

## **9H-Quinolino[3,2,1-k]phenothiazine: A New Electron-Rich Fragment for Organic Electronics**

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# 9H-Quinolono[3,2,1-k]phenothiazine: A new electron-rich fragment for organic electronics

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Since the last 30 years, the fantastic emergence of organic electronics is assignable to the design, the synthesis and the study of thousands of organic semi-conductors (OSCs) with specific properties fitting with the different electronic devices.<sup>[1-4]</sup> Since the discovery of the 'host/guest concept' in Phosphorescent Organic Light Emitting Diodes (PhOLEDs),<sup>[5]</sup> the design of OSCs usable as host materials for blue emitting phosphors has been an intense research field worldwide.<sup>[4]</sup> In addition to its high thermal/morphological stability and its high triplet energy ( $E_T$ ), an ideal host should also possess a bipolar character with HOMO and LUMO energy levels adapted to the Fermi levels of the electrodes.<sup>[6-9]</sup> However, there is a certain antimony between a high  $E_T$  (essential to insure efficient energy transfers) and a short HOMO/LUMO gap (essential to insure efficient charge injection) and the best trade-off should be found for the ideal host. If the nature of the linkage<sup>[10]</sup> between the electron rich and the electron poor units is of key importance to control the  $E_T$  as well as the HOMO/LUMO gap, their intrinsic properties remain nevertheless the most important characteristics. In this context, the widely known electron rich carbazole fragment has led to impressive breakthroughs and is nowadays an important building block in organic electronics.<sup>[11,12]</sup> Around the molecular structure of carbazole, many other carbazole-like fragments have been developed in the recent years sometimes surpassing the performance of the carbazole fragment in electronics. Thus, indolocarbazole (IC),<sup>[13]</sup> phenylacridine (PA)<sup>[14-16]</sup> and more recently indoloacridine (IA)<sup>[17]</sup> are nowadays key building blocks, widely used in all organic electronic devices.

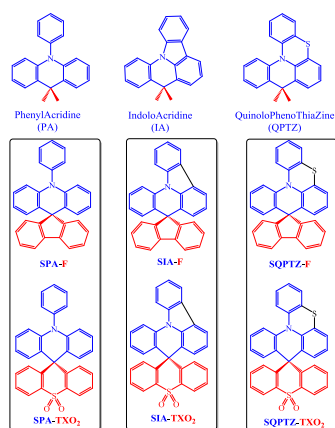
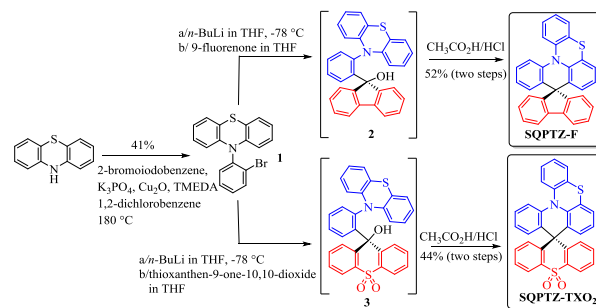


Chart 1: Phenylacridine (PA, left),<sup>[14, 18]</sup> indoloacridine (IA, middle)<sup>[17]</sup> and quinolophenothiazine (QPTZ, right) based semi-conductors

As the future of organic electronics strongly relies on the

discovery of novel elementary molecular fragments, such as for example the new bithiophenesulfonamide building block recently reported by Marks, Facchetti *et al.*,<sup>[19]</sup> we wish to report herein a new promising electron rich core, namely the quinolophenothiazine (QPTZ). The QPTZ unit can be seen as a two ortho-bridged triphenylamine or as a pentacyclic fused ring amine. The QPTZ fragment is hence an IA core in which a sulphur atom has been incorporated within the biphenyl linkage. This sulphur atom is expected to have a major role in the electronic properties with the raising of the HOMO energy level and of the spin-orbit coupling. Thus, this molecular fragment, described more than 35 years ago,<sup>[20]</sup> is nevertheless to the best of our knowledge, almost absent from literature with only one example reported without any device application.<sup>[21]</sup> Herein, the potential of the QPTZ fragment has been evaluated through the synthesis of two spiro-configured dyes, Spiroquinolophenothiazine-Fluorene (**SQPTZ-F**) and Spiroquinolophenothiazine-dioxothioxanthene (**SQPTZ-TXO<sub>2</sub>**) possessing a D-Spiro-A design (Scheme 1).<sup>[14-16]</sup> These semi-conductors have been studied in detail, incorporated as host materials in blue PhOLEDs and compared to their structurally related parents possessing a PA (**SPA-TXO<sub>2</sub>**/**SPA-F**,<sup>[18]</sup> Chart 1-left) or an IA core (**SIA-TXO<sub>2</sub>**/**SIA-F**,<sup>[17]</sup> Chart 1-middle) previously reported in literature. It turns out that with a combination of a moderate triplet energy and a high HOMO energy level, the QPTZ fragment appears to lead to strongly more efficient blue PhOLEDs than those reported with the IA and PA fragments. This first electronic application displays the fantastic potential of this fragment in organic electronics.



Scheme 1: Synthesis of **SQPTZ-F** and **SQPTZ-TXO<sub>2</sub>**

**SQPTZ-F** and **SQPTZ-TXO<sub>2</sub>** were readily obtained through a short and highly efficient synthetic approach by coupling the halogeno-aryl 10-(2-bromophenyl)-10H-phenothiazine **1**, obtained from the copper catalysed Goldberg coupling of phenothiazine and 2-bromiodobenzene (Scheme 1), with either 9-fluorenone (for **SQPTZ-F**) or 9H-thioxanthen-9-one-10,10-dioxide (for **SQPTZ-TXO<sub>2</sub>**). The corresponding fluoreneol **2** and thioxanthenol **3** respectively obtained were not isolated and were involved in an intramolecular electrophilic substitution to provide the two targets **SQPTZ-TXO<sub>2</sub>** and **SQPTZ-F** with good yields. These short and efficient synthetic approaches allow a gram-scale preparation, key feature for electronic applications.

X-ray diffraction data (Figure 1) show that **SQPTZ-F** crystallizes in a monoclinic system,  $P2_1/a$ , revealing an asymmetric unit containing two molecules (Mol1/Mol2, see Fig. S1-S4/S7-S10 in SI) whereas **SQPTZ-TXO<sub>2</sub>** crystallizes in a triclinic system  $P-1$  with also two molecules (Mol1/Mol2, see Fig. S13-S16/S19-S22 in SI). A rough value of the molecular radius of each molecule

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(approximately obtained by the distance from the spiro carbon atom to the farthest carbon atom) has been evaluated at 6.7 Å for **SQPTZ-F** and at 6.8-6.9 Å for **SQPTZ-TXO<sub>2</sub>**. For both molecules, this radius is imposed by the QPTZ unit, which is larger than the other aromatic units (F or TXO<sub>2</sub>). These radii (involved in the Lippert-Mataga formalism used to determine the polarity of the excited states, Figure 3D) are smaller than those reported for structurally analogues **SIA-TXO<sub>2</sub>/SIA-F** (6.95 to 7.02 Å)<sup>[17]</sup> and **SPA-TXO<sub>2</sub>/SPA-F** (7.14 Å).<sup>[14]</sup> As the molecular radius is governed by the QPTZ unit, its decrease comes from a more intense deformation of the QPTZ unit compared to that of its structurally related analogues, IA and PA fragments, which are not deformed at all. Indeed, the significant deformation of the QPTZ core mainly finds its origin in the deformation of the phenothiazine unit (deformation angle from 37.4° to 46.9°) and to a lesser extent in that of the acridine units (deformation angle between 15.1 and 32.9°). This strong deformation of the QPTZ core is a key feature to maintain a high E<sub>T</sub> (see below). It should be stressed that these important structural features lead to very short intramolecular C/C contacts (shorter than the sum of their Van der Waals radii, see Fig. S5, S11, S17 and S23 in SI). A very short intramolecular C/C distance of ca 3.0 Å is especially detected between the two carbons in β position of the nitrogen atom highlighting the strong steric congestion between the phenothiazine and the acridine backbones.

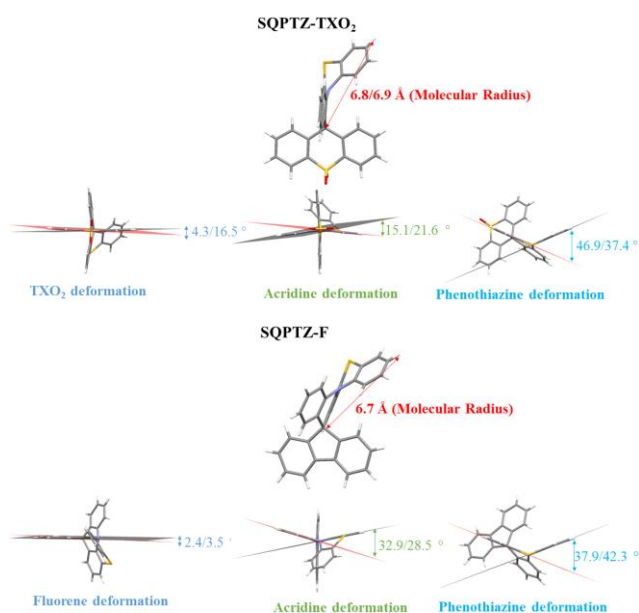


Figure 1. Molecular structures of **SQPTZ-TXO<sub>2</sub>** (top) and **SQPTZ-F** (bottom) exemplified by one molecule (*Mol 1*) with extreme values of the different parameters for the 2 molecules present in the asymmetric unit

The electron rich character of the QPTZ fragment is the most important feature and has been investigated by cyclic voltammetry (Figure 2). In oxidation, the two compounds remarkably present **two reversible single electron oxidation processes** with peak potentials E<sup>1</sup>/E<sup>2</sup> (vs SCE) recorded at 0.89/1.61 V for **SQPTZ-F** and at 1.0/1.64 V for **SQPTZ-TXO<sub>2</sub>**. As TXO<sub>2</sub> and F moieties are oxidised through irreversible processes, these two oxidation waves have been assigned to two **single** electron transfers involving the QPTZ core. Thus, the first oxidation of the QPTZ fragment remarkably occurs at a less anodic potential than that of the PA (E<sup>1</sup> > 1.11 V in **SPA-TXO<sub>2</sub>**)<sup>[17]</sup> and that of the IA core (E<sup>1</sup> = 1.33 V for **SIA-TXO<sub>2</sub>** and 1.19 V for **SIA-F**).<sup>[14]</sup> This strong electron rich character finds hence its origin in the electron donating behaviour of the intracyclic sulphur atom, which fully drives the HOMO energy level of these dyes (Figure 4). Another important feature deserves

to be stressed out. Indeed, the reversibility of the two first oxidation waves clearly shows the efficient delocalization of the charges within the QPTZ core, which is not the case for the structurally analogues based on the IA or on the PA core (Figure 2-right).<sup>[14, 17]</sup> Thus, the sulphur atom has a significant impact on the electrochemical properties not only in terms of potential but also in the stability of the cationic species. From the onset potential pointed at 0.9 V, we determined the HOMO energy level of **SQPTZ-TXO<sub>2</sub>** at -5.3 eV, significantly higher than that of **SPA-TXO<sub>2</sub>** (-5.42 eV) and **SIA-TXO<sub>2</sub>** (-5.61 eV). Similarly, the HOMO energy level of **SQPTZ-F** (-5.16 eV) is more than 0.3 eV higher than that of **SIA-F** (-5.48 eV) confirming the strong electron-rich nature of the QPTZ fragment.<sup>[14]</sup> One can also note that the HOMO energy level of **SQPTZ-F** is higher than that of **SQPTZ-TXO<sub>2</sub>**, translating the non-negligible influence of the acceptor unit on the HOMO level via the spiro bridge. Thus, the HOMO energy levels of QPTZ based molecules are high and should favour the hole injection within an electronic device.

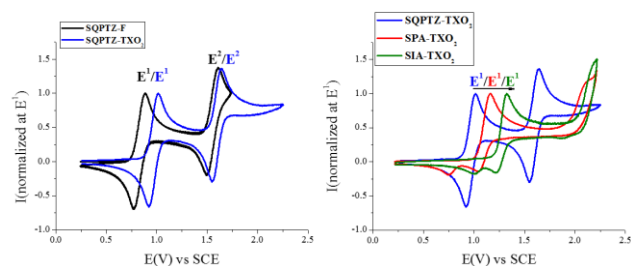


Figure 2. Cyclic voltammetry (CH<sub>2</sub>Cl<sub>2</sub> + Bu<sub>4</sub>NPF<sub>6</sub> 0.2 M, sweep-rate 100 mV.s<sup>-1</sup>) of **SQPTZ-F** and **SQPTZ-TXO<sub>2</sub>** (left) and **SIA-TXO<sub>2</sub>**, **SPA-TXO<sub>2</sub>** and **SQPTZ-TXO<sub>2</sub>** (right).

In reduction, neither **SQPTZ-F** nor **SQPTZ-TXO<sub>2</sub>** present any distinguishable reduction wave before the accessible limit of the present electrolytic solution (see Fig. S33 in SI). We determined their LUMO energy levels from the onset potential of their reduction at -1.85 and -2.06 eV for **SQPTZ-F** and **SQPTZ-TXO<sub>2</sub>**, highlighting the stronger electron-poor character of TXO<sub>2</sub> compared to that of F in accordance with the above mentioned results on HOMO energies. The electrochemical gaps ( $\Delta E^{el}$ ) were found to be 3.31 eV for **SQPTZ-F** and 3.24 eV for **SQPTZ-TXO<sub>2</sub>**. Thus, **SQPTZ-TXO<sub>2</sub>** possesses a  $\Delta E^{el}$  significantly contracted by 0.2/0.3 eV compared to that of the PA and IA counterparts ( $\Delta E^{el}$  **SPA-TXO<sub>2</sub>** = 3.43 eV,  $\Delta E^{el}$  **SIA-TXO<sub>2</sub>** = 3.56 eV) due to its high HOMO energy level. As the same gap contraction is detected between **SIA-F** and **SQPTZ-F** (3.60 vs 3.31 eV *resp.*), this clearly highlights the strong impact of the QPTZ fragment to contract the gap, independently of the spiro-connected unit. Geometry optimization of the two compounds was performed using density functional theory (DFT, B3LYP/6-31G(d)). The HOMO and LUMO levels follow the same tendencies than those obtained by electrochemical studies with notably a higher HOMO energy level for **SQPTZ-F** (-5.03 eV) than for **SQPTZ-TXO<sub>2</sub>** (-5.30 eV). The HOMO levels are spread on the QPTZ fragment (Figure 4), confirming a first electron transfer from this core. The calculated LUMO levels are respectively lying at -0.92 and -1.17 eV for **SQPTZ-F** and **SQPTZ-TXO<sub>2</sub>** and are spread out on the fluorene of **SQPTZ-F** and on the TXO<sub>2</sub> of **SQPTZ-TXO<sub>2</sub>** (Figure 4). There is hence a complete electronic separation between the HOMO and the LUMO, which will have important repercussions on the photophysical processes (see below). This orbital separation is a key point to gather within a single dye the HOMO/LUMO energy levels of the constituting building blocks, a short  $\pi$ -conjugation pathway and a high E<sub>T</sub>.

Absorption spectra of **SQPTZ-TXO<sub>2</sub>** and **SQPTZ-F** are presented in Figure 3A. **SQPTZ-F** presents an UV-vis spectrum with maxima recorded at 262, 296 and 308 similar to that of its

building unit fluorene (265, 290, 301 nm) at the exception of the long tail observed between 320 and 360 nm. In the light of TD-DFT, the main absorption band centred at 308 nm is assigned to a transition with two contributions (HOMO/LUMO+4 and HOMO/LUMO+5,  $f=0.117$ , see Fig. S37 in SI) implying only the QPTZ fragment, the one at 296 nm is attributed to fluorene-fluorene transition (HOMO-1/LUMO,  $f=0.067$ ), and the large high energy band at 262 nm to phenyl-phenyl transitions. More importantly, the low energy tail is noticeably large due to the flexibility induced by the presence of the sulphur atom within the QPTZ fragment and is assigned to HOMO/LUMO+1 and HOMO/LUMO+2 transitions (Figure 4-left). Indeed, the calculated absorption spectrum shows that the first excited state corresponds to an HOMO-LUMO transition but with a very weak oscillator strength ( $f=0.0002$ ), not detectable experimentally. This is a crucial point in the D-spiro-A design, which finds its origin in the spatial separation of HOMO and LUMO levels (HOMO delocalized on the QPTZ core and LUMO on the fluorene core) leading to through-space forbidden transitions. Oppositely, the HOMO/LUMO+1 and HOMO/LUMO+2 transitions possess oscillator strengths of ca 0.02, being ca 100 times more probable than the HOMO/LUMO transition (Figure 4-left). These transitions involve almost exclusively the QPTZ core which hence governs the optical gap  $\Delta E^{\text{opt}}$  of **SPQZT-F** (ca 3.3/3.4 eV). Despite an almost identical  $\Delta E^{\text{opt}}$ , the absorption spectrum of **SPQZT-TXO<sub>2</sub>** is slightly different in the low energy region with a broader tail. As observed for **SPQZT-F**, the HOMO/LUMO transition of **SPQZT-TXO<sub>2</sub>** possesses a very weak oscillator strength ( $f=0.003$ , Figure 4-right) due to the spatial separation of HOMO (centred on QPTZ) and LUMO (centred on TXO<sub>2</sub>) and is not experimentally observed. Thus, the main low-energy transition is an HOMO/LUMO+1 transition ( $f=0.042$ , see Fig. S39 in SI). There is nevertheless a significant difference between **SPQZT-F** and **SPQZT-TXO<sub>2</sub>**, which is linked to the different localization of the LUMO+1 orbital: on the QPTZ core for the former and on the TXO<sub>2</sub> core for the latter. This difference confers to the HOMO/LUMO+1 transition of **SPQZT-TXO<sub>2</sub>** a notable charge transfer character, which is not the case for **SPQZT-F** (see solvatochromic experiments Figure 3D).

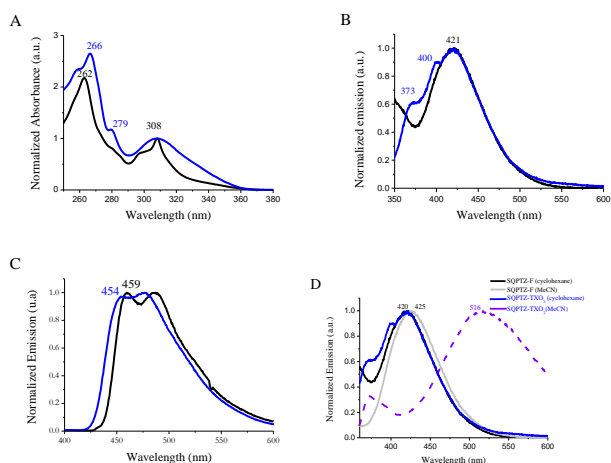


Figure 3. Optical properties of **SPQZT-F** (black line) and **SPQZT-TXO<sub>2</sub>** (blue line); **A**: UV-vis absorption spectra (cyclohexane), **B**: Emission spectra (cyclohexane) at rt under argon ( $\lambda_{\text{exc}}=330$  nm), **C**: Emission spectra at 77K (2-MeTHF,  $\lambda_{\text{exc}}=315$  nm). **D**: Emission spectra in cyclohexane and acetonitrile.

**SPQZT-TXO<sub>2</sub>** and **SPQZT-F** display almost no fluorescence and the emission spectra were recorded in absence of oxygen (under argon, Figure 3B). Thus, the emission of **SPQZT-F** and that of **SPQZT-TXO<sub>2</sub>** appear very similar with a main large and unresolved band at 421 nm. The quantum yields (QY), measured

in presence of oxygen, appear very low ( $\text{QY} < 0.05$  for both). Thus, QPTZ fragment, due to the insertion of the intracyclic sulphur atom, is very weakly fluorescent in opposition to the IA core, which is highly fluorescent ( $\text{QY}\sim 0.4$ ).<sup>[17]</sup> This feature can be assigned to internal heavy atom effects as recently shown for phenothiazinium dyes and highlights the significant impact of the sulphur atom on the electronic properties of the QPTZ fragment.<sup>[22]</sup> Solvatochromic experiments allow a deeper understanding of the photophysical properties through the determination of the polarity of the excited states (Figure 3, bottom-right). Indeed, the emission maximum of **SPQZT-F** in polar MeCN is recorded at 425 nm, being only very weakly shifted by 4 nm, compared to that in apolar cyclohexane (421 nm), in accordance with a transition occurring with only one fragment involved, namely QPTZ. **SPQZT-TXO<sub>2</sub>** displays a very different behaviour with an impressive red shift of 98 nm between cyclohexane (421 nm) and MeCN (516 nm) highlighting a strong photoinduced Intramolecular Charge Transfer (ICT) character due to the different localization of the HOMO and LUMO+1 levels.  $\Delta\mu$  values of 6.05 D and 21.51 D have been evaluated using the Lippert-Mataga formalism (the dipole moments at the ground state obtained through DFT calculations were: 1.6 and 5.1 D for **SPQZT-F** and **SPQZT-TXO<sub>2</sub>**, see Fig. 27-29 in SI), translating the different polarity of the two dyes at the excited state ( $\mu^*=7.5$  and 26.6 D resp.). Thus, the strong solvatochromic effect and the corresponding high  $\mu^*$  observed for **SPQZT-TXO<sub>2</sub>** are indicative of a significant photoinduced ICT whereas the negligible solvatochromic effect and the corresponding low  $\mu^*$  observed for **SPQZT-F** are indicative of a very weak photoinduced ICT. Thus, the acceptor unit can drastically change the electronic properties of QPTZ based dyes.

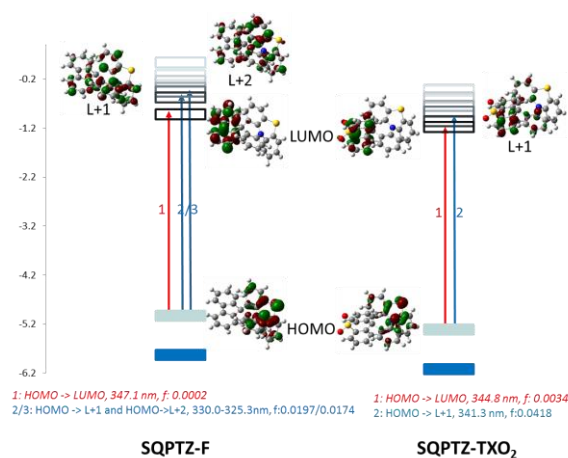


Figure 4. The first calculated electronic transitions of **SPQZT-F** (Left) and **SPQZT-TXO<sub>2</sub>** (Right) by TD-DFT B3LYP/6-31G(d), shown with an isovalue of 0.04 [ $e \text{ bohr}^{-3}$ ]<sup>1/2</sup>.

The phosphorescent spectra (Figure 3C), recorded at 77K, appear large and weakly structured and contrarily to the fluorescence spectra, are relatively intense. As the fluorescence of the two molecules is low, the intersystem crossing (ISC) between  $S_1$  and  $T_1$  is fast and the phosphorescence contribution at 77K is very intense. This is a remarkable characteristic of the QPTZ fragment, which results from the incorporation of the sulphur atom which enhances the ISC by vibronic spin-orbit coupling.<sup>[22]</sup> The transition of highest energy presents a maximum at 459 nm for **SPQZT-F** and 454 nm for **SPQZT-TXO<sub>2</sub>** giving respective triplet energy  $E_T$  of 2.70 and 2.73 eV. QPTZ based compounds possess hence an  $E_T$  of ca 2.7 eV, smaller than those of PA (3.08 eV for **SPA-TXO<sub>2</sub>** and 3.02 eV for **SPA-F**) and IA analogues (2.93 eV **SIA-TXO<sub>2</sub>** and 2.87 eV for **SIA-F**) indicating a different localization of the triplet exciton in IA, PA and QPTZ



based molecules. In summary, thanks to the QPTZ fragment, the two dyes possess an HOMO/LUMO gap relatively small as well as an  $E_T$  high enough to be used with the blue FIrpic phosphor ( $E_T=2.62$  eV<sup>[23]</sup>). This combination appears appealing for the future design of host materials based on this fragment.

The potential of the QPTZ fragment in electronics has been finally evaluated using **SQPTZ-F** as host in a blue PhOLED (10% FIrpic). The performance of these devices has been compared to those of structurally related **SIA** and **SPA** analogues with exactly the same device architecture. Indeed, in order to accurately compare the performance of the hosts, the device architecture should be strictly identical (see PhOLED architecture in SI). Thus, **SQPTZ-F** displays a very high EQE of 15.8% at  $J=10$  mA/cm<sup>2</sup> with corresponding Current Efficiency (CE) of 32.8 cd.A<sup>-1</sup> and Power Efficiency (PE) of 21.7 lm.W<sup>-1</sup> (see Fig. S40 in SI). These performances are ca 50% higher than those recorded with the IA counterpart **SIA-F** (EQE of 10.7% at  $J=10$  mA/cm<sup>2</sup> with corresponding CE of 24.2 cd.A<sup>-1</sup> and PE of 13.3 lm.W<sup>-1</sup>) (see Fig. S42 in SI). In addition, the turn on voltage  $V_{on}$  (at 1 cd.m<sup>-2</sup>) of **SQPTZ-F** based device appears to be very low, 2.7 V, for a blue PhOLED and significantly lower than that of **SIA-F** recorded at 3.5 V. This feature clearly translates a better charge injection within the **SQPTZ-F**/FIrpic emitting layer than within the **SIA-F**/FIrpic one. Similarly, **SQPTZ-TXO<sub>2</sub>** emits light at a low voltage of 2.8 V and displays a very high EQE of 15.8% (10 mA/cm<sup>2</sup>) with corresponding CE and PE of 33.1 cd.A<sup>-1</sup> and 17.6 lm.W<sup>-1</sup> (see Fig. S41 in SI), clearly highlighting that these very high performances are induced by the QPTZ fragment. The electroluminescent spectra of all devices (see Fig. S40-41 in SI) are identical, exclusively showing the emission of the blue dopant at 472 and 496 nm close to the photoluminescence of pure FIrpic (468/495 nm in solution in CH<sub>2</sub>Cl<sub>2</sub>), indicating efficient energy transfers.<sup>[23]</sup> It is also important to mention that **SQPTZ-F** and **SQPTZ-TXO<sub>2</sub>** displays a better performance and a lower turn on voltage than structurally related IA and PA compounds reported in literature. Thus, **SIA-TXO<sub>2</sub>** (EQE of 12.7 % at 10 mA/cm<sup>2</sup> with corresponding CE of 27.7 cd.A<sup>-1</sup>, PE of 14.5 lm.W<sup>-1</sup> and a  $V_{on}$  of 2.9 V)<sup>[17]</sup> and **SPA-TXO<sub>2</sub>** (EQE of 11.4% at 10 mA/cm<sup>2</sup> with CE of 28.4 cd.A<sup>-1</sup>, LE of 14.4 lm.W<sup>-1</sup> and a  $V_{on}$  of 2.9 V)<sup>[15]</sup> both display a lower performance and a higher  $V_{on}$  than that exposed above. Thus, switching from IA or PA core to QPTZ allows a strong EQE increase as well as a decrease of the  $V_{on}$ . Some recent universal hosts based on the indolocarbazole backbone, namely diphenyl-indolocarbazole (EQE=10.4%) and phenylpyridine-indolocarbazole (EQE=13.4%)<sup>[13]</sup> also display lower performance for FIrpic based devices than those reported herein, indicating the strong potential of the QPTZ core in electronics.

In summary, this work reports the first example of a new molecular fragment, namely QPTZ, used in electronics. The QPTZ fragment can be easily synthesized, presents a high HOMO level, -5.2/5.3 eV and a moderate  $E_T$  value of ca 2.7 eV. This combination leads to high-efficiency blue PhOLEDs displaying higher performance (increase by 50%) than those reported with the PA and IA analogues. This preliminary result highlights the potential of this fragment in the field of PhOLEDs. We believe that the future development of QPTZ based materials is appealing for other organic devices and can advance the field of organic semi-conducting materials.

#### Acknowledgment:

**Keywords:** Quinolophenothiazine · PhOLED · organic semi-conductors · spiro compounds

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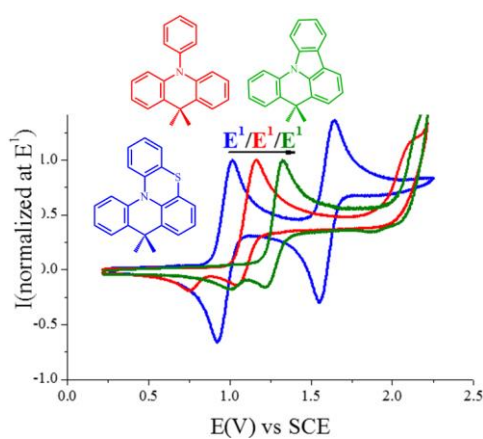
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Page – Page

9H-Quinolino[3,2,1-k]phenothiazine:  
A new electron-rich fragment for  
organic electronics



We report a novel electron-rich fragment for organic electronic applications, namely the quinolophenothiazine (QPTZ). The sulphur atom of QPTZ fragment plays a major role in the electronic properties (increasing the HOMO energy level, modulate the spin-orbit coupling...) and leads to highly efficient blue Phosphorescent Organic Light Emitting diode, indicating the strong potential of this new molecular fragment in organic electronics.