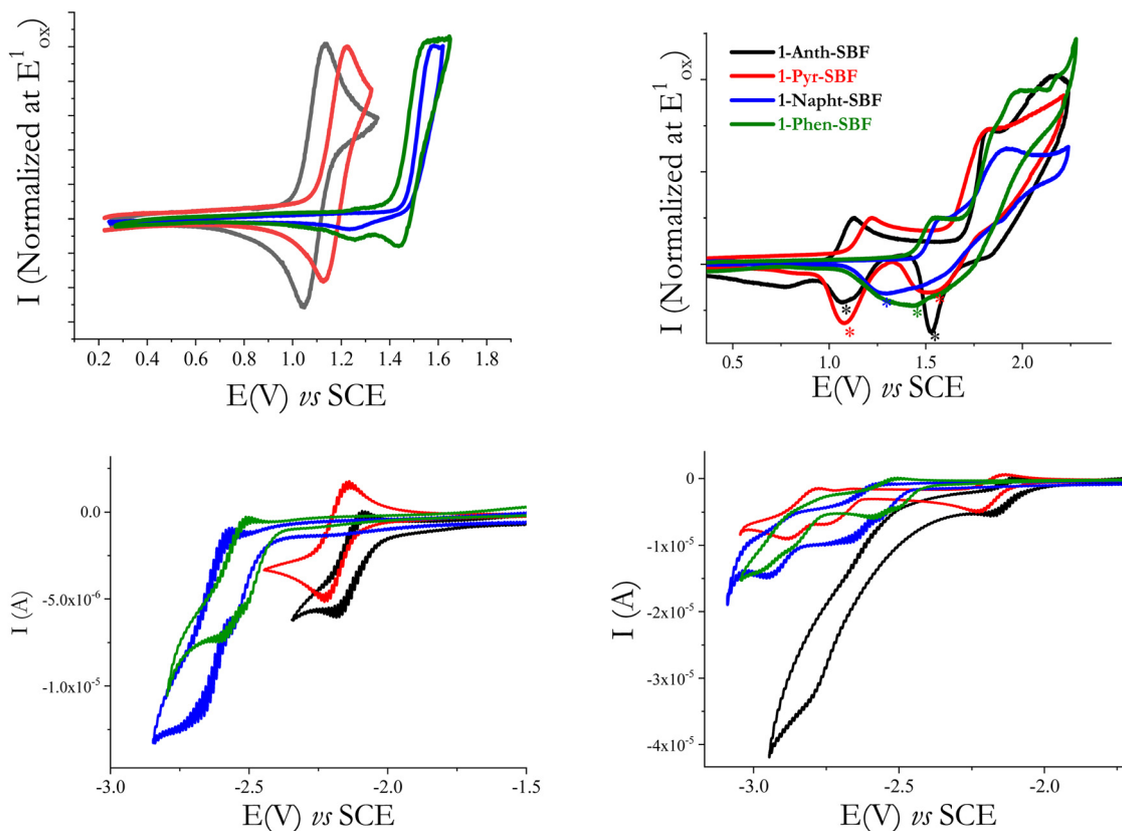


Fig. 6 Cyclic voltammograms of **1-Naphth-SBF** (blue line), **1-Phen-SBF** (green line), **1-Anth-SBF** (black line) and **1-Pyr-SBF** (red line): (top-left) up to their first oxidation wave or (top-right) between 0.5 and 2.3/2.4 V in CH_2Cl_2 0.2 M + Bu_4NPF_6 and (bottom-left) to their first reduction wave or (bottom-right) up to their highest reduction step in DMF 0.1 M + Bu_4NPF_6 sweep-rate: 100 mV s^{-1} and a platinum disk working electrode.

pyrene units (a high oscillator strength of 0.082 is noted for this transition in the case of **1-Anth-SBF**). Moreover, both the HOMO and LUMO energies are calculated with a large energy difference from their respective HOMO-1 and LUMO+1 orbital levels. This is a different behaviour from what was observed above for **1-Naphth-SBF** and **1-Phen-SBF**. It should be noted that the substitution of anthracene with SBF increases both the HOMO and LUMO levels, to a lesser extent for the LUMO, which induces a red shift for **1-Anth-SBF** compared to anthracene (15 nm according to the calculations in agreement with experimental data, 13 nm). Similar observations can explain the red shift of **1-Pyr-SBF** compared to that of pyrene (theoretical calculations: 15 nm vs. experimental data: 12 nm).

From the onset absorption wavelength, the optical energy gaps were calculated as 3.94 eV for **1-Naphth-SBF** and 3.93 eV for **1-Phen-SBF**; both are significantly lower than those of their corresponding building block (4.25 eV for naphthalene and 4.16 eV for phenanthrene) and both are slightly lower than that of **SBF** (3.97 eV). This indicates that (i) the SBF unit mainly drives the optical gap due to the short π -conjugation pathways of both naphthalene and phenanthrene and (ii) there is a small but not null impact of the substituent. The optical gaps of **1-Anth-SBF** (3.10 eV) and **1-Pyr-SBF** (3.47 eV) are significantly contracted and are both lower than that of **SBF**, due to the extended π -system of the substituent. These gaps are also

contracted compared to those of their building units anthracene (3.26 eV) and pyrene (3.64 eV). The optical gaps are then measured as follows: naphthalene > phenanthrene > pyrene > anthracene (4.25, 4.16, 3.64, and 3.26 eV, respectively) showing the different π -conjugation pathways in accordance with the above-stated conclusions. Hence, for all compounds, one can note that the C1-substituted compounds display a shorter optical gap than their constituting building units with however a different magnitude. Through-space interactions between the substituent and its cofacial fluorene are at the origin of this behaviour. Thus, the optical gap is driven by the most conjugated compound and tuned by the strength of the intramolecular interactions.

The emission spectra recorded in cyclohexane are presented in Fig. 3-right. Compared to the fluorescence of their related aryl units, 1-aryl-SBFs display emission spectra red shifted and roughly less defined. Superposition of optimized geometries of the ground state (S_0) and the first singlet excited state (S_1) indicates, except for **1-Anth-SBF**, a significant decrease (*ca* 20/25°) of the dihedral angle between the fluorene and its substituent (Fig. 5). Therefore, this planification provides an increase of the π -conjugation extension between the two fragments at the excited state. Thus, in fluorescence, both through-space and through-bond interactions seem to be involved. For **1-Anth-SBF**, the dihedral angle remains very high and, in this



case, through-space interaction seems to be involved in the red shift observed compared to anthracene fluorescence. We can finally note that the quantum yields of the substituents are not significantly affected, except in the case of pyrene, by their incorporation at C1 of the SBF.

Electrochemistry. The electrochemical properties of the four 1-aryl-SBF were studied in $\text{CH}_2\text{Cl}_2 + \text{Bu}_4\text{NPF}_6$ 0.2 M using cyclic voltammetry (CV) and compared to those of **1-Ph-SBF**,⁸ and **SBF**⁴⁵ (Fig. 6). As shown in Fig. 6 top, **1-Anth-SBF** is the easiest to oxidize in the series with a reversible wave possessing a maximum at $E_{\text{ox}}^1 = 1.12$ V, followed by **1-Pyr-SBF**, which also displays a reversible wave with $E_{\text{ox}}^1 = 1.22$ V. For these two compounds, the first electron transfer is located on the pendant substituent as shown above by the density of the HOMO (Fig. 4).

The first oxidation wave is reversible for **1-Anth-SBF** and **1-Pyr-SBF**, while it is not reversible for **1-Napht-SBF** and **1-Phen-SBF**. When oxidized at higher potential values (Fig. 6-top-right), the four compounds present other oxidation waves and are involved in electrodeposition processes which are indicated by the appearance of new redox systems (marked with a * in Fig. 6-top right), by their growth along recurrent sweeps and by the coverage of the electrode surface by an insoluble and electro-active deposit (see for each compound the detailed study of the

electrodeposition process in the ESI[†]). Such an electrodeposition process is classically observed with fluorene or SBF based compounds^{46–48} and more generally with bridged oligophenylenes (indenofluorene⁴⁹ or carbazole⁵⁰). The HOMO levels, determined from the onset of the first oxidation potential, decrease from $-5.43/-5.52/-5.85$ to -5.90 eV from **1-Anth-SBF/1-Pyr-SBF/1-Phen-SBF** to **1-Napht-SBF**. The same trend is observed by theoretical calculations from $-5.40/-5.52/-5.88$ to -5.91 eV from **1-Anth-SBF/1-Pyr-SBF/1-Phen-SBF** to **1-Napht-SBF** (Table 1). In order to check the exact influence of the pendant substituent, the HOMO energy levels of the constituting building blocks have also been measured and a similar trend was observed: -5.46 eV for anthracene, -5.54 eV for pyrene, -5.89 eV for phenanthrene and -5.94 eV for naphthalene. For the four compounds, the HOMO is slightly higher (by 0.02–0.04 eV) than the one of the corresponding substituents. This can be, in principle, assigned to two features, which have the same effect on the HOMO energy level. First, the electronic coupling between the fluorene and its substituent increases the conjugation length and in turn the HOMO energy. Second, the cofacial arrangement found in 1-aryl SBF induces through-space $\pi-\pi$ interactions between the fluorene and its facing pendant substituent, which are known to decrease the oxidation potential, increasing in turn the HOMO energy.^{51–53}

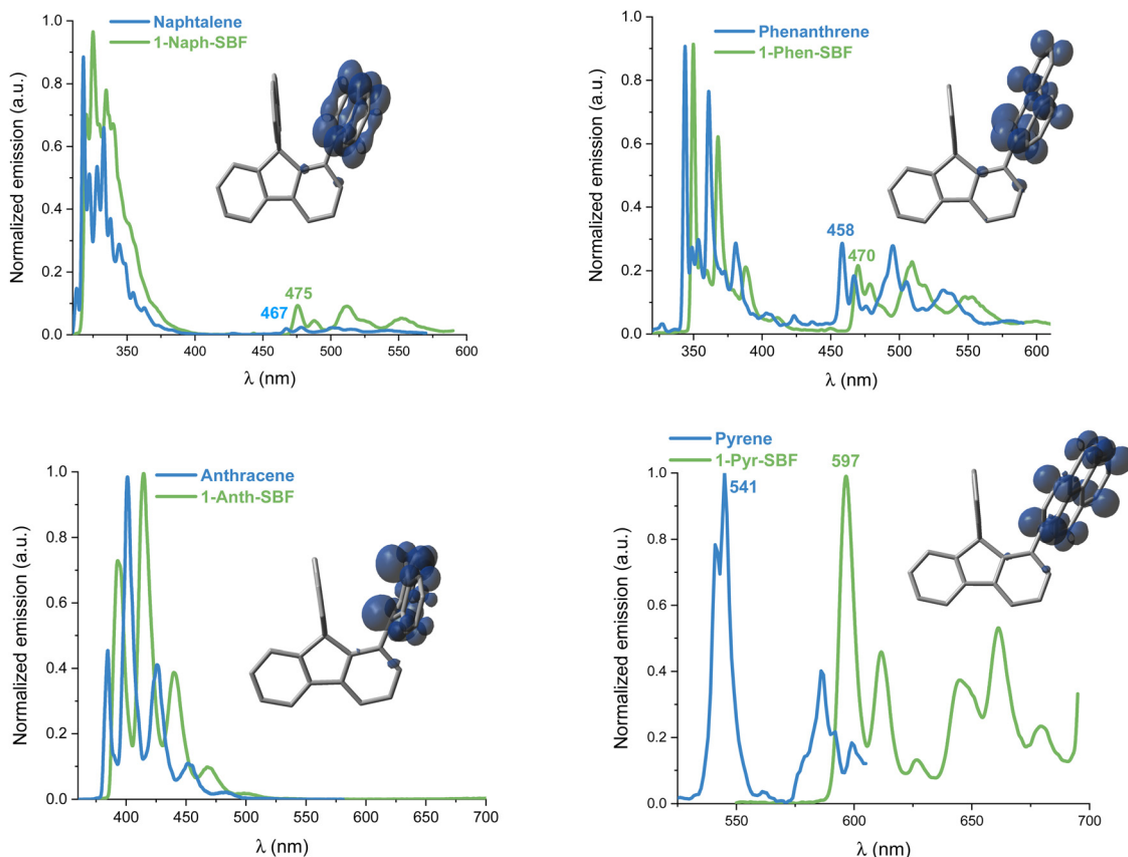


Fig. 7 Emission spectra of **1-Napht-SBF/Naphtalene** (top-left, $\lambda_{\text{ex}} = 300$ nm), **1-Phen-SBF/Phenanthrene** (top-right, $\lambda_{\text{ex}} = 310$ nm), **1-Anth-SBF/Anthracene** (bottom-left, $\lambda_{\text{ex}} = 350$ nm), and **1-Pyr-SBF/Pyrene** (bottom-right, $\lambda_{\text{ex}} = 310/340$ nm) at 77 K (2-Me-THF). Inset: spin density distribution (SDD) of the triplet of 1-aryl-SBF (isovalue = 0.004).



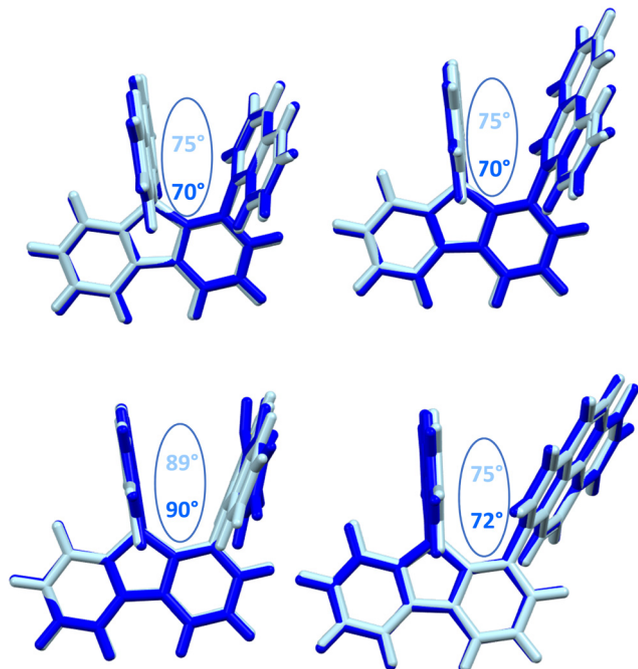


Fig. 8 Superposition of optimized geometries: S_0 (ground state, light blue) and T_1 (first triplet excited state) blue of **1-Napht-SBF** (top-left), **1-Phen-SBF** (top-right), **1-Anth-SBF** (bottom-left) and **1-Pyr-SBF** (bottom-right). On each structure, the modeled dihedral angle between the substituted fluorene and the substituent in the S_0 and T_1 states is presented.

As the extension of conjugation is weak at the C1 position due to the high dihedral angle (especially in the HOMO as previously reported^{8,22,45}), the cofacial arrangement and the resulting interactions are surely the main parameters involved in the increase of the HOMO energy. This was confirmed by plotting the electrostatic potential surfaces, which show an electron dense region between the face of the unsubstituted fluorene and the substituent (see the ESI[†]).

In reduction, recorded in DMF (see Fig. 6-Bottom), the first reduction wave of both **1-Napht-SBF** and **1-Phen-SBF** is irreversible and occurs at -2.72 V for **1-Napht-SBF** and -2.60 V for **1-Phen-SBF**. The two other C1-substituted SBFs, **1-Pyr-SBF** and **1-Anth-SBF**, are reduced in a first reversible process peaking at higher potential values, -2.22 and -2.18 V, respectively. The LUMO energy levels calculated from the onset reduction potentials decrease from -1.97 eV for **1-Napht-SBF** to -2.0 eV for **1-Phen-SBF**, -2.29 eV for **1-Pyr-SBF** and -2.36 eV for **1-Anth-SBF** following the same trend as their the respective aryl units ($-1.87/-1.95/-2.31$ and -2.42 eV for naphthalene/phenanthrene/pyrene and anthracene). The same conclusions as those for the HOMO can be then drawn. This is in accordance with the electron density of the LUMO, which is mainly distributed over the substituent. Thus, one can note that, compared to **SBF** (LUMO: -1.74 eV), the LUMO energy levels of the four C1-substituted SBFs are significantly more modified than their HOMO energy levels. Similar effects have been observed for C1-based dimers.⁴⁵ The electrochemical gaps calculated from the HOMO (in DCM) and the LUMO (in DMF) following the

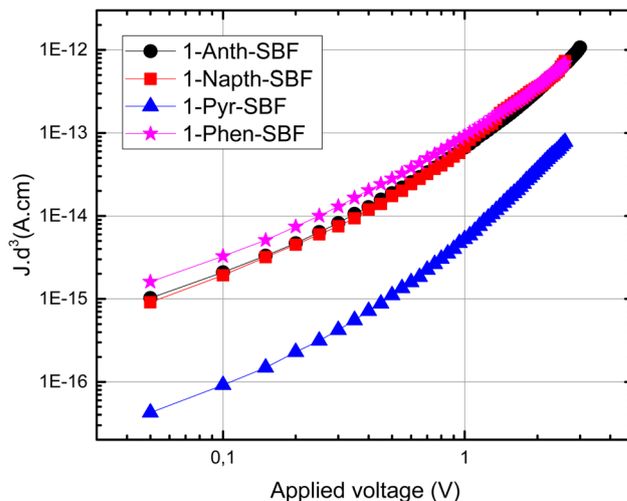


Fig. 9 SCLC device (left): thickness-scaled current–voltage characteristics of **1-Napht-SBF**, **1-Phen-SBF**, **1-Anth-SBF** and **1-Pyr-SBF** hole-only SCLC devices (right): The dotted lines indicate the SCLC regime and the continuous ones indicate the Ohmic regime.

trend of the substituent ($3.93/3.85/3.23/3.07$ eV for **1-Napht-SBF/1-Phen-SBF/1-Pyr-SBF/1-Anth-SBF**) are in accordance with those obtained by theoretical calculations (Table 1) and confirm the above-mentioned conclusions on the impact of the substituent.

The influence of the nature of the substituent on the triplet state energy level (E_T) is also of chief importance in the design of organic materials when hosting a phosphor in PhOLEDs is the goal.⁴⁵ For the last 20 years, the field of host materials for PhOLEDs has been driven by this feature. Since the first report, in 2017, of a C1-substituted SBF as the host material in a blue PhOLED,⁸ the C1-substituted scaffold has become one of the most efficient molecular platforms to reach high performance devices.^{1,22,23,54}

At 77 K, the phosphorescence contributions of the emission spectra of both **1-Napht-SBF** and **1-Phen-SBF** are similar with a first band centred at 475/470 nm leading to E_T values of 2.61 and 2.64 eV, respectively (Fig. 7). The E_T values are slightly lower than for their corresponding substituents, 2.65 and 2.71 eV respectively, in accordance with a triplet exciton spread out exclusively on the substituent (Fig. 7, inset). The small decrease can be assigned to the through-space interaction as detailed above. Interestingly, the E_T value is significantly decreased compared to that of **SBF**, evaluated at $E_T = 2.89$ eV, highlighting the significant influence of the substituent on the T_1 state. This is a different result to what was reported for structurally related C1-substituted SBFs previously reported in the literature such as **1-Phenyl-SBF** for which the E_T is maintained very high, 2.86 eV, as a very small π -conjugated unit, *i.e.* the phenyl unit, is attached. Thus, the E_T can be easily tuned by the grafted substituent. As the π -conjugation of the attached substituent increases, there is a drastic change in the corresponding phosphorescence contributions. Indeed, in **1-Pyr-SBF**, the E_T is significantly decreased to 2.07 eV, with a shift of the first emission band at 597 nm (the SDD of the triplet is exclusively centred on the pyrene unit). The E_T of pyrene,



measured 2.29 eV, is higher than that **1-Pyr-SBF** in accordance with the above-mentioned conclusion. Finally, with anthracene grafted in **1-Anth-SBF**, there is no phosphorescence contribution detected in the range as anthracene with its three fused phenyl ring provides a highly conjugated system. Radiative deactivation of the triplet state has also been measured and appears to be very slow under these experimental conditions: the phosphorescence decay was measured and the lifetimes of the T_1 state were found to be 2.7 and 3.6 s for **1-Napht-SBF** and **1-Phen-SBF**, respectively. In **1-Pyr-SBF**, the lifetime is significantly shorter, 0.6 s, highlighting the more important contribution of the pyrene fragment.

Superposition of optimized geometries of the ground state (S_0) and the first triplet excited state (T_1) shows that the high rigidity and the particular molecular arrangement of the 1-SBF scaffold strongly prevents reorganization at the first triplet excited state oppositely to what was observed for S_1 (see above) (Fig. 8). For the ground state, the substituted-fluorene dihedral angle is very high maintaining the π -conjugation breaking between the two fragments (Table 1). The E_T decrease of 1-SBFs compared to their building units can then not be assigned to an increase of the π -conjugation between the substituent and the fluorene but more to the cofacial arrangement.

Charge transport properties

Finally, as the C1 position of SBF and related fluorene compounds has been successfully used in the OLED field with very high efficiencies either in phosphorescence or in TADF,^{23,33} the charge transport properties were investigated using space charge limited current (SCLC) devices (see the device architecture in Fig. 9) to measure the out-of-plane hole mobilities. The whole series of C1-substituted SBF compounds exhibit a hole mobility of *ca* 10^{-6} cm² V⁻¹ s⁻¹, which are intrinsically low values but in line with other reports for materials constructed on SBF scaffolds (3D shape). In fact, compared with very high performance host materials for PhOLEDs recently reported, the present mobilities are in the same range.²³

The small differences observed within the C1-substituted SBF molecules suggest that the nature of the substituents does not significantly influence the π -overlaps between neighbouring molecules in the solid state, up to a certain level of steric hindrance. Indeed, one can nevertheless note that **1-Pyr-SBF** displays a mobility of one order of magnitude lower than its analogues, despite the pyrene flat and extended π -system which can act as the π -stacking platform. This probably results from the high steric hindrance imposed by the pyrene-substituent whose conjugation plane appears to be slightly more tilted as regards to the cofacial fluorene (high slippage angle).

Conclusions

The present work reports the synthesis and the study of a series of organic semi-conductors constructed on an emerging molecular fragment, namely the C1-substituted SBF. This scaffold is nowadays one of the most efficient materials for

phosphorescence and TADF OLEDs and deserves particular attention. However, this fragment has been, to date, used only with small substituents introduced at the C1 position. In the present work, extended π -conjugated systems, namely naphthalene, anthracene, phenanthrene and pyrene, were introduced and their effects on the electronic properties were investigated. We show that the substituent attached mainly drives the optical and electrochemical properties, which are nevertheless modulated by the π - π intramolecular interactions occurring with the cofacial fluorene. Thanks to the structural study based on the work of Janiak and coworkers, we demonstrate that the particular arrangement of the C1 SBF scaffold induces moderate to strong cofacial interactions with the pendant substituent attached. The impact of this cofacial interaction appears to be different as a function of the substituent and of the property investigated: HOMO/LUMO energy levels, absorption, fluorescence, and phosphorescence. As tuning intramolecular interactions is of great interest notably to control the ΔE_{S-T} in TADF OLEDs, this study shows how the fragment grafted at C1 has a key role in this interaction. This study will help to further design organic semi-conductors based on this more and more used molecular scaffold.

Conflicts of interest

There are no conflicts to declare.

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