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The Dark State of the Thiele Hydrocarbon: Efficient Solvatochromic Emission from a Non-polar Centrosymmetric Singlet Diradicaloid

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Singlet diradicaloid • polyhalogenated trityl radicals • Thiele hydrocarbon • double exciton state • solid state fluorescence

ABSTRACT: In this work, a comprehensive investigation of the photoinduced processes and mechanisms, which are linked to the luminescence of a novel non-perchlorinated Thiele hydrocarbon (TTH), is presented. Despite the comparable diradical character of TTH ($\gamma_0 = 0.32 - 0.44$) and unsubstituted Thiele hydrocarbon (TH) ($\gamma_0 = 0.30$), the polyhalogenated species is inert and photostable, showing an intense deep-red/NIR fluorescence (PLQY = 0.84 in toluene), even at room temperature and in solid state (PLQY = 0.19). TTH displays a large Stokes shift (307 nm in benzonitrile) and solvatochromic behavior, which resulted unusual for a centrosymmetric, non-polar and low-conjugated species. These outstanding emission features were interpreted through quantum-chemical calculations indicating that its fluorescence arises from the low-lying dark doubly-excited zwitterionic state, typically found at low excitation energies in diradicaloids, acquiring dipole moment and intensity by state mixing *via* twisting around the strongly elongated exocyclic CC bonds of the excited *p*-quinodimethane (*p*QDM) core, with a mechanism similar to sudden polarization occurring in olefins. Such mechanism was derived from ns and fs transient absorption measurements.

INTRODUCTION

Singlet diradicaloids (SDs) are emerging as a promising class of molecular materials due to their potential multiple applications in several fields.^{1,2} Many examples of SDs have been reported as conductive materials,^{3,4} semiconductors in organic field-effect transistors (OFETs),^{5–8} singlet-fission species for organic photovoltaics (OPV),^{9–11} NIR absorbing chromophores and dyes for non-linear optics (NLO).^{12,13} A widely underestimated property of these versatile compounds is their luminescence.^{11,14–16} This is quite surprising, since the first ever synthesized SD, the Thiele hydrocarbon (TH) (Figure 1),¹⁷ was described as fluorescent in solution, slightly

oxygen and light sensitive.¹⁸ Probably, its photosensitivity and essentially quinoidal ground state configuration limited the interest in the luminescence properties own by the simplest *para*-quinodimethane (*p*QDM) diradicaloid. Actually, most of the synthetic efforts for developing new and stable TH derivatives, were focused on the enhancement of the diradical character (described with the diradical index γ_0 , ranging from zero to one, moving from a closed-shell species to a pure diradical)¹⁹ and mainly consisted in the introduction of heteroatoms in the TH backbone and the extension of the π -system.^{20–25} However, the luminescence arising from open-shell species is currently intensively investigated, focusing on the development of highly fluorescent polyhalogenated trityl radicals.²⁶

TTM and **PTM** (**Figure 1**) adducts have been proficiently used as functional units in several molecular materials for NLO,^{27,28} circularly polarized luminescence (CPL),^{29–31} magneto^{32,33} and electro luminescence.^{34,35} In particular, **TTM** was the first polyhalogenated triaryl radical exhibiting excimer emission when dispersed into rigid hosts,³⁶ while its push-pull derivatives led to record device performance when used as emitters in organic light-emitting diodes (OLEDs), reaching the physical limit of 100% of internal quantum efficiency.³⁷ The key for this success is their extraordinary stability, mainly ensured by the six *ortho* chlorine atoms, which exert a shielding effect on the odd electron localized on the *ipso*-C.³⁸ Interestingly, perchlorination was already used as synthetic approach for the stabilization of *p*QDM diradicaloids.^{39,40} In 1991, Castañer *et al.* reported the synthesis and characterization of the perchlorinated Thiele hydrocarbon (**PTH**) (**Figure 1**).⁴⁰ The authors expected that the twist induced by the bulky chlorine atoms

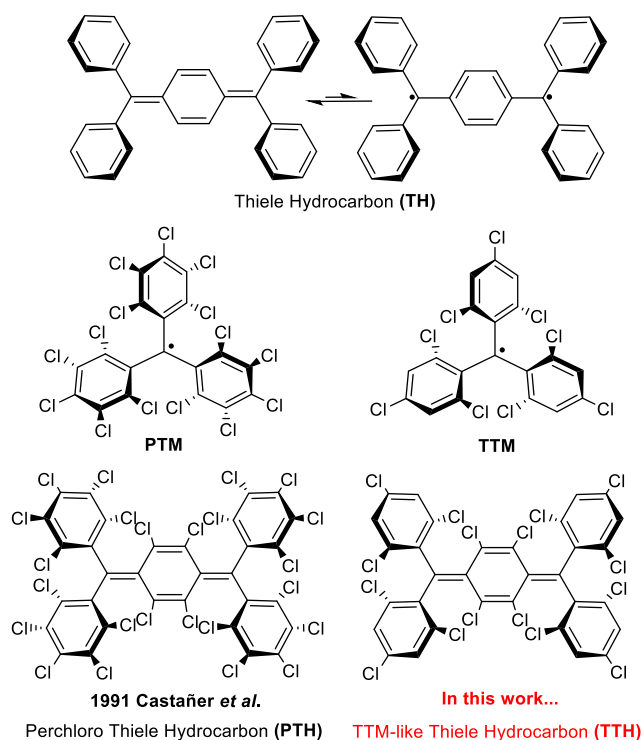
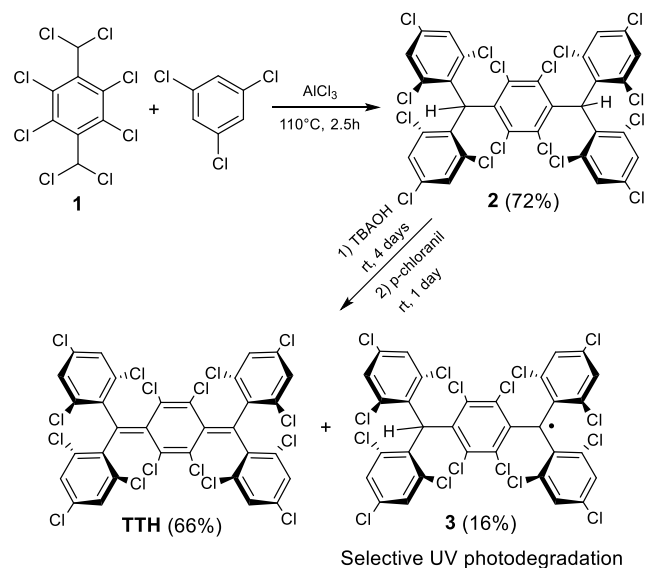


Figure 1. Molecular structures Thiele hydrocarbon (**TH**), **PTM** and **TTM** radicals, perchloro Thiele hydrocarbon (**PTH**), and TTM-like Thiele hydrocarbon (**TTH**).

nearby the exocyclic double bonds of the perchlorinated *p*-xylylene bridge, would favor a triplet electronic configuration. In contrast, the electron paramagnetic resonance (EPR) and UV-Vis characterizations depicted a quinoidal structure, consistent with the one of the unsubstituted **TH**. A further study by Domingo *et al.* focused instead on the internal charge transfer (ICT) properties of the **PTH** mixed-valence compound (anion-radical).⁴¹ Oddly, both works did not mention any luminescence feature for the perchlorinated diradicaloid. Based on these findings, we synthesized a novel TTM-like Thiele hydrocarbon (**TTH**) (**Figure 1**), namely the 2,2',2'',2'''-(perchlorocyclohexa-2,5-diene-1,4-diylidene)bis(methanediylidene)tetrakis-(1,3,5-trichlorobenzene), aiming to obtain a highly fluorescent and inert *p*QDM diradicaloid, suitable to be easily characterized and processed. In fact, in analogy with what already reported for polyhalogenated trityl radicals, **TTM** derivatives exhibit superior photoluminescence quantum yields (PLQYs) compared with **PTM** ones.^{34,42} This aspect, together with the elevated molecular rigidity of **TTH** given by the coexistence of exocyclic

double bonds and *ortho* halogens, should guarantee high values of PLQY. Moreover, the absence of chlorine atoms in *meta* positions, undeniably affects the molecule processability, improving its solubility and favoring the thermal evaporation. The interest in the synthesis of highly fluorescent *p*QDM diradicaloids goes beyond their possible applications in optoelectronics or NLO,^{43,44} since the study of their luminescence can provide new fundamental insights into the nature of the emitting excited state^{14,45} and its correlation with the ground-state open-shell character.⁴⁶ Furthermore, their diradical character is attracting much interest for the possibility of controlling spin states through conformational changes.^{47–49}



Scheme 1. Reaction scheme for the synthesis of **TTH**.

RESULTS AND DISCUSSION.

Synthesis and purification. **TTH** was synthesized with a similar procedure to the one used for **PTH** and briefly described in **Scheme 1** (details in SI). An excess of tetrabutylammonium hydroxide (TBAOH) was used for the conversion of **2** into the dianionic species, then, after the oxidation with *p*-chloranil, a mixture of **TTH** and mono radical **3** was obtained.⁵⁰ The radical nature of **3** was suggested by the presence of an intense absorption band at 382 nm, typical of polychlorinated trityl radicals (**Figure 2a**),³⁶ and was confirmed by high-resolution mass analysis (data not shown). It was not possible to purify the product *via* crystallization nor through column chromatography on silica gel since both species showed almost the same retention factor. Therefore, to isolate **TTH**, its exceptional photostability was exploited. In fact, polyhalogenated trityl radicals such as **TTM** (**Figure 1**), suffer of a severe photosensitivity (**Figure 2b**), that is usually reduced by coupling the radical core, that behaves as electron-acceptor (A) moiety, with an electron-donor (D) group, generating in this way a push-pull derivative.^{51–53} As in the case of **TTM**, the impurity **3** behaved as a photosensitive species due to the lack of a charge-transfer stabilization of the radical. The absorption spectrum in CHCl₃ of a mixed fraction containing both **TTH** and **3**, before and after the irradiation with a UV lamp at 365 nm, is shown in **Figure 2a**. The absorption band at 382 nm, belonging to **3**, was rapidly quenched while the band at 502 nm of **TTH** remained almost unaffected by the prolonged photoexcitation. After the complete photodegradation of **3**, **TTH** was easily isolated through column chromatography on silica gel (See SI).

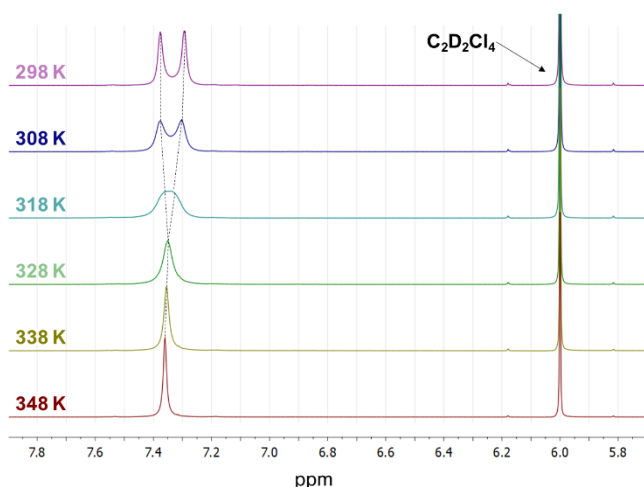


Figure 4. $^1\text{H-NMR}$ spectra of **TTH** in $\text{C}_2\text{D}_2\text{Cl}_4$ varying the temperature in the range 298 – 348 K.

High temperature $^1\text{H-NMR}$ characterization. The quinoidal structure of **TTH** was confirmed by $^1\text{H-NMR}$ characterization, performed in deuterated 1,1,2,2-tetrachloroethane ($\text{C}_2\text{D}_2\text{Cl}_4$) varying the temperature in the 298 – 348 K range. As can be seen in **Figure 4**, no thermal-accessible triplet states were detected increasing the temperature, while the temperature effect on the free rotation of phenyl substituents was clearly seen. In fact, while at 298 K the spectrum displayed two singlets at 7.37 ppm and 7.29 ppm, due to steric restrictions to the free rotation of phenyl substituents (confirming the high rigidity of the molecule at RT), starting from 328 K these signals coalesced to a single peak at 7.36 ppm, indicating their free rotation. Beyond its impressive photostability, **TTH** showed an elevated thermal stability, compatible with thermal evaporation processing. The thermogravimetric analysis (TGA) in nitrogen atmosphere displayed no degradation up to 330 °C (**Figure S8**), a lower value compared with **PTH** for which a temperature of 395 °C was reported.⁴⁰ A similar behavior is seen for poly-halogenated trityl radicals, for which their thermal stability is directly correlated with the corresponding melting points and, in turn, the molecular weight.⁵⁶

Electrochemical characterization. **TTH** shows four reversible redox couples related to the formation of the radical-anion ($E_{1/2} = -1.047 \text{ V vs. Fc/Fc}^+$), anion-anion ($E_{1/2} = -1.572 \text{ V vs. Fc/Fc}^+$), radical-cation ($E_{1/2} = +1.030 \text{ V vs. Fc/Fc}^+$) and cation-cation ($E_{1/2} = +1.278 \text{ V vs. Fc/Fc}^+$) species (**Figure 5a**). This result is impressive considering the small dimensions of the *p*QDM derivative.^{57–59} The quinoidal stabilization of the diradicaloid determines a slightly reduced electron-acceptor character compared to both **TTM** and **PTH**.^{41,60} The electrochemical bandgap (**Figure 5b**) of 2.08 eV resulted 0.1 eV smaller than the optical one. The amphoteric redox behavior of **TTH** with four reversible redox couples, makes this diradicaloid a promising candidate for energy storage applications^{61,62} or air-stable organic field-effect transistor (OFET),^{6,10} especially considering the ambipolar behavior that is usually displayed by singlet diradicaloids.⁶³

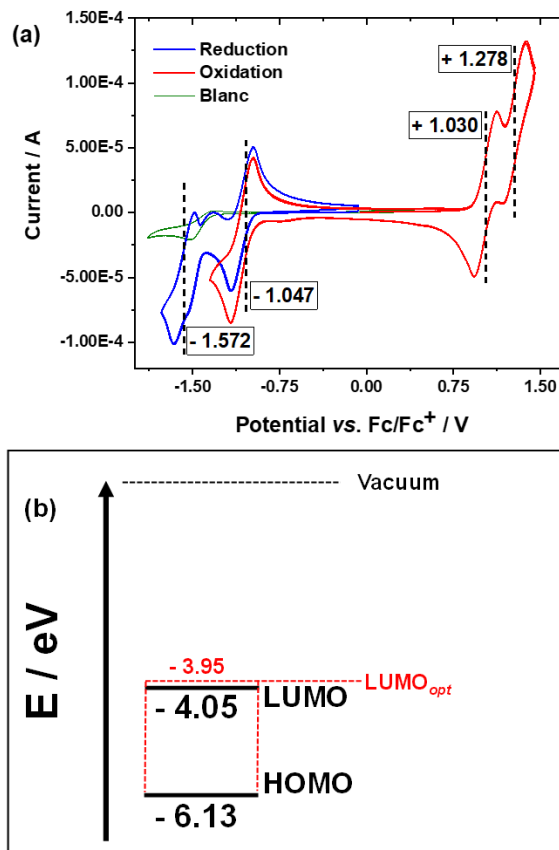


Figure 5. a) Cyclic voltammogram of a 0.5 mM solution of **TTH** with 0.1M of $\text{N}(\text{Bu})_4\text{PF}_6$, in CH_2Cl_2 , at the scan rate of 0.1 V/s; b) Representation of the HOMO and LUMO orbitals calculated through the empirical equations $E_{\text{HOMO}} = - [E_{(\text{ox vs Fc/Fc}^+)} + 5.1\text{V}]$ and $E_{\text{LUMO}} = - [E_{(\text{red vs Fc/Fc}^+)} + 5.1\text{V}]$,⁶⁴ LUMO_{opt} energy level was estimated as follows: $\text{LUMO}_{\text{opt}} = \text{HOMO} + E_{\text{gopt}}$, in which E_{gopt} is obtained from the λ_{onset} .

Spectroscopic Characterization. The absorption spectra of **TTH** in different solvents are reported in **Figure 6a**, and the main optical properties are summarized in **Table 2**. The absorption maximum is peaked around 500 nm, slightly depending on the solvent. The molar extinction coefficient in CHCl_3 was $48200 \text{ M}^{-1}\text{cm}^{-1}$ (502 nm), a value two times higher than the one reported for **PTH** ($23150 \text{ M}^{-1}\text{cm}^{-1}$ at 508 nm).⁴⁰ Despite the similar spectral region, **TTH** exhibits a sharper onset and a more symmetric band-shape compared to **PTH**. Furthermore, although the transition in the UV is already more intense for **TTH** ($13100 \text{ M}^{-1}\text{cm}^{-1}$ at 326 nm for **TTH** vs. $7800 \text{ M}^{-1}\text{cm}^{-1}$ at 335 nm for **PTH**),⁴¹ such band is still far less intense compared to that in the visible. As already mentioned, **TTH** showed an intense fluorescence in the deep-red/NIR spectral region, with a broad luminescence peaked at 691 nm in cyclohexane, having a Stokes shift of 193 nm (0.695 eV) and a photoluminescence quantum yield (PLQY) of 0.69 (detail in SI). The emission was bathochromically shifted increasing the polarity of the solvent, with the emission maximum at 813 nm in benzonitrile (BCN), reaching a Stokes shift value of 307nm (0.925 eV). This red shift was followed by a slowing of the recombination dynamics (**Figure 6b**). The PLQY values do not follow the same monotonically trend increasing the solvent polarity, in fact extremely high PLQYs were

detected in chloroform (PLQY = 0.83) and toluene (PLQY = 0.84), with a maximum of emission around 720 nm. In order to investigate the viscosity dependent optical properties of **TTH**, different mixtures of hexane and paraffin were employed (Table S5). Upon an increase in viscosity, the absorption (Figure S10) as well as fluorescence (Figure 6c) maxima were subject to minor bathochromic shifts. As such, the strong solvent dependent shifts seen for the fluorescence maxima are exclusively attributed to a polarity induced solvatochromism. This contrast the observations made with more extended and flexible derivatives.⁶⁵ As the paraffin/hexane ratio increases, fluorescence decay lifetimes are subtly shortened (Figure S11, Table S5) compared to trends observed when increasing the solvent polarity (Table 2). Interestingly though, the PLQYs show an opposite trend, namely increasing with decreasing fluorescence lifetime. We rationalize such an observation by a non-radiative deactivation that is hampered at conical intersections between nearby geometries. At the forefront are rotational barriers that rise upon increasing the overall viscosity. At the same time, restrictions of internal motion favor radiative deactivations at certain molecular geometries. This is likely to compensate the overall singlet excited state lifetime as it is inversely proportional to the sum of both radiative and non-radiative rate constants. Our considerations contribute to explain the high PLQYs in benzonitrile or even in the solid state and the short lifetimes. Relative to THF the PLQYs are higher, and the lifetimes are shorter. In the nonpolar regime, even subtle solvent changes exert sizeable changes on the optical features. A leading example is cyclohexane versus toluene. Even a slightly better stabilization of the zwitterionic character is seen to increase the transition dipole moment and, in turn, to favor the radiative decay. The immediate consequence is a higher PLQY. Interestingly, **TTH** displays an intense fluorescence even in the solid state (Figure S9), unlike what happens with trityl radicals, which suffer a severe aggregation caused quenching (ACQ).

Table 2. Main optical properties of **TTH** in different solvents

Solvent	λ_{abs} (nm)	ϵ ($M^{-1}\text{cm}^{-1}$)	λ_{em} (nm)	$\tau^{[a]}$ (ns)	PLQY
CycloHex	498	44450	691	51.2 ± 0.07	0.69 ^[b]
Hex	496	45300	691	58.2 ± 0.09	0.62 ^[b]
Toluene	503	43900	716	24.7 ± 0.03	0.84 ^[b]
CHCl ₃	501	48200	723	22.6 ± 0.03	0.83 ^[b]
Chloro-benzene	505	43250	745	17.1 ± 0.03	0.53 ^[b]
DCM	502	45820	765	11.9 ± 0.02	0.48 ^[b]
THF	501	46060	785	5.3 ± 0.01	0.10 ^[b]
Benzonitrile	506	42600	813	4.2 ± 0.01	0.14 ^[b]
Film	513	N.A.	703	4.7 ± 0.2	0.19 ^[c]

[a] The lifetimes have been obtained by best fitting the decay profiles with a mono-exponential function. [b] Values obtained integrating the emission profile up to 900 nm. [c] The value is obtained integrating on the whole emission spectrum (up to 900 nm) obtained by simulating the extension of the experimental data (see SI).

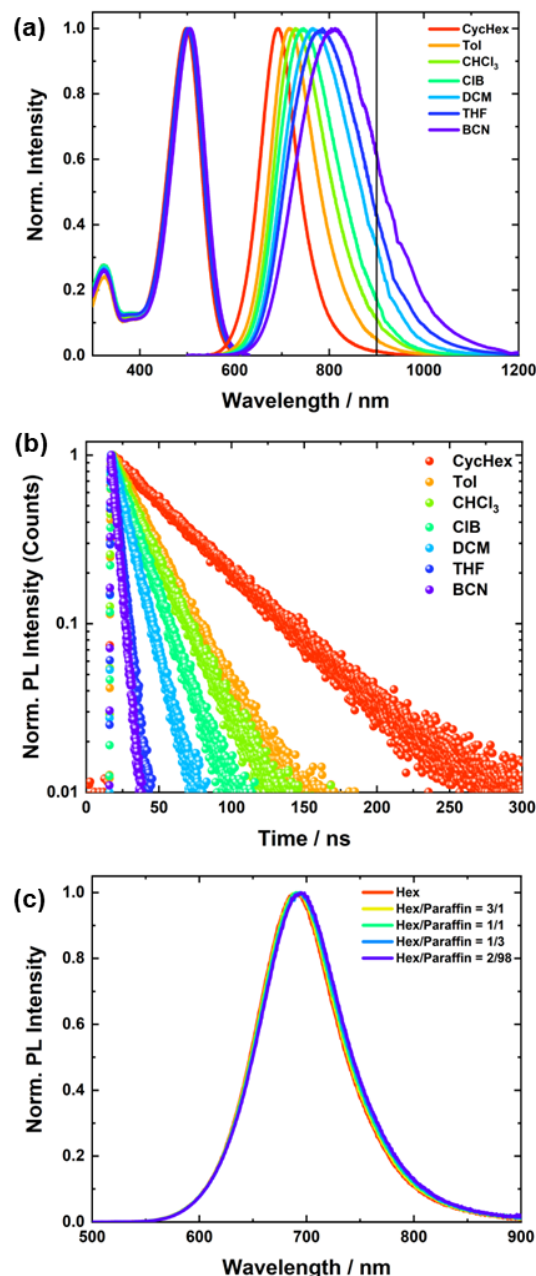


Figure 6. a) Normalized absorption and emission spectra of **TTH** in different solvents; b) normalized TCSPC decay profiles of **TTH** measured in different solvents; c) Normalized emission spectra of **TTH** in different hexane:paraffin mixtures.

In fact, thin films of **TTH** spin-coated (1000 rpm for 60 s) from a chloroform solution (4 mg/ml), show an emission peaked at 703 nm with a PLQY of 0.19 (detail in SI), a value comparable with the state of the art for deep-red/NIR emitters, which makes **TTH** a promising fluorophore for bioimaging and OLEDs.^{66,67} Despite the structural analogies between **TTH** and **TTM**, no excimer emission was detected in the case of the diradicaloid even when processed as thin film. The possibility to obtain long-wavelength emission without an extended conjugated system, and a remarkable solvatochromism from a non-polar centrosymmetric derivative, opens to new strategies for the molecular design of fluorophores in the NIR region. In fact, similar properties were observed in the case of tetraaryl *p*QDM derivatives having a strong push-pull character.⁴⁴

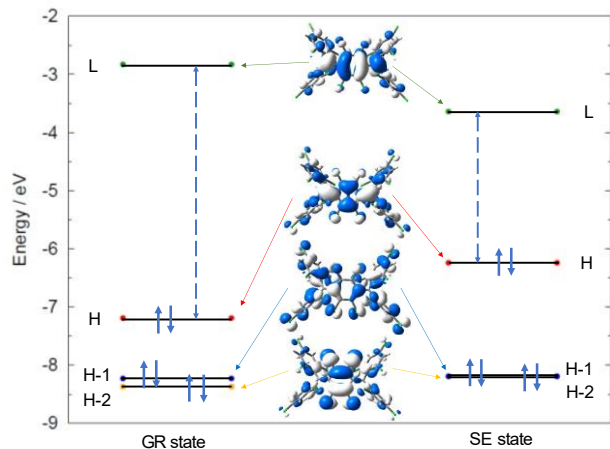


Figure 7. Relevant frontier molecular orbital energies and shapes of **TTH** (M06-2X/def2-SVP +D3 level) at the ground (GR) and singly excited (SE) state optimized geometries showing the remarkable reduction of H/L gap at the excited state geometry, implying an energy decrease of the doubly excited configuration and consequently of the DE state.

Since **TTH** lacks any push-pull feature that might simply explain its emission properties, quantum-chemical calculations aiming to provide new insights about the physical nature of its emitting state, were performed.

Quantum-chemical Calculations. It is well known that in diradicaloids, beside the excited state dominated by the singly excited $H \rightarrow L$ configuration (SE state, bright), one distinctive signature is the presence of a “dark” low-lying excited singlet state dominated by the doubly excited ($H,H \rightarrow L,L$) electronic configuration (DE state)⁶⁸, featuring an ionic (or zwitterionic)

character similarly to the SE state.^{69,70} Such state, described as two local triplet states coupled as an overall singlet in polyenes,^{71,72} has been shown to be the lowest excited state of several conjugated systems displaying a medium to large diradical character^{73–75} and can be expected to be a low-lying state also for **TTH**. While the SE state can be correctly captured by TDDFT calculations and accounts for the main band in the absorption spectrum (**Figure S13**), the DE state demands for highly correlated wavefunction methods.⁷³ Notably, the equilibrium structure of the SE state is markedly diradicaloid, as demonstrated by the elongated CC exocyclic bonds, and by phenyl twisting angles almost identical to those of **TTM** (**Figure S14**). Furthermore, the H/L gap is strongly reduced compared to the ground state geometry (dashed double-arrows in **Figure 7**), implying an increasing weight of the $H,H \rightarrow L,L$ excitation in both ground and excited states as documented by DFT-MRCI and CASSCF calculations (additional details in the SI). Indeed, at the SE optimized geometry there is a dramatic difference between the low-lying excited states predicted at TD-DFT and CASSCF/NEVPT2 level (**Figure 8**), with the latter showing a low-lying DE excited state just above the SE (note that the DE state becomes the lowest according to CASPT2 calculations (**Figure S15**)). Interestingly, the optimized geometry of the DE state displays even more remarkably elongated exocyclic CC bonds than the SE state (**Figure S16**), implying a small barrier for twisting. To assess the role of twisting around such CC bonds, CASSCF/NEVPT2 calculations were carried out on a smaller *p*QDM derivative, the fully-fluorinated tetracyano-quinodimethane **F4TCNQ** (**Figure S17**) featuring a similar low-lying DE state and

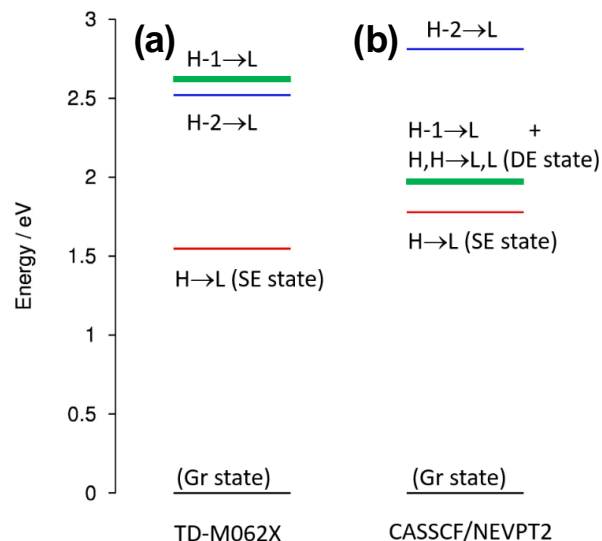


Figure 8. Comparison between low-lying excited states of **TTH** predicted at the equilibrium structure of the SE state: a) TD-M06-2X/def2-SVP calculations; b) CASSCF/NEVPT2 calculations. The thick green line is the “dark” state strongly affected by electron correlation.

documented luminescence properties (maximum value of PLQY = 0.04 in cyclohexane).⁷⁶ Twisting around one exocyclic CC bond of **F4TCNQ** generates asymmetric excited conformers featuring non-negligible dipole moments as a result of sudden polarization,^{77–79} leading to a solvent-sensitive dipolar structure with red-shifted emission driven by the mixing of the DE and SE ionic states. Calculations show that such twisted excited state may be preferentially stabilized in polar solvents (**Figure S17**). At the same time, twisting and SE/DE mixing, confers an increased transition dipole moment, thus favoring radiative decay and accounting for the observed luminescence properties. In contrast to the excited state, the ground state is remarkably destabilized for increasing torsional angles, thereby red-shifting the emission and increasing the Stokes shift. These evidences, applied to **TTH**, suggest that, after relaxation to the SE state *via* planarization of the *p*QDM core (**Figure 9a**) and fast internal conversion to the lower lying DE state, the long exocyclic CC bond length afforded in the ionic DE state, along with its proximity to the SE state, assist the intra- molecular torsion leading ultimately to a charge separated twisted excited state whose stabilization will be preferential in polar solvents (**Figure 9b**). Such twisted, charge separated (zwitterionic) excited states are well known in olefins and justified by sudden polarization,^{77–79} a mechanism explaining, in this case, the observed solvent dependent Stokes shift of **TTH**. Thus, the extreme elongation of the exocyclic CC bonds in the excited DE state of **TTH**, rather unusual for other more extended *p*QDM diradicaloids, makes it comparable to short olefins. While Herzberg-Teller mechanism has been invoked to explain photoluminescence from the doubly excited “dark” state corresponding to the entangled triplet pair ¹(TT) that is the immediate product of singlet fission in oligoacene films⁸⁰ (other conjugates systems displaying some diradical character), here we propose that luminescence from the formally “dark” state of **TTH** diradicaloid is driven by twisting-induced mixing of SE and DE states.

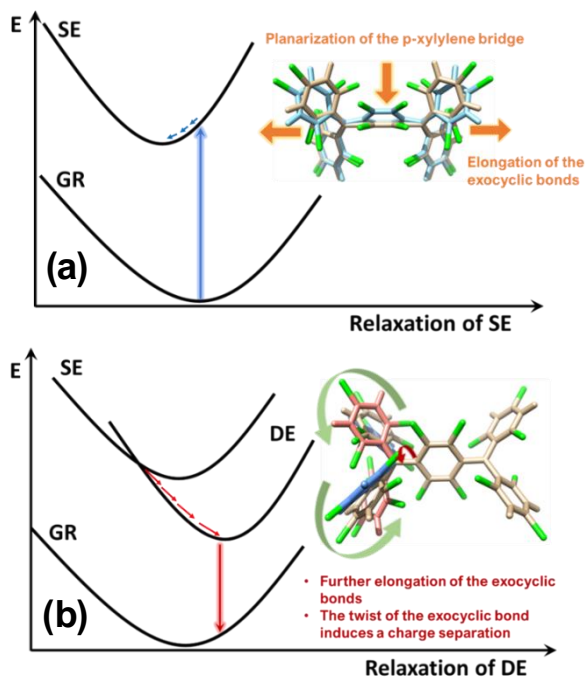


Figure 9. Schematic 2D representation of the potential energy surfaces (PES) of the relevant electronic states of **TTH** involved in the photoinduced processes as inferred from CASSCF/NEVPT2 calculations. (a) main relaxation coordinate in the SE state involving planarization of the *p*QDM core; (b) subsequent internal conversion to the DE state and relaxation along the CC exocyclic twisting coordinate (red and green arrows in the molecular structure on the right)

Transient absorption (TA) spectroscopy. We elaborated on the existence and role of the excited DE state by performing fs and ns TA spectroscopy by photoexciting close to the absorption band edge at 550 nm. Our fs TA experiments unambiguously show the involvement of two excited states in the photophysical deactivation cascade (see **Figures S18-S24**). fs TA experiments (**Figure 10**) reveal the primarily populated bright singlet excited (SE) state. The spectroscopic signatures thereof include excited state absorption (ESA) maxima at around 580 nm and in the range from 1000 to 1150 nm. In addition, ground state bleaching (GSB) at ca. 505 nm and stimulated emission centered around 700 nm were noted. A difference in solvent polarity exerts only a subtle impact on ESA, GSB, and stimulated emission. All of them follow the trend seen in steady-state absorption maxima. Solely, the 1000 to 1150 nm ESA displays an opposing trend. In stark contrast, the subsequently formed lower-lying excited state signature, which we assign to the dark DE state, reveals a strong solvent polarity dependence. On one hand, DE exhibits similar solvent-insensitive TA characteristics seen for SE. This speaks for a twisting-induced state mixing between SE and DE and reflects their singlet multiplicities. On the other hand, two additional ESAs evolve at 700 and 1400 nm upon changing the solvent environment. None of them were present in the TA spectra of SE. Spectro-electrochemical absorption experiments (**Figure S25**) reveal absorptions, which are due to the one-electron reduced or oxidized form of TTH, which are in excellent agreement with these ESAs. This observation supports the notion of a twisted DE with charge separated character. Stabilization thereof in polar media is derived from an intensity gain of the corresponding ESA. Interestingly, the internal conversion, by which SE transforms into DE and are found to depend on the solvent viscosity rather than solvent polarity. The corresponding SE lifetimes are obtained from a sequential global fitting procedure and ranged

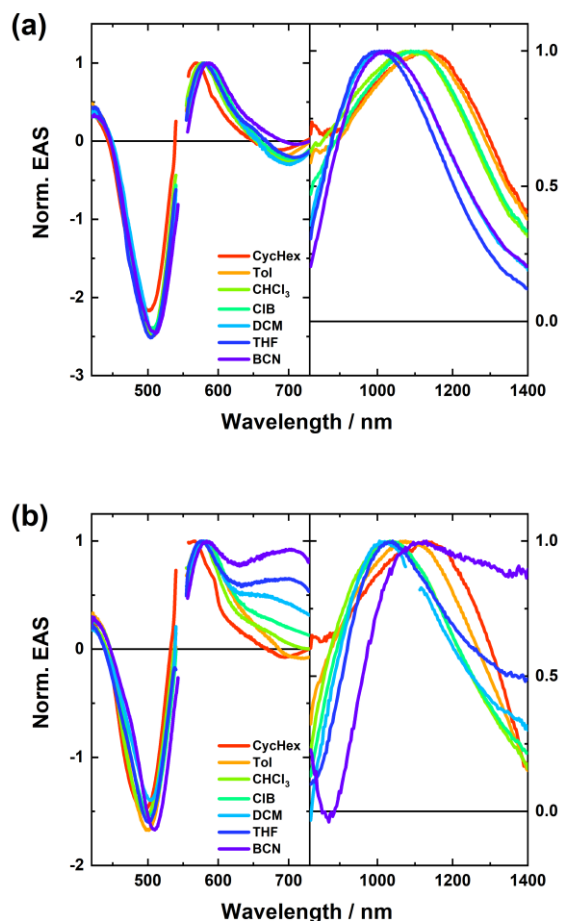


Figure 10. Normalized evolution associated spectra (EAS) of the deconvoluted species obtained via sequential global fitting of the fs TA raw data measured in different solvents upon photoexcitation at 550 nm. (a) EAS of the excited SE state; (b) EAS of the relaxed excited DE state.

from 2.0 ps for DCM to 8.4 ps for BCN (a summary of all lifetimes is found in **Table S9** in the SI). An internal conversion that is viscosity dependent emphasizes the predicted planarization of the *p*QDM bridge and the simultaneous elongation of the exocyclic CC bonds, which goes ahead of the interstate transition. This is succeeded by a rather slow relaxation along the potential energy surface to reach the minimum of the potential energy surface together with a reorganization of the solvent shell. Lifetimes in the range of 100 to 300 ps infer a dependence on solvent polarity and viscosity. This is in line with the quantum-mechanically predicted structural rearrangements along the CC exocyclic twisting coordinate which accompany the transformation towards the relaxed zwitterionic DE (*vide supra*). Its decay exceeds, however, the time-window of fs-TA experiments and, in turn, necessitates ns TA experiments to record the full deactivation dynamics (**Figures S26-S33**). Importantly, the zwitterionic DE decays monoexponentially on the μ s timescale to recover the ground state directly, that is, without the involvement of any other electronic state. The underlying lifetimes show an excellent agreement with the TCSPC results. In other words, the zwitterionic DE is solvatochromatic emissive. By virtue of solvent stabilization, the energy gap with the electronic ground state as a reference point is reduced. Immediate consequences are steadily shorter lifetimes and lower PLQYs as the solvent polarity is reduced all the way to benzonitrile.

Temperature dependent spectroscopy. To gather additional evidence for a mechanism that is based on twisting zwitterionic DE states, temperature dependent absorption and fluorescence experiments were performed in 2-methyltetrahydrofuran. With respect to steady state absorptions of **TTH** (Figure S34), a decrease in temperature down to 80 K induces only subtle shifts. Here, a redistribution of vibrational and rotational energy levels is predominantly operative. This is, however, in stark contrast to the steady state fluorescence (Figure 11a). When going from room temperature to 160 K, the 775 nm fluorescence maximum shifts bathochromically. Internal motions, which are restricted by the low temperatures, aid in the stabilization of the twisted DE state. Going beyond 160 K, that is, to 80 K, the trend is reverted and the maximum shifts hypsochromically all the way to 622 nm. Now, even twisting around the elongated exocyclic CC bonds is inhibited. It restricts the formation of a transition dipole moment as a consequence of sudden polarization. Time-resolved fluorescence spectroscopy in the ns regime (Figures 11b and 11c) independently confirms the aforementioned. At 100 K, a shift of the fluorescence maximum from 630 to 695 nm takes place over the course of excited state deactivation. A global fit of the data yields three lifetimes: 0.4, 1.5, and 6.5 ns. These results reflect the internal conversion from SE to DE as well as structural relaxations. A considerable change is noted at 80 K. As a matter of fact, only a minor bathochromic shift from 595 to 600 nm and two rather than three lifetimes are derived: 2.7 and 11.0 ns. Now, restrictions of internal motions prevent twisting the exocyclic bonds. This renders a transition towards the zwitterionic DE impossible and the fluorescence stems exclusively from the unrelaxed/relaxed SE.

CONCLUSIONS

In this work, a comprehensive investigation encompassing synthesis, diverse spectroscopic characterizations, and computational investigations have been used to describe and model, for the first time, the photoinduced processes leading to the luminescence of **TTH**, a partially chlorinated derivative of the Thiele Hydrocarbon. From the synthetic point of view, we demonstrated that polychlorination is a powerful and simple synthetic tool for obtaining neutral, photostable and inert *p*QDM diradicaloids. Interestingly, **TTH** showed a long-wavelength emission in the deep-red/NIR region without an extended π -system and push-pull character. In addition, extremely high values of PLQY were observed, which is quite uncommon for *p*QDM derivatives, especially considering its solid-state emitting behavior. The large Stokes shift and remarkable solvatochromism of **TTH** luminescence was interpreted with the help of correlated multi-reference quantum-chemical calculations indicating that the low-lying DE ionic “dark” state, typically found at low excitation energies in diradicaloids, is responsible for light emission. In **TTH**, such zwitterionic state becomes the lowest energy excited state and acquires dipole moment and intensity *via* twisting around the elongated exocyclic CC bonds in the excited *p*QDM core, with a mechanism involving mixing with the nearby SE state, similar to sudden polarization occurring in olefins. Such mechanism has been confirmed by fs and ns TA measurements. Because the proximity of the SE and DE states can be tuned by chemical substitution, this study paves the way toward modulation of the luminescence properties in diradicaloids by chemical design,¹⁶ making polyhalogenated *p*QDMs extremely interesting open-shell species for photonics and optoelectronics. Interestingly, no solvatochromism, no neat film emission, and lower PLQYs have been detected in the case of a more extended polyhalogenated Müller’s hydrocarbon, clearly showing the relevance of the moderate radical character in design *p*QDM fluorophores.⁸¹

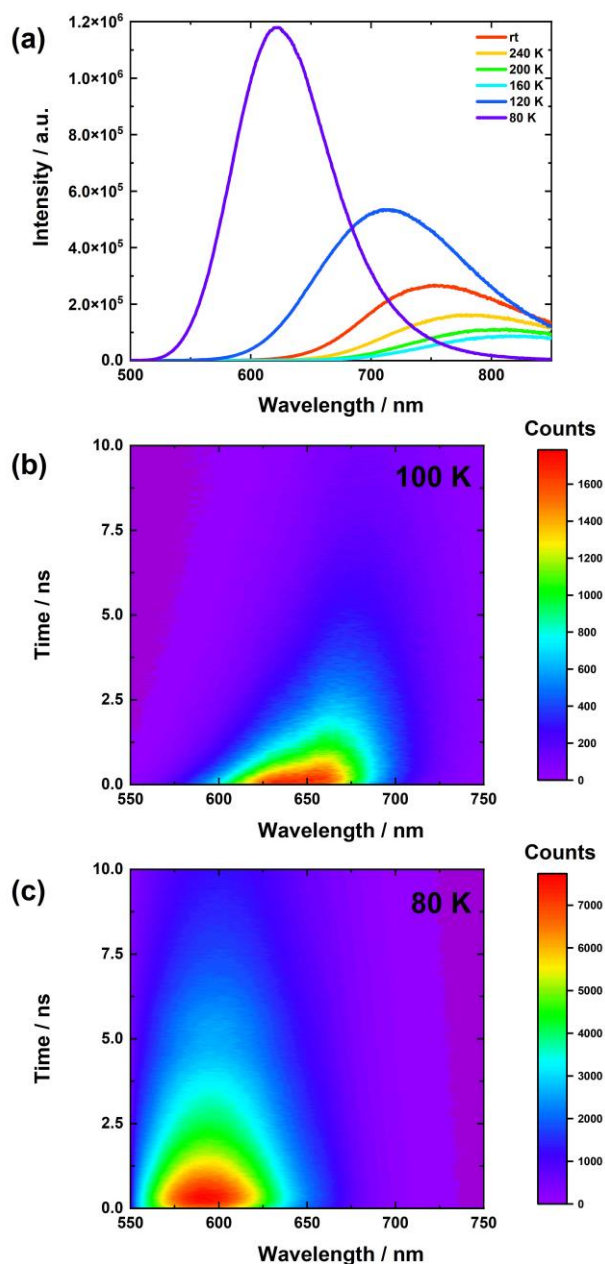


Figure 11. a) Steady state fluorescence spectra of **TTH** in 2-methyltetrahydrofuran obtained at various temperatures; b) ns time-resolved fluorescence plot of **TTH** obtained at 100 K; c) ns time-resolved fluorescence plot of **TTH** obtained at 80 K.

In addition to their exceptional and unique luminescence properties, **TTH** exhibits four reversible redox couples, which make it suitable for energy storage and electronic applications, or even as photo-electrocatalytic species for organic synthesis and biomass valorization.^{82–84}

ASSOCIATED CONTENT

Supporting Information. Additional experimental details, material and methods, synthetic, spectroscopic, crystallographic and computational details, NMR spectra for all compounds, TGA data, TCSPC fittings, fs and ns TA spectra in different solvents and

temperature dependent absorption spectra. This material is available free of charge via the Internet at <http://pubs.acs.org>.”

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Author Contributions

The authors declare no competing interests.

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SYNOPSIS TOC: Intense solvatochromic emission has been observed from an inert Thiele Hydrocarbon, both in solution and as neat thin film. Quantum-chemical calculations and transient absorption measurements show that this emission arises from a low-lying doubly-excited “dark” zwitterionic state, ascribable to the diradical character of the species. In addition, this derivative shows an impressive photo- and electrochemical stability, which is uncommon for such diradicaloid systems.

