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# Linear and nonlinear optical properties of a quadrupolar *carbo*-benzene and its benzenic parent: The *carbo*-merization effect

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## ABSTRACT

Herein, the optical properties of thiophene-functionalized quadrupolar carbo-benzenes and a benzenic parent, of generic structure Th-C=C-[core]-C=C-Th, Th =  $R_2C_4HS$ , are comparatively investigated. Beyond the previously unknown dioctylthienylethynylbenzene (core  $= p-C_6H_4$ , R = nOct), two bis-dialkylthienylethynyl-carbobenzenes (core =  $C_{18}$ Ph<sub>4</sub>, R = nOct, nBu) are envisaged for the unique "carbo-aromatic" character of the  $C_{18}$ macrocycle. The three targets were synthesized from the corresponding ethynylthiophenes in 47, 20 and 10% yield, respectively, then characterized by classical methods such as NMR spectroscopy, and X-ray crystallography for one of the carbo-benzenes. Regarding linear and nonlinear optical properties, our results show that the carbomerization induces a significant shift to lower energies of the one-photon electronic excitations accompanied by an 8-fold increase of the molar extinction coefficient compared to the parent molecule. Intriguingly, these excitations lead to a broad band of photoluminescence comprising decay transitions of the type  $S_1 \rightarrow S_0$  but also of the type  $S_2 \rightarrow S_0$ . This phenomenon of emission from higher excited states, which is contrary to Kasha's rule, is assigned to - or revealed by - a reduction of the internal conversion efficiency between  $S_2$  and  $S_1$ . Two-photon induced transitions are also enhanced, the two-photon absorption cross-section ( $\sigma_{2PA}$ ) being in average five times larger for the carbo-benzenes than for their benzene parent in the wavelength range 650–950 nm, with a maximum of  $\sigma_{2PA} = 1430$  GM (1 GM =  $10^{-50}$  cm<sup>4</sup> s/photon). Beyond a moderate nonlinearity, this comparative study provides quantitative insights about the way *carbo*-merization or insertion of a  $\pi$ -conjugated macrocycle between chromophoric functions (here thiophene rings) can tune optical properties of organic molecules. The optical properties of the bis-dialkylthienylethynyl-carbo-benzenes are also discussed in regard of recent reports on organic chromophores based on other types of  $\pi$ -conjugated macrocyclic cores.

## 1. Introduction

During the last three decades,  $\pi$ -conjugated organic molecules and polymers have emerged as functional materials for diverse applications in the fields of optoelectronics and photonics [1a,1b, 1c]. Typically, the optical and electronic properties of these materials are tailored by

designing molecular architectures with effective  $\pi$ -conjugated backbones, appropriate donor (D) and acceptor (A) electroactive substituents, and auxiliary functions. Classically, the  $\pi$ -conjugation in molecules and polymers is brought by combination of unsaturated linear chains and/or small carbo- or hetero-cycles. Recently, a special type of  $\pi$ -conjugated backbone based on shape-persistent macrocyclic

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architectures have gained significant attention for two-photon absorption (2 PA) or other nonlinear optical effects [2a,2b, 2c,2d], optoelectronics [3a,3b, 3c], near-infrared luminescence [4a,4b], sensing of anions [5], photodynamic therapy [6], redox- or photo-responsive switches [7a,7b] and liquid crystallinity [8]. In such compounds, the number of  $\pi$ -electrons varies with the size of the macrocycle and in some cases intriguing electronic and optical properties are achieved even without the use of electroactive substituents [9]. Over the years, various  $\pi$ -conjugated macrocyclic architectures have been investigated, including macrocyclic annulenes [10], cyclic oligophenylenes and belt-shaped paraphenylenes [11a,11b, 11c]. macrocvclic oligophenylene-ethynylenes [12a,12b, 12c], porphyrins and expanded porphyrins [13a,13b, 13c], and cyclic oligothiophenes [14]. More macrocyclic architectures recently, novel exhibiting aggregation-induced emission (AIE) properties have been designed [15]. Oligofuran-based macrocycles [16], polyradicaloids [17], hexagonal ring carbo-mers of benzene [18] and many others have been also considered.

The term carbo-mer designates "carbon enriched" molecules formally generated by systematic insertion of C<sub>2</sub> units into each (or a selected part) of the covalent bonds of a representative Lewis structure of any parent molecule [19]. At the Lewis-Gillespie level of representation, the carbo-merization process preserves the topology, symmetry, shape and  $\pi$ -electronic resonance of the parent molecule, while its bonding size is approximately expanded by a factor of three. Ring carbo-mers of benzene, simply called "carbo-benzenes", constitute the most exemplified series of carbo-mers [20a,20b, 20c,20d], and have been studied both theoretically, in particular for their aromaticity [21a,21b, 21c,21d,21e], and experimentally, e. g. for redox [18,22], conductive [23], and mesogenic properties [24]. It was also demonstrated that carbo-benzenes exhibit unique chromophoric [18,25a,25b,25c] and nonlinear optical properties. In particular, the two-photon absorption (2 PA) by the quadrupolar representatives (p-difluorenyl)tetraphenyl-carbo-benzene 1a and (p-difluorenylethynyl)tetraphenyl-carbo-benzene 1b (Fig. 1) was recently evaluated, with a 2 PA cross-section  $\sigma_{2PA} = 656$  GM at 800 nm for 1b [26]. This first attempt to assess the optical response of carbo-benzenes at infrared wavelengths showed their relevance as nonlinear materials. In this context, the understanding of the structure-optical properties relationships in *carbo*-mer series is still limited and awaits further investigation.

In this work, the effects of carbo-merization are evaluated through the comparison of the optical properties of the quadrupolar carbo-benzenes 2a,b with those of the benzenic parent molecule 2P, in which the central  $C_{18}$  macrocycle is replaced by a  $C_6$  benzene ring (Fig. 1). In these molecules, the thienyl substituents, largely used in the design of 2 PA chromophores, behave as electronic donors that are anchored to the C<sub>18</sub> macrocycle through an acetylenic linker. Dialkylthienylethynyl substituents were selected for two reasons: i) two alkyl chains are attached to the thiophene ring with the view of enhancing the solubility of the targeted carbo-benzenes 2a,b to allow sufficiently high concentrations for the measurement of  $\sigma_{\rm 2PA}$  by the Z-scan technique (*ca*  $10^{-3}$  -  $10^{-2}$  mol  $L^{-1}$ ), and ii) the ethynyl linkers serve to both facilitate the synthesis key step, and increase the length of the  $\pi$ -conjugated system, thus a priori favoring high 2 PA efficiency, as previously evidenced by the comparison between 1a and 1b (Fig. 1). The linear photophysical properties (UV-vis absorption, emission, excitation) and nonlinear optical properties (2 PA) of the core-carbo-mers 2a and 2P are here reported in detail.

## 2. Results and discussion

## 2.1. Synthesis

The synthesis of the bis-dialkylthiophenylethynyl-*carbo*-benzenes **2a,b** required the preparation of known key precursors, namely the [6] pericyclynedione **3** [27a,27b, 27c], a classical intermediate in the synthesis of most quadrupolar *carbo*-benzenes [20], and the dialkylthienylacetylenes **4a,b**.

Though a synthesis of **4b** was previously published [28], **4a** and **4b** were not prepared following exactly the reported methodology. First, the described nickel-catalyzed Kumada coupling of commercial 3, 4-dibromothiophene with octyl- or butyl-magnesium bromide led to mixtures of dialkylthiophenes **5a**,**b** and monoalkylthiophenes **5a**',**b**' in a 95:5 ratio (Scheme 1). In both cases, the separation of the two products by silica gel column chromatography appeared problematic because of their similar low polarity. The direct formylation of the mixtures **5a**,**b** 



Fig. 1. Quadrupolar *carbo*-benzenes 1a and 1b previously studied [26] for their 2 PA properties (*top*); bis-dithientlethynyl-*carbo*-benzenes 2a,b and parent molecule 2P considered in this work (*bottom*).



Scheme 1. Synthesis of the dialkylthinylethynylacetylenes 4a,b.

and 5a',b' leading to inseparable mixtures of aldehydes 6a,b and 6a',b', the preparation of more polar intermediates was envisioned with the view to facilitating removal of the monoalkyl-thiophene contaminants. The mixtures of 5a,b and 5a',b' were thus treated with one equivalent of *n*-butyllithium, followed by addition of paraformaldehyde, to give mixtures of the alcohols 7a,b and 7a',b' which could then be separated by silica gel column chromatography. The pure alcohols 7a and 7b, isolated with 83% and 95% yield, respectively, were then oxidized by treatment with MnO<sub>2</sub> to give the corresponding aldehydes 6a and 6b almost quantitatively. Finally, the aldehydes were converted to the terminal alkynes **4a** and **4b** by a Corey-Fuchs reaction, in two steps *via* the dibrominated intermediates **8a,b** [29].

The *carbo*-benzenes 2a,b were then prepared from 3 and 4a,b by using the classical two-step process developed for the synthesis of other quadrupolar *carbo*-benzenes [20]. First, two equivalents of the lithium dialkylthienylacetylide of 4a or 4b, deprotonated with *n*-butyllithium, were added to the diketone 3, leading to the pericyclynediols 9a,b bearing the two thienylethynyl substituents. The last reductive



Scheme 2. Two-step synthesis of the carbo-benzenes 2a,b from the known [6]pericyclynedione 3.

aromatization step was performed with  $SnCl_2$  alone to avoid the reduction and hydrochlorination of the exocyclic triple bonds, affording the *carbo*-benzenes **2a** and **2b** in 35% and 20% yield respectively (Scheme 2). In the presence of  $SnCl_2$  and HCl, an inseparable mixture of chromophoric products was formed from **9a**, among which **2a'** could be identified by X-ray diffraction analysis (see SI, Figure S1). Such a hydrochlorination side-reaction was previously observed for [6]pericyclynediols bearing indolylethynyl donor groups, and its occurrence here with donor thienylethynyl groups is thus consistent [25c]. The unexpected reduction of the other triple bond to a *trans*-substituted double bond can be compared to a previously reported  $SnCl_2$ -mediated reduction of a *spC-spC* bond of a *carbo*-meric C<sub>18</sub> ring to a *trans*-CH=CH bond [18].

The *carbo*-chromophores **2a**,**b** were isolated as violet solids, soluble in most of the conventional organic solvents while giving intense violet solutions. **2a** was characterized by NMR and UV–vis spectroscopy, and by mass spectrometry. Its structure was also determined by X-ray diffraction analysis of a single crystal deposited upon slow evaporation of a dichloromethane solution at room temperature (Fig. 2 and Section 2.2) [30].

The parent molecule **2P** was prepared by a Sonogashira coupling of two equivalents of **4a** with diiodobenzene, and isolated with 47% yield (Scheme 3).

## 2.2. X-ray crystallography

In the X-ray crystal structure of **2a** [30], the macrocycle is quasi planar, the maximum deviation from planarity being of 0.02 Å, with almost coplanar phenyl substituents (maximal torsion angle of 12.9°). The shape of the  $C_{18}$  ring is however quite distorted from a regular hexagon, the three  $sp^2C \dots sp^2C$  diagonal distances varying from 7.69 to 7.87 and 8.33 Å, while the internal bond angles vary from 116.9 to 121.8°, with two concave PhC<sup>...</sup>C·..·C(C<sub>2</sub>Th) edges.

In the crystal packing, the molecules of **2a** are shifted one from the other, but the mean planes of two successive macrocycles are parallel, albeit without normal overlapping between them. The distance between the mean planes of two successive molecules is of 3.26 Å. As one of the octyl groups is bent, some H atoms of the chain point towards the center of the closest *carbo*-benzene ring, the shortest distance measured between the centroid of the  $C_{18}$  ring and a H atom of the octyl chain being of 2.21 Å.

## 2.3. Electrochemistry

The electrochemical properties of the *carbo*-benzene **2b** were investigated by square-wave (SWV) and cyclic voltammetry (CV; Table 1). Four reduction and three oxidation waves were observed by SWV, the two first reductions at -0.62 and -1.02 V/SCE being reversible.

The first reduction is observed at a quite low potential (in absolute value), which is consistent with the extended  $\pi$ -conjugated system of **2b** *a priori* associated with a low LUMO level, and similar to the corresponding potential previously reported for **1b** (see Fig. 1) [26].

The first oxidation, observed at 1.13 V/SCE, is a non-reversible process, as observed for 1b, which is first oxidized irreversibly at 1.17 V/SCE.

## 2.4. Optical properties

## 2.4.1. Linear optical properties

Effects of *carbo*-merization on optical properties can be evaluated by comparing the photoinduced electronic transitions of **2P** and **2a** (Fig. 3). While 2P exhibits an ordinary absorption band with a peak at 350 nm assigned to a  $\pi \rightarrow \pi^*$  transition, **2a** presents the characteristic features of carbo-benzenes [18], with a sharp and intense band, in this case at 495 nm (for the simplicity of discussion this main band is denoted as M in Fig. 3), a shoulder at 542 nm, and a secondary band with a peak at 608 nm (denoted as S band). The effect of dioctylthienylethynyl substitution at the para positions of the C<sub>18</sub> macrocycle (C1 and C10 positions in 2a) is clearly distinguishable by comparison with the absorption spectrum of the previously reported para-dialkyl-substituted analogue 1c (Supplementary Information, Figure S2) with a much shorter  $\pi$ -conjugation extent [31]. The overall effect of substitution of the carbo-benzene ring with thiophenylethynyl units produces a bathochromic shift of ca 50 nm of both M and S bands as compared to 1c accompanied by a moderate spectral broadening of the M band and vanishing of minor peaks between the M and S bands. The peak of maximum absorption for 2a is practically at the same wavelength (495 nm) as the one observed for the fluorenylethynyl-substituted carbo-benzene 1b [26] but with a ca 30% reduction of the molar extinction coefficient  $\varepsilon$ . Nevertheless,  $\varepsilon$  increases from 2.8  $\times$  10  $^4$   $M^{-1}$  cm  $^{-1}$  in 2P to 2.25  $\times$  10  $^5$   $M^{-1}$  cm  $^{-1}$  in 2a due to the expansion of the  $\pi$ -conjugation. For organic molecules in general, the magnitude of  $\varepsilon$  is a signature of their polarizability at optical frequencies, which correlates with the degree of  $\pi$ -conjugation within the backbone and with substituents. For reference, Table 2 presents the



Fig. 2. Molecular view of the X-ray crystal structure of 2a (H atoms were omitted for clarity). See note [30], CCDC 2027203.



Scheme 3. Synthesis of the parent molecule 2P of 2a.

Table 1
SWV and CV data for the <i>carbo</i> -benzene <b>2b</b> .

Reduction			Oxidation
$E_{1/2}^{a,b} (\Delta E_p)$	<i>RI</i> p <sup>c</sup>	$E_{\rm p}^{\rm redd}$	$E_{\rm p}^{\rm oxd}$
- 0.62 (69)	0.99	- 1.52	1.13
- 1.02 (59)	0.76	- 1.74	1.75
			1.89

<sup>a</sup> Measurements performed at room temperature in DCM; supporting electrolyte:  $[nBu_4N][PF_6]$  (0.1 M); working electrode: Pt; reference electrode: saturated calomel electrode (SCE; 0.242 V *versus* the hydrogen electrode); scan rate: 0.2 V s<sup>-1</sup>, unless otherwise noted.

<sup>b</sup> Half-wave potential,  $E_{1/2} = (E_p^{\text{red}} + E_p^{\text{ox}})/2$ , in V/SCE.

<sup>c</sup> Peak current ratio  $RI_p = |I_p^{ox}/I_p^{red}|$ .

<sup>d</sup>  $E_p$  values measured from CV in V/SCE.

optical properties and the molecular structure of some representative  $\pi$ -conjugated macrocycles. In these examples the number of  $\pi$ -electrons (*m*) in a conjugated circuit inscribed in the corresponding macrocycle varies from 18 to 180, producing molar extinction coefficients in the range  $0.2-6.39 \times 10^5 \,\mathrm{M^{-1}\ cm^{-1}}$ . In spite of a relatively low out-of-plane  $\pi_{\rm xy}$  electron count m = 18 (unconjugated with the 12 in-plane  $\pi_{\rm xy}$  electrons of the 6 *spC-spC* bonds), the  $\varepsilon$  values of *carbo*-benzenes are thus among the largest. According to the parity of m/2, the macrocycles of Table 2 can be qualitatively identified as aromatic *vs* antiaromatic using the well-known m = 4n + 2 *vs* 4n Hückel rule.

It should be noted that characteristic features of the absorption spectra of *carbo*-benzenes, i.e., a sharp and intense **M** band followed by a secondary **S** band of weaker intensity, are not observed in other families of  $\pi$ -conjugated macrocycles (as those exemplified in Table 2) since they generally exhibit broad and structureless absorptions, except in the porphyrinoids whose spectra present the well-known Soret (B Band) and Q bands [(13a–c),45]. The main band of the absorption spectra of substituted *C*<sub>2</sub>-symmetrical *carbo*-benzenes was previously interpreted [26] as additive and subtractive combinations of excitations from the two highest occupied molecular orbitals (HOMO-1, HOMO) to the two lowest unoccupied molecular orbitals (LUMO, LUMO+1), as schematically shown in Fig. 4. This approach invokes the Gouterman four-orbital model developed to explain the origin of the Soret and Q bands in the absorption spectra of porphyrins [45].

Fig. 3 presents the photoluminescence (PL) spectra of **2P** and **2a** under excitation at their main absorptions, 350 and 497 nm, respectively. The PL spectrum of the parent molecule **2P** exhibits a band with a relatively small Stokes shift of 36 nm and noticeable vibronic structure. In the case of the *carbo*-benzene **2a**, the broad PL band is distorted due to absorption inner effects from the **S** band. Inner filter effects occurred inevitably in all the PL measurements of **2a** given its high  $\varepsilon$  and poor emission properties. Nevertheless, PL characterization of **2a** is worthwhile since it can provide information on the excited states that lead to radiative decays in *carbo*-benzenes. For example, radiative decays occur not only at longer wavelengths (lower energies) than the low-lying band **S**, but also at shorter wavelengths (higher energies). The matching between the PL excitation spectra and the absorption spectrum shown in



Fig. 3. Absorption, PL (with excitation wavelength  $\lambda_{exc}$ ) and excitation spectra (with detection of PL at the wavelength  $\lambda_{det}$ ) obtained for a) **2P** and b) **2a** in THF solutions prepared at  $8.43\times 10^{-5}$  M.

Fig. 3b clearly indicates that the electronic transitions corresponding to the **M** and **S** bands contribute to the PL emission. Furthermore, significant PL still occurs at wavelengths with small shifts with respect to the excitation, suggesting small geometric changes between the excited and ground states. This is not surprising as we consider the planar and quite rigid monocyclic *carbo*-benzene ring, identified previously as both aromatic and flexible ("*carbo*-aromatic") [20c]. Small Stokes shifts are also observed in other planar conjugated systems such as the more rigid pentacyclic porphyrins and expanded porphyrins with residual

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#### Table 2

Comparative properties for various  $\pi$ -conjugated macrocycles. The molecular structure of each of these macrocycles is presented in Table S1.

π-conjugated macrocycle	$\lambda_{\rm abs}~({\rm nm})$	$\lambda_{\rm em}$ (nm)	$\varepsilon \times 10^5  (\mathrm{M}^{-1}  \mathrm{cm}^{-1})$	Conjugated $\pi$ -electron count m and Hückel aromatic character <sup>a</sup>	$\sigma_{2PA (GM)}$	$\lambda_{\max}(2PA)$	Ref.
					$\sigma_{2PA}/MW$	()	
Thienylene-phenanthrylene	385	495	1.51	56 antiaromatic	-	-	[32]
Tetraphenylethylene based	330	520	-	non-aromatic	-	-	[15]
Oligofuran based	401	530	1.5	32 antiaromatic	-	-	[16]
BODIPY-phenylacetylene	364	527	0.26	32 antiaromatic	-	_	[33]
Phenanthrylene-ethynylidene	462	437	-	non-aromatic	-	-	[34]
Expanded 1,3-Dithiolane [5]radialene	403		1.2	non-aromatic	_	_	[35]
Triphenylamine phenylacetylene	380	430	-	non-aromatic	1300	650	[36]
					19.7		
					0.81		
Triangle-shaped annulene	332	389	-	18 aromatic	11.5	372	[37]
					0.64		
					0.031		
Triangular –[D–π–A] <sub>3</sub> –	375	469	2.0	66 aromatic	2800	750	[38]
					42.42		
					0.85		
Phenylacetylene-based macrocycle	413	486	1.1	non-aromatic	125	820	[39]
					4.2		
					0.069		
Giant expanded oligothiophenes Thiophene [30]mer	480	559	6.18	180 antiaromatic	107 800	710	[14,40]
					598		
					16.30		
Zinc(II) porphyrin	471	_	-	18 aromatic	3520	1360	[41]
					195		
					2.66		
Hydrocarbon-substituted metal-free porphyrins	440	660	-	18 aromatic	42 600	800	[42]
					2366		
					38.33		
Pentapyrrolic expanded porphyrin-isosmaragdyrin	712	716	-	22 aromatic	2900	1280	[43]
					131.8		
					4.81		
Cyclo [8]pyrrole expanded porphyrin	427	1496	1.2	30 aromatic	3030	1600	[44a, 44b]
					101		
					3.12		
[26]Hexaphyrin expanded porphyrin	567	1036	4.2	26 aromatic	9890	1200	[44a, 44b]
					380		
					6.77		
Quadrupolar carbo-benzene 2a	496		2.25	18 aromatic	1430	650	This work
					83.3		
					2.12		

<sup>a</sup> Number *m* of conjugated  $\pi$ -electrons inscribed exclusively in the circuit of the corresponding macrocycle.

flexibility [44]. Unexpectedly, the observation that PL can occur directly from the excitation of the **M** band is in contraposition with Kasha's rule [46] since it is assumed that such excitation leads to the quasi-degenerated exited state  $S_2$  (see Fig. 4 for the Jablonsky diagram proposed for 2a in analogy with the Gouterman four-orbital model). PL from higher excited states  $(S_2 \rightarrow S_0)$  is a quite rare phenomenon that mainly occurs in small organic compounds [47] although it has also been reported in macrocycles such as [18]annulene C<sub>18</sub>H<sub>18</sub> [48], akin to carbo-benzene also identifiable as the most symmetric dodecadehydro-[18] annulene C<sub>18</sub>H<sub>6</sub>, BODYPI-based macrocycles [33], metalloporphyrins [49a,49b], and theoretically predicted in cyclo[n]thiophenes [50]. Anomalous emission had been also already observed for a dichromic carbo-cyclohexadiene [18] having a green PL band appearing between its blue and red absorption bands. However, this PL band was not explicitly identified as a radiative decay of the type  $S_2 \rightarrow S_0$ . The accepted interpretation for anomalous PL in organic molecules is that it can occur when the energy gap  $\Delta E(S_2 - S_1)$  is sufficiently large to reduce the Franck-Condon factor [51], the latter expressing the overlap integral of vibrational wavefunctions involved in the internal conversion (IC) process between  $S_2$  and  $S_1$ , to a level where the rate of radiative  $S_2 \rightarrow S_0$ transitions competes with the IC rate. Azulene, a polar hydrocarbon molecule, is a well-known prototype that exhibits efficient PL from  $S_2$ due to a very large  $\Delta E(S_2 - S_1) \sim 10\ 000\ \mathrm{cm}^{-1}$  [47]. The carbo-chromophore **2a** also exhibits significant  $\Delta E(S_2 - S_1) \sim 3673 \text{ cm}^{-1}$ . This value is even larger for the *carbo*-cyclohexadiene reported in Ref. [18],  $\Delta E(S_2 - C_2)$ 

 $S_1$ ) ~ 6272 cm<sup>-1</sup>. This suggests that the fluorescence decay  $S_2 \rightarrow S_0$  in **2a** could be assigned to the relatively wide separation between the **M** and **S** bands, which decreases, to some extent, the IC efficiency between  $S_2$  and  $S_1$ , thus increasing the probability of bypassing  $S_1$  during relaxations. It must be noted that the conformation of the macrocycle determines the density and shape of vibronic levels, and, in turn, the Franck–Condon factor. In consequence, the fluorescence in *carbo*-benzenes originates simultaneously from both the first and second excited states ( $S_1 \rightarrow S_0$  and  $S_2 \rightarrow S_0$ ) as depicted in Fig. 4. All this characterization of the linear optical properties for **2P** and **2a** was obtained in diluted solutions such that self-assembly effects were discarded.

As a consequence of *carbo*-merization, **2a** exhibited significantly longer fluorescence lifetime than its parent molecule **2P**. Fig. 5 presents the PL decays of the two molecules using the technique of timecorrelated single photon counting (TCSPC); in this figure, the dotted lines denote the instrument response, the solid lines the lifetimes of **2P** (black) and **2a** (blue). The average lifetime measured in **2a** was 5.04 ns, while the measurement for **2P** gave a significantly shorter value (1.02 ns), just barely different from the instrument response function. Although radiative decays from  $S_2$  and  $S_1$  should imply differentiated lifetimes, no wavelength dependence of the PL was observed, the resolution of the TCSPC technique probably being insufficient to allow their discrimination. Further investigation using techniques of higher temporal resolution will thus be necessary.





Excitations between pairs of molecular orbitals that produce excited states in *carbo*-benzens

Fig. 4. Jablonsky diagram (left) and Gouterman four-orbital model (right) for 2a.



**Fig. 5.** Lifetime measurements of **2a** (*blue line*) and **2P** (*black line*). As reference, the response function of the instrument (*dotted lines*) is presented. The emission lifetime of **2a** increases 5 times in comparison to the lifetime of **2P**. The lifetime of **2a** was monitored at 550 nm and **2P** at 414 nm. Inset: Plots in semilog scale. (For interpretation of the references to color in this figure legend, the reader is referred to the Web version of this article.)

## 2.4.2. Nonlinear optical properties

Comparison of one- and two-photon excitations of **2P** and **2a** provides insight into the effects of *carbo*-merization on the energy level configuration and nonlinear polarizability. To measure  $\sigma_{2PA}$  in **2P**, the TPEF technique was used since that molecule fluoresces appreciably (quantum efficiency of 22%) and this technique is of higher sensitivity than Z-scan for molecules with small  $\sigma_{2PA}$  values. TPEF spectra of **2P** were recorded from 680 nm to 780 nm in a THF solution (8.43 × 10<sup>-5</sup> M), as shown in Figure S3, and used to calculate the  $\sigma_{2PA}$  spectrum presented in Fig. 6. In the case of **2a**, it was not possible to measure  $\sigma_{2PA}$  through the TPEF technique because of its very weak fluorescence. The  $\sigma_{2PA}$  spectrum of **2a** (THF solution at 8.43 × 10<sup>-2</sup> M) was thus acquired from 650 nm to 950 nm through the Z-scan technique. To ensure a fair

comparison of the nonlinearities obtained for the carbo-meric and parent molecules through different techniques, the  $\sigma_{2PA}$  values of **2P** in solution  $(8.43 \times 10^{-2} \text{ M in THF})$  were also determined by Z-scan at three different wavelengths (black dots in Fig. 6a), showing the agreement between the two techniques. In spite the high concentration of the solutions used in Z-scan experiments, self-assembly effects in 2P and 2a were not detected. For instance, light scattering, an affect that is typically produced in presence of self-assembled molecules, was not observed in the tested solutions. Of course, self-assembly could be present if the molecules are deposited in solid films, but such studies were beyond our present work. Larger nonlinearities are observed in 2a with respect to **2P**, with a *ca* 7-fold increase in  $\sigma_{2PA}$  at the infrared (IR) range of wavelengths (which is, in fact, the region of interest for practical applications of the 2 PA process). We can see that the onset of detectable nonlinear absorption in 2a takes place at a significantly longer wavelength (950 nm, as shown in Figure S4) compared with 2P (780 nm). Out of the IR range, the maximum values of  $\sigma_{2PA}$  were 506 GM for **2P** at 680 nm and 1430 GM for 2a at 650 nm, with a tendency to increase at shorter wavelengths for both molecules. It is also illustrative to evaluate the enhancement of nonlinearities by comparison of the ratio between  $\sigma_{2PA}$  and the number *m* of  $\pi$ -electrons inscribed inclusively in the  $\pi$ -conjugated circuit. For instance, at IR wavelengths we have the relation  $\sigma_{2PA}/m$  (2a) ~ 2 ×  $\sigma_{2PA}/m$  (2P). Thus, the overall linear and nonlinear characterization of 2a shows that carbo-merization, though preserving the shape, symmetry and topology of a parent molecule, can induce enhancements of the polarizability ( $\varepsilon$ ) and hyperpolarizability  $(\sigma_{2PA})$  due to a wider  $\pi$ -conjugation extent and an improvement of the optical effects per  $\pi$ -conjugated electron. Table 2 presents  $\sigma_{2PA}/m$  values for a variety of macrocyclic chromophores. The carbo-benzene 2a shows intermediate nonlinearities as compared to its counterparts. A similar trend is observed when  $\sigma_{2\text{PA}}$  is normalized by the molecular weight. Note that in some series of macrocycles of increasing number m, an enhancement of nonlinearities is observed: giant oligothiophenes with m = 72 and 180 showed  $\sigma_{2PA}/m \sim 209$  and  $\sim 598$  GM, respectively [14]. In line with these observations, it would thus be of interest, at least at the theoretical level, to study the effect of enhancement or saturation of  $\sigma_{2PA}$ in carbo-mers of larger size than 2a, for instance in carbo-benzenes of second generation exhibiting 30 cyclically  $\pi$ -conjugated electrons [20b].

The excited states accessible by two-photon transitions in substituted



**Fig. 6.** a) Spectra of the 2 PA cross section for **2P** and **2a** measured in THF solution by TPEF and Z-scan techniques at the concentration of  $8.43 \times 10^{-5}$  and  $8.43 \times 10^{-2}$  M, respectively. 2 PA cross sections were also measured for **2P** by Z-scan technique is a THF solution at  $8.43 \times 10^{-2}$  M; b) Optical limiting at 800 nm for **2a** emphasizing that, at the relatively low intensity of 20 GW/cm<sup>2</sup>, its transmission diverges from a linear behavior.

carbo-benzenes were examined. In Fig. 6a, the total energy for a pair of infrared photons  $(2\hbar\omega)$  exciting **2a** is higher than the energy corresponding to the one-photon peak at the M band. This blue-shifting of the 2 PA resonance with respect to one-photon resonance is expected in symmetric molecules like 2a because one-photon and two-photon transitions are mutually exclusive according to Laporte's rule [51]. In the four-orbital model used to construct the excited states diagram of 2a, the quasi-degenerate states  $3A_u$  and  $4A_u$  of the  $\boldsymbol{M}$  band have an ungerade parity and are thus two-photon forbidden from the ground state of gerade parity (Ag). A strategy to enhance the nonlinearities at IR wavelengths of equivalent energies to the M band would thus be to lower the symmetry of carbo-benzenes by modifying the positions of the substituents of the C18 hexagonal core, strategy that was successfully applied with other macrocyclic molecules to bring 2 PA states closer in energy to one-photon allowed states [52]. Of course, not only the symmetry of the molecule but also the nature and position of the substituents and  $\pi$  connectors play complementary roles that have to be globally taken into account in the design of new carbo-chromophore targets.

Finally, the 2 PA properties of the *carbo*-benzene **2a** allowed implementation of optical limiting, an application of utility for the protection of sensors, optical elements and human eyes. Measurements were performed at 800 nm with pulses of 80 fs and intensities ranging from 0.1 to 250 GW/cm<sup>2</sup>. Even at energies as low as 20 GW/cm<sup>2</sup>, **2a** begins to deviate from the linear regime getting to a maximum deviation at 60 GW/cm<sup>2</sup> (Fig. 6b). For intensities as high as 250 GW/cm<sup>2</sup> no damage or degradation was detected in the sample, making **2a** appropriate for optical damage control [1c].

#### 3. Conclusions

To get insights into the underlying optical features that emerge in a given  $\pi$ -conjugated molecular topology when a C<sub>6</sub> benzene ring is replaced by a C18 aromatic macrocycle, the one- and two-photonpromoted electronic transitions were characterized in a quadrupolar carbo-benzene, namely 2a, and compared with those of its parent molecule 2P. In passing, but not less importantly, the synthesis and extensive characterization of the previously unknown carbo-benzene 2a, homologue 2b, and parent molecule 2P are disclosed. In the linear regime, a notorious increment in the polarizability was observed, whereas, in the nonlinear regime, a higher 2 PA activity per  $\pi$ -electron was determined. The presence of the hexagonal carbo-benzene ring in the conjugated backbone introduces anomalous PL due to simultaneous radiative decays  $S_1 \rightarrow S_0$  and  $S_2 \rightarrow S_0$ . In view of these results, the C<sub>18</sub>  $\pi$ -conjugated macrocycle can be considered as a versatile platform on which substituents can be attached at various positions in dipolar, quadrupolar or octupolar geometries, the two latter being relevant for potential applications involving 2 PA as an initial process, or possibly for energy transfer from either the  $S_1$  or  $S_2$  excited states. Finally, the results also suggest that lowering the symmetry of substituted carbo-benzenes, through replacement of some peripheral phenyl groups by other substituents, might allow tuning the peak of 2 PA to longer wavelengths.

## 4. Experimental section

## 4.1. General

THF, Et<sub>2</sub>O, pentane and DCM were dried with a PureSolv-MD-5 Innovative Technology system for the purification of solvents. All other reagents were used as commercially available. In particular, solutions of *n*-BuLi were 1.6 or 2.5 M in hexane, solutions of EtMgBr were 3.0 M in THF, solution of *n*-OctMgBr were 2 M in Et<sub>2</sub>O, solution of *n*-BuMgCl were 2.0 M in THF. All the reactions were carried out under an argon atmosphere by using Schlenk tube and vacuum line techniques. Column chromatography was carried out on silica gel (60 Å, C.C. 70–200 µm). Silica gel TLC plates (60F254, 0.25 mm) were observed under UV light and/or by treatment with a solution of phosphomolybdic acid (20%) in ethanol. The following analytical instruments were used: <sup>1</sup>H and <sup>13</sup>C NMR spectroscopy: Avance 300, Avance 400 and Avance 400 HD spectrometers; MS: TSQ 7000 Thermo Electron and Voyager DE-ST Perseptive Biosystems spectrometers; UV/vis: PerkinElmer UV–vis Win-Lab Lambda 35; FT-IR: PerkinElmer 1725.

NMR chemical shifts are given in ppm, with positive values to high frequency relative to the tetramethylsilane (TMS) reference. Coupling constants, *J*, are in Hz, UV–vis extinction molar coefficients,  $\varepsilon$ , are in M<sup>-1</sup> cm<sup>-1</sup>, wavelengths,  $\lambda$ , in nm.

## 4.2. Voltammetry

Voltammetric measurements were carried out with an Autolab PGSTAT100 potentiostat controlled by GPES 4.09 software. Experiments were performed at room temperature in a homemade, air-tight, three-electrode cell connected to a vacuum/argon line. The reference electrode consisted of a SCE separated from the solution by a bridge compartment. The counter electrode was a platinum wire of about 1 cm<sup>2</sup> apparent surface. The working electrode was a Pt microdisk (0.5 mm diameter). The supporting electrolyte, [*n*Bu<sub>4</sub>N][PF<sub>6</sub>], was used as received (Fluka, 99% electrochemical grade) and simply degassed under argon. The solution used in the electrochemical study was  $10^{-3}$  M in

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*carbo*-benzene and 0.1 M in supporting electrolyte. Before each measurement, the solution is degassed by bubbling with argon, and the working electrode was polished by using a polishing machine (Presi P230). Typical instrumental parameters for recording SW voltammograms were: SW frequency, f = 20 Hz; amplitude,  $E_{sw} = 20$  mV; scan increment dE = 0.5 mV.

## 4.3. Crystallography

Data collections were performed on a high flux microfocus Rigaku FRX rotating anode at the copper  $k_{\alpha}$  wavelength equipped with a Dectris Pilatus 200 K hybrid detector at 130 K. The structure was solved using Superflip [53a], and refined by full-matrix least-squares procedures on *F* using the programs of CRYSTALS [53b]. Atomic scattering factors were taken from the International tables for X-ray Crystallography [53c]. All non-hydrogen atoms were refined anisotropically. Hydrogen atoms were refined using a riding model. Absorption corrections were introduced using the program MULTISCAN.

## 4.4. Synthesis and characterization

• 2-(2-{10-[2-(3,4-dialkylthien-2-yl)ethynyl]-4,7,13,16-tetraphenylcyclooctadeca-1,2,3,7,8,9,13,14,15-nonaen-5,11,17-triyn-1-yl}ethynyl)-3,4-dialkylthiophene (**2a**, **2b**).

To the solution of **9a** or **9b** (0.025 mmol) in 20 mL of dry DCM at -78 °C was added anhydrous SnCl<sub>2</sub> (20 eq.), and the resulting mixture was allowed warming to room temperature and stirred at this temperature for 3 h. Then the mixture was filtered through a pad of silica gel using DCM until complete elution of colored fractions. The resulting colored solutions were joined and concentrated under vacuum and the residue were washed with pentane to give **2a** or **2b** as a dark violet solid (**2a**: 35%, **2b**: 20%).

**2a**: <sup>1</sup>**H NMR** (CD<sub>2</sub>Cl<sub>2</sub>, 400 MHz):  $\delta$  0.61 (t, J = 7.1 Hz, 6H), 0.94 (t, J = 6.6 Hz, 6H), 1.01–1.13 (m, 8H), 1.17–1.29 (m, 4H), 1.32–1.58 (m, 24H), 1.65 (quint, J = 7.2 Hz, 4H), 1.81 (quint, J = 7.6 Hz, 4H), 1.99 (quint, J = 8.7, Hz, 4H), 2.75 (t, J = 7.8 Hz, 4H), 3.28 (t, J = 7.7 Hz, 4H), 7.26 (s, 2H), 7.73 (t, J = 7.5 Hz, 4H), 7.96 (t, J = 7.8 Hz, 8H), 9.43 (d, J = 7.5 Hz, 8H).

 $^{13}$ C {<sup>1</sup>H} NMR (CD<sub>2</sub>Cl<sub>2</sub>, 100 MHz):  $\delta$  14.1, 14.3, 22.9, 23.2, 29.3, 29.6, 29.7, 29.8, 30.0, 30.0, 30.1, 30.1, 30.4, 30.9, 32.1, 32.4, 85.1, 94.0, 97.6, 105.1, 114.0, 118.7, 119.4, 120.0, 124.8, 130.3, 130.3, 130.4, 139.4, 143.7, 150.2.

**HRMS** (MALDI-TOF/DCTB): m/z calcd for  $C_{86}H_{90}S_2$  [M\*]+1186.6484, found: 1186.6416.

**UV–Visible** (CHCl<sub>3</sub>):  $\lambda_{max} = 497 \text{ nm} (\varepsilon = 225 \text{ 000 L mol}^{-1} \text{ cm}^{-1}).$ 

**2b:** <sup>1</sup>**H NMR** (CDCl<sub>3</sub>, 300 MHz):  $\delta$  1.06–1.13 (m, 12H), 1.57 (m, 4H), 1.72–2.01 (m, 12H), 2.75 (t, J = 7.8 Hz, 4H), 3.28 (t, J = 7.6 Hz, 4H), 7.22 (s, 2H), 7.75 (t, J = 7.3 Hz, 4H), 7.97 (t, J = 7.7 Hz, 8H), 8.46 (d, J = 7.3 Hz, 8H).

**HRMS** (MALDI-TOF/DCTB): *m*/*z* calcd for C<sub>70</sub>H<sub>58</sub>S<sub>2</sub> [M\*]+ 962.3943, found: 962.3980.

• 2-(2-{4-[2-(3,4-dioctylthien-2-yl)ethynyl]phenyl}ethynyl)-3,4-dioctylthiophene (**2P**).

To a degassed solution of 1,4-diiodobenzene (48 mg, 0.143 mmol) and 4a (100 mg, 0.301 mmol) in a mixture of dry THF/Et<sub>3</sub>N (5 mL, 4/1) under an argon atmosphere were added copper(I) iodide (3 mg, 0.0143 mmol) and bis(triphenylphosphine)palladium(II) dichloride (5 mg, 0.00715 mmol). The mixture was heated in a sealed tube at 75 °C for 12 h before cooling to room temperature and concentration under vacuum. The residue was purified by SiO<sub>2</sub> column chromatography (pentane: EtOAc = 10 : 1) to give **2P** as a waxy solid (49 mg, 47%).

<sup>1</sup>**H** NMR (CDCl<sub>3</sub>, 400 MHz):  $\delta$  0.84–0.95 (m, 12H), 1.22–1.47 (m, 40), 1.56–1.69 (m, 8H), 2.51 (t, *J* = 7.6 Hz, 4H), 2.71 (t, *J* = 7.6 Hz, 4H), 6.88 (s, 2H), 7.42–7.48 (s, 4H).

<sup>13</sup>C {<sup>1</sup>H} NMR (CDCl<sub>3</sub>, 100 MHz):  $\delta$  14.1, 22.7, 28.3, 29.1, 29.3, 29.3, 29.4, 29.5, 29.5, 29.6, 29.8, 30.0, 31.9, 31.9, 85.2, 94.6, 118.3,

121.7, 123.0, 131.0, 142.2, 147.3.

**HRMS** (MALDI-TOF/DCTB): *m*/*z* calcd for C<sub>50</sub>H<sub>74</sub>S<sub>2</sub> [M\*]<sup>+</sup> 738.5232, found: 738.5288.

• 2-ethynyl-3,4-dialkylthiophene (4a,b).

*n*-BuLi (1.2 eq.) was added at -78 °C to a solution of dibromoolefin **8a** or **8b** (13 mmol,1 eq.) in pentane (300 mL). The temperature was increased up to *ca* -40 °C and stirring was continued for 30 min. Then, the mixture was cooled again to -78 °C and treated with a saturated aqueous solution of NH<sub>4</sub>Cl. After extractions of the aqueous layer with diethyl ether, the combined organic layers were washed with brine, dried over MgSO<sub>4</sub> and evaporated under reduced pressure to give the expected product **4a** or **4b** as a yellow liquid (**4a**: 99%, **4b**: 91%).

**4a**: <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>)  $\delta$  0.90–0.93 (m, 6H), 1.29–1.41 (m, 24H), 1.55–1.63 (m, 4H), 2.51 (t, J = 8.0 Hz, 2H), 2.66 (t, J = 8.0 Hz, 2H), 3.43 (s, 1H), 6.84 (s, 1H).

 $^{13}$ C {<sup>1</sup>H} NMR (CDCl<sub>3</sub>, 100 MHz)  $\delta$  14.1, 22.7, 28.1, 29.0, 29.3, 29.3, 29.4, 29.5, 29.5, 29.6, 29.8, 30.0, 31.9, 31.9, 82.6, 117.3, 121.4, 141.9, 148.1.

**HRMS** (DCI/CH<sub>4</sub>): *m*/*z* calcd for C<sub>22</sub>H<sub>37</sub>S [M − H]<sup>+</sup> 333.2616, found: 333.2603.

IR (cm<sup>-1</sup>):  $\nu$  = 3313, 2954, 2923, 2854, 2101, 1465, 1155, 875, 744, 655, 575.

**4b**: the obtained data were in accordance with those previously described [28].

• **3,4-dialkylthiophenes** (5a, 5b) were prepared following described procedures. Physical properties of the obtained products are in accordance with those previously reported [28,54].

• 3,4-dialkyl-2-thiophenocarboxaldehyde (6a, 6b).

A solution of the alcohol **7a** or **7b** (1 eq., 15 mmol) in DCM (300 mL) was treated with  $MnO_2$  (10 eq.) at 0 °C. The mixture was stirred at 0 °C for 10 min, then 1.5 h at room temperature, before being filtered through Celite<sup>®</sup>. The solution was concentrated under vacuum to give the expected spectroscopically pure aldehydes **6a** or **6b** as a yellow oil (**6a**: 95%, **6b**: 94%)

**6a**: the obtained data were in accordance with those previously described [54].

**6b**: the obtained data were in accordance with those previously described [55].

• 2-Hydroxymethyl-3,4-dialkylthiophene (7a,7b).

*n*-BuLi (1.2 eq.) was added at 0 °C to a solution of the **6a**, **6a**' or **6b**, **6b**' mixture (1 eq., 16 mmol) in THF (50 mL). After 45 min of stirring, the mixture was brought to room temperature and a suspension of paraformaldehyde (1.8 eq.) in THF (10 mL) was added. The mixture was stirred for 3 h at room temperature, and poured into ice and water. Then, the pH was adjusted to 6 by addition of concentrated HCl. After extractions of the aqueous layer with diethyl ether, the combined organic layers were washed with water, dried over MgSO<sub>4</sub> and evaporated under reduced pressure. After purification by silica gel chromatography (pentane: diethyl ether, **98** : 2), the alcohols **7a** or **7b** were obtained as a yellow liquid (**7a**: 83%, **7b**: 95%)

**7a**: t he obtained data were in accordance with those previously described [54].

**7b**: the obtained data were in accordance with those previously described [56].

• ((1,1-Dibromo) -2-ethene) -3,4-dialkylthiophene (8a,8b).

A solution of aldehyde **6a** or **6b** (1 eq., 15 mmol) in DCM (200 mL) was added at 0 °C to a mixture of PPh<sub>3</sub> (4 eq.) and CBr<sub>4</sub> (2 eq.) in DCM (50 mL). The resulting mixture was stirred for 5 min before being concentrated to dryness. The residue was purified by silica gel chromatography (pentane: EtOAc 100 : 1) to give the expected product **8a** or **8b** as a yellowish liquid (**8a**: 91%, **8b**: 66%).

**8a**: <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>)  $\delta$  0.96 (t, J = 8.0 Hz, 6H), 1.34–1.40 (m, 22H), 1.64–1.70 (m, 2H), 2.53–2.60 (m, 4H), 7.04 (s, 1H), 7.68 (s, 1H).

<sup>13</sup>C {<sup>1</sup>H} NMR (CDCl<sub>3</sub>) δ 14.2, 22.8, 27.4, 28.8, 29.3, 29.4, 29.4, 29.6, 29.7, 29.7, 29.8, 31.0, 32.0, 85.9, 121.7, 129.9, 131.9, 141.8,

143.2.

HRMS (DCI/CH<sub>4</sub>): m/z calcd for C<sub>22</sub>H<sub>36</sub>OSBr<sub>2</sub> [M] 490.0904, found: 490.0904.

**8b**: <sup>1</sup>H NMR (300 MHz, CDCl<sub>3</sub>) δ 0.97 (m, 6H), 1.39–1.62 (m, 12H), 2.50–2.60 (m, 4H), 7.03 (s, 1H), 7.64 (s, 1H).

<sup>13</sup>C {<sup>1</sup>H} NMR (75 MHz, CDCl<sub>3</sub>): δ 13.9, 14.0, 22.6, 22.7, 27.2, 28.4, 31.9, 33.1, 85.9, 121.7, 129.8, 131.8, 141.8, 143.2.

**HRMS** (DCI/CH<sub>4</sub>): *m*/*z* calcd for C<sub>14</sub>H<sub>20</sub>OSBr<sub>2</sub> [M] 377.9639, found: 377.9652.

• 1,10-bis[2-(3,4-dialkylthien-2-yl)ethynyl]-4,7,13,16-tetramethoxy-4,7,13,16-tetraphenylcyclooctadeca-2,5,8,11,14,17-hexayne-1,10-diol (**9a**, **9b**).

To the solution of **4a** or **4b** (3 eq.) in dry THF (10 mL) at -78 °C was added dropwise a *n*-BuLi solution (1.6 M in hexane, 2.5 eq.). Stirring was continued at -78 °C for 30 min, then at room temperature for 1 h. After cooling to -78 °C, a solution of **3** (0.15 mmol) in dry THF (2 mL) was added dropwise and the mixture was allowed warming slowly to room temperature overnight. The reaction was treated with a saturated aqueous solution of NH<sub>4</sub>Cl and the aqueous layer was extracted with Et<sub>2</sub>O. The combined organic layers were dried with MgSO<sub>4</sub>, concentrated under vacuum and purified by SiO<sub>2</sub> column chromatography (pentane: EtOAc 10 : 1) leading to the diol **9a** or **9b** as a waxy solid (**9a**: 58%, **9b**: 48%).

**9a**: <sup>1</sup>**H NMR** (CDCl<sub>3</sub>, 300 MHz): δ 0.84–0.94 (m, 12H), 1.18–1.65 (m, 24H), 2.49–2.67 (8H), 2.93–3.21 (m, 2H), 3.41–3.68 (m, 12H), 6.92 (s, 2H), 7.28–7.42 (m, 12H), 7.74–7.82 (m, 8H).

<sup>13</sup>C {<sup>1</sup>H} NMR (CDCl<sub>3</sub>, 100 MHz): δ 14.1, 22.7, 28.3, 28.9, 29.2, 29.4, 29.5, 29.7, 29.8, 30.0, 30.1, 31.9, 53.6, 54.9, 71.9, 78.3, 80.4, 84.4, 90.5, 116.1, 123.0, 126.5, 128.5, 129.1, 139.2, 142.2, 149.0.

**HRMS** (MALDI-TOF/DCTB): m/z calcd for  $C_{86}H_{90}S_2$  [M]<sup>+</sup> 1186.6484, found: 1186.6460.

**9b**: <sup>1</sup>H NMR (CDCl<sub>3</sub>, 300 MHz): δ 0.88–0.98 (m, 12H), 1.27–1.59 (m, 16H), 2.48–2.62 (m, 8H), 3.52–3.54 (m, 12H), 6.88 (s, 2H), 7.29–7.33 (m, 12H), 7.74 (m, 8H).

**MS** (MALDI-TOF/DCTB): *m*/*z* [M - Na]<sup>+</sup> 1143.7 [M - K]<sup>+</sup> 1159.7.

## 4.5. Linear and nonlinear spectroscopic characterization

Spectra of linear absorption, emission, excitation, and lifetime of fluorescence were obtained with the use of a fluorometer (FS5, Edinburgh Instruments). Emission was also obtained by pumping the sample with short optical pulses (100 kHz repetition rate) from an optical parametric amplifier (Orpheus, Light Conversion, Lithuania) pumped at 1030 nm by a mode-locked oscillator (Pharos-Model PH1-20-0200-02-10, Light Conversion, Lithuania). To minimize absorption inner effects, the excitation and recollection of emission was produced in the front face of a 1 mm-thick cuvette containing a solution of the sample. The emission analyzed with an imaging spectrograph (Shamrock 193i, Andor Technology) and detected with a multichannel detector. The  $\sigma_{2PA}$ was obtained by Z-scan method with 80 fs (800 nm) laser pulses delivered by a Ti:Sa amplifier (Libra HE from Coherent) at 1 KHz. At the position Z = 0 the beam waist was 20  $\mu$ m. THF solutions were tested at the concentration of  $8.43 \times 10^{-2}$  M in a 1 mm quartz cell. For wavelengths other than 800 nm in the range 650–950 nm, the Z-scan method was implemented with an optical parametric amplifier (TOPAs, from Light Conversion). The nonlinear coefficient  $\beta$  was calculated from the equation [57a,57b]:

$$T(z) = 1 - \frac{1}{2\sqrt{2}}\beta \frac{I_0 L_{eff}}{1 + (Z/Z_0)^2}$$
(1)

where  $I_0$  is the peak intensity,  $L_{eff}$  is the effective length of the sample with  $L_{eff} = [1 - exp(-\alpha_0 L)]/\alpha_0$ , where L is the sample thickness and  $\alpha_0$  the absorption coefficient, Z the sample position and  $Z_0 = \left(\pi\omega_0^2/\lambda\right)$  the Rayleigh range. To determine  $\omega_0$  the knife edge method was used.

The Z-scan apparatus was calibrated with traces in the close aperture scheme using CS<sub>2</sub> as standard nonlinear refractive material with  $n_2 = 2 \times 10^{-15} \text{ cm}^2/\text{W}$  [58]. Finally,  $\sigma_{2PA} = \frac{\hbar\omega}{N}\beta$ , where  $\hbar\omega$  is the photon energy and N the density of molecules in the solution.

Two-photon excited fluorescence (TPEF) was implemented with 140 fs pulses from a Ti:Sa oscillator (Chameleon ultra, from Coherent) in the range 680–1080 nm. THF solutions were tested at a concentration of  $8.43 \times 10^{-5}$  M. Laser beam was focused in the frontal face of a 1 cm quartz cell with a 35 mm focal length lens. TPEF signal was detected in the perpendicular direction to the excitation with a hand-held spectrometer (Ocean optics, 2000+) and using a low pass filter to block the excitation. The  $\sigma_{2PA}$  value was obtained using the equation [59]:

$$\sigma_{TPA} = \sigma_R \frac{F_s C_R \Phi_R \eta_R}{F_R C_R \Phi_S \eta_S}$$
(2)

where, F is the integrated area of the emission, C is the molar concentration (mol. L<sup>-1</sup>),  $\phi$  is the fluorescence QY, and  $\eta$  is the refraction index of the solvent, the subscripts S and R denoting sample and reference, respectively. The Rodhamine 6G at a concentration of  $1 \times 10^{-5}$  M was used as reference for calibration.

## 4.6. Optical limiting

For optical limiting measurements, the same laser source as the Z-scan experiments was used at 800 nm. The setup consisted in a laser beam hitting a 1 mm quartz cell containing a solution of **2a** at a concentration of  $8.43 \times 10^{-2}$  M in THF.

## Author contribution

Rodrigo M. Barba-Barba: Investigation (general optical studies, linear and nonlinear); Formal analysis; Visualization; Writing - original draft. Marwa Chammam: Investigation (general chemical studies and synthesis); Formal analysis; Visualization; Writing - original draft. Gabriel Ramos-Ortiz: Conceptualization; Supervision; Investigation; Formal analysis; Funding acquisition; Resources; Project administration; Writing - review & editing. Dymytrii Listunov: Investigation (chemical synthesis). Jayaramakrishnan Velusamy: Investigation (nonlinear optical characterization). Mario Rodriguez: Conceptualization (molecular design). Ramon Carriles: Investigation (linear optical characterization). Carlos Silva: Validation; Resources, Funding acquisition; Writing - review & editing. Carine Duhayon: Investigation (X-ray crystallography). Brice Kauffmann: Investigation (X-ray crystallography). Valérie Maraval: Conceptualization; Supervision; Formal analysis; Funding acquisition; Resources; Project administration; Writing - review & editing. Remi Chauvin: Conceptualization; Supervision; Formal analysis; Funding acquisition; Resources; Writing - review & editing.

## Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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## Appendix A. Supplementary data

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