## **NJC**



View Article Online PAPER



Cite this: New J. Chem., 2023, **47**, 1234

# nonlinear optical properties, a theoretical investigation<sup>†</sup>

Bis-TTF-Ge derivatives: promising linear and

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In this work, based on bis(tetrathiafulvalenyldithio)-germane (bis-TTF-Ge), 37 compounds (denoted T0-T36) are designed by the introduction of donor and acceptor groups at different substituent positions. The ground state electronic structures, reactivity indices, electronic transition, charge transfer properties (charge, distance and dipole moment), hyper-Rayleigh scattering and depolarization ratios (static and dynamic) of these compounds are fully investigated using DFT and TD-DFT calculations. The quantum calculations were performed at the CAM-B3LYP/6-311g(d,p) level and using the sum-over-states (SOS) approach in both static and dynamic cases. The chemical hardness (n) of the two substituted bis-TTF-Ge derivatives at the **b** position has a smaller value than those with the same substituent at positions **d**, **e**, **a** and c. In both static and dynamic regimes the investigations show that the bis-TTF-Ge derivatives substituted with donor or acceptor groups have larger first hyperpolarizability than T0. Our work predicts that the introduction of the substituent group at the  $\mathbf{b}$  position can increase the hyperpolarizability values more than those at the other positions, and the NO2, NO and COCN acceptor groups lead to the largest values. For example, the dynamic  $\beta_{HRS}^{2=1064}$  value of T25 is about 145 times larger than that of T0 and about 11 times that in the static regime. Interestingly, the  $\beta_{HRS}^{\lambda=1064}$  of bis-TTF-Ge substituted with the NO<sub>2</sub> group increases with the number of NO2 in the TTF fragment. The large nonlinear optical (NLO) origin of bis-TTF-Ge substituted with NO2, NO and COCN is attributed to charge transfer from the TTF to the acceptor group at the second TTF fragment (overlap  $S_r$  index = 0). The studied bis-TTF-Ge substituted with  $NO_2$ compounds exhibits the possibility of being an excellent second-order NLO material.

Received 25th July 2022, Accepted 2nd December 2022

DOI: 10.1039/d2nj03671a

rsc.li/njc

## Introduction

Nonlinear optics (NLO) is the study of the interaction of high intensity light with matter. In linear optics the relation between polarization (P) and electric field (E) is linear and the frequency is kept. For an intense laser light, this linearity no longer holds and one gets different kinds of frequencies; this frequency transformation has been observed experimentally by Franken,1 and the intense light induces a change in the refraction index of the material (optical Kerr effect), which leads to a phenomenon called the self-phase modulation.<sup>2</sup> The induced polarization at the macroscopic level is generally written as3

$$\Delta P = \gamma^{(1)}E + \gamma^{(2)}E^2 + \gamma^{(3)}E^3 + \cdots$$

 $\chi^{(1)}$ ,  $\chi^{(2)}$  and  $\chi^{(3)}$  are the first, second and third order susceptibility tensors, respectively.  $\chi^{(1)}$  belongs to the domain of linear optics, whereas  $\chi^{(2)}$  and  $\chi^{(3)}$  quantifies the second and third order NLO effects, respectively.

Since the discovery of lasers by Maiman and Collins in 1960<sup>4,5</sup> great interest has been devoted to the discovery and design of new materials with excellent nonlinear optical properties. This is due to their important applications in different areas such as in photonic devices such as molecular switching, frequency doubling, telecommunications and optical information processing.6-12

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<sup>†</sup> Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d2nj03671a

Paper

Numerous techniques for developing new high-performance NLO materials have been proposed, including diradical characterization, 13 synthesis of molecular structures of electrondonor (D)-π-electron-acceptor (A) (D-π-A), 14 reinforcing pushpull effects, 15,16 designing octupolar compounds, 17,18 multidecker sandwich complexes, 19 asymmetric coordination complexes, 20-23 excess electrons, 24-29 etc. Among the studied NLO materials the inorganic compounds have attracted exceptional interest because of their physicochemical stability and thermal stability. 30,31 For example, tetrathiafulvalene (TTF) derivatives present excellent second- and third-order nonlinear optical responses, and they are a very active field in contemporary chemistry since shortly after their first synthesis in the early 1970s by Wudl and coworkers. 32,33 These properties are derived from the large  $\pi$  conjugated systems, the strong electron-donor character and the existence of polarizable sulfur atoms, leading to charge density redistribution within the molecule. 34-36

The combination of the TTF group and electron acceptor (A) units that are covalently linked by a  $\pi$ -conjugated bridging group is highly desirable and classified as excellent push-pull organic molecules (D- $\pi$ -A). So far, TTF- $\pi$ -A as promising NLO materials have been widely studied by many scientific researchers, because these compounds exhibit intramolecular charge transfer (ICT) and have low energy-gaps. 37-39

On the other hand, the binding of the TTF group with transition metals (TM) through covalent linkages or via electrostatic interactions allows the development of new metal complexes with  $\pi$ -conjugated ligands and can present a respectable class of NLO materials. 36,40-42 For example, in 2015, Liu and coworkers studied static and dynamic HRS hyperpolarizability of a series of extended tetrathiafulvalene (exTTF) and TTF derivatives of metal cations (TM: Ni<sup>2+</sup>, Cu<sup>2+</sup>, Mg<sup>2+</sup>, Zn<sup>2+</sup> and Cd<sup>2+</sup>) and they found that the TM complexes containing the redox-active TTF unit and the pyridine group exhibit large hyperpolarizability values.<sup>43</sup> In addition, the nonlinear optical responses of several square-planar metal TTF complexes<sup>44,45</sup> have been studied theoretically and experimentally and have shown the important role of the ligand in perturbing the electron distribution in these complexes and its effect on both the ground and the excited states and, thus, on the linear and nonlinear optical properties. 46-51 In recent years growing attention has been expressed concerning germanium, and this interest is due to its high carrier mobility, good stability so that Ge is the most logical candidate to substitute carbon; moreover, it has also potential applications in photodetectors and nonlinear optical fields.<sup>52-60</sup>

Ueda et al.61 reported spiro molecules: bis(tetrathiafulvalenyldithio)-germane (here named bis-TTF-Ge T36, see Fig. 1). This asymmetric compound which contains large  $\pi$  systems should be a potential second-order NLO material. In this work, our aim is to study the effects of acceptor/donor substituents (NO, CN, COCN, CHO, NO<sub>2</sub>, NH<sub>2</sub> and NMe<sub>2</sub>) and the substituent positions (a, b, c, d and e; see Fig. 1) on the reactivity indices, and the linear and nonlinear optical properties of bis-TTF-Ge compounds. Also in this study, we focus to offer new perspectives with respect to the origin of nonlinear optical materials for title compounds. Furthermore, to the best of

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R1=R2=R3=R4=H
                                 T18 R1=R4=CHO R2=R3=H
    R1= NO, R2=R3=R4=H
                                 T19 R1=R2=R3=CHO, R4=H
    R1= R2= NO R3=R4=H
T2
                                 T20 R1=R2=R3=R4= CHO
    R1=R4=NO, R2=R3=H
                                 T21 R1= NO2, R2=R3=R4=H
T3
T4
    R1=R2=R3=NO, R4=H
                              b
                                 T22 R1= R2= NO2, R3=R4=H
    R1=R2=R3=R4= NO
                                 T23 R1=R4=NO2, R2=R3=H
T5
    R1= CN, R2=R3=R4=H
                                 T24 R1=R2=R3=NO2, R4=H
    R1= R2= CN, R3=R4=H
                                 T25 R1=R2=R3=R4= NO2
T7
    R1=R4=CN, R2=R3=H
                                 T26 R1= NH2, R2=R3=R4=H
Т9
    R1=R2=R3=CN, R4=H
                              b
                                 T27 R1= R2= NH2, R3=R4=H
T10 R1=R2=R3=R4= CN
                                 T28 R1=R4=NH2 R2=R3=H
T11 R1= COCN, R2=R3=R4=H
                                 T29 R1=R2=R3=NH2, R4=H
T12 R1= R2= COCN, R3=R4=H
                                 T30 R1=R2=R3=R4= NH2
T13 R1=R4=COCN, R2=R3=H
                                 T31 R1= NMe2, R2=R3=R4=H
T14 R1=R2=R3=COCN, R4=H
                                 T32 R1= R2= NMe2, R3=R4=H
T15 R1=R2=R3=R4= COCN
                                 T33 R1=R4=NMe2, R2=R3=H
T16 R1= CHO, R2=R3=R4=H
                                 T34 R1=R2=R3=NMe2 R4=H
T17 R1= R2= CHO, R3=R4=H
                                 T35 R1=R2=R3=R4= NMe2
                                 T36 R1=R2=R3=R4=Me
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Fig. 1 Chemical structures of the studied bis-TTF T0 to T36, where a, b, c. d and e are the substituent position (a: one substituent at R1; b: two substituents at R1 and R2; c: two substituents at R1 and R4; d: three substituents R1. R2 and R3: e: four substituents).

our knowledge, there are no computational studies that address in detail the effect of the number and the nature of substituent groups in compounds made of two TTF groups linked by spiro germanium on the NLO properties.

## Computational details

The geometries of the bis-TTF-Ge derivatives were fully optimized using Coulomb-attenuating exchange-correlation density functional method CAM-B3LYP<sup>62</sup> using the 6-311g(d,p)<sup>63,64</sup> basis set. CAM-B3LYP is a long-range corrected functional developed to handle the inaccuracies of the non-Coulomb part of the exchange functional at long distances. Quantum chemical calculations were performed using the Gaussian 09 program with the TIGHT SCF convergence and ultra-fine integration grid. 65-67 No symmetry constraints were applied and the local minima were confirmed on potential energy surface by frequency calculations of the ground state for bis-TTF-Ge compounds at the same level.

Chemical reactivity analysis takes advantage of its ground in DFT through a functional Taylor expansion of energy that introduces several reactivity descriptors from energy derivatives of chemical significance.<sup>68</sup> According to finite difference approximation and Koopmans' theorem<sup>69-71</sup> the electronic chemical potential  $(\mu)$  and global hardness  $(\eta)$  can be evaluated from the frontier orbital energies HOMO and LUMO ( $\varepsilon_{\rm H}$  and  $\varepsilon_{\rm L}$ , respectively).

Electronic chemical potential  $(\mu)$  is the tendency of an atom or molecule to attract electrons; it is given by the following equation:

$$\mu = \frac{1}{2}(\varepsilon_{\rm H} + \varepsilon_{\rm L}) = -\chi \tag{1}$$

 $\chi$  is the electronegativity.

The chemical hardness  $(\eta)$  expresses the resistance of a system to change its number of electrons; it is calculated using the relation

$$\eta = \varepsilon_{\rm L} - \varepsilon_{\rm H} \tag{2}$$

It is important to note that the global hardness ( $\eta$ ) measures the stability of a system in terms of resistance to electron transfer and the chemical potential  $(\mu)$  characterizes the escaping tendency of electrons from the equilibrium system.

Global electrophilicity index ( $\omega$ ), introduced by Parr et al., <sup>72</sup> is calculated from the hardness and chemical potential:

$$\omega = \frac{\mu^2}{2\eta} \tag{3}$$

This index expresses the ability of a molecule to accept electrons from the environment.

Furthermore, to investigate and understand the electronic transition and optical properties, the electron excitation energies, oscillator strengths, for the studied compounds were calculated at the TD-CAM-B3LYP level.

The hyper-Rayleigh scattering (HRS) technique involves the detection of the incoherent scattering second harmonic generation at frequency  $2\omega$  from a laser with frequency  $\omega$  incidence. In HRS, we can evaluate the first hyperpolarizability  $\beta_{HRS}$  and its depolarization ratios DR, through the relation<sup>6,73,74</sup>

$$\langle \beta_{\rm HRS} \rangle = \sqrt{\{\langle \beta_{ZZZ}^2 \rangle + \langle \beta_{XZZ}^2 \rangle\}}$$
 (4)

$$DR = \frac{\langle \beta_{ZZZ}^2 \rangle}{\langle \beta_{YZZ}^2 \rangle}$$
 (5)

where  $\langle \beta_{ZZZ}^2 \rangle$  and  $\langle \beta_{XZZ}^2 \rangle$  are orientational averages of the  $\beta$ tensor without assuming Kleinman's conditions:

$$\langle \beta_{ZZZ}^{2} \rangle = \frac{1}{7} \sum_{i}^{x,y,z} \beta_{iii}^{2} + \frac{4}{35} \sum_{i \neq J}^{x,y,z} \beta_{iij}^{2} + \frac{2}{35} \sum_{i \neq J}^{x,y,z} \beta_{iii} \beta_{ijj} + \frac{4}{35} \sum_{i \neq J}^{x,y,z} \beta_{jii} \beta_{iij}$$

$$+ \frac{4}{35} \sum_{i \neq J}^{x,y,z} \beta_{iii} \beta_{jji} + \frac{1}{35} \sum_{i \neq J}^{x,y,z} \beta_{Jii}^{2} + \frac{4}{105} \sum_{i \neq J \neq k}^{x,y,z} \beta_{iij} \beta_{jkk}$$

$$+ \frac{1}{105} \sum_{i \neq J \neq k}^{x,y,z} \beta_{jii} \beta_{jkk} + \frac{4}{105} \sum_{i \neq J \neq k}^{x,y,z} \beta_{iij} \beta_{kkj} + \frac{2}{105} \sum_{i \neq J \neq k}^{x,y,z} \beta_{ijk}^{2}$$

$$+ \frac{4}{105} \sum_{i \neq J \neq k}^{x,y,z} \beta_{ijk} \beta_{jik}$$

$$\begin{split} \left<\beta_{XZZ}^{2}\right> &= \frac{1}{35} \sum_{i}^{x,y,z} \beta_{iii}^{2} + \frac{4}{105} \sum_{i \neq J}^{x,y,z} \beta_{iii} \beta_{ijj} - \frac{2}{35} \sum_{i \neq J}^{x,y,z} \beta_{iii} \beta_{jji} \\ &+ \frac{8}{105} \sum_{i \neq J}^{x,y,z} \beta_{iij}^{2} + \frac{3}{35} \sum_{i \neq J}^{x,y,z} \beta_{ijj}^{2} - \\ &+ \frac{1}{35} \sum_{i \neq J \neq k}^{x,y,z} \beta_{ijj} \beta_{ikk} - \frac{2}{105} \sum_{i \neq J \neq k}^{x,y,z} \beta_{iik} \beta_{jjk} + \frac{2}{35} \sum_{i \neq J \neq k}^{x,y,z} \beta_{ijk}^{2} \\ &- \frac{2}{105} \sum_{i \neq J \neq k}^{x,y,z} \beta_{iij} \beta_{jkk} - \frac{2}{105} \sum_{i \neq J \neq k}^{x,y,z} \beta_{ijk} \beta_{jik} \end{split}$$

Previous works have pointed out that CAM-B3LYP, adopted in this work, can provide an absorption spectrum close to experimental results and has been proven to be efficient in calculating the static and dynamic NLO responses. 7,75-77 Frequency-dependent hyperpolarizabilities were calculated using an incident wavelength of 1064 nm.

The electron density difference maps (EDDM) corresponding to the crucial excited states can be accurately evaluated as follows:

$$\Delta \rho(r) = \rho_{\rm ex}(r) - \rho_{\rm GS}(r)$$

where  $\rho_{\rm ex}(r)$  and  $\rho_{\rm GS}(r)$  are defined as the electronic densities associated with the excited and ground states, respectively.

Following the procedure proposed by Bahers et al. 78 the excited states of interest were examined using charge-transfer indices (CT) including charge-transfer distance ( $d^{CT}$ ). The product of these quantities  $d^{\text{CT}}$  and  $q^{\text{CT}}$  (transferred charge) gives the variation in the dipole moment between the ground and the excited states  $(\Delta \mu_{0 \to n} = d^{\text{CT}} \times q^{\text{CT}})$ .

On the other hand, the overlap distribution between the holes and electrons can be expressed as

$$S_{\rm r}(r) = \sqrt{
ho^{
m hole}(r)
ho^{
m ele}(r)}$$

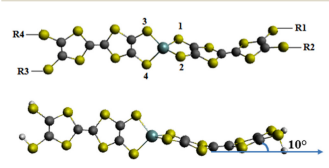
To characterize the overlapping extent of holes and electrons,  $S_r$ index is defined as

$$S_{\rm r}({\rm index}) = \int S_{\rm r}(r) {\rm d}r \equiv \int \sqrt{\rho^{\rm hole}(r)\rho^{\rm ele}(r)} {\rm d}r$$

In our work, the crucial excited state wavefunctions have been obtained by natural transition orbitals (NTOs) in terms of "excited particle" to "empty hole" of the electronic transition density matrix. The CT indices have been determined using the MULTIWFN program. 79

### Results and discussion

The optimized geometries of bis-TTF linked by a germane noted Ti (i = 0-36) compounds obtained from DFT quantum chemical calculations is shown in Fig. 2 along with the atom numbering schemes. As can be seen from the figure, the optimized structure of the bis-TTF-Ge is tetrahedral, in which the Ge-S (S is S1, S2, S3 and S4) average length is 2.259 Å and the  $\angle$  S3-Ge-S4 and  $\angle$  S1-Ge-S2 angles are  $\sim$  97° whereas  $\angle$  S1-Ge-S3 and  $\angle$  S2-Ge-S4 are about 115°. On the other hand,



Optimized molecular geometry of bis-TTF-Ge compounds

the TTF groups deviate slightly from the perfect planar geometry by 10°, showing a slightly distorted geometry. One of the reasons for this distortion is a steric repulsion between the sulphur atoms and the d orbitals from the germanium atom.

On the other hand we can observe that the TTF fragments in the title compounds are distorted into a boat conformation as shown in Fig. 2 in their ground state and this shape agrees well with the available studies. For instance, Demiralp et al. in their study of organic donors (X) containing the TTF group found that X<sup>+</sup> always has a planar TTF region but that the neutral is distorted into a boat conformation.80 They suggested that the off-planar geometry is caused by ring strain, which is developed when the H atoms of TTF are substituted by bulky ligands. 48 Furthermore, Linker et al. in their study of ethylenedioxy-TTF (EDO-TTF) indicated that a neutral derivative has a boat-shaped conformation, whereas the positively charged EDO-TTF is planar.81

#### Reactivity

The values of the HOMO and LUMO frontier orbital energies  $(E_{\text{HOMO}})$  and  $E_{\text{LOMO}}$ , respectively), the electronegativity  $(\gamma)$ , the chemical hardness ( $\eta$ ) and the electrophilicity index ( $\omega$ ), calculated for each bis-TTF-Ge compound, are given in Table 1.

The electronegativities  $(\chi = -\mu)$  of the T1 to T25 bis-TTF-Ge which are substituted with the acceptor group increases with the number of acceptor groups in the title compounds. These results indicate that the trend of the electrons to leave the equilibrium systems increases from the four-acceptor substituted (e) to onesubstituted (a) bis-TTF-Ge. The reverse trend is observed for the bis-TTF-Ge with the electron donor R (NH<sub>2</sub> and NMe<sub>2</sub>), so it can be concluded that the bis-TTF-Ge with four electron donors (e) have the largest tendency to donate electrons than d, b, c and a, respectively.

On the other hand, we observe that the substituent position and the number of the substitution play an important role in the stability of studied bis-TTF-Ge. In each group with the same substituent, the substituent position **b** has a smaller hardness than the d, e, a and c substituent positions. The CHO and CN substituents in bis-TTF-Ge compounds show a decrease according to the following order: a, d, c and e. The ordering of the chemical hardness at the a position is T1 (NO) < T21 (NO<sub>2</sub>) < T11 (COCN) < T31 (NMe2) < T26 (NH2) < T6 (CN) < T16 (COH). The smallest  $\eta$  values are obtained for acceptor substituents and similar trends for the substituent positions b, c, d and e. Therefore, the hardness of bis-TTF-Ge can be effectively reduced when substituent positions are substituted with acceptor groups. According to the theoretical scale of electrophilicity proposed by Domingo et al.<sup>82</sup> of global electrophilicity power ( $\omega$ ) the title compounds can be assigned to a strong electrophile class (2.51 to 6.70 eV).

#### **Electronic transitions**

The TD-DFT calculation results of absorption spectra parameters such as excitation energy ( $\Delta E$ ), absorption wavelength  $(\lambda)$  and corresponding oscillator strength (f), charge transfer parameters  $(q^{\text{CT}}, d^{\text{CT}}, \text{ and } \Delta\mu_{0 \to i})$  and  $S_{\text{r}}$  index are calculated at

Table 1 Energy HOMO and LUMO (eV), electronegativity (χ, eV), chemical hardness  $(\eta, eV)$  and electrophilicity index  $(\omega, eV)$  of the Ti (i = 0 to 36)compounds

	$\mathrm{T}i$	$E_{ m HOMO}$	$E_{ m LUMO}$	η	χ	ω
	T0	-5.311	-2.577	2.733	3.944	2.845
a	T1	-5.329	-3.495	1.834	4.412	5.307
b	T2	-5.341	-3.776	1.564	4.558	6.644
c	T3	-5.361	-3.505	1.855	4.433	5.296
d	T4	-5.374	-3.790	1.583	4.582	6.631
e	T5	-5.430	-3.798	1.631	4.614	6.523
a	T6	-5.388	-2.708	2.680	4.048	3.057
b	T7	-5.429	-2.786	2.642	4.107	3.192
c	T8	-5.670	-2.838	2.832	4.254	3.195
d	T9	-5.710	-2.914	2.795	4.312	3.326
e	T10	-5.860	-2.985	2.874	4.423	3.402
a	T11	-5.377	-2.957	2.419	4.167	3.589
b	T12	-5.429	-3.374	2.054	4.401	4.715
c	T13	-5.600	-3.000	2.600	4.300	3.555
d	T14	-5.701	-3.416	2.284	4.559	4.548
e	T15	-5.855	-3.437	2.418	4.646	4.463
a	T16	-5.361	-2.657	2.703	4.009	2.973
b	T17	-5.400	-2.728	2.672	4.064	3.091
c	T18	-5.527	-2.738	2.788	4.132	3.062
d	T19	-5.528	-2.746	2.782	4.137	3.076
e	T20	-5.710	-2.878	2.831	4.294	3.256
a	T21	-5.382	-3.009	2.373	4.195	3.708
b	T22	-5.412	-3.790	1.622	4.601	6.523
c	T23	-5.624	-3.054	2.570	4.339	3.663
d	T24	-5.653	-3.834	1.819	4.744	6.184
e	T25	-5.838	-3.798	2.039	4.818	5.691
a	T26	-5.247	-2.576	2.671	3.912	2.864
b	T27	-5.019	-2.450	2.569	3.734	2.714
c	T28	-5.199	-2.502	2.697	3.851	2.749
d	T29	-4.996	-2.413	2.583	3.704	2.656
e	T30	-4.971	-2.363	2.607	3.667	2.578
a	T31	-5.203	-2.536	2.667	3.870	2.807
b	T32	-5.031	-2.451	2.579	3.741	2.713
c	T33	-5.179	-2.495	2.684	3.837	2.743
d	T34	-4.989	-2.409	2.580	3.699	2.651
e	T35	-4.964	-2.323	2.641	3.644	2.513
e	T36	-5.291	-2.715	4.003	2.575	3.111

the CAM-B3LYP/6-311g(d,p) level and summarized in Table S1 (ESI†).

The simulated spectra and the electron density difference maps (EDDMs) are provided in Fig. 3 and Fig. S1, S2 (ESI†), respectively. In general, the absorption spectra of the title compounds are characterized by a strong band (B3) at 267 nm, two weak bands (B1 and B2) and two moderate absorption shoulders (S1 and S2).

The bis-TTF-Ge with NO electron acceptor substituents have five absorption bands, and their absorption wavelength range extends to 750 nm. T1, T2 and T3 show electronic transitions at 727, 680-637 and 654 nm, respectively, with small oscillator

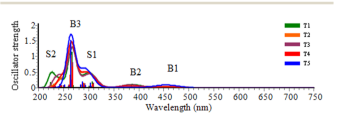


Fig. 3 Calculated UV-vis absorption spectra of bis-TTF-Ge T1-T5 compounds (B: bands; S: shoulders).

strength ( $f \le 0.001$ ), mainly arising from TTF rings 1, 2, and 3 to NO, from TTF rings 1-3 to NO (at positions R1 and R2) and from rings 5 and 6 to NO (at position R3) respectively. The HOMO → LUMO excitations of T1, T2 and T3 are located at 307, 456 and 418 nm, respectively, mainly ascribed to a  $\pi \to \pi^*$ charge transfer (CT) character. The electronic transitions of T5 are assigned as a CT from rings 1, 2, 5, and 6 to four NO units (677, 637 and 455 nm) and from rings 1, 2, 5, and 6 to the 4S-Ge unit (at 400 and 289 nm). For T4, the electronic transition appears in the 676, 455 and 316 nm from rings 1-2 to NO (at positions R1 and R2) and from rings 5-6 to S-NO (positions R1 and R2)  $\pi \rightarrow \pi^*$  CT character, besides the electronic transition at 290 nm (f = 0.132) from rings 5-6 to NO (at position R3) and to the 4S-Ge unit.

For the T6 to T10 compounds, the HOMO → LUMO transition at  $\sim$  399 nm is a charge transfer from rings 6 and 5 to 4S-Ge. The bis-TTF-Ge T6 is also ascribed to the same CT character at 292 nm (f = 0.141). For T8 the absorption bands located at 304 and 243 nm, with large oscillator strength, mainly arise from rings 1 and 2 to 4S-Ge and CN (at position R3). For bis-TTF-Ge T10, it involves three CT characteristic bands at 363, 279 and 241 nm from rings 1-3 to C-S-COCN units attached to ring 1, from TTF rings to 4S-Ge (f = 0.198) and from ring 6-5 to C-S-COCN (at R1 and R2 positions) and C3-C4 atoms attached to ring 6 (f = 0.129), respectively. For T7 and T9, the absorption at 276 nm (f = 0.234and 0.191, respectively) have a mixed character of charge transfer from rings 5 and 6 to S-CN units at the R1 and R2 positions of type  $n \to \pi^*$  and  $\pi \to \pi^*$  electronic transition.

For T14 and T15, the electronic transition at 374 and 351 nm, respectively, is mainly ascribed to the  $n \to \pi^*$  and  $\pi \to \pi^*$  CT mixed character from rings 4, 5 and 6 to S-COCN unit at position R3. The transition at 400 nm of T14 and T15 mainly comes from rings 1-3 to S-COCN (at R1 and R2 positions). The electronic transition of T13 at 398 and 359 nm arises from rings 5 and 6 to 4S-Ge and from rings 1, 2, 5 and 6 to two COCN units CT, respectively.

For T16, T17 and T19 the HOMO → LUMO transition at  $\sim$  401 nm is CT from rings 6 and 5 to the 4S-Ge unit. For T16 and T19 the HOMO-2  $\rightarrow$  LUMO and HOMO-1  $\rightarrow$  LUMO+4 excitations are CT from rings 1 and 2 to 4S-Ge unit and from rings 1-3 to 4-6, respectively (see Table S1 and Fig. S2, ESI†). Bis-TTF-Ge T18 exhibits absorption at 398 nm (HOMO → LUMO) with a charge transfer from rings 1, 2, 5 and 6 to the 4S-Ge unit. The T20 compound has three absorption bands at 397, 304 and 278 nm with a charge transfer character from rings 6 and 5 to 4S-Ge, from rings 6 and 5 to the four units S-CHO and from rings 1, 2, 5 and 6 to the 4S-Ge unit, respectively.

The bis-TTF-Ge with NH<sub>2</sub> and NMe<sub>2</sub> electron donors show three main absorption bands. The most intense absorption band of Ti (i = 26 to 35) is at 269 nm, with the largest oscillator strength, mainly coming from intra-rings charge transfer (ICT). The band at  $\sim$  404 nm is formed by a charge transfer from rings 1 and 2 to 4S-Ge unit from HOMO → LUMO. The absorption band at  $\sim$  310 nm shows that the electronic transitions have a mixed character of charge transfer from rings 1-3 to rings 4-6 of type  $n \to \pi^*$  and  $\pi \to \pi^*$  electronic transition.

For T21, the electronic transition at 434 and 295 nm is assigned as a CT from rings 1 and 2 to S-NO2 and from rings 5 and 6 to S-NO2, respectively. For T23, there is a charge transfer from rings 1, 2, 5 and 6 to S-NO2 (at the R4 position) and to the 4S-Ge unit.

For T25, it is interesting to find that there is a CT from rings 2 and 3 to two units C-S-NO2 (at R1 and R2 positions), from rings 1-2 and 5-6 to the 4S-Ge unit and from rings 4-6 to the C-S-NO<sub>2</sub> unit (at R1 and R2 positions) at 524, 394 and 325 nm, respectively; besides n,  $\pi \to \pi^*$  mixed CT from 1–3 to 4–6 rings and from rings 2-3 and 5-6 to C-S-NO<sub>2</sub> (at ring 6). For T22, the electronic transition arises from rings 1-2 to S-NO2 and from rings 4-6 to S-NO<sub>2</sub> (at R1 and R2 positions), from rings 5-6 to 4S-Ge and from rings 1-2 to 4S-Ge. For bis-TTF-Ge T24, the electronic transition is mainly ascribed to a n,  $\pi \to \pi^*$  mixed character from rings 2-3 to S-NO<sub>2</sub> (at R1 and R2 position), from rings 4-6 to S-NO<sub>2</sub> (at R1 and R2 positions) and to S-NO<sub>2</sub> at the R3 position.

TD-DFT calculations on bis-TTF-Ge T0-T36 compounds show that the wavelength of the largest oscillator strength located at  $\sim 266$  nm, are due to n  $\rightarrow \pi^*$  and  $\pi \rightarrow \pi^*$  mixed character type intra-TTF electron transition (see Table S1 and Fig. S1, ESI†). Furthermore, these excitations show a relatively small  $d^{CT}$  and a large  $S_r$  index, indicating that the hole and electron are very close. For example, for T0, T36, T13, T18, T23 and T16 the electron density difference maps (EDDM) of this transition, shown in Fig. 4, reveals a significant intramolecular CT (ICT) and, for all compounds, is reported in the ESI† in Fig. S2.

On the other hand, the TD-DFT results show that the electronic transition HOMO → LUMO for T2, T4, T11, T12, T21, T22 and T24 (R = NO, COCN and NO<sub>2</sub>; position is  $\mathbf{a}$ ,  $\mathbf{b}$  and  $\mathbf{d}$ ) is a strong non-local excitation as indicated by the overlapping index between the hole and the electron which is zero  $(S_r = 0,$ Table S1, ESI†).

Overall, T22, T24 and T25 show red-shift wavelength absorption compared to T23 and T12. On the other hand, the absorption

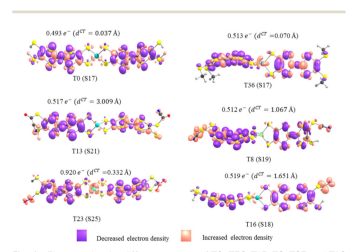


Fig. 4 Electron density difference maps of T0, T36, T13, T8, T23 and T16 compounds from the ground state to the crucial excited state S0  $\rightarrow$  Sn (S17, S17, S2, S19, S25 and S18 respectively), plotted using 0.008 a.u. isovalues

wavelengths of T6 to T10 are close to T16 to T20. Furthermore, this study displays that the bis-TTF-Ge with electron-acceptor substituents (NO2 and NO) shows a red-shift wavelength absorption compared to the Ti with donor electron substituents. In general, from this calculation, we can conclude that the electronic transition characters and the absorption wavelength of Ti compounds can be tuned through the introduction of the electron donors or electron acceptor substituents at different positions.

#### Nonlinear optical properties

In current years, quantum chemical calculations have become a widely accepted method for predicting new NLO materials. 20,21 For the NLO calculation we carried out a test on T36 with the 6-311++G and 6-311G(d,p) basis set, but did not find significant differences (see Fig. S3, ESI†). This is no surprise because of the quasi linear shape of the molecules and the 6-311G(d,p) basis is a good choice for the title compounds. On the other hand, we calculated the dynamic hyperpolarizabilities at the  $\lambda$  value of 1064 nm ( $\omega$  = 0.04282 a.u.), which is sufficiently apart from the observed lowest-energy absorption of the title compounds (the  $\lambda_{\text{Max}}$  values are in the range 250–270 nm). It is known that it is difficult to compare the calculated dynamic first hyperpolarizabilities to the experimental data measured under resonance conditions.83 However the tendency is very similar between experimental and theoretical values,77 and one has hyperpolarizabilities (experimental) > hyperpolarizabilities (dynamic) > hyperpolarizabilities (static) and our calculation shows the same tendency: hyperpolarizabilities (dynamic) > hyperpolarizabilities (static).

The second-order nonlinear responses calculated in the static ( $\lambda = \infty$ ) and dynamic ( $\lambda = 1064$  nm) regimes for the 37 derivatives are summarized in Table 2. The reported quantities include the hyper-Rayleigh scattering ( $\beta_{HRS}$ ) hyperpolarizabilities and the associated depolarization ratio (DR).

For bis-TTF-Ge T26, T28, and T30 their  $\beta_{HRS}^{\lambda=\infty}$  values are close to the value of T31, T33 and T35 (~111 a.u.) which indicates that when the NH<sub>2</sub> (at a, c and e positions) is replaced by the  $\mathrm{NMe_2}$  group the effect on the  $\beta_{\mathrm{HRS}}$  value is not significant. On the other hand, the  $\beta_{\rm HRS}^{\lambda=\infty}$  value calculated for T26, T28, T30-T31, T33 and T35 are relatively close to the value of T0 (75 a.u.). We can observe that the dynamic regime has the same behavior as the static regime for NH2 and NMe2 donor groups. Furthermore, the hyperpolarizability  $\beta_{\rm HRS}^{\lambda}$  ( $\lambda = \infty$ , 1064 nm) of T27 and T29 are three times larger than that of the T26, T28 and T30. For T32 and T34 the  $\beta_{HRS}^{\lambda}$  values are about twice that of T31, T33 and T35. We can observe that the introduction of donor substituents (NH2 and NMe2) at b and d positions can increase the hyperpolarizability of bis-TTF-Ge. For T1-T5 (acceptor NO unit) the  $\beta_{HRS}^{\lambda}$  (static and dynamic) of T2 is found to be about twice larger than that of T1, T3, T4, and T5.

For bis-TTF-Ge T11–T12, the  $\beta_{HRS}^{\lambda}$  values are larger than those of T13, T14 and T15, respectively. Comparing  $\beta_{HRS}^{\lambda=1064}$  values of T11-T12 and T0, it can be seen that the T11-T12 values are about 11 times larger than that of T0. For bis-TTF-Ge substituted by CN and CHO units, the  $\beta_{HRS}$  (static and dynamic) values are also larger in the substitution at **b** and **a** positions than for **d**, **c** and **e**.

**Table 2** Static ( $\lambda = \infty$ ) and dynamic ( $\lambda = 1064$  nm) HRS first hyperpolarizability ( $\beta_{HRS}$ , a.u.) and depolarization ratio (DR), as well as calculated dipole moment (D, Debye), calculated at the CAM-B3LYP/6-311g(d,p) level in Ti (i = 0-36) compounds

	Τi	μ	$DR^{\lambda=\infty}$	$eta_{ m HRS}^{\lambda=\infty}$	$\mathrm{DR}^{\lambda=1064}$	$eta_{ m HRS}^{\lambda=1064}$
	ТО	2.048	1.503	75	1.407	137
a	T1	0.797	3.817	440	4.629	1266
b	T2	0.955	5.247	782	7.147	3397
c	T3	0.014	1.862	314	1.720	1015
d	T4	0.702	3.505	464	5.108	2469
e	T5	1.005	2.34	317	1.541	1140
a	T6	4.492	3.679	263	4.278	501
b	T7	7.508	5.729	416	6.576	742
c	T8	1.436	2.053	152	1.812	246
d	T9	4.290	3.907	231	3.067	383
e	T10	5.674	1.781	173	1.407	390
a	T11	4.271	4.731	644	5.215	1566
b	T12	6.626	5.387	627	7.022	1374
c	T13	1.070	2.316	371	2.322	858
d	T14	3.643	1.915	324	1.859	807
e	T15	3.440	2.739	234	1.845	629
a	T16	3.294	2.693	180	3.256	360
b	T17	5.625	3.913	265	5.131	576
c	T18	1.829	1.688	116	1.676	206
d	T19	2.623	2.633	150	3.137	331
e	T20	1.501	1.633	104	1.326	271
a	T21	4.106	3.92	713	4.27	2859
b	T22	6.091	4.499	800	1.145	9638
c	T23	2.939	2.378	709	1.902	2726
d	T24	2.583	2.527	829	1.754	12 600
e	T25	2.836	1.952	761	0.749	19810
a	T26	4.838	2.447	103	1.942	185
b	T27	3.384	4.077	327	3.741	602
c	T28	1.694	2.793	118	1.914	194
d	T29	2.717	4.596	295	4.054	581
e	T30	4.024	1.538	115	1.422	183
a	T31	4.919	2.813	110	2.158	189
b	T32	2.567	4.724	192	4.061	369
c	T33	0.912	4.829	110	2.265	173
d	T34	2.014	6.843	185	5.122	364
e	T35	1.388	3.059	111	2.779	158
e	T36	4.024	1.538	116	1.422	183

For T6-T10, T11-T15 and T16-T20 compounds, we observe that the  $\beta_{\rm HRS}^{\lambda=\infty}$  and  $\beta_{\rm HRS}^{\lambda=1064}$  values of bis-TTF-Ge substituted by the acceptor COCN group are larger than that of bis-TTF-Ge substituted by CN and COH group at the same substituent position, respectively.

In static and dynamic regimes, the compounds T21-T25 exhibit larger hyperpolarizability values than the other bis-TTF-Ge compounds, which show that the introduction of the acceptor  $NO_2$  group can effectively enhance  $\beta_{HRS}^{\lambda}$  values. This observation is in line with Gong et al.'s study of helicenes84 and linear [3]spirobifluorenylene. 85 On the other hand, the  $\beta_{HRS}^{\lambda=\infty}$  decreases slightly from T29  $\approx$  T27 to T25, T21 and T23, respectively, indicating that the substituent position (number) of NO2 substitutions in bis-TTF-Ge have little effect on the  $\beta_{HRS}^{\lambda=\infty}$  values. On the other hand, the  $\beta_{\rm HRS}^{\lambda=1064}$  values decrease in the order T25 > T24 > T22 > T21  $\approx$  T23, indicating that the  $\beta_{\rm HRS}^{\lambda=1064}$  increases with the number of NO<sub>2</sub> groups in the bis-TTF-Ge compound (see Fig. 5). The  $\beta_{HRS}^{\lambda=1064}$  value of T25 is about 7 times larger than that of T23. Furthermore, the  $\beta_{HRS}^{\lambda=\infty}$  value calculated for T0 (and T36) is about 11 times smaller than that of T21–T25 and  $\beta_{\rm HRS}^{\lambda=1064}$  of T25, T24, T22 and T21 is 145, 92, 70 and 20 times larger than that of T0, respectively.

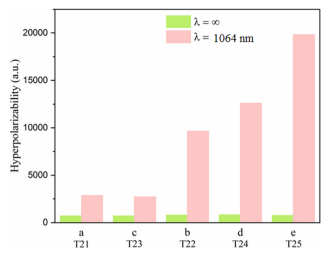


Fig. 5 Static and dynamic hyperpolarizability values for T21-T25 calculated at the CAM-B3LYP/6-311q(d,p) level.

From this study, we observed that the substituent position **b** has a stronger effect on the enhancement the  $\beta_{HRS}^{\lambda}$  value than the a, c, d and e substituent positions. Comparing T0 and the bis-TTF-Ge substituent position **b**, it is found that the  $\beta_{HRS}^{\lambda=\infty}$  values of T2, T12, T7, T17, T27 and T32 is about 10, 8, 6, 4, 4 and 3 times larger than that of T0, respectively, and the  $\beta_{HRS}^{\lambda=1064}$  value of these compounds is about 25, 10, 5, 4, 4 and 3 times larger than that of T0, respectively. We conclude that the highest hyperpolarizability value can be obtained by the introduction NO2, NO or COCN acceptor groups at the **b** position for bis-TTF-Ge.

The experimental study of Bao et al.,86 shows that the different numbers (mono-, di-, tri-, and tetra-) of ester substituents as well as their position (trans and cis) have a significant influence on the NLO properties of tetraphenylporphyrins. They indicate that the compounds with four ester groups present a small value of hyperpolarizability. In the static regime, the same results are observed for title compounds, excepting the bis-TTF-Ge substituted with NO<sub>2</sub> group (see Fig. 5 and Table 2).

Generally, in the dynamic regime, the hyperpolarizability is 2-5 times larger than that in the static regime, excepting T22, T24 and T25, where  $\beta_{HRS}^{\lambda=1064}$  is about 26, 15 and 12 times larger than the  $\beta_{HRS}^{\lambda=\infty}$ , respectively. The static hyperpolarizability of T1 to T35 is about 2-11 times larger than that of T0.

On the other hand, the TD-DFT results of T22, T24 and T25 show that the  $\lambda_{0\rightarrow 1}$  (between 524 and 554 nm) corresponds to the two-photon resonance value (1048-1108 nm). This value is very close to near-resonant wavelength (1064 nm) causing resonance. The two-photon resonance value of T25 is 1048 nm and that of T22 and T24 is 1108 and 1096 nm, respectively. The resonance value of T25 is the closest to the near-resonant wavelength of 1064 nm and that of T22 and T24 is the most shifted from the near-resonant wavelength. This is the reason why the value of  $\beta_{\rm HRS}^{\lambda=1064}$  of T25 is larger than the value of  $\beta_{HRS}^{\lambda=1064}$  of T24 and T22, respectively. Accordingly, the wavelength resonance is one of the reasons why the value of  $\beta_{\rm HRS}^{\lambda=1064}$  is much larger than that of  $\beta_{\rm HRS}^{\lambda=\infty}$ .

On the other hand, the urea molecule is one of the prototypical compounds used in the study of the second-order NLO response of molecules. Therefore it is used frequently as a threshold value for comparative purposes. 87,88 The HRS hyperpolarizability of urea was calculated at the same level of theory. From our results we observe that the all bis-TTF-Ge derivative showed values for the static and dynamic HRS hyperpolarizability higher than the T0 ( $\beta_{HRS}^{\lambda=\infty}$  = 75 a.u. and  $\beta_{HRS}^{\lambda=1064}$  = 137 a.u.) and urea  $(\beta_{HRS}^{\lambda=\infty} = 38 \text{ a.u. and } \beta_{HRS}^{\lambda=1064} = 40 \text{ a.u.}).$ 

The depolarization ratio (DR) is an essential parameter for displaying the contribution of the hyper-Rayleigh scattering response. As revealed by the static and dynamic DR values collected in Table 2, the NLO responses of T1, T2, T4, T6, T7, T9, T11, T12, T17, T21, T22, T27, T29, T32, T33 and T34 compounds display a dominant dipolar character, with the exception of T9, T33 compounds ( $\lambda = 1064$  nm) dominated by their octupolar component. On the other hand, in our studied bis-TTF-Ge T3, T5, T8, T10, T13, T15, T18, T20, T23, T24, T25, T26, T28, T30, T31, T35 and T36 the octupolar component is dominant. From this study we observed that the (static and dynamic) DR values are more strongly sensitive to the substitution positions than to the nature of the substituent (donor and acceptor). This reveals that the a, b and d substitution positions lead to a dominant dipolar nature of the NLO response; the **b** substitution positions show a larger DR value in the static and dynamic regimes. On the other hand, the title compounds with c and e substitution positions are octupolar molecules. For the e substitution positions, the DR value ( $\lambda = \infty$ , 1064 nm), close to 1.5 (T10, T18, T20, T30, T36 and T0) characterizes typical octupolar systems. Furthermore, we observed that the DR value at  $\lambda = 1064$  nm of T0, T10, T20, T22, T25, T30 and T36 is less than 1.5, indicating that those compounds are close to resonance.

On the other hand, we studied the correlation between the energy gaps (hardness) of the title compounds and their hyperpolarizability in the static regime. The results show that  $\beta_{\rm HRS}^{\infty}$  values are increased with the decrease of the bis-TTF-Ge energy gaps (the correlation factor is 0.832, see Fig. S4, ESI†). Taking T24 as an example, the energy gap of T24 is the smallest (1.819 eV) but its hyperpolarizability value is the largest ( $\beta_{HRS}^{\infty}$  = 829 a.u.), which is in good agreement with the literature. 89,90

In order to understand the nonlinear optical origins of studied compounds, the excitation energy  $(\Delta E_{0\rightarrow n})$  and wavelength  $(\Delta \lambda_{0 \to n})$ , oscillator strengths  $(f_{0 \to n})$ , transfer distance charges ( $d^{CT}$ ), charge transfer ( $q^{CT}$ ) and transition dipole moment  $(\Delta \mu_{0 \to n})$  calculated for the crucial singlet excited states are summarized in Table 3, with more details in Table S1 (ESI†), and the corresponding electron density difference maps (EDDMs) are shown in Fig. S2 (ESI†).

The charge transfer involves both  $n \to \pi^*$  and  $\pi \to \pi^*$ transitions, the main factor being the conjugation of the p electrons through the delocalization domain.

From the EDDM of bis-TTF-Ge substituted with acceptor groups (T1-T5 (R: NO), T11-T15 (R: COCN) and T21-T25 (R:NO<sub>2</sub>)), it can be observed that there is obvious CT from the TTF fragment to the substituted groups (R) at the other TTF

T36

3.083

401

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**Table 3** Vertical transition energy ( $\Delta E_{0 \to n}$ , eV) and wavelength ( $\Delta \lambda_{0 \to n}$ , nm), oscillator strengths ( $f_{0 \to n}$ , dimensionless), charge transfer ( $q^{\text{CT}}$ , |e|), charge transfer distance ( $d^{CT}$ , Å), and dipole moment variation ( $\Delta\mu_{0\rightarrow n}$ , D) associated with the S0  $\rightarrow$  Sn (S<sub>0 $\rightarrow$ n</sub>) transition, as calculated at the CAM-B3LYP/6-311g(d,p) in Ti compounds

BSLTP/0-SIIg(a,p) iii 17 compounds								
Τi	$S_{0\rightarrow n}$	$\Delta E_{0 \to n}$	$\Delta \lambda_{0 \to n}$	$f_{0 \to n}$	$q^{\mathrm{CT}}$	$d^{\mathrm{CT}}$	$\Delta\mu_{0\to n}$	
T0	$S_{0 \rightarrow 1}$	3.096	400	0.003	0.821	0.035	0.054	
T1	$S_{0\rightarrow 1}$	1.706	727	0.001	0.891	8.636	36.946	
	$S_{0\rightarrow7}$	3.218	385	0.079	0.707	9.586	32.559	
T2	$S_{0\rightarrow 13}$	3.892	318	< 0.001	0.999	16.977	81.427	
	$S_{0\rightarrow 16}$	4.303	307	0.126	0.618	6.649	19.741	
Т3	$S_{0\rightarrow 1}$	1.896	654	0.001	0.888	11.586	49.439	
	$S_{0\rightarrow 13}$	3.999	310	0.162	0.508	2.906	7.095	
T4	$S_{0\rightarrow 17}$	3.921	316	< 0.001	0.997	17.038	81.612	
m-	$S_{0\rightarrow 20}$	4.010	309	0.125	0.576	6.199	17.151	
T5	$S_{0\rightarrow 1}$	1.831	677	< 0.001	0.805	6.489	25.050	
TC	$S_{0\rightarrow 5}$	2.725	455	0.046	0.794	1.452	5.526	
T6	$S_{0\rightarrow 1}$	3.070	404	0.003 0.167	0.872	6.729	28.067	
T7	$S_{0\rightarrow7}$	4.003 3.051	307 406	0.107	$0.631 \\ 0.871$	7.18 6.634	20.478 27.747	
1 /	$S_{0\rightarrow 1}$ $S_{0\rightarrow 7}$	4.028	308	0.002	0.623	5.961	17.829	
Т8	$S_{0\rightarrow 1}$	3.132	396	0.004	0.792	1.963	7.467	
10	$S_{0\rightarrow7}$	4.074	304	0.429	0.513	0.320	0.784	
Т9	$S_{0\rightarrow 5}$	3.421	362	0.042	0.781	4.721	17.702	
	$S_{0\rightarrow 12}$	4.304	288	0.178	0.593	3.380	9.545	
T10	$S_{0\rightarrow 1}$	3.086	402	0.005	0.747	6.418	23.035	
	$S_{0\rightarrow 13}$	4.305	288	0.204	0.594	4.507	12.859	
T11	$S_{0 \rightarrow 13}$	4.238	293	0.141	0.601	4.082	11.783	
T12	$S_{0 \rightarrow 20}$	4.200	295	0.205	0.708	7.483	25.392	
T13	$S_{0\rightarrow 17}$	4.309	288	0.764	0.575	1.699	4.684	
T14	$S_{0\rightarrow 9}$	3.790	327	0.100	0.752	4.598	16.609	
T15	$S_{0\rightarrow 8}$	3.528	351	0.041	0.912	6.937	30.399	
T16	$S_{0\rightarrow 1}$	3.082	402	0.003	0.838	6.540	26.862	
TT4 7	$S_{0\rightarrow7}$	4.020	308	0.221	0.610	3.628	10.635	
T17	$S_{0\rightarrow 1}$	3.065	404	0.003	0.869	6.778	28.212	
T18	$S_{0\rightarrow7}$	4.032	307	0.173	0.650	8.709	27.195 4.946	
110	$S_{0\rightarrow 1}$ $S_{0\rightarrow 7}$	3.119 $4.018$	398 309	0.003 $0.295$	$0.800 \\ 0.558$	1.281 $0.723$	1.945	
T19	$S_{0\rightarrow 7}$ $S_{0\rightarrow 1}$	3.117	398	0.004	0.831	5.726	22.867	
115	$S_{0\rightarrow 8}$	4.091	303	0.172	0.669	5.704	18.314	
T20	$S_{0\rightarrow 1}$	3.125	397	0.004	0.786	3.243	4.816	
	$S_{0\rightarrow7}$	4.074	304	0.386	0.581	0.315	0.879	
T21	$S_{0\rightarrow 1}$	2.854	434	0.083	0.831	5.372	21.438	
	$S_{0\rightarrow 14}$	4.207	295	< 0.001	0.999	15.939	76.256	
T22	$S_{0\rightarrow 6}$	3.421	362	< 0.001	0.969	15.746	73.084	
	$S_{0\rightarrow 11}$	3.971	312	0.113	0.883	6.229	26.408	
T23	$S_{0\rightarrow 1}$	2.867	432	0.123	0.829	3.650	14.512	
T24	$S_{0\rightarrow 2}$	2.869	432	0.034	0.828	5.318	22.476	
	$S_{0\rightarrow 8}$	3.603	344	< 0.001	0.999	15.685	75.300	
T25	$S_{0\rightarrow 2}$	2.366	524	0.036	0.383	7.626	30.538	
TOC	$S_{0\rightarrow 9}$	3.811	325	< 0.001 0.003	0.916 $0.875$	13.204	58.126 26.428	
T26	$S_{0\rightarrow 1}$	3.083 4.030	402 308	0.003	0.875	6.448 $4.420$	11.286	
T27	$S_{0\to7} \atop S_{0\to1}$	3.020	411	0.002	0.330	6.741	28.401	
127	$S_{0\rightarrow 7}$	3.993	311	0.135	0.744	8.137	29.066	
T28	$S_{0\rightarrow 1}$	3.050	406	0.002	0.852	6.211	25.277	
T29	$S_{0\rightarrow 1}$	3.026	410	0.002	0.874	6.746	28.277	
	$S_{0\rightarrow7}$	4.002	309	0.151	0.738	8.838	31.299	
T30	$S_{0\rightarrow 1}$	3.039	408	0.002	0.873	6.537	27.177	
	$S_{0\rightarrow7}$	4.016	309	0.146	0.734	6.854	24.012	
T31	$S_{0\rightarrow 1}$	3.081	402	0.003	0.870	6.972	29.120	
	$S_{0\rightarrow7}$	4.023	308	0.319	0.593	7.924	22.565	
T32	$S_{0 \rightarrow 1}$	3.027	410	0.002	0.893	7.276	31.218	
	$S_{0\rightarrow7}$	3.998	310	0.201	0.749	9.054	32.562	
T33	$S_{0\rightarrow 1}$	3.089	401	0.003	0.843	4.628	18.658	
mo 1	$S_{0\rightarrow7}$	4.021	308	0.420	0.522	3.447	8.593	
T34	$S_{0\rightarrow 1}$	3.026	410	0.003	0.896	7.178	30.898	
Т2 =	$S_{0\rightarrow7}$	3.981	311	0.219	0.689	8.247	27.285	
T35	$S_{0\rightarrow 1}$	3.055	406	0.003	0.869	5.083 $2.059$	21.208	
ma c	$S_{0\rightarrow7}$	3.989	311	0.315	0.568	2.059	5.619	

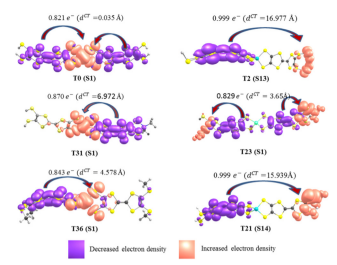


Fig. 6 Electron density difference maps of T0, T2, T21, T22, T31 and T36 compounds from the ground state to the crucial excited state S0  $\rightarrow$  Sn (S1, S13, S1, S1, S1 and S14, respectively), plotted using 0.008 a.u. isovalues.

fragment. These transitions are dominated by an electronic excitation from the HOMO to the LUMO and have high charge transfer parameters  $(q^{\text{CT}}, d^{\text{CT}})$  and  $\Delta \mu_{0 \to n}$  character which is the necessary condition to obtain a high NLO response. On the other hand, the CT characters of T6-T10, T16-T20 and T26-T36 are from the TTF fragment (rings) to the 4S-Ge unit and these transitions are also associated with a HOMO → LUMO excitation with fewer charge transfer parameters than the other compounds (see Table 3, Fig. S2 and Table S1, ESI†). For instance, the dipole moment variation of T21 (76.26 D) is due to a transferred excitation charge  $q^{\text{CT}} = 0.999 |e|$  and a larger associated CT distance ( $d^{\text{CT}} = 15.94 \text{ Å}$ ,  $S_{\text{r}} = 0$ ) than that of T31  $(q^{\text{CT}} = 0.87 | e|, d^{\text{CT}} = 6.97 \text{ Å}; S_r = 0.367 \text{ and } \Delta\mu_{0 \to 1} = 29.12 \text{ D}).$  The relatively small  $d^{\rm CT}$  and  $\Delta\mu_{0\to n}$  values are obtained for bis-TTF-Ge with the substitution donor or acceptor at position c. This can be attributed to the CT from the TTF to the substituted group on the same fragment; see for example T23 in Fig. 6. In contrast, T0 undergoes a moderate CT excitation ( $q^{\text{CT}}$  = 0.821 |e|) with a weak CT distance and small dipole moment variation ( $d^{CT} = 0.035 \text{ Å}$  and  $\Delta \mu_{0 \to 1} = 0.054 \text{ D}$ ) (see Fig. 6). Furthermore, the title compounds show a nice correlation between HRS hyperpolarizability and the largest dipole moment variation (Fig. 7). Also, there is a good correlation between  $\beta_{HRS}^{\lambda=\infty}$  and  $\beta_{HRS}^{\lambda=1064}$ , except for T2, T4 and T5 (Fig. 8).

#### Two-level model analyses for hyperpolarizability

The sum-over-states (SOS) method is one of the most commonly used methods for theoretical estimation of hyperpolarizability. 91-96 The full form of the SOS formula of component ABC of hyperpolarizability is defined as

$$\begin{split} \beta_{xyz}(-\omega_{\sigma};\omega_{1},\omega_{2}) = & \hat{P}[x(-\omega_{\sigma}),y(\omega_{1}),z(\omega_{2})] \\ \times \sum_{i\neq 0} \sum_{j\neq 0} \frac{\mu_{0i}^{x}\overline{\mu_{ij}^{y}}\mu_{j0}^{z}}{(\Delta_{i}-\omega_{\sigma})(\Delta_{j}-\omega_{2})} \end{split}$$

0.843

4.578

18.492

0.003

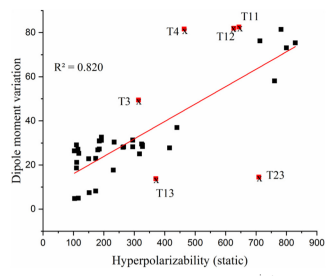


Fig. 7 Correlation between HRS hyperpolarizability  $(\beta_{\text{HRS}}^{\lambda=\infty})$  and dipole moment variation ( $\Delta\mu_{0\rightarrow i}$ ), (T3, T4, T11, T13 and T23 are exceptions)

where 
$$\mu^x_{ij} = \langle i | \hat{\mu}^x | j \rangle$$
,  $\overline{\mu^x_{ij}} = \mu^x_{ij} - \mu^x_{00} \delta_{ij}$ ,  $\omega_{\sigma} = \sum_i \omega_i$ ,  $\omega$  is the

energy of external fields, and  $\omega = 0$  corresponds to the static electric field;  $\Delta_i$  stands for excitation energy of state i with respect to the ground state (0).  $\hat{P}$  is the permutation operator that acts on the *xyz* indices of the  $\beta$ -components.  $\mu_{ij}^x$  is the *x* component of the transition dipole moment between state i and state i.

Molecular NLO responses of material is intimately related to the electronic absorption properties and through the two-level model established by Oudar and Chemla the static hyperpolarizabilities  $\beta$  can be expressed as  $^{97,98}$ 

$$\beta_0 \propto \frac{f \times \Delta \mu}{\Delta E^3}$$

where  $\Delta E$ , f and  $\Delta \mu$  are the excitation energy, oscillator strength and difference of dipole moment between the ground state and

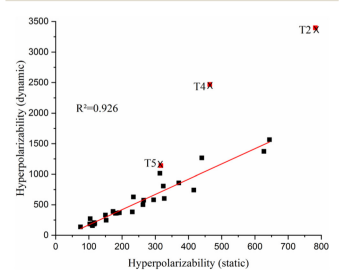
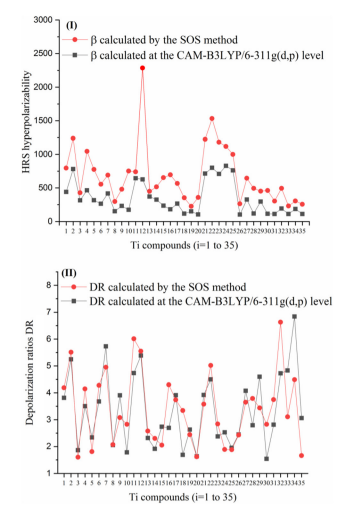


Fig. 8 Correlation between static and dynamic hyperpolarizability, T2, T4 and T5 are exceptions



(I) Calculated static first-hyperpolarizabilities of Ti compounds by using the SOS method and at the CAM-B3LYP/6-311g(d,p) level; (II) depolarization ratios DR of Ti compounds calculated via the SOS method and at the CAM-B3LYP/6-311q(d,p) level

the excited state, respectively. According to this relation, it is observed that the static hyperpolarizability  $(\beta_0)$  is inversely proportional to  $\Delta E^3$ , demonstrating that lower excited energy  $(\Delta E)$ , greater dipole moment  $(\Delta \mu)$  and oscillator strength (f) will lead to the largest  $\beta_0$  of compounds.

Furthermore, it is well known that the molecular NLO properties calculated using the SOS method are closely related to the number of excited states;<sup>99</sup> therefore in our study we plotted the relationship between the first hyperpolarizability  $(\beta^{SOS})$  and 100 excited states (Fig. S5, ESI†). Generally, these graphs demonstrate that 100 excited states are sufficient for the convergence of the  $\beta^{SOS}$  value. As shown in Table S2 and Fig. S5 (ESI†) the introduction of the substituent group at the b position can increase the hyperpolarizability values more than at the other positions, and the NO2, NO and COCN acceptor groups, lead to the largest values. This prediction agrees well with hyperpolarizability calculated by CAM-B3LYP/6-311g(d,p).

In addition, the two-level analysis discloses that the excited state S7 makes a dominant contribution to the hyperpolarizability value of compounds T6, T7, T8, T16-T18, T20 and T26-

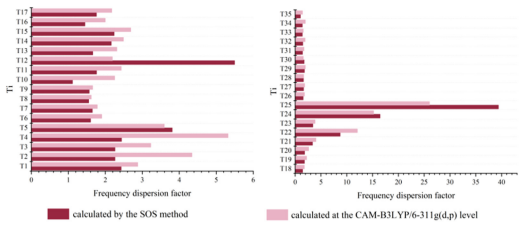


Fig. 10 Variation of the frequency dispersion factor of bis-TTF-Ge derivatives calculated by SOS method and at the CAM-B3LYP/6-311q(d,p) level

T35; the crucial excited states S12, S13, S12, S13, S20 and S17 with largest hyperpolarizability in compounds T9, T10, T19, T11, T12, and T13 respectively; these transitions can be assigned to the CT from TTF fragment (rings) to the 4S-Ge unit (see Tables S1 and S2, ESI†). For systems T1-T5 the dominant contribution to hyperpolarizability values can be assigned to S7, S18, S23, S22 and S5, respectively, and are mainly made up of CT from TTF fragment to the NO groups at the other TTF fragment. For compounds T21 to T25 the excited states S1, S11, S1, S2 and S2, respectively, with CT from the TTF fragment to NO<sub>2</sub> groups make large contributions to the hyperpolarizability value. For compounds, T14 to T15 the crucial excited state is S9 and S8 respectively, with CT from the TTF fragment to COCN groups have (see Tables S1, S2 and Fig. S2, ESI†). On the other hand, SOS calculations show that the local excitation transition at  $\sim 266$  nm of title compounds (noted in italic in Table S2, ESI†) exhibit a small hyperpolarizability value and a large dipole moment variation. These results agree well with the character of the electronic transition that we have noted before (in the section of nonlinear optical properties and Table 3).

As can be seen from Table S2 and Fig. 9, as a whole, the hyperpolarizability  $\beta_{HRS}^{SOS}$  and ratio DR<sup>SOS</sup> values (calculated DR<sup>SOS</sup> ratio and  $\beta_{HRS}^{SOS}$  based on the hyperpolarizability tensor derived by the SOS formula) qualitatively reproduce the general trend of the HRS hyperpolarizabilities and DR character (Table 2) with a few exceptions resulting from the limitation of the SOS method. On the other hand, it is worth stressing that the SOS method obviously overestimates the hyperpolarizability and DR value of Ti compounds (Fig. 9(I) and (II)).

Furthermore, the frequency dispersion factor (FDF) between static and dynamic at a definite wavelength  $\lambda = 1064$  nm depicted by the ratio  $\beta^{1064}/\beta^{\infty}$ , namely FDF<sub>HRS</sub> and FDF<sub>HRS,SOS</sub>, are calculated from the results in the Table 2 and Table S2 (ESI†), respectively and are presented in Fig. 10. From this Fig. 10, as expected, both FDF<sub>HRS</sub> and FDF<sub>HRS,SOS</sub> follow the same trend and in addition, the FDF<sub>HRS</sub> values are quite close to the FDF<sub>HRS,SOS</sub> values, with the exception of T2, T4, T12, T22 and T25. On the other hand, we can observe that the dispersion of optical nonlinearity of T24, T25 and T22 have maximum

frequency dispersion factor at  $\lambda = 1064.8$  nm, confirming their great nonlinear optical properties, in contrast to the other compounds (dispersion factor from  $\sim 1$  to 5.5). It is important to highlight that the SOS method consistently produces large hyperpolarizability and small FDF values of compound bis-TTF-Ge. Indeed, all the compounds containing NO<sub>2</sub> substituent(s) (and, to a smaller extent, containing NO) are much more hyperpolarizable at 1064 nm frequency than in static regime, whereas with other substituents, the difference is not significant. This may be related to the NO<sub>2</sub> lone pairs, which may be more mobile through the molecular systems.

To summarize, the large HRS hyperpolarizabilities origin of bis-TTF-Ge substituted with acceptor groups come from the CT from the TTF fragment towards the acceptor group located on the second TTF unit. In the meantime, the values of the CT from the TTF to the acceptor group located on the same TTF fragment are weak, whereas the NLO origin of bis-TTF-Ge substituted with donor originates a CT from TTF fragment to the 4S-Ge unit.

#### Conclusions

The structural properties, reactivity parameters, and linear and nonlinear optical properties of 37 bis-TTF-Ge derivatives differing by a donor and acceptor substitution on the TTF fragment have been investigated by using DFT and TD-DFT calculations. These calculations were performed at the CAM-B3LYP/ 6-311g(d,p) level and using the sum-over-states (SOS) approach in both static and dynamic regimes.

Our results indicate that the bis-TTF-Ge hardness can effectively be reduced when substituent positions are substituted with an acceptor group (NO, NO2, and COCN). Furthermore, the **b** substituent position leads to a smaller hardness than the **d**, **e**, a and c substituent positions. The electronic excitation properties of the title compounds can be described as a mixed transition of intra-TTF charge transfer (ICT) and charge transfer (CT) from TTF to the 4S-Ge unit, TTF to the acceptor group (NO2, NO and COCN) and TTF to TTF. For example, the electronic transition at ~266 nm is a local excitation with a

relatively small  $d^{ct}$  and large  $S_r$  indexes. The transition HOMO → LUMO of bis-TTF-Ge with the acceptor substituent (R = NO, COCN and NO<sub>2</sub>) at positions a, b and d is a non-local excitation with large dipole moment variation and a small  $S_r$  index  $(S_r \sim 0)$ . In the case of bis-TTF-Ge with a donor substituent (R: NH2 and NMe<sub>2</sub>) at positions **a**, **b** and **d**, the transition HOMO  $\rightarrow$  LUMO from the TTF fragment to 4S-Ge shows moderate values of the transition dipole moment and  $S_r$  indexes ( $S_r = \sim 0.3$ ) when we compare it with the results of an acceptor substituent.

All bis-TTF-Ge derivatives showed values for the static and dynamic HRS hyperpolarizability higher than T0 and urea.

The quantum chemical calculations indicate that the position and the nature (donor/acceptor) of the substituent have strong effects on the NLO responses. The introduction of a substituent group at the b position induces a greater increase of the first hyperpolarizability and the acceptor NO2, NO and COCN groups show a large value of HRS hyperpolarizability which indicate that the acceptor group has a stronger effect on the second order NLO than the donor group. On the other hand, the compounds with c and e substitution positions are octupolar molecules and a, b and d substitution positions lead to a dominant dipolar nature of the NLO response.

The large first hyperpolarizability origin comes from the CT from the TTF fragment towards the acceptor group located on the second TTF unit. On the other hand, the  $\beta_{HRS}^{\lambda=\infty}$  values of compounds T21-T25 are high and relatively close, indicating that the introduction of the acceptor NO2 group at any substituent position can effectively enhance the first hyperpolarizability. Interestingly, the  $\beta_{HRS}^{\lambda=1064}$  of bis-TTF-Ge substituted with the NO<sub>2</sub> group increases with the number of NO2 in the TTF fragment.

Finally, it is important to emphasize that the intrinsic asymmetric geometry of the bis-TTF-Ge can tune the second order NLO response through the nature, position and number of the substituent group at the TTF fragment, as well as by the  $S_r$  index.

On the other hand, our study suggests that these compounds can be used as new materials for developing nonlinear second order photonic devices. Furthermore, the bis-TTF-Ge compounds substituted with the an acceptor group may become excellent candidates for second order NLO materials, which will also promote the evolution of high technology applications.

#### Author contributions

All the authors discussed the results. Dalila Kamli: calculations achievements, data interpretation, draft writing. Douniazed Hannachi: project initiation, conceptualization, calculations supervision, data interpretation, draft writing. Djamila Samsar: calculations, data interpretation, draft writing. Henry Chermette: calculations, data interpretation, draft writing.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors gratefully acknowledge the GENCI/CINES for HPC resources/computer time (Project cpt2130), and the PSMN of the ENS-Lyon for computing resources.

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