


 Cite this: *New J. Chem.*, 2022, 46, 13431

How do structural factors determine the linear and non-linear optical properties of fluorene-containing quadrupolar fluorophores? A theoretical answer†‡

 Safa Gam,^{id abc} Sabri Messaoudi,^{id bcd} Jean-François Halet^{id *ae} and Abdou Boucekine^{id *a}

A large series of push–push and pull–pull quadrupolar fluorophore derivatives with conjugated rods made from arylene–vinylene (PV) or arylene–ethynylene (PE) building blocks, bearing different electron-releasing or electron-withdrawing end-groups, were theoretically investigated using DFT and TD–DFT computations. These compounds, which exhibit good transparency in the visible region, are very promising for various applications, especially for optical limitation. Their absorption and photoluminescence as well as their two-photon absorption (TPA) properties in the near infrared (NIR) region were systematically investigated in order to derive structure–property relationships. The results indicate that (i) for all the studied compounds, the lowest excited state reached by the TPA is the S_2 state, corresponding to the HOMO–1 to LUMO electronic transition, (ii) push–push molecules are found to be more efficient than pull–pull molecules, (iii) an increase of the strength of the donor terminal group (OPh, OMe, NH_2 , NPh_2 , and NBu_2) enhances the TPA cross-section, and (iv) replacement of PV by PE always leads to an increase of TPA cross-sections in the NIR region.

 Received 10th March 2022,
 Accepted 30th May 2022

DOI: 10.1039/d2nj01192a

rsc.li/njc

Introduction

Over the years, a large number of π -conjugated organic molecules with large two-photon absorption properties have been developed.^{1–25} This was the case, for instance, of a series of push–push and pull–pull derivatives prepared and characterized by Mongin, Blanchard-Desce and colls,^{13,20,22} Jen and colls,¹⁴ Belfield and colls^{18,19} or Painelli and colls²⁴ containing conjugated rods made from phenylene–vinylene (PV) or

phenylene–ethynylene (PE) building blocks. More specifically, these quasi one-dimensional quadrupolar biphenyl derivatives, *i.e.*, symmetrical conjugated molecules, bear either electron-releasing ($X = \text{OPh}$, OMe, NH_2 , NPh_2 , and NBu_2) or electron-withdrawing ($X = \text{SO}_2\text{CF}_3$ and CN) end-groups attached at the *para* position to conjugated PV or PE connectors (Scheme 1). Interestingly, such elongated rod-like or banana-shaped fluorophores exhibit enhanced TPA cross-sections (σ_2 , expressed in $\text{GM} = 1 \times 10^{-50} \text{ cm}^4 \text{ s photon}^{-1} \text{ molecule}^{-1}$) in the visible red–near IR region (*i.e.*, 700–1300 nm, a spectral window very

^a Univ Rennes, CNRS, Institut des Sciences Chimiques de Rennes (ISCR) – UMR 6226, F35000 Rennes, France. E-mail: Abdou.Boucekine@univ-rennes1.fr, Jean-Francois.Halet@univ-rennes1.fr

^b Faculty of Sciences of Bizerte FSB, University of Carthage, 7021 Jarzouna, Tunisia

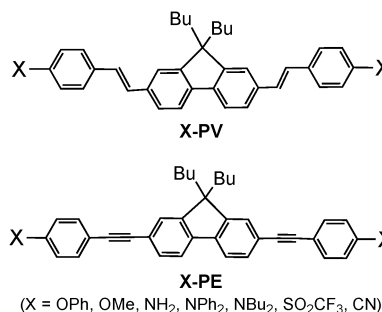
^c Laboratory of Materials, Molecules and Applications, IPEST, University of Carthage, Sidi Bou Said Road, B.P. 51, 2070 La Marsa, Tunisia

^d Department of Chemistry, College of Science, Qassim University, Buraidah 51452, Saudi Arabia

^e CNRS – Saint-Gobain – NIMS, IRL 3629, Laboratory for Innovative Key Materials and Structures (LINK), National Institute for Materials Science (NIMS), Tsukuba, 305-0044, Japan

† Dedicated to Dr Christian Bruneau for his important contribution to molecular chemistry.

‡ Electronic supplementary information (ESI) available: Optimized geometries and Cartesian coordinates, UV-visible spectra, TD-DFT results, frontier MO diagrams, and OPA and TPA computational detailed results. See DOI: <https://doi.org/10.1039/d2nj01192a>



Scheme 1 X-PV and X-PE series of compounds with different electron donor ($X = \text{OPh}$, OMe, NH_2 , NPh_2 , and NBu_2) or acceptor groups ($X = \text{CN}$ and SO_2CF_3).



appealing for biological applications).^{13,14,18,19,22} Moreover, the fluorene core exhibits high thermal and photochemical stability, making fluorene-containing compounds promising in areas such as laser applications.^{7,8,10,16}

From an experimental viewpoint, it is often challenging to fully characterize the electronic processes responsible for the properties of such compounds. Fortunately, it is possible to obtain crucial information on their molecular photo-physical properties, namely those relevant to excitation and emission phenomena and the nature of the involved excited states, thanks to quantum chemical computations.^{26–41} In this context, we carried out standard density functional theory (DFT) and time-dependent density functional theory (TD-DFT) computations to first explore the ground state (GS) characteristics, *i.e.*, the geometric and electronic structures of the compounds shown in Scheme 1. In the second step, we investigated their optical properties aiming to propose an assignment of the observed absorption bands and emission spectra using TD-DFT simulations. Then, we theoretically investigated their TPA properties, trying to understand and rationalize the structural origin of their large non-linear optical (NLO) responses.

It is noteworthy that systems bearing strong donor or acceptor terminal groups can also lead to a reduced fluorescence quantum yield and/or a pronounced red-shift of the one-photon absorption band.^{1,42} So, the aim of our work was to (i) compare the effect of the PV or PE connectors on the absorption/emission properties and on their TPA response, (ii) study the influence of the strength of the donor/acceptor terminal group on the considered properties, and (iii) evaluate the effect of the replacement of the biphenyl core by a bi-thiophene one. The purpose of these systematic structural variations was to derive complete structure–TPA relationships and to determine the appropriate combination of the core, linker (double or triple bond) and connector (PV, PE) for optimized properties of TPA/luminescence and/or TPA/transparency. Herein, we report and discuss the main results that were obtained.

Computational details

DFT and TD-DFT calculations were performed using the Gaussian 09⁴³ and Gaussian 16⁴⁴ programs. Solvent effects (toluene) were taken into account using the Polarizable Continuum Model (PCM).^{45–48} All the ground state (GS) geometries (thereafter S_0) were optimized without any symmetry constraint and were checked to be genuine minima on the potential energy surface (PES), *i.e.*, the computed frequencies of their normal modes of vibration were all real. Calculations were carried out employing the PBE0 functional⁴⁹ and the 6-31+G(d) standard basis set. TD-DFT computations were carried out first at the same level of theory using the optimized GS geometries to investigate the absorption spectral properties of the considered molecules. It was found (*vide infra*) that calculations carried out at the CAM-B3LYP/6-31+G(d) level⁵⁰ showed more satisfying agreement with experiments with respect to the UV-vis spectra than

with respect to the PBE0/6-31+G(d) ones. The optimized geometries and vibration frequencies of the first triplet state (T_1) were obtained using spin-unrestricted calculations, whereas the excited S_1 state geometries were optimized using TD-DFT computations. The molecular orbitals (MO), and the simulated UV-vis and fluorescence spectra were plotted using the GaussView 5.0 program.⁵¹

The TPA properties were computed using the damped cubic response theory of Lasse, Jensen and colls³⁶ implemented in the Amsterdam Density Functional (ADF2018 and ADF2019)^{52,53} program packages. Unless otherwise stated, these response properties were calculated using the statistical average of orbital model exchange–correlation potentials (SAOPs) with a DZP basis set in the gas phase. Solvent effects were not included in the TPA calculations. The SAOP potential model^{54,55} was chosen due to its correct coulombic behavior of the potential at long distances, which is important for the description of response properties. It should be kept in mind that geometry optimizations cannot be carried out using the SAOP model and that the geometries used for the TPA calculations were those optimized at the PBE0 level in solution. Relevant SAOP/DZP MOs were plotted using the ADFView program.⁵⁶

In previous studies, we showed that SAOP/DZP calculations lead to TPA cross-sections that were in satisfying agreement with the experimental measurements.^{57,58} The excited state lifetime was included in the theory using a damping parameter of 0.0034 atomic unit (a.u.) (~ 0.1 eV or ~ 800 cm^{-1}),⁵⁹ which was previously found to be suitable for TPA computations.^{36,60} The full TPA simulated spectra were plotted point by point after the computation of the cross-sections for different laser energies.

The TPA cross-sections (σ^{TPA}) were calculated from the imaginary part of the third-order hyperpolarizability γ as

$$\sigma^{\text{TPA}}(\omega) = \frac{N\pi^3\alpha_f^2\hbar^3\omega^2}{15e^4}\gamma^{\text{TPA}} \quad (1)$$

with

$$\gamma^{\text{TPA}} = \sum_{\alpha\beta} \text{Im}[\gamma_{\alpha\alpha\beta\beta}(-\omega; \omega, \omega, -\omega) + \gamma_{\alpha\beta\beta\alpha}(-\omega; \omega, \omega, -\omega) + \gamma_{\alpha\beta\alpha\beta}(-\omega; \omega, \omega, -\omega)] \quad (2)$$

where α_f is the fine structure constant (not to be confused with the α -index of the summation in eqn (2)), ω is the photon energy, \hbar is the reduced Planck's constant, and e is the elementary charge. The integer value N is related to the experimental setup and in this work, the value $N = 4$ was used for all simulated TPA spectra.^{35,36} The cross-section values (σ^{TPA}) are given in Goepfert–Mayer (GM) ($1 \text{ GM} = 10^{-50} \text{ cm}^4 \text{ s photon}^{-1} \text{ molecule}^{-1}$).⁶¹ σ^{TPA} values were first obtained in atomic units (a.u.) and then converted $(0.529177 \times 10^{-8} \text{ cm a.u.}^{-1})^4 \times (2.418884 \times 10^{-17} \text{ s a.u.}^{-1})$ into conventional units ($\text{cm}^4 \text{ s photon}^{-1} \text{ molecule}^{-1}$). All other properties are expressed in a.u. The expression of γ^{TPA} given in eqn (2) which avoids any negative TPA intensities caused by the pure



one-photon processes and allows for appropriate σ^{TPA} calculations in the presence of one- and two-photon double-resonance effects,^{62,63} was used for all TPA simulations.

Results

Ground state geometries

The different possible geometric conformations of the studied compounds (Scheme 1) were first explored. They are displayed in Fig. S1 (ESI†). Different energy minima were found for all of them (Table 1). The **OPh-PV** and **OMe-PV** molecules exhibit four thermodynamically stable forms labeled a–d that differ either by the orientation of the terminal group (OPh, OMe, respectively) or by the orientation of the two butyl groups. Forms a and b are close in energy and significantly more stable than forms c and d. This indicates that forms a and b might at least coexist in solution. Regarding **NH₂-PV**, **NPh₂-PV**, **NPh₂-PE**, **CN-PV** and **NBu₂-PV**, only two forms were computed to be thermodynamically stable (a and b), showing different orientations of the two butyl groups with respect to the π -conjugated backbone. Finally, for the **SO₂CF₃-PV** and **SO₂CF₃-PE** molecules, three forms were found (a, b and c with a and b very close in energy) that differ either by the orientation of the triflate terminal group (SO₂CF₃) and/or by the orientation of the two butyl groups. In all cases, form a is the most stable.

In the most stable configurations (form a), the two butyl groups are perpendicular to the plane of the molecules, and the peripheral arms built from the *para*-substituted PV or PE moieties adopt a planar structure relatively to the central fluorene core (Table S1, ESI†). Here after, unless specified, results which are presented and discussed concern only these most stable configurations.

Absorption properties

The photo-physical properties of some of the fluorophores considered in Scheme 1, as well as their absorption and emission features including fluorescence quantum yields and lifetimes, were measured experimentally in toluene solution.^{13,20,22} All of them display an intense absorption in the near UV-vis blue region with typical molar extinction coefficients ranging from 60 000 to 180 000 mol⁻¹ L cm⁻¹. TD-DFT calculations were first

Table 2 Experimental and theoretical UV-vis absorption wavelengths (λ_{max} (nm)) for the most stable form (a) of the **X-PV** and **X-PE** series of compounds in toluene

Compound	$\lambda_{\text{max}}^{\text{abs}}$		Exp.	Ref.
	PBE0	CAM-B3LYP		
OPh-PV	434	385		
OMe-PV	429	381	381	13
NH₂-PV	440	387	392	13
NPh₂-PV	479	408	411	20
NPh₂-PE	446	382	389	22
NBu₂-PV	470	407	415	13
CN-PV	447	392		
SO₂CF₃-PV	451	395		
SO₂CF₃-PE	420	371	372	13

carried out considering the most stable form (a) of all compounds with different electron donor (X = OPh, OMe, NH₂, NPh₂, and NBu₂) or acceptor (X = CN and SO₂CF₃) groups in order to assign the different absorption bands observed experimentally in toluene. Note that starting from the optimized geometries obtained at the PBE0/6-31+G(d) level, vertical TD-DFT computations were carried out using both the PBE0 and CAM-B3LYP functionals to simulate the UV-vis absorption spectra of the molecules in toluene. As can be seen for the lowest energy absorption band (Table 2), CAM-B3LYP calculations show perfect agreement with the experiments,^{13,20,22} whereas PBE0 overestimates systematically the experimentally measured wavelengths by *ca.* 50–60 nm. Thereafter, only the CAM-B3LYP results are discussed. Complete results are given in Table 3. The simulated absorption spectra of some of them (X = OPh, NH₂, NPh₂, CN, and SO₂CF₃) are shown in Fig. 1, and the molecular orbitals involved in the corresponding electronic excitations are displayed in Fig. 2–4.

As can readily be seen in Table 3, the lowest energy absorption band calculated for each compound with different donors is broad and intense with a maximum very close to the visible region, at 385 nm ($f_{01} = 2.58$), 387 nm ($f_{01} = 2.84$), 408 nm ($f_{01} = 3.10$) and 382 nm ($f_{01} = 3.51$) for **OPh-PV**, **NH₂-PV**, **NPh₂-PV** and **NPh₂-PE**, respectively. Indeed, the absorption band energy depends very slightly upon (i) the nature of the peripheral donor groups (a small red shift is observed upon the increase of the donor properties), and (ii) the nature of the conjugated arms (C=C or C≡C) tethered to the fluorene core as can be seen comparing **NPh₂-PV** and **NPh₂-PE** (a small blue shift is noticed for the latter with respect to the former). Similarly, with acceptor groups, the computed lowest energy absorption band is broad and intense with a maximum very close to the visible as well at 392 nm ($f_{01} = 2.95$), 395 nm ($f_{01} = 2.90$) and 371 nm ($f_{01} = 2.89$) for **CN-PV**, **SO₂CF₃-PV** and **SO₂CF₃-PE**, respectively. Again, a slight blue shift is observed when the vinyl group is replaced by an ethynyl one. These little changes in the absorption band energy for these different compounds can be interpreted with a glance at the energy and nodal properties of their HOMO and LUMO which are mostly involved in the corresponding electronic excitations. They are all overall delocalized over the entire molecular π -conjugated backbone with a poor contribution of the peripheral donor/acceptor groups (see Fig. 2–4 for a

Table 1 PBE0/6-31+G(d) relative energies (ΔE , kcal mol⁻¹) of different geometrical forms of the **X-PV** and **X-PE** series of compounds

Compound	ΔE			
	a	b	c	d
OPh-PV	0.00	0.19	3.06	3.20
OMe-PV	0.00	0.25	3.51	3.25
NH₂-PV	0.00	3.12		
NPh₂-PV	0.00	3.15		
NPh₂-PE	0.00	3.08		
NBu₂-PV	0.00	4.08		
CN-PV	0.00	3.13		
SO₂CF₃-PV	0.00	0.13	4.53	
SO₂CF₃-PE	0.00	0.17	4.33	



Table 3 CAM-B3LYP results (in toluene) for the most stable form (a) of the **X-PV** and **X-PE** series of compounds (S_n is the excited state number, E_{0n} is the transition energy (in eV), λ_{calc} is the wavelength (in nm) and f_{0n} is the oscillator strength of the n -th excited state)

Compound	S_n	E_{0n}	λ_{calc}	f_{0n}	Main MO transition percentage
OPh-PV	1	3.22	385	2.58	HOMO → LUMO (87%)
	2	3.89	319	0.28	HOMO-1 → LUMO (47%)
	8	4.83	257	0.24	HOMO-2 → LUMO (22%)
OMe-PV	17	5.52	225	0.32	HOMO-1 → LUMO+1 (19%)
	1	3.26	381	2.71	HOMO → LUMO (87%)
	8	4.88	254	0.35	HOMO-2 → LUMO (32%)
	15	5.52	225	0.31	HOMO-1 → LUMO+2 (24%)
NH₂-PV	28	6.10	203	0.27	HOMO → LUMO+14 (17%)
	1	3.20	387	2.84	HOMO → LUMO (85%)
	7	4.72	262	0.37	HOMO-2 → LUMO (35%)
	18	5.43	227	0.24	HOMO-1 → LUMO+1 (24%)
NPh₂-PV	18	5.43	227	0.24	HOMO-1 → LUMO+2 (31%)
	27	5.88	211	0.15	HOMO-3 → LUMO (31%)
	1	3.04	408	3.10	HOMO → LUMO (76%)
	2	3.48	356	0.17	HOMO-1 → LUMO (48%)
NPh₂-PE	28	5.47	227	0.35	HOMO → LUMO+1 (38%)
	1	3.25	382	3.51	HOMO → LUMO (72%)
	7	4.33	287	0.56	HOMO-1 → LUMO+6 (44%)
	27	5.56	223	0.28	HOMO-4 → LUMO (31%)
NBu₂-PV	29	5.60	221	0.17	HOMO-8 → LUMO (21%)
	1	3.05	407	2.88	HOMO → LUMO (83%)
	2	3.59	345	0.18	HOMO-1 → LUMO (59%)
	19	5.39	230	0.25	HOMO → LUMO+1 (35%)
CN-PV	19	5.39	230	0.25	HOMO-4 → LUMO (28%)
	29	5.67	217	0.18	HOMO-1 → LUMO+2 (15%)
	1	3.16	392	2.95	HOMO-6 → LUMO (14%)
	6	4.72	263	0.24	HOMO → LUMO (86%)
SO₂CF₃-PV	9	5.13	242	0.22	HOMO-1 → LUMO+1 (27%)
	13	5.52	225	0.29	HOMO-3 → LUMO (39%)
	1	3.14	395	2.90	HOMO → LUMO+4 (28%)
	6	4.76	261	0.23	HOMO-6 → LUMO+1 (15%)
SO₂CF₃-PE	9	5.11	243	0.25	HOMO → LUMO (86%)
	23	6.15	202	0.40	HOMO-1 → LUMO+1 (29%)
	1	3.35	371	2.89	HOMO → LUMO+2 (22%)
	13	5.67	219	0.52	HOMO-3 → LUMO (44%)
SO₂CF₃-PE	29	6.35	195	0.78	HOMO-1 → LUMO+6 (15%)
	30	6.39	194	0.42	HOMO → LUMO (84%)
	13	5.67	219	0.52	HOMO-3 → LUMO+5 (34%)
					HOMO-3 → LUMO+5 (17%)

few representatives and Fig. S2–S10, ESI†). At higher energy, all compounds exhibit absorptions in the range 220–320 nm with a shoulder at a higher wavelength (Fig. 1, Table 3 and Table S2, ESI†). These absorptions are also very slightly bathochromically or hypsochromically shifted depending on the increasing electron releasing or withdrawing properties of the *para*-substituent.

Emission and fluorescence

The emission spectra of part of the series (**OMe-PV**, **NH₂-PV**, **NPh₂-PV**, **NPh₂-PE**, **NBu₂-PV** and **SO₂CF₃-PE**) were experimentally obtained under excitation at the wavelength of maximum absorption.^{13,20,22} In order to investigate theoretically these emission spectra, geometries of the first singlet excited states

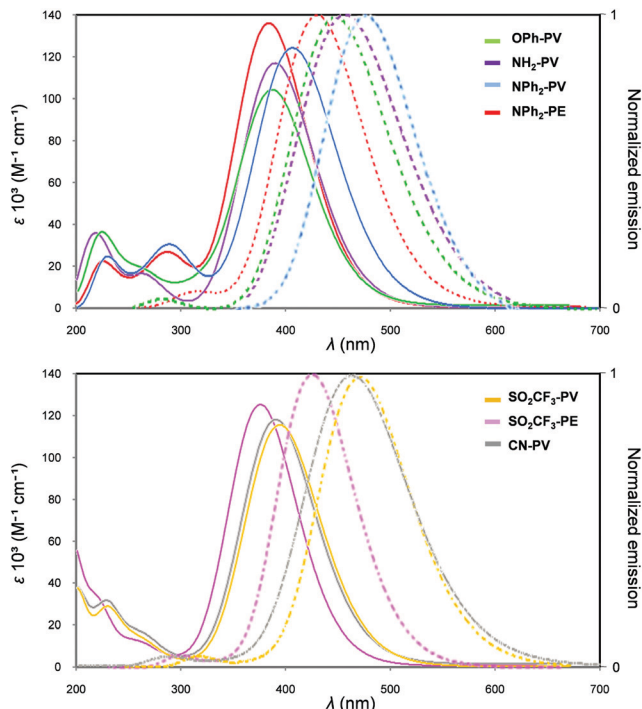


Fig. 1 CAM-B3LYP/6-31+G(d) simulated UV-vis absorption (solid line) and emission (dashed line) spectra of some examples of the **X-PV** and **X-PE** series of compounds with donor ($X = \text{OPh}, \text{NH}_2, \text{NPh}_2$, top) or acceptor ($X = \text{CN}, \text{SO}_2\text{CF}_3$, bottom) groups in toluene.

S_1 of all conformers of the whole series optimized at the CAM-B3LYP/6-31+G(d) level were considered. Results, which are given in Table 4, indicate that the computed fluorescence wavelengths of the different forms of a given compound such as forms a–d for the **OPh-PV** and **OMe-PV** molecules hardly differ. All compounds exhibit maxima emission peaks in the blue region in the range of 425–473 nm in toluene (see the computed fluorescence spectra of selected compounds in Fig. 1). As can be seen, the experimental and theoretical values are in satisfying agreement, the computed wavelengths being systematically higher than the experimentally measured ones by *ca.* 25 nm, the highest emission wavelength being for **NBu₂-PV** and the lowest one for **SO₂CF₃-PE**.

The simulated UV absorption and fluorescence emission spectra show that these molecules exhibit large Stokes shifts (Table 4), which might be due to a large difference between the Franck–Condon state and the excited state of emission of the molecules. High values of Stokes shifts ensure that there is no re-absorption of the emitted radiation. This must depend on the nature of the different electron releasing and withdrawing groups as well as the butyl groups attached to the biphenyl core. This greatly affects the intensity of absorption and emission of the target compounds resulting from a better π -extended conjugation of the synthesized compounds. The large Stokes shift provides significant information about the configuration rearrangement of molecules. This is illustrated in Fig. 5, which shows the atom-by-atom superposition of the geometry of the S_0 state on the geometry of the S_1 state for molecules **SO₂CF₃-PV** and **SO₂CF₃-PE**.



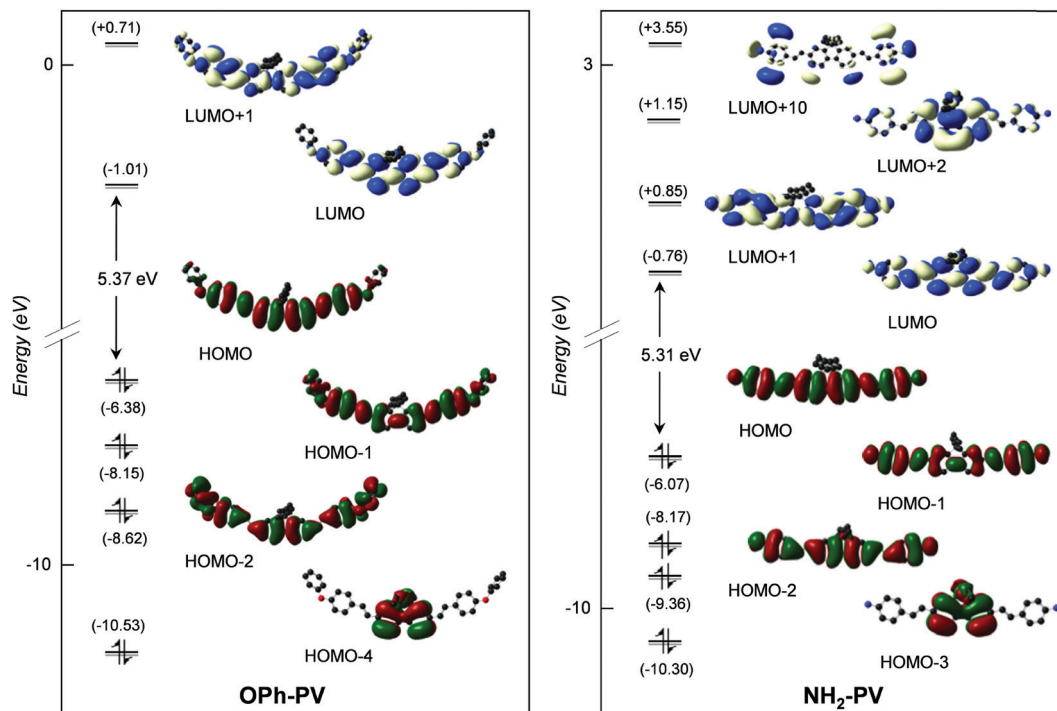


Fig. 2 Frontier MO diagrams of **OPh-PV** (left) and **NH₂-PV** (right) (contour isodensity values: ± 0.015 (e bohr^{-3/2})). Hydrogen atoms are omitted for clarity.

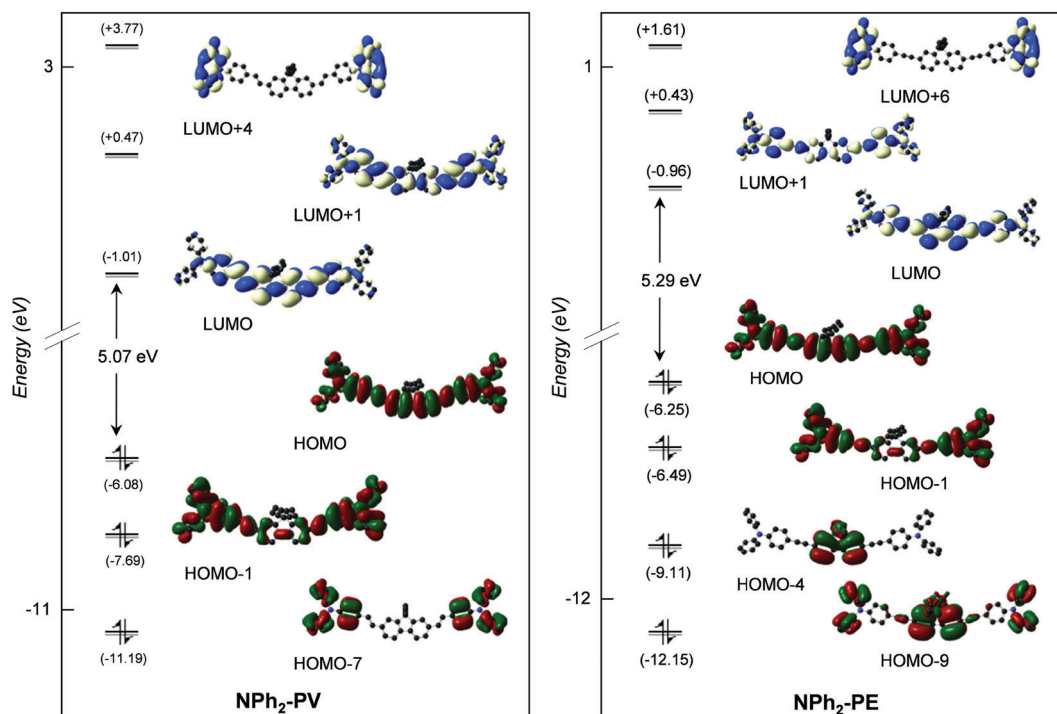


Fig. 3 Frontier MO diagrams of **NPh₂-PV** (left) and **NPh₂-PE** (right) (contour isodensity values: ± 0.015 (e bohr^{-3/2})). Hydrogen atoms are omitted for clarity.

The **SO₂CF₃-PV** molecule with the largest Stokes shift (3811 cm⁻¹) exhibits a higher displacement of the triflate moiety (SO₂CF₃) and the butyl groups compared to the **SO₂CF₃-PE** molecule with a smaller Stokes shift (3480 cm⁻¹).



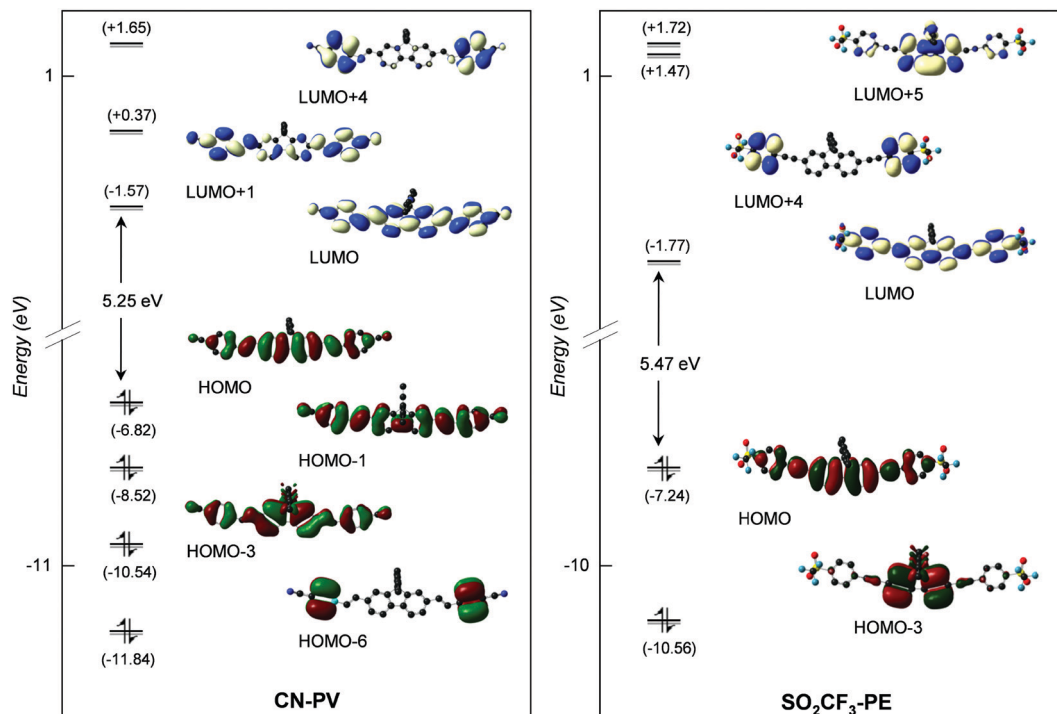


Fig. 4 Frontier MO diagrams of **CN-PV** (left) and **SO₂CF₃-PE** (right) (contour isodensity values: ± 0.015 (e bohr⁻³)^{1/2}). Hydrogen atoms are omitted for clarity.

Table 4 CAM-B3LYP/6-31+G(d) photo-physical properties of the **X-PV** and **X-PE** series of compounds, including experimental and computed absorption wavelengths (λ^{abs} , nm), fluorescence wavelengths (λ^{em} , nm) in toluene, experimental and computed Stokes shift ($\Delta\omega$, cm⁻¹)

Compound	Exp.			Ref.	Calc.			
	$\lambda_{\text{max}}^{\text{abs}}$	$\lambda_{\text{max}}^{\text{em}}$	$\Delta\omega$		Form	$\lambda_{\text{max}}^{\text{abs}}$	$\lambda_{\text{max}}^{\text{em}}$	$\Delta\omega$
OPh-PV					a	385	451	3801
					b	380	447	3945
					c	381	449	3976
					d	379	446	3963
OMe-PV	381	415	2200	13	a	381	448	3926
					b	380	448	3995
					c	378	447	4084
NH₂-PV	392	433	2400	13	d	377	446	4103
					a	387	455	3862
					b	386	454	3881
					a	408	471	3279
NPh₂-PV	411	454	2305	20	b	406	471	3400
					a	382	430	2922
NPh₂-PE	389	421	1954	22	b	380	428	2951
					a	407	473	3428
NBu₂-PV	415	457	2200	13	a	404	471	3521
					b	391	460	3839
CN-PV					a	392	462	3736
					b	395	465	3811
SO₂CF₃-PV					a	395	464	3764
					b	396	464	3701
					c	371	426	3480
SO₂CF₃-PE	372	404	2200	13	a	370	425	3498
					b	370	425	3489
					c	370	425	3489



Fig. 5 Superimposed S_0 (red)/ S_1 (black) state geometries of **SO₂CF₃-PV** (left) and **SO₂CF₃-PE** (right).

of the vertical S_1 state and that of the ground state S_0 ($\Delta\rho(r) = \rho_{S_1}(r) - \rho_{S_0}(r)$)⁶⁴ were computed to get information about the direction and the intensity of the charge transfer in these molecules. They are illustrated in Fig. 6 for **NH₂-PV**, **NPh₂-PV**, **NPh₂-PE** and **CN-PV**.

This density difference shows a very short charge transfer distance d^{CT} between the positive and negative charge centroids due to the symmetry of the molecules. On the other hand, the amount of charge Q^{CT} transferred between S_0 and S_1 is important in the four cases. Fig. 6 also clearly reveals that the charge transfer is higher in the case of the stronger electron donating terminal group **NPh₂** with respect to **NH₂**, namely 0.515 e vs. 0.447 e . Interestingly, the charge transfer is larger in **NPh₂-PE** (0.537 e) than in **NPh₂-PV** (0.515 e). The amount of charge Q^{CT} transferred between S_0 and S_1 in **CN-PV** containing the electron acceptor terminal group **CN** is also important (0.481 e). This confirms that the fluorene core can also somewhat act as an electron-donating moiety depending on the peripheral counterparts.

Overall, conclusions reached for the absorption properties of this series of compounds can be extended to the emission

To ascertain the nature of the charge transfer induced upon excitation, the differences between the electron density



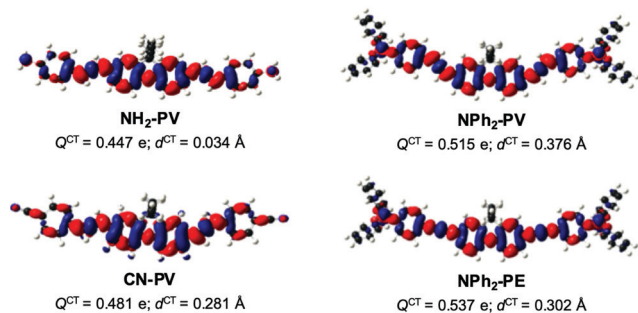


Fig. 6 Plots of the change in total density upon excitation from the ground state S_0 to the first singlet excited state S_1 for **NH₂-PV**, **NPh₂-PV**, **NPh₂-PE** and **CN-PV**. Red and blue colors indicate increase and decrease of electron density, respectively (contour isodensity value: 0.0002 e bohr⁻³). Q^{CT} and d^{CT} are the amount of transferred charge and the distance between the centroids of the positive and negative charges, respectively.

properties, *i.e.*, replacing a double bond by a triple bond induces a blue shift of the emission bands. On the other hand, increasing the electron releasing or withdrawing properties of the peripheral groups leads to a red shift of the emission bands, indicative of a more pronounced either core-to-periphery or periphery-to-core intramolecular charge transfer.

Another question is why does the considered series of fluorene derivatives not exhibit any phosphorescence in toluene? The relative energies of the different ground state S_0 , singlet excited states S_1 and S_2 , and triplet excited state T_1 were computed for the whole series. They are shown in Fig. 7 for **OPh-PV**, **NH₂-PV**, **NPh₂-PV**, and **NPh₂-PE**. Large energy differences ranging from 0.77 to 1.07 eV are computed between S_1 and T_1 states for these molecules. Obviously, this large difference must prevent any intersystem crossing and consequently, phosphorescence is not observed.

Two-photon absorption

The TPA properties of the whole **X-PV** and **X-PE** series of compounds were computed at the SAOP/DZP level under vacuum. One-photon absorption (OPA) spectra were also computed at

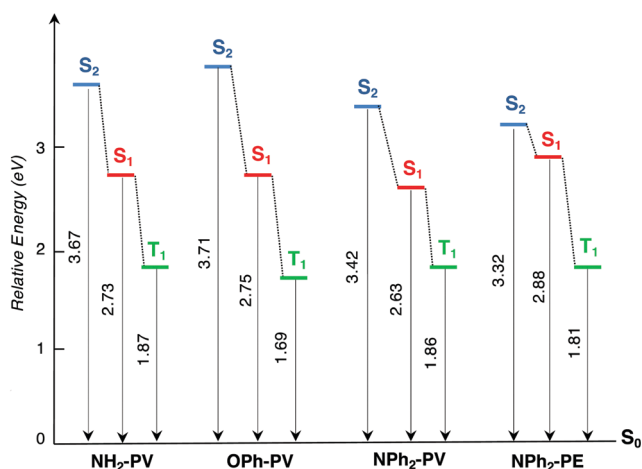


Fig. 7 Relative energy (eV) of the excited states S_1 , S_2 and T_1 with respect to that of S_0 for compounds **NH₂-PV**, **OPh-PV**, **NPh₂-PV**, and **NPh₂-PE** computed at the CAM-B3LYP/6-31+G(d) level in toluene.

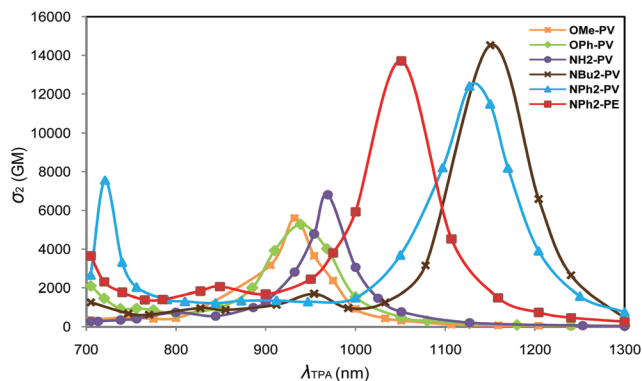


Fig. 8 Plots of the SAOP/DZP computed TPA cross-sections σ_2 vs. wavelength (λ^{TPA}) for **OMe-PV** (orange line), **OPh-PV** (green line), **NH₂-PV** (purple line), **NBu₂-PV** (brown line), **NPh₂-PV** (blue line) and **NPh₂-PE** (red line).

this level of theory for comparison. The geometries used for these calculations were those optimized at the PBE0/6-31+G(d) level in toluene. For quadrupolar systems, the excited S_2 state is generally one-photon forbidden (or at least approximately forbidden in the case of small symmetry distortions) but is two-photon allowed. TPA energies correspond then to the half of the OPA excitation energies to the S_2 state. The TPA simulated spectra, *i.e.*, the plot of the cross-sections as a function of the laser wavelength are displayed in Fig. 8 and 9. For the whole series, the TPA-allowed excitation generally involves a HOMO-1 \rightarrow LUMO transition (Fig. S11, ESI[†]). Experimental^{13,20,22} and theoretical OPA, TPA maximum energies and cross-sections are summarized in Table 5. More detailed information (σ_2 , OPA and TPA energies ω) is provided in Tables S3–S26 (ESI[†]).

As can be seen in Table 5, the computed cross-sections are much higher than the ones experimentally measured. Such a large deviation between the measured and SAOP/DZP computed TPA cross-sections is generally expected and has been discussed in the literature.^{57,58} Indeed, the main reason is that SAOP calculations (i) underestimate the HOMO–LUMO gaps and (ii) neglect solvent effects. Nevertheless, the theoretical

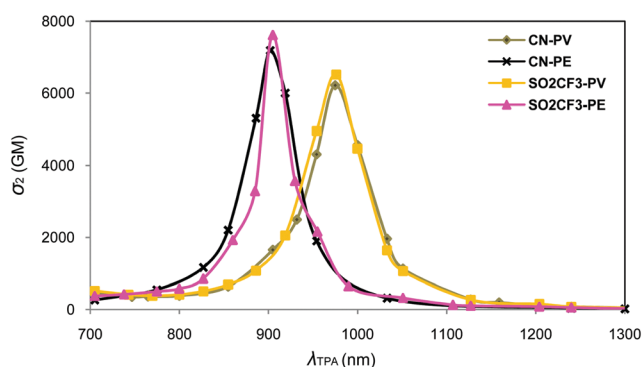


Fig. 9 Plots of the SAOP/DZP computed TPA cross-sections σ_2 vs. wavelength (λ^{TPA}) for **CN-PV** (grey line), **CN-PE** (black line), **SO₂CF₃-PV** (yellow line), and **SO₂CF₃-PE** (pink line).



Table 5 Experimental and computed (for the most stable conformer a) OPA and TPA energies ω (eV), $2\lambda^{\text{OPA}}$ (nm), $\lambda_{\text{max}}^{\text{TPA}}$ (nm) wavelengths, and cross-sections σ_2 (GM) of the **X-PV** and **X-PE** series of compounds

Compound	Exp.			Ref.	Calc. ^d				
	$2\lambda_{\text{max}}^{\text{OPA}}$	$\lambda_{\text{max}}^{\text{TPA}}$	σ_2		ω^{OPA}	ω^{TPA}	$2\lambda^{\text{OPA}}$	$\lambda_{\text{max}}^{\text{TPA}}$	σ_2
OMe-PV	762	705	110	13 ^a	2.65	1.33	936	932	5599
OPh-PV					2.63	1.32	942	939	5266
NH₂-PV	784	705	400	13 ^a	2.55	1.28	972	969	6813
NH₂-PE					2.79	1.40	889	886	7483
NBu₂-PV	830	740	1260	13 ^a	2.12	1.08	1170	1150	14 520
NPh₂-PV	—	745	570	20 ^b	2.17	1.10	1142	1127	12 401
NPh₂-PE	782	720	980	22 ^c	2.35	1.18	1056	1051	13 710
CN-PV					2.53	1.27	980	975	6220
CN-PE					2.73	1.375	908	902	7192
SO₂CF₃-PV					2.54	1.27	976	976	6523
SO₂CF₃-PE	744	730	68	13 ^a	2.73	1.37	908	905	7615
NBu₂-PV-SO₂CF₃					2.55	1.28	972	969	3322
NPh₂-PE-CPDT					2.25	1.13	1102	1097	15 241

^a Experimental values for the TPA cross-sections were obtained *via* two-photon excited fluorescence (TPEF) in toluene using a mode-locked Ti:sapphire laser delivering 80 fs pulses at 80 MHz. The highest TPA cross-section was measured in the 700–1000 nm range. ^b TPA cross-sections were measured by using the two-photon-induced fluorescence method in toluene using a mode-locked Ti:sapphire laser delivering 120 fs pulses at 76 MHz in the wavelength range of 710–1000 nm. ^c TPA maximal values in dichloromethane determined by TPEF in the femto-second regime. TPEF cross-sections were measured relative to fluorescein in 0.01 M aqueous NaOH over the range 715–980 nm, with the appropriate solvent-related refractive index corrections. ^d Highest TPA cross-section calculated at the SAOP/DZP level in the 700–1300 nm range.

results respect the trend experimentally observed for **OMe-PV**, **NH₂-PV**, **NPh₂-PV** and **NBu₂-PV**.^{13,20} Additionally, the increase in intensity of the maximum TPA cross-sections is well correlated with the increase of the electron-donor properties of the end-groups, *i.e.*, **OPh-PV** < **OMe-PV** < **NH₂-PV** < **NPh₂-PV** < **NBu₂-PV** (Table 5 and Fig. 8). We note also that a red shift of both one- and two-photon absorption bands of push–push derivatives occurs, as illustrated in Fig. 8 and observed from Table 5. Moreover, as observed experimentally, comparing the computed TPA cross-sections of both **NPh₂-PV** and **NPh₂-PE** (6813 *vs.* 7483 GM) shows that replacing in the conjugated bridge a carbon–carbon double bond by a triple one slightly increases the NLO response in agreement with the experimental results.^{20,22} This result is also confirmed by comparing the **CN-PV** and **CN-PE** systems (6220 *vs.* 7192 GM).

To summarize, the extension of the π -conjugated connectors and the increase in the strength of the donor terminal groups enhance not only the photoluminescence properties, but also allow obtaining rather good two-photon absorbers on the edge of the NIR domain.

The simulated spectra obtained for the molecules containing electron-withdrawing end-groups (SO₂CF₃ and CN) are shown in Fig. 9. Comparable energies and intensity of the cross-sections are observed for **SO₂CF₃-PV** and **CN-PV**. A blue shift and a decrease of the intensity are observed relative to the molecules containing electron-releasing groups (see above). As observed earlier for the latter systems, the replacement of the vinyl group (**SO₂CF₃-PV**) by an alkyne one (**SO₂CF₃-PE**) is

accompanied with a substantial blue shift and a slight increase of the TPA cross-sections in the NIR domain (Fig. 9). A similar effect is observed with the electron-withdrawing CN group. It turns out that the nature of the connector unit (C=C or C≡C) in these conjugated rods plays a non-negligible role in tuning the TPA spectra and influencing the TPA cross-section magnitude. Regardless of the nature of the end-groups (donor or acceptor), the core moiety or the length of the conjugated bridges, replacement of a double bond (C=C) by a triple bond (C≡C) always leads to a blue shift in energy and an increase and broadening of the TPA cross-sections in the NIR region. As a result, all fluorophores built from vinylene linkers instead of ethynylene ones exhibit lower TPA cross-sections in the whole red-NIR region. This result was not expected since higher energies gaps are often accompanied by a lowering of the cross-section. This is of particular importance when imaging applications are concerned because (i) improved penetration depth is achieved when shifting to higher wavelength (due to reduction of scattering losses) and (ii) spectral broadening offers much more flexibility in terms of two-photon excitation (allowing for a wider choice of laser sources).

We found it interesting to study the linear and non-linear optical properties of an unsymmetrical donor–bridge–acceptor molecule, namely the push–pull system **NBu₂-PV-SO₂CF₃**, containing the NBu₂ electron donor group on one hand and the electron acceptor SO₂CF₃ group on the other hand as terminal groups. The TPA spectrum of **NBu₂-PV-SO₂CF₃** exhibits a band with a maximum value at 969 nm and a cross-section at 3322 GM (Table 5 and Table S26, ESI†). Such a push–pull system has a lower cross-section than both the pull–pull **NBu₂-PV** (14 520 GM at 1150 nm) and the push–push **SO₂CF₃-PV** (6523 GM at 976 nm) species. Indeed, this confirms that quasi one-dimensional quadrupolar systems, *i.e.*, symmetrical conjugated molecules bearing electron-releasing or electron-withdrawing end-groups, have been found to be more efficient than push–pull systems in terms of TPA.¹³

Finally, we may wonder what is the effect of the fluorene group present in all molecules that we studied on the TPA response? For that, the two phenyl groups of the **NPh₂-PE** core

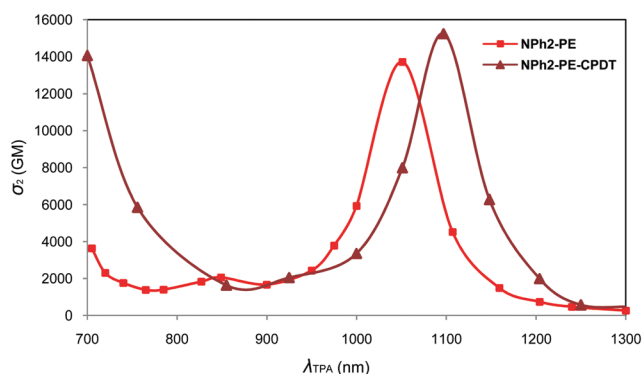


Fig. 10 Plots of the SAOP/DZP computed TPA cross-sections σ_2 *vs.* wavelength (λ^{TPA}) for **NPh₂-PE** and **NPh₂-PE-CPDT**.



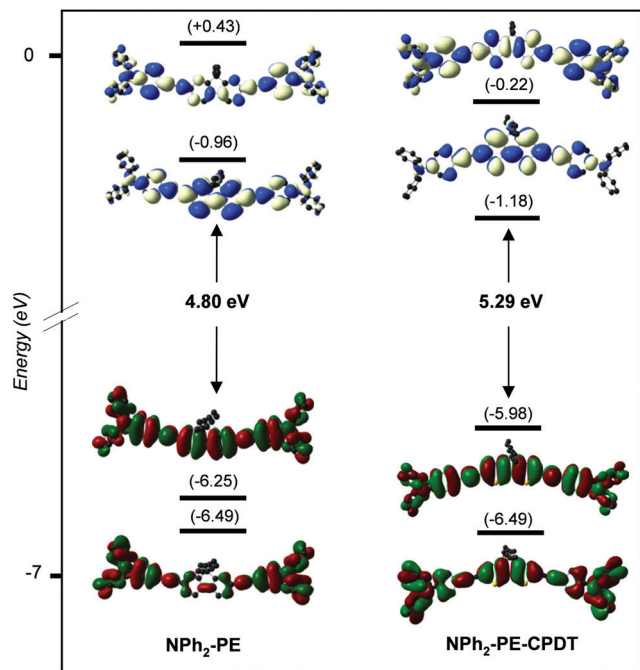


Fig. 11 CAM-B3LYP/6-31+G(d) frontier MO diagrams of **NPh₂-PE** (left) and **NPh₂-PE-CPDT** (right). Contour isodensity values: ± 0.015 ($e \text{ bohr}^{-3/2}$). Hydrogen atoms are omitted for clarity.

Table 6 Computed photo-physical properties (absorption and emission wavelengths λ (nm and eV (in brackets)), OPA and TPA energies ω (eV), and cross-sections σ_2 (GM)) of **NPh₂-PE** and **NPh₂-PE-CPDT**^a

Compound	$\lambda_{\text{max}}^{\text{abs}}$	$\lambda_{\text{max}}^{\text{em-elect}}$	E_{gap}^b	ω^{OPA}	ω^{TPA}	σ_2
NPh₂-PE	382 (3.25)	430 (2.88)	3.06	2.35	1.18	13 710
NPh₂-PE-CPDT	454 (2.73)	543 (2.28)	2.51	2.25	1.13	15 241

^a CPDT = 4*H*-cyclopenta[2,1-*b*:3,4-*b'*]dithiophene. ^b The electronic gap (E_{gap}) is calculated from the absorption and emission maxima energy difference.

were replaced by two thiophene rings (**NPh₂-PE-CPDT**). Their TPA spectra and CAM-B3LYP/6-31+G(d) frontier MO diagrams are shown in Fig. 10 and 11, respectively, and their main photo-physical properties are given in Table 6. Corresponding SAOP/DZP frontier MO diagrams are shown in Fig. S12 (ESI[†]) – they are qualitatively comparable to the CAM-B3LYP/6-31+G(d) ones. This replacement leads to a significant increase and a red shift of the TPA cross-section. As seen before, the TPA energy corresponds to half of the OPA excitation energy to state S_2 . For the previous quadrupolar systems, this excitation allowed by TPA generally corresponds to a HOMO–1 \rightarrow LUMO transition (see Fig. S12, ESI[†]). With the dithiophene core, the transition to the S_2 state is HOMO \rightarrow LUMO+1. The blue shift in energy observed with **NPh₂-PE-CPDT** is due to the destabilization of the HOMO which is shifted higher in energy by 0.29 eV, with respect to that of **NPh₂-PE**. These results demonstrate that the nature of the central conjugated core significantly influences the TPA spectra also and governs the TPA cross-section magnitude of this type of molecules.

Conclusion

This systematic theoretical study of the absorption, photoluminescence and two-photon absorption properties of a large series of quadrupolar fluorophores, made of a fluorene core bearing conjugated bridges and electron donor or acceptor end-groups, allowed us to extract some structure–property relationships of great significance for both spectral tuning and amplification of molecular TPA in the NIR spectral domain. First, their different possible geometrical configurations were optimized using DFT computations. Then, TD-DFT calculations were carried out showing absorption at low wavelengths for all compounds around 400 nm or below, due to HOMO–LUMO transition of rather large energy, with high oscillator strength in agreement with the available experimental data. The computed fluorescence wavelengths are also in good agreement with the experiments. The presence of the fluorene core and the phenylene–vinylene (PV) or phenylene–ethynylene (PE) conjugated bridges leads to high efficiency for light harvesting. The influence of each moiety constituting these quadrupolar structures (cores, linkers, connectors and end-groups) was investigated. Regarding the non-linear optical properties, the theoretical results confirm that the excited state (S_2) reached by the two-photon absorption of lowest energy corresponds to the HOMO–1 \rightarrow LUMO transition for all compounds. It is found that an increase of the strength of the donor terminal group (OPh, OMe, NH₂, NPh₂, and NBu₂) enhances the TPA cross-section. Planarization of the fluorene core always leads to an increase in the TPA cross-sections. The nature of the linkers (PV or PE) and the phenylene units is of particular importance as well. Small changes in the structures may have important implications in terms of molecular engineering for specific applications, since the TPA properties (in terms of cross-sections, position of the maximum and band width), as well as the one-photon and the photoluminescence characteristics can be considerably affected. Interestingly, replacement of the fluorene core by a bithiophene one could enhance two-photon absorption. With these findings in hand, quadrupolar fluorophores, combining important peak TPA cross-sections and broad TPA bands in the whole 700–1300 nm range, could thus be obtained.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

S. G. and S. M. are grateful to the Tunisian Ministry of Higher Education and Scientific Research for financial support. GENCI-IDRIS and GENCI-CINES are acknowledged for computing time allocation (grant 2020-2021-080649). The authors thank Dr Olivier Mongin (University of Rennes) for his valuable comments.



References

- M. Albota, D. Beljonne, J.-L. Brédas, J. E. Ehrlich, J.-Y. Fu, A. A. Heikal, S. E. Hess, T. Kogej, M. D. Levin, S. R. Marder, D. McCord-Maughon, J. W. Perry, H. Röckel, M. Rumi, G. Subramaniam, W. W. Webb, X.-L. Wu and C. Xu, *Science*, 1998, **281**, 1653–1656.
- L. Ventelon, L. Moreaux, J. Mertz and M. Blanchard-Desce, *Chem. Commun.*, 1999, 2055–2056.
- M. Rumi, J. E. Ehrlich, A. A. Heikal, J. W. Perry, S. Barlow, Z. Hu, D. McCord-Maughon, T. C. Parker, H. Röckel, S. Thayumanavan, S. R. Marder, D. Beljonne and J.-L. Brédas, *J. Am. Chem. Soc.*, 2000, **122**, 9500–9510.
- M. Blanchard-Desce, *C. R. Phys.*, 2002, **3**, 439–448.
- O. Mongin, L. Porrès, L. Moreaux, J. Mertz and M. Blanchard-Desce, *Org. Lett.*, 2002, **4**, 719–722.
- M. G. Silly, L. Porrès, O. Mongin, P.-A. Chollet and M. Blanchard-Desce, *Chem. Phys. Lett.*, 2003, **379**, 74–80.
- K. D. Belfield, M. V. Bondar, A. R. Morales, O. Yavuz and O. V. Przhonska, *J. Phys. Org. Chem.*, 2003, **16**, 194–201.
- K. D. Belfield, A. R. Morales, J. M. Hales, D. J. Hagan, E. W. Van Stryland, V. M. Chapela and J. Percino, *Chem. Mater.*, 2004, **16**, 2267–2273.
- C. D. Entwistle and T. B. Marder, *Chem. Mater.*, 2004, **16**, 4574–4585.
- S. Yao and K. D. Belfield, *J. Org. Chem.*, 2005, **70**, 5126–5132.
- Z. Yuan, C. D. Entwistle, J. C. Collings, D. Albesa-Jové, A. S. Batsanov, J. A. K. Howard, N. J. Taylor, H. M. Kaiser, D. E. Kaufmann, S.-Y. Poon, W.-Y. Wong, C. Jardin, S. Fathallah, A. Boucekkine, J.-F. Halet and T. B. Marder, *Chem. Eur. J.*, 2006, **12**, 2758–2771.
- M. Charlot, N. Izard, O. Mongin, D. Riehl and M. Blanchard-Desce, *Chem. Phys. Lett.*, 2006, **417**, 297–302.
- O. Mongin, L. Porrès, M. Charlot, C. Katan and M. Blanchard-Desce, *Chem. – Eur. J.*, 2007, **13**, 1481–1498.
- Y. Tian, C. Y. Chen, Y. J. Cheng, A. C. Young, N. M. Tucker and A. Y. Jen, *Adv. Funct. Mater.*, 2007, **17**, 1691–1697.
- C. D. Entwistle, J. C. Collings, A. Steffen, L. O. Pålsson, A. Beeby, D. Albesa-Jové, J. M. Burke, A. S. Batsanov, J. A. K. Howard, J. A. Mosely, S. Y. Poon, W. Y. Wong, F. Ibersiene, S. Fathallah, A. Boucekkine, J.-F. Halet and T. B. Marder, *J. Mater. Chem.*, 2009, **19**, 7532–7544.
- K. D. Belfield, M. V. Bondar, C. O. Yanez, F. E. Hernandez and O. V. Przhonska, *J. Mater. Chem.*, 2009, **19**, 7498–7502.
- C. Katan, S. Tretiak and J. Even, Nanophotonics III, in: *Proc. SPIE*, ed. D. L. Andrews, J.-M. Nunzi and A. Ostendorf, 2010, vol. 7712, p. 77123D.
- S. Yao, H.-Y. Ahn, X. Wang, J. Fu, E. W. Van Stryland, D. J. Hagan and K. D. Belfield, *J. Org. Chem.*, 2010, **75**, 3965–3974.
- C. D. Andrade, C. O. Yanez, L. Rodriguez and K. D. Belfield, *J. Org. Chem.*, 2010, **75**, 3975–3982.
- C. Rouxel, M. Charlot, Y. Mir, C. Frochot, O. Mongin and M. Blanchard-Desce, *New J. Chem.*, 2011, **35**, 1771–1780.
- A. R. Morales, A. Frazer, A. W. Woodward, H.-Y. Ahn-White, A. Fonari, P. Tongwa, T. Timofeeva and K. D. Belfield, *J. Org. Chem.*, 2013, **78**, 1014–1025.
- Z. Pokladek, N. Ripoche, M. Betou, Y. Trolez, O. Mongin, J. Olesiak-Banska, K. Matczyszyn, M. Samoc, M. G. Humphery, M. Blanchard-Desce and F. Paul, *Chem. – Eur. J.*, 2016, **22**, 10155–10167.
- S. A. Kurhuzenkau, A. W. Woodward, S. Yao, K. D. Belfield, Y. O. Shaydyuk, C. Sissa, M. V. Bondar and A. Painelli, *Phys. Chem. Chem. Phys.*, 2016, **18**, 12839–12846.
- S. A. Kurhuzenkau, M. Y. Colon Gomez, K. D. Belfield, Y. O. Shaydyuk, D. J. Hagan, E. W. Van Stryland, C. Sissa, M. V. Bondar and A. Painelli, *J. Phys. Chem. C*, 2018, **122**, 5664–5672.
- J. He, F. Rauch, A. Friedrich, D. Sieh, T. Ribbeck, I. Krummenacher, H. Braunschweig, M. Finze and T. B. Marder, *Chem. – Eur. J.*, 2019, **25**, 13777–13784.
- F. Terenziani, C. Katan, E. Badaeva, S. Tretiak and M. Blanchard-Desce, *Adv. Mater.*, 2008, **20**, 4641–4678.
- K. A. Nguyen, P. N. Day and R. Pachter, *Theor. Chem. Acc.*, 2008, **120**, 167–175.
- L. K. Yan, A. Pomogaeva, F. L. Gu and Y. Aoki, *Theor. Chem. Acc.*, 2010, **125**, 511–520.
- G. L. C. Moura and A. M. Simas, *J. Phys. Chem. C*, 2010, **114**, 6106–6116.
- X.-T. Liu, L.-Y. Zou, A.-M. Ren, J.-F. Guo, Y. Sun, S. Huang and J.-K. Feng, *Theor. Chem. Acc.*, 2011, **130**, 37–50.
- Y. Zhao, J.-F. Guo, A.-M. Ren and J.-K. Feng, *Theor. Chem. Acc.*, 2011, **128**, 265–274.
- G. Argouarch, R. Veillard, T. Roisnel, A. Amar, A. Boucekkine, A. Singh, I. Ledoux and F. Paul, *New J. Chem.*, 2011, **35**, 2409–2411.
- D. W. Silverstein and L. Jensen, *J. Phys. Chem.*, 2012, **136**, 064111.
- I. H. Nayyar, A. E. Masunov and S. Tretiak, *J. Phys. Chem. C*, 2013, **117**, 18170–18189.
- M. T. Beerepoot, D. H. Friese, N. H. List, J. Kongsted and K. Ruud, *Phys. Chem. Chem. Phys.*, 2015, **17**, 19306–19314.
- H. Zhongwei, J. Autschbach and L. Jensen, *J. Chem. Theory Comput.*, 2016, **12**, 1294–1304.
- M. M. Alam and C. Daniel, *Theor. Chem. Acc.*, 2016, **135**, 41.
- M. Abe, Y. Chitose, S. Jakkampudi, P. T. T. Thuy, Q. Lin, B. T. Van, A. Yamada, R. Oyama, M. Sasaki and C. Katan, *Synthesis*, 2017, 3337–3346.
- O. Sengul, E. B. Boydas, M. Pastore, W. Sharmouk, P. C. Gros, S. Catak and A. Monari, *Theor. Chem. Acc.*, 2017, **136**, 67.
- M. T. Beerepoot, M. M. Alam, J. Bednarska, W. Bartkowiak, K. Ruud and R. Zalesny, *J. Chem. Theory Comput.*, 2018, **14**, 3677–3685.
- A. Triadon, A. N. Ndimba, N. Richey, A. Amar, A. Boucekkine, T. Roisnel, M. P. Cifuentes, M. G. Humphrey, M. Blanchard-Desce and O. Mongin, *New J. Chem.*, 2018, **42**, 11289–11293.
- S.-J. Chung, M. Rumi, V. Alain, S. Barlow, J. W. Perry and S. R. Marder, *J. Am. Chem. Soc.*, 2005, **127**, 10844–10845.
- M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, B. Mennucci, G. A. Petersson, H. Nakatsuji, M. Caricato, X. Li, H. P. Hratchian, A. F. Izmaylov, J. Bloino, G. Zheng,



- J. L. Sonnenberg, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. Bearpark, J. J. Heyd, E. Brothers, K. N. Kudin, V. N. Staroverov, T. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, N. Rega, J. M. Millam, M. Klene, J. E. Knox, J. B. Cross, V. Bakken, C. Adamo, J. Jaramillo, R. Gomperts, R. E. Stratmann, O. Yazyev, A. J. Austin, R. Cammi, C. Pomelli, J. W. Ochterski, R. L. Martin, K. Morokuma, V. G. Zakrzewski, G. A. Voth, P. Salvador, J. J. Dannenberg, S. Dapprich, A. D. Daniels, O. Farkas, J. B. Foresman, J. V. Ortiz, J. Cioslowski and D. J. Fox, *Gaussian 09, Revision D.01*, Gaussian, Inc., Wallingford CT, 2013.
- 44 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman and D. J. Fox, *Gaussian 16, Revision C.01*, Wallingford, CT, 2016.
- 45 M. Cossi, G. Scalmani, N. Rega and V. Barone, *J. Chem. Phys.*, 2002, **117**, 43–54.
- 46 V. Barone, M. Cossi and J. Tomasi, *J. Chem. Phys.*, 1997, **107**, 3210–3221.
- 47 J. Tomasi, B. Mennucci and R. Cammi, *Chem. Rev.*, 2005, **105**, 2999–3094.
- 48 S. Miertuš, E. Scrocco and J. Tomasi, *Chem. Phys.*, 1981, **55**, 117–129.
- 49 C. Adamo and V. Barone, *J. Chem. Phys.*, 1999, **110**, 6158–6170.
- 50 T. Yanai, D. P. Tew and N. C. Handy, *Chem. Phys. Lett.*, 2004, **393**, 51–57.
- 51 R. Dennington, T. Keith and J. Millam, *GaussView, version 5.0*, 2009.
- 52 ADF, Amsterdam Density Functional, SCM Computing & Modelling, Theoretical Chemistry, Vrije Universiteit, Amsterdam, The Netherlands, <http://www.scm.com>.
- 53 G. teVelde, F. M. Bickelhaupt, E. J. Baerends, C. Fonseca Guerra, S. J. A. van Gisbergen, J. G. Snijders and T. Ziegler, *J. Comput. Chem.*, 2001, **22**, 931–967.
- 54 P. R. Schipper, O. V. Gritsenko, S. J. van Gisbergen and E. J. Baerends, *J. Phys. Chem.*, 2000, **112**, 1344–1352.
- 55 L. Jensen, P. T. van Duijnen and J. G. Snijders, *J. Chem. Phys.*, 2003, **119**, 12998–13006.
- 56 ADFGUI 2012, SCM, Amsterdam, The Netherlands, <http://www.scm.com>.
- 57 A. Amar, A. Elkechai, J.-F. Halet, F. Paul and A. Boucekkine, *New J. Chem.*, 2021, **45**, 15074–15081.
- 58 A. Amar, A. Boucekkine, F. Paul and O. Mongin, *Theor. Chem. Acc.*, 2019, **138**, 105.
- 59 L. Jensen, J. Autschbach and G. C. Schatz, *J. Chem. Phys.*, 2005, **122**, 224115.
- 60 H. Zhongwei, J. Autschbach and L. Jensen, *J. Chem. Phys.*, 2014, **141**, 124305.
- 61 M. Göppert-Mayer, *Ann. Phys.*, 1931, **9**, 273–294.
- 62 J. Pérez Moreno and M. G. Kuzyk, *J. Chem. Phys.*, 2005, **123**, 194101.
- 63 J. Pérez-Moreno, K. Clays and M. G. Kuzyk, *J. Chem. Phys.*, 2008, **128**, 084109.
- 64 T. Le Bahers, C. Adamo and I. Ciofini, *J. Chem. Theory Comput.*, 2011, **7**, 2498–2506.

