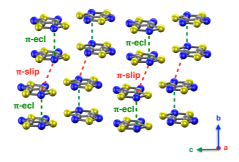
Reorganization of intermolecular interactions in the polymorphic phase transition of a prototypical dithiazolyl-based bistable material ¶

Tommaso Francese^{a,b}, Fernando Mota^a, Mercè Deumal^a, Juan J. Novoa^a, Remco W.A. Havenith^{b,c}, Ria Broer^b, Jordi Ribas-Arino^{a,*}

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Abstract:

The spin transitions undergone by several molecular crystals of dithiazolyl (DTA) radicals make this type of radicals promising candidates for future sensors and memory devices. Here, we present a systematic computational study of the intermolecular interactions existing in the two polymorphs of the neutral radical 1,3,5-trithia-2,4,6-triazapentalenyl in order to elucidate the origin of the difference in energy between those two polymorphs involved in its spin transition and to understand the crystal packing of this prototype of bistable materials. The π - π interactions between radicals are the main driving force for the crystal packing of both polymorphs, which comprises π -stacks of radicals. Among the interstack interactions, the strongest ones are those mediated by six- and four-center S···N bridges. The difference in energy between polymorphs, in turn, is mainly controlled also by the intrastack π - π intermolecular interactions, and the interstack S···S contacts instead of the S···N contacts. Since the supramolecular motifs herein identified as important for the crystal packing and/or for the energy difference between polymorphs (and, thus, for the spin transition temperature) are common to other members of the DTA family, our results provide valuable information to understand better the structure and properties of other switchable DTA-based materials.



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[¶] Dedicated to Jean-Paul Malrieu on the occasion of his 80th birthday, for his contribution to the modeling and the understanding of magnetic properties.

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The spin transitions undergone by several molecular crystals of dithiazolyl (DTA) radicals make this type of radicals promising candidates for future sensors and memory devices. Here, we present a systematic computational study of the intermolecular interactions existing in the two polymorphs of the neutral radical 1,3,5-trithia-2,4,6-triazapentalenyl in order to elucidate the origin of the difference in energy between those two polymorphs involved in its spin transition and to understand the crystal packing of this prototype of bistable materials. The π - π interactions between radicals are the main driving force for the crystal packing of both polymorphs, which comprises π -stacks of radicals. Among the interstack interactions, the strongest ones are those mediated by six- and four-center S···N bridges. The difference in energy between polymorphs, in turn, is mainly controlled also by the intrastack π - π intermolecular interactions, and the interstack S···S contacts instead of the S···N contacts. Since the supramolecular motifs herein identified as important for the crystal packing and/or for the energy difference between polymorphs (and, thus, for the spin transition temperature) are common to other members of the DTA family, our results provide valuable information to understand better the structure and properties of other switchable DTA-based materials.

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Introduction

Thiazyl-based radicals have been intensively investigated during the last decades because they can be used as versatile building blocks for molecular materials with interesting magnetic and/or electric properties. 1,2,3,4,5,6,7,8 These radicals have also become main actors in the field of switchable molecular materials, i.e., materials whose physical properties can be altered by means of external stimuli, such as temperature and light.^{9,10,11} Indeed, the family of dithiazolyl (DTA) neutral radicals has furnished multiple examples of metal-free compounds capable of undergoing structural phase transitions that entail a drastic change in the magnetic response of the material when it is subjected to changes of temperature 12,13,14,15,16,17,18,19,20, ^{21,22,23} and, in some cases, when it is irradiated with light.^{19,24} It should be mentioned that the closely-related family of dithiadiazolyl radicals has also provided two examples of switchable magnetic materials in recent years. 25,26 In some of the DTA phase transitions^{12-15,17,23}, the transition temperature in the warming cycle is found to be higher than the transition temperature in the cooling cycle. This hysteretic behavior gives rise to a loop in the magnetic susceptibility versus temperature plot, yielding a temperature range of magnetic bistability wherein the crystal can be observed in two different states depending on its immediate history. This intriguing property, which can also be observed in transition-metal based spin-crossover compounds 27,28,29 and in other purely organic materials 30,31,32, makes these radicals potential candidates for future sensors and memory devices.^{9,33}

All the phase transitions of DTAs that have been reported so far involve the formation/cleavage of dimers of radicals. In many cases $^{12-16,18,22}$, the dimers present in the crystal structure are cofacial π -dimers, which are held toghether by means of a long, multicenter bond 34,35 (alternativaly called "pancake" bond 36,37,38 , 39). These cofacial, eclipsed π -dimers are always found in the low-temperature (LT) phases of this family of switchable DTA compounds, where the cofacial, eclipsed π -dimers pile up giving rise to distorted π -stacks of DTA radicals that contain slipped pairs of cofacial π -dimers. Conversely, in the high-temperature (HT) phases of this family of switchable DTA compounds, the radicals pile up giving rise to regular π -stacks with a uniform distance between the slipped

radicals. The distinct difference in magnetic response between the LT and HT phases (LT phases are typically diamagnetic, while the HT phases exhibit weak paramagnetism) originates in the large changes in the magnetic exchange couplings between adjacent radicals in the π -stacks upon phase transition.^{40,41}

The transition temperature is a key property of any switchable material. Since the electronic energy difference between phases is one of the most important factors controlling the transition temperature, the identification of the elements that govern the energy difference between phases within a family of materials is a mandatory exercise in the quest of new materials with tailored transition temperatures. Here, we present a detailed computational study aimed at identifying the intermolecular interactions that play a key role in defining the energy difference between phases in switchable materials based on the formation/dissociation of eclipsed π -dimers between DTA radicals. As a model system of these materials, our study will focus on the 1,3,5-trithia-2,4,6-triazapentalenyl (TTTA) neutral radical 43 , which has become a prototypical example of molecule-based bistable materials on account of its spin transition with a wide hysteresis loop encompassing room temperature (see Figure 1) that can be induced both with temperature and absorption of light 13,24,40,44,45,46,47,48,49,50,51,52,53,54,55

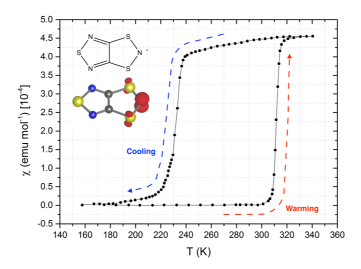


Figure 1. Temperature dependence of the paramagnetic susceptibility of the TTTA compound in the warming and cooling cycles. The insets show the molecular structure of a TTTA neutral radical and its spin density, which shows that the unpaired electron is delocalized over the S-N-S atoms of the dithiazolyl ring.

The hysteretic phase transition of TTTA involves a LT diamagnetic phase and a HT paramagnetic phase. The triclinic ($P\overline{1}$ space group) LT polymorph, which is the single polymorph observed on cooling below the bistability range, presents distorted π -stacks of radicals comprising slipped pairs of cofacial π -dimers. As displayed in Figure 2a, two types of intermolecular interactions between neighboring radicals are present in these distorted stacks: the long, multicenter bond between radicals in the eclipsed π -dimers (hereafter referred to as π -ecl interaction), and the interaction between slipped pairs of radicals (π -slip interaction). Conversely, the monoclinic ($P2_1/c$ space group) HT polymorph, which is the single polymorph observed on heating above the bistability range, presents regular π -stacks of radicals, where each molecule exhibits a slipped overlap with its two adjacent molecules along the stacking direction (π -slip interaction in Figure 3a).

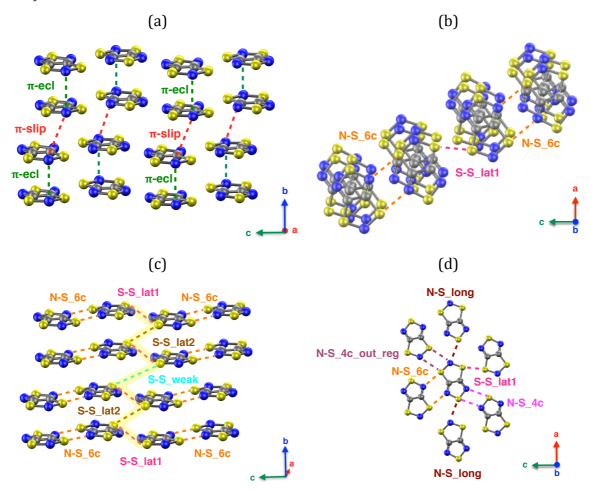


Figure 2. Different views of the crystal packing of the LT polymorph of TTTA (CCDC refcode = SAXPOW06) at room temperature (300K) along (a) , (b) , (c) and (d) . The different types of intermolecular interactions between radicals are marked (see main text for contact nomenclature).

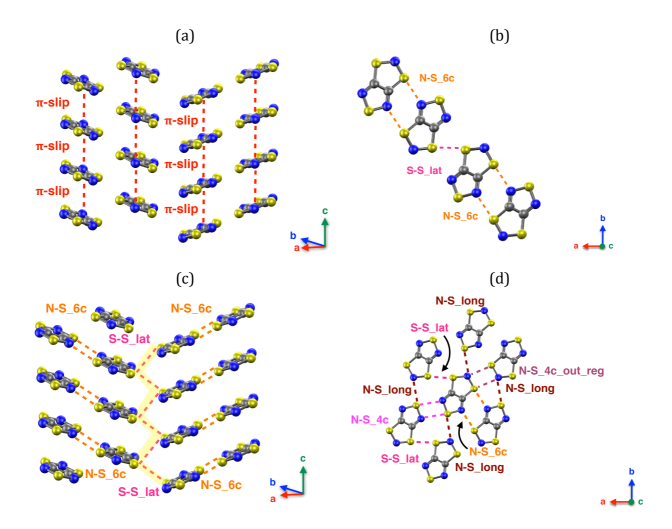


Figure 3. Different views of the crystal packing of the HT polymorph of TTTA (CCDC refcode = SAXPOW05) at room temperature (300K) along (a) , (b) , (c) and (d) . The different types of intermolecular interactions between radicals are marked (see main text for contact nomenclature).

The π -stacks of both LT and HT polymorphs of TTTA form 2D layers, where the π -stacks are laterally linked by a series of intermolecular 6-center S···N bridges (which will be referred to as N-S_6c interactions) and intermolecular S···S contacts (S-S_lat interactions, see Figures 2b and 3b). The 2D layers in LT contain one single molecular plane orientation (Figure 2c), whereas the 2D layers in HT contain two distinct molecular plane orientations (Figure 3c). The S-S_lat contacts in the 2D layers of both LT and HT phases define a zigzag pattern along the stacking direction (b axis) (Figures 2c and 3c). While the zigzag pattern in HT is regular (i.e., all the shortest interstack S···S contacts have the same distance), the zigzag pattern of LT features three different S-S_lat contacts: S-S_lat1, S-S_lat2 and S-S_weak, the

latter having the longest S···S distance (see Figure 2c). The different 2D layers of both phases of the TTTA crystal are linked by a series of interstack contacts, as shown in Figures 2d and 3d: 4-center S···N bridges involving radicals in the sample plane (N-S_4c interactions), 4-center S···N bridges involving radicals of out-of-registry stacks (N-S_4c_out_reg interactions), and S···N contacts (N-S_long interactions).

In this computational work, we will rationalize the different stability of the LT and HT polymorphs of TTTA on the basis of differences in intermolecular interactions between radicals. Specifically, the intermolecular interactions playing a leading role in establishing the energy difference between polymorphs and, thus, the transition temperature will be determined. In addition, we will identify which are the stronger intermolecular interactions and, thus, the interactions that drive the crystal packing of the polymorphs. Finally, the nature of the key intermolecular interactions present in the polymorphs will be evaluated. Since many of the intermolecular interactions herein considered are also present in other DTA-based crystals, the conclusions that will be drawn for TTTA will also be relevant for the other members of the family of DTA-based switchable materials.

Computational details

The analysis of the origin of the different stability of the two polymorphs of TTTA was performed by means of a set of single-point electronic structure calculations using different model systems: i) supercells containing 32 TTTA radicals for the solid-state calculations, ii) isolated π -stacks and isolated pairs of π -stacks of radicals, and iii) isolated dimers of radicals and isolated radicals. The single-point calculations of all these model systems were done using the relative atomic coordinates as directly extracted from the X-ray structures of the LT and HT polymorphs recorded at 300 K.¹³ These structures (whose diffraction data were collected using a Rigaku AFC5S four-circle diffractometer for HT, and on a Rigaku R-AXIS-IV imaging-plate system for LT) were obtained after isolating samples of the HT and LT phases by annealing at 40°C and at liquid nitrogen temperature for several hours, respectively.¹³ The supercells employed for the solid-state

calculations include 8 stacks of radicals, each of them containing 4 radicals. The lattice parameters of these supercells are collected in Table 1. In the calculations of the isolated π -stacks and isolated pairs of π -stacks, each stack comprised 4 radicals and periodic boundary conditions were considered along the stacking direction (the supercell vectors associated with the stacking direction were the same ones as those employed in the solid-state calculations). In a recent work⁵⁴, we demonstrated that the regular π -stacking motif (···A···A···A···A···)_n of the HT phase of TTTA is not a potential energy minimum but the average structure arising from a dynamic inter-conversion between two degenerate dimerized configurations: (···A-A···A-A···)_n \leftrightarrow (-A···A-A···A-)_n. We also demonstrated that the regular π -stacking motif is a minimum in the free energy surface of the system at 300 K. Consequently, the regular stacks of the X-ray recorded structure of the HT polymorph properly represent this phase and are thus adequate for the analysis herein presented.

The single-point electronic structure studies for all model systems were carried out with plane wave pseudopotential calculations using the PBE exchange-correlation functional 56,57 within the spin unrestricted formalism, together with Vanderbilt ultrasoft pseudopotentials 58 , and Γ -point sampling of the Brillouin zone, as implemented in the QUANTUM ESPRESSO package. 59 The semiempirical dispersion potential introduced by Grimme 60 (D2 version) was added to the conventional Kohn-Sham DFT energy in order to properly describe the Van der Waals interactions between the different TTTA radicals. The plane wave basis set was expanded at a kinetic energy cutoff of 35 Ry.

Table 1. Supercell parameters employed in the calculations of the LT and HT polymorphs of TTTA. The a, b and c parameters are given in Angstrom. The α , β and γ angles are given in degree.

	а	b	С	α	β	γ
LT ¹	15.062	20.046	14.048	100.598	96.978	77.638
HT	18.888	14.844	15.063	90.0	104.628	90.0

¹ The (a,b,c) supercell parameters of LT and HT are multiples of the (a,b,c) parameters defining the unit cells of the LT-300 and HT-300 X-ray resolved structures, respectively. The relations between the supercell (sc) parameters and the unit cell (uc) parameters are the following ones: $a_{\rm LT,sc} = 2 \cdot a_{\rm LT,uc}$; $b_{\rm LT,sc} = 2 \cdot b_{\rm LT,uc}$; $c_{\rm LT,sc} = 2 \cdot c_{\rm LT,uc}$; $a_{\rm HT,sc} = 2 \cdot a_{\rm HT,uc}$; $b_{\rm HT,sc} = 4 \cdot b_{\rm HT,uc}$; $c_{\rm HT,sc} = c_{\rm HT,uc}$

Some of the plane wave pseudopotential calculations of isolated π -stacks and isolated pairs of π -stacks and the plane wave pseudopotential calculations of isolated pairs of radicals were benchmarked against all-electron PBE-D2 calculations with the cc-pVTZ basis set^{61,62} using the CRYSTAL09 code.^{63,64} The DFT calculations of isolated pairs of radicals, in turn, were validated by means of NEVPT2 ^{65,66}/aug-cc-pVTZ calculations using the ORCA code.⁶⁷ These NEVPT2 calculations were carried out after CASSCF calculations using a minimum active space (i.e., 2 electrons in 2 orbitals). In all the calculations, an antiferromagnetic coupling between the spins of adjacent radicals was assumed, in agreement with previous studies.^{40,54}

The interaction energy decomposition analysis performed for a selected subset of pairs of radicals was carried out using an Energy Decomposition Analysis⁶⁸ method that can be applied within the DFT framework⁶⁹, as implemented in the GAMESS suite of programs.⁷⁰ In the method herein employed the interaction energy is decomposed into electrostatic, exchange, repulsion, polarization, and dispersion terms.⁶⁹ The nature of the bonding between radicals was also evaluated using the "Atoms in Molecules" (AIM) methodology.⁷¹ Specifically, bond critical points (BCPs) between pairs of radicals were located with the AIMAll⁷² software, which allows for a fully automated analysis of the topological features of the Laplacian of the electron density distribution, including all critical points. The molecular wavefunction data required by the AIMAll code was obtained by means of calculations performed with Gaussian09.⁷³

Results and Discussion

The presentation of the results is organized as follows. We will first analyze the intermolecular interactions in isolated pairs of radicals (Subsection 1). Then, we will investigate the different stability of the two polymorphs of TTTA on the basis of the energetics of isolated π -stacks of radicals and isolated pairs of π -stacks of radicals (Subsection 2). Finally, we will explore the nature of the intermolecular

interactions that play a key role in determining the different stability of the two polymorphs of TTTA (Subsection 3).

1. Intermolecular interactions in isolated pairs of TTTA radicals

In this subsection, we analyze the interaction energy between radicals in different isolated pairs of TTTA radicals in order to i) identify the strongest intermolecular interactions in the TTTA polymorphs, i.e., the interactions that are key driving forces in the crystal packing, ii) provide insights into the origin of the different stability of the two phases of TTTA, and iii) validate the DFT methodology used in the calculations carried out with periodic boundary conditions (subsection 2).

The set of isolated pairs of radicals that have been considered for the LT and HT polymorphs are displayed in Figures 4 and 5, respectively. The results collected in

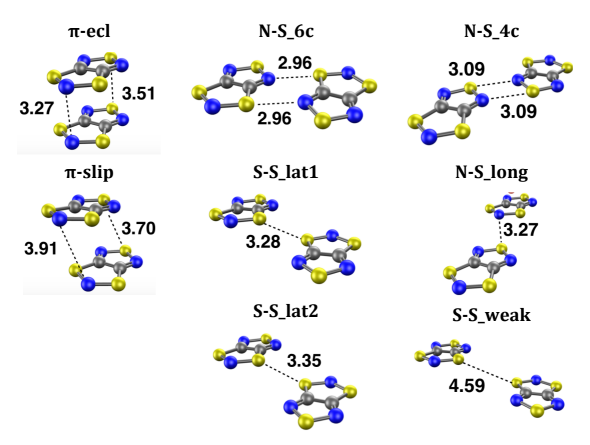


Figure 4. Different pairs of radicals in the LT polymorph of the TTTA crystal. The set of pairs considered in our study include those pairs with the shortest intermolecular contacts (distances are given in Å).

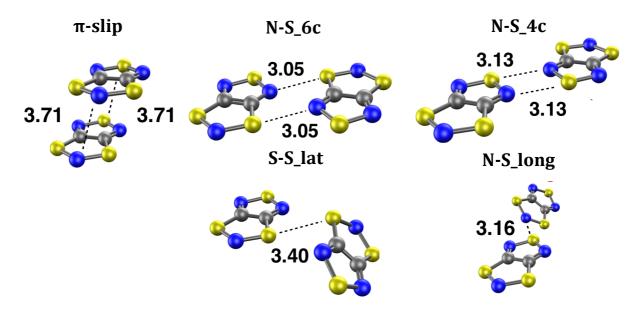


Figure 5. Different pairs of radicals in the HT polymorph of the TTTA crystal. The set of pairs considered in our study include those pairs with the shortest intermolecular contacts (distances are given in Å).

Table 2. Interaction energies^a ($E_{int,LT}$ given in kcal mol⁻¹) between pairs of radicals in the LT polymorph of TTTA, ordered according to stability.

Pair ^b	PBE-D2/USP-PW ^c	PBE-D2/cc-PVTZd	NEVPT2/aug-cc-PVTZ ^e
π-ecl	-7.66	-9.01	-16.04
π-slip	-4.81	-5.64	-11.13
N-S_6c	-5.10	-5.72	-8.22
N-S_4c	-3.77	-4.36	-6.63
S-S_lat1	-2.13	-2.44	-3.73
S-S_lat2	-2.31	-2.60	-3.86
N-S_long	-1.35	-1.67	-2.81
S-S_weak	-0.57	-0.73	-0.98

^a Each interaction energy was computed as $E(\text{pair}) - E(\text{radical}_1) - E(\text{radical}_2)$, where E(pair) is the energy of the pair of radicals and $E(\text{radical}_1)$ and $E(\text{radical}_2)$ are the energies of the isolated radicals forming the dimer; ^b The structures of the different types of pairs of radicals are displayed in Figure 4; ^c Plane wave pseudopotential DFT calculations carried out with Quantum Espresso; ^d All-electron DFT calculations using Gaussian basis sets carried out with Crystal; ^e Correlated wave function all-electron calculations carried out with ORCA.

Tables 2 and 3 reveal that the strongest intermolecular interaction between radicals in both polymorphs is found in the pairs exhibiting a π - π interaction (be it either of the π -ecl or the π -slip type of interaction). It then follows that the intermolecular interactions along the π -stacks are the most attractive interactions in the crystals of TTTA. Among the other types of interaction between radicals (i.e.

Table 3. Interaction energies^a (E_{int,HT} given in kcal mol⁻¹) between pairs of radicals in the HT polymorph of TTTA, ordered according to stability.

Pairb	PBE-D2/USP-PWc	PBE-D2/cc-PVTZd	NEVPT2/cc-PVTZe
π-slip	-5.04	-5.91	-11.42
N-S_6c	-4.83	-5.50	-7.75
N-S_4c	-3.65	-4.21	-6.48
N-S_long	-1.61	-1.94	-3.31
S-S lat	-2.24	-2.54	-4.05

^a Each interaction energy was computed as $E(\text{pair}) - E(\text{radical}_1) - E(\text{radical}_2)$, where E(pair) is the energy of the pair of radicals and $E(\text{radical}_1)$ and $E(\text{radical}_2)$ are the energies of the isolated radicals forming the dimer; ^bThe structures of the different types of pairs of radicals are displayed in Figure 5; ^cPlane wave pseudopotential DFT calculations carried out with Quantum Espresso; ^dAll-electron DFT calculations using Gaussian basis sets carried out with Crystal; ^eCorrelated wave function all-electron calculations carried out with ORCA.

Table 4. Difference in energies ($\Delta E_{int}(HT-LT)$ given in kcal mol⁻¹) between pairs of TTTA radicals of the HT polymorph and pairs of radicals of the LT polymorph.^a

НТ	LT	PBE-D2/USP-PWb	PBE-D2/cc-PVTZ ^c	NEVPT2/cc-PVTZd
π-slip	π-ecl	2.58	2.96	4.77
π-slip	π-slip	-0.27	-0.41	-0.14
N-S_6c	N-S_6c	0.24	0.09	0.63
N-S_4c	N-S_4c	0.09	0.02	0.31
S-S_lat	S-S_lat1	-0.14	-0.21	-0.13
S-S_lat	S-S_lat2	0.04	-0.05	-0.03
S-S_lat	S-S_weak	-1.70	-1.94	-2.91
N-S_long	N-S_long	-0.29	-0.40	-0.34

^a The difference in energy between two different pairs (pair1 and pair2) was computed as: ΔE (pair1, pair2)= E (pair1, HT) – E (pair2, LT); ^b Plane wave pseudopotential DFT calculations carried out with Quantum Espresso; ^c All-electron DFT calculations using Gaussian basis sets carried out with Crystal; ^d Correlated wave function all-electron calculations carried out with ORCA.

the interactions that connect π -stacks), the N-S_6c and N-S_4c interactions are significantly stronger than the others.

According to the differences in energy between pairs of radicals gathered in Table 4, the different stability of the LT and HT phases of TTTA is mainly controlled by the two following types of interaction: the interactions along the π -stacks (π -slip^{HT} and π -ecl^{LT}) and the interstack S···S contacts (S-S_lat^{HT} and S-S_weak^{LT}). The former type of interactions favor the LT polymorph because the transformation of the π -ecl interactions into π -slip interactions in going from LT to HT results in a much less attractive interaction (~4.7 kcal mol-1, according to NEVPT2 calculations) and,

thus, in a lower stability of HT. The interstack S···S contacts, instead, favor the HT polymorph due to the disruption of the regular zigzag pattern of S···S contacts in going from HT to LT. As a result of this disruption, one out of every four S···S contacts increases its distance (i.e., one out of every four S-S_lat^{HT} interactions transforms itself into a S-S_weak^{LT} interaction), thereby decreasing significantly its associated interaction energy (by about 2.9 kcal mol⁻¹, according to NEVPT2 calculations).

The results collected in Tables 2, 3 and 4 also demonstrate that PBE-D2 calculations (both with plane waves and cc-PVTZ basis sets) properly capture the main trends observed in the more accurate NEVPT2 calculations. Although the interaction energies and the differences in interaction energy obtained at the PBE-D2 level are smaller than those obtained at the NEVPT2 level, the conclusions that can be drawn from the PBE-D2 calculations are fully consistent with the conclusions drawn from the NEVPT2 calculations. It thus follows that the PBE-D2 methodology is adequate for describing the intermolecular interactions between TTTA radicals and, thus, adequate for the calculations considering periodic boundary conditions that will be presented in the next subsection.

2. Intermolecular interaction model systems considering periodic boundary conditions

In this subsection we shall further investigate the origin of the different stability of the two polymorphs of TTTA by means of PBE-D2 calculations of isolated π -stacks and isolated pairs of π -stacks.

Let us set the stage by first establishing the difference in lattice cohesive energy⁷⁴ between the two phases of TTTA. According to plane wave pseudopotential PBE-D2 solid-state calculations carried out using supercells of 32 radicals, the LT phase of TTTA is 0.92 kcal mol⁻¹ more stable (given per TTTA molecule) than the HT phase (*cf.* Table 5). The agreement between this value and the transition enthalpy of 0.6 kcal mol⁻¹ measured in DSC experiments⁴⁵ is remarkable taking into account that we are dealing with very small energy differences. The error associated with

Table 5. Differences in energy, ΔE (given in kcal mol⁻¹), between the HT and LT polymorphs of TTTA and between an isolated π-stack of HT and an isolated π-stack of LT.

ΔE polymorphs ^a	ΔE isolated π -stacks
0.92	1.08^{b}

 $^{^{\}rm a}$ The differences in energy are calculated as $E({\rm HT})$ – $E({\rm LT})$, and are given per TTTA radical. The values were obtained by means of plane wave pseudopotential DFT calculations carried out with Quantum Espresso.

the computational estimation of lattice energies is commonly 1 kcal mol⁻¹ or slightly above ⁷⁵. Nevertheless, the computational estimation of differences between the lattice energies of two polymorphs of TTTA within the framework of the same level of theory is perfectly meaningful.

The calculations carried out on isolated π -stacks, in turn, show that an isolated π -stack of LT is 1.08 kcal mol⁻¹ more stable than an isolated π -stack of HT. The fact that these two values are so similar demonstrates that the intra- π -stack interactions are the leading factor behind the different stability of the two polymorphs of TTTA. In fact, the difference in energy between a π -stack of LT and a π -stack of HT can be accurately predicted on the basis of the differences in energy gathered in Table 4. If we assume that the energy of a given column (given per radical) is determined exclusively by the energy of the TTTA radical and the sum of interaction energies between this radical and its nearest neighbors, the expression of the energies of the π -stacks of LT and HT can be written as:

$$E_{stack}^{LT} = E_{rad} + \frac{1}{2}E_{int}^{\pi - ecl} + \frac{1}{2}E_{int}^{\pi - slip, LT}$$
 (Eq. 1)

$$E_{stack}^{HT} = E_{rad} + E_{int}^{\pi - slip,HT}$$
 (Eq. 2)

where E_{rad} is the the energy of a single radical and E_{int} refers to the interaction energies between radicals. Note that in the expressions given above we distinguish the π -interaction between slipped radicals in LT from that of HT because they do not present the same interaction energy (*cf.* Tables 2, 3 and 4). On the basis of Eqs.

^b This value, obtained with Quantum Espresso, was further corroborated with all-electron calculations using a cc-pVTZ basis set (carried out with CRYSTAL09), which furnished a very similar difference in energy of 1.27 kcal mol⁻¹.

1 and 2, the difference in energy between an isolated π -stack of HT and isolated π -stack of LT can be expressed as:

$$E_{stack}^{HT} - E_{stack}^{LT} = \frac{1}{2} \left(E_{int}^{\pi - slip,HT} - E_{int}^{\pi - ecl} \right) + \frac{1}{2} \left(E_{int}^{\pi - slip,HT} - E_{int}^{\pi - slip,LT} \right)$$
 (Eq. 3)

The difference in energy between the HT and LT π -stacks that results from inserting the values of the first two entries of the "PBE-D2/USP-PW" column of Table 4 in the equation above is 1.15 kcal mol⁻¹. The close correspondence between this value and the value obtained from the PBE-D2 calculations of isolated π -stacks (1.08 kcal mol⁻¹) demonstrates that the difference in energy between the HT and LT isolated π -stacks is essentially governed by the short-range two-body interactions between neighboring radicals, primarily by the difference in energy between the π -ecl dimers of LT and the π -slip dimers of HT.

Upon comparing the difference in lattice energy between the HT and LT polymorphs (0.92 kcal mol⁻¹) with the difference in energy between the HT and LT isolated π -stacks (1.08 kcal mol⁻¹), it is concluded that the former is largely dominated by the latter and that the interactions between π -stacks make a nonnegligible contribution to the energy gap between HT and LT, specifically by decreasing it. In view of the non-negligible contribution of the interstack interactions, we shall now study them in detail. The interaction energies between π -stacks in both HT and LT polymorphs (*cf.* Table 6) follow the same trend as that observed in the interaction energies for the isolated pairs of radicals (Tables 2 and 3). Indeed, the strongest interaction between π -stacks is found for the stacks that interact through the N-S_6c interactions, which feature the largest interaction energy among the interstack contacts. The other interstack contacts are, in decreasing order of interaction strength, those mediated by N-S_4c contacts, lateral S···S contacts and N-S long contacts (cf. Table 6). As can be seen in Tables 2 and 3, the interaction energies of isolated pairs of radicals have the same ordering. However, the values of the interaction energies between π -stacks in Table 6 are considerably larger than the values that would be predicted on the basis of the interaction energies of isolated pairs of radicals (note that the values of Table 6 are given per radical, while the values of Tables 2 and 3 are given per pairs of radicals).

Table 6. Interaction energies (E_{int}) and differences in interaction energies (ΔE_{int}) (given in kcal mol⁻¹) between π -stacks of radicals in both HT and LT polymorphs.^a

Type of interaction	$E_{int}(HT)$	$E_{int}(LT)$	$\Delta E_{ m int}^{ m d}$
N-S_6c	-3.46	-3.34	-0.11
N-S_4c	-2.29	-2.41	0.12
S-S_lat ^b	-2.13	-1.86	-0.27e
N-S_long	-1.06	-1.17	0.11
N-S_4c_out_reg ^c	-1.38	-1.48	0.10

^a The interaction energies are given per TTTA radical. The values reported were obtained by means of plane wave pseudopotential DFT calculations carried out with Quantum Espresso.

This implies that the interaction energy between π -stacks cannot be expressed solely in terms of a sum of pairwise interaction energies between nearest neighbors, contrary to the intra- π -stack interaction energies.

When considering the difference in interaction energies between the π -stacks of HT and the π -stacks of LT ($\it cf$. Table 6), it is observed that all types of interactions result in small but not negligible relative stabilizations of either one or the other polymorph. Among all types of lateral interactions, the lateral S···S contacts are those that lead to a larger stabilization of one polymorph relative to the other one. Specifically, the lateral S···S contacts lead to a significant relative stabilization of the HT polymorph because the interaction energy between π -stacks through this type of contacts at HT is 0.27 kcal mol⁻¹ more stable than at LT. This result is in line with the intermolecular interactions evaluated for isolated pairs of radicals (see previous subsection), which showed that the increase of the S···S distance in going from the S-S_lat interaction at HT to the S-S_weak interaction at LT entailed a considerable weakening (by 0.85 kcal mol⁻¹, $\it cf$. Table 4)⁷⁶ of the intermolecular

^b Note that the S-S_lat-type interaction in HT involves only one type of interaction between pairs of radicals (the S-S_lat interaction, see Fig. 3c). Conversely, the S-S_lat-type interaction in LT involves three different types of interaction between radical pairs, namely: S-S_lat1, S-S_lat2 and S-S_weak (see Figure 2c).

^c The "N-S_4c_out_reg" type of interaction present in these columns resembles the type of interaction present in the "N-S_4c" case. However, in the former case the π -stacks are out-of-registry, whereas in the latter they are in-registry (see Figures 2d and 3d).

^d For a given type of interaction, the difference in interaction energy (ΔE_{int}) was obatined via: $\Delta E_{\text{int}} = E_{\text{int}}(\text{HT}) - E_{\text{int}}(\text{LT})$.

^e This value, obtained with Quantum Espresso, was further corroborated with all-electron calculations using a cc-pVTZ basis set (carried out with CRYSTAL09), which furnished a very similar difference in interaction energy of -0.30 kcal mol⁻¹.

interaction between radicals. It should be mentioned, though, that the relative stabilization of 0.27 kcal mol $^{-1}$ is somewhat smaller than the value that might have been expected on the basis of the results obtained for the isolated pairs. As explained in the previous paragraph, this is due to the fact that the interaction energy between π -stacks cannot be expressed solely in terms of a sum of pairwise interaction energies between nearest neighbors.

3. Evaluation of the nature of the key intermolecular interactions

In this subsection we shall gain further insight into the energetic differences between the LT and HT polymorphs of TTTA by means of an analysis of the intermolecular bond critical points (BCPs) present in the different pairs of radicals and an energy decomposition analysis of the interaction energy for the most relevant pairs. Before presenting the results of these analyses, it is worth mentioning that the intermolecular interactions in the crystal induce very small changes in the electron density of a TTTA radical, as shown by the electron density difference isosurfaces shown in Figure 6. The deformation of the electron density, which is mainly localized in the S-N-S fragment of the dithiazolyl ring, is slightly larger in the LT polymorph than in the HT polymorph.

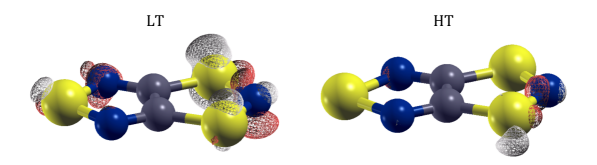


Figure 6. Electron density difference isosurfaces at an isovalue of $0.0015 \ e/Å^3$ for a TTTA radical in the LT (*left*) and in the HT (*right*) polymorphs; grey/red isosurfaces indicate charge depletion/accumulation. The electron density differences have been computed by subtracting the electron density of a given TTTA radical in the solid state (either in the LT or HT polymorphs) and the electron density of the same TTTA radical in the gas phase.

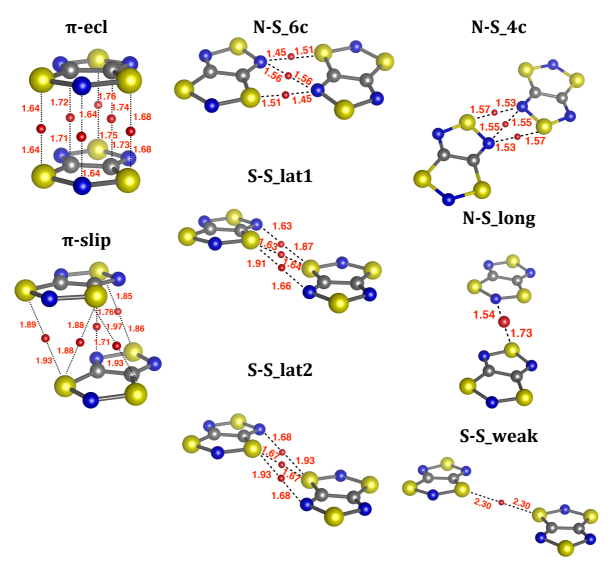


Figure 7. Bond critical points (BCP are marked as pink spheres) associated with several interatomic contacts of the different types of intermolecular interactions between radicals in the LT polymorph of TTTA. The images show the distances (given in Å) between the bond critical points and the atoms.

The number of BCPs (Figures 7 and 8) and the electron density at these BCPs (Table 7) correlate well with the strength of interaction energies of the isolated pairs of radicals (cf. Tables 2 and 3). The π -ecl interaction at LT, which is the most attractive interaction, features the largest number of BCPs (six), together with the largest value of the density on some of these critical points. In going from the π -ecl interaction of LT to the π -slip interaction of HT, not only the number of BCPs decreases (from six to five) but also the corresponding electron densities.

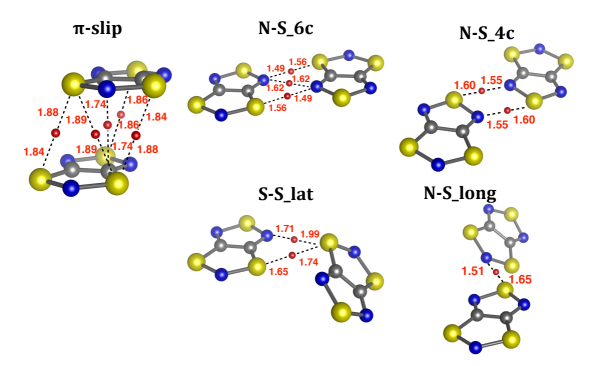


Figure 8. Bond critical points (BCP are marked as pink spheres) associated with several interatomic contacts of the different types of intermolecular interactions between radicals in the HT polymorph of TTTA. The images show the distances (given in Å) between the bond critical points and the atoms.

Interestingly, the BCP between the nitrogen atoms of the dithiazolyl rings, which are the atoms that formally hold the unpaired electron of the radical, disappears upon transformation of the π -ecl(LT) interaction into the π -slip(HT) interaction. Among all lateral interactions, the N-S_6c type of interactions exhibit both the largest number of BCPs and the largest values of electron density, which is consistent with the fact that these are the strongest lateral interactions. The analysis of BCPs also reflects the strong weakening of the lateral S···S contact in going from the S-S_lat interaction in HT the to S-S_weak interaction in LT. Even though the elongated S···S contact in S-S_weak(LT) still features a BCP, the data in Table 7 shows that its associated value of the electron density is one order of magnitude smaller than the corresponding value in the BCP associated with the S...S contact in the S-S_lat(HT) interaction. It is worth mentioning that the computed values of the electronic density in the BCPs associated with the S···S contacts of HT and the shorter S···S contacts of LT are very similar to the experimental value determined for the S···S contacts in TiS₂⁷⁷ and the computed values for other organic crystals featuring S···S chalcogen interactions. 78,79

Table 7. Values of the electronic density (in atomic units, i.e, $e ext{ a}_0^{-3}$) at the bond critical points associated with the intermolecular contacts in the LT and HT polymorphs of TTTA. For each type of intermolecular interaction (see Figs. 6 and 7), the value of the electronic density is reported for all the interatomic contacts presenting a bond critical point.^a

Н	T	LT		
π-s	slip	π-ecl		
$S(DTA)\cdots S(DTA)$	$0.0064^{ m b}$	S(DTA)···S(DTA)	$0.0146^{\rm b}$	
$N(DTA)\cdots N(DTA)$	_C	$N(DTA)\cdots N(DTA)$	0.0054	
$C(DTA)\cdots C(DTA)$	0.0044	$C(DTA)\cdots C(DTA)$	0.0045^{b}	
$S(TDA)\cdots S(TDA)$	0.0064	S(TDA)···S(TDA)	0.0096	
N-S	_6c	N-S	_6c	
N(TDA)···S(DTA)	$0.0112^{\rm b}$	N(TDA)···S(DTA)	$0.0134^{\rm b}$	
$N(TDA)\cdots N(TDA)$	0.0054	$N(TDA)\cdots N(TDA)$	0.0067	
N-S	_4c	N-S_4c		
$N(TDA)\cdots S(TDA)$	0.0088^{b}	N(TDA)···S(TDA)	0.0095^{b}	
$N(TDA)\cdots N(TDA)$	-	$N(TDA)\cdots N(TDA)$	0.0074	
N-S_	long	N-S_long		
$N(DTA)\cdots S(TDA)$	0.0092	N(DTA)···S(TDA)	0.0079	
S-S_lat		S-S_weak		
$S(DTA)\cdots S(DTA)$	0,0093	S(DTA)···S(DTA)	0.0008	
S(DTA)···N(TDA)	0.0038	S(DTA)···N(TDA)	-	
		S-S_lat ₁ / S-S_lat ₂		
		S(DTA)···S(DTA)	0.0106	
		S(DTA)···N(TDA)	0.0047^{b}	

^a The interatomic contacts are identified on the basis of the atoms of the heterocyclic rings (DTA and TDA refer to the dithiazolyl and thiadiazole rings, respectively) involved in the contact.

The evaluation of the nature of the key intermolecular interactions behind the different stability of the two polymorphs of TTTA was carried out by means of an energy decomposition analysis (EDA). Since the difference in stability between polymorphs is primarily dictated by the energy difference between the π -ecl(LT) and π -slip(HT) interactions and the energy difference between the S-S_weak(LT) and S-S_lat(HT) interactions, the EDA was carried out for these types of interaction only.

A comparison between the interaction energy components of the π -ecl dimers of LT and the π -slip dimers of HT reveals that all the attractive components of the interaction energy of the former are larger than those of the latter. The largest differences are found for the polarization and dispersion components (*cf.* Tables 8 and

^b This value results from taking the average of the density values of different bond critical points of the same type (see Figures 6 and 7).

^c There is no critical point associated with this contact.

Table 8. Energy decomposition analysis of the interaction energy of π -slip and π -ecl-type interactions in TTTA. All the components of the interaction energy are given in kcal mol⁻¹.

	π-slip (HT)	π-slip (LT)	π-ecl (LT)
Electrostatic	-3.23	-3.03	-7.30
Exchange	-2.74	-2.46	-6.35
Repulsion	15.28	14.09	33.07
Polarization	-3.95	-3.55	-11.46
DFT dispersion ^a	-5.77	-5.32	-10.81
Grimme dispersionb	-5.63	-5.47	-6.59
Total dispersion ^c	-11.40	-10.79	-17.40

^a This accounts for the dispersion that the PBE functional is able to recover by itself.

Table 9. Difference between the various components of the interaction energy of π -slip and π -ecl-type interactions in TTTA. All the differences are given in kcal mol⁻¹.

	$E(\pi\text{-slip, HT})$ - $E(\pi\text{-ecl, LT})$	$E(\pi\text{-slip, HT})$ - $E(\pi\text{-slip, LT})$
Electrostatic	4.07	-0.20
Exchange	3.61	-0.28
Repulsion	-17.79	1.19
Polarization	7.51	-0.40
DFT dispersion	5.04	-0.45
Grimme dispersion	0.96	-0.16
Total dispersion	6.00	-0.61

Table 10. Energy decomposition analysis of the interaction energy of S-S_lat and S-S_weak type interactions in TTTA. All the components of the interaction energy are given in kcal mol⁻¹.

	S-S_lat (HT)	S-S_weak (LT)	ΔE^a
Electrostatic	-1.61	0.02	-1.63
Exchange	-1.66	-0.12	-1.54
Repulsion	7.38	0.10	7.28
Polarization	-2.81	-0.36	-2.45
DFT dispersion ^b	-2.12	0.04	-2.16
Grimme dispersion	-1.83	-0.39	-1.44
Total dispersion	-3.95	-0.35	-3.60

^a Given a component of the interaction energy, this column collects the difference between the value of this component in S-S_lat(HT) and in S-S_weak(LT).

9), in line with the known fact that pancake bonding between π -radicals is dominated by SOMO–SOMO bonding interaction and dispersion interaction.^{34,38,80}. It is also worth mentioning that the electrostatic interaction between radicals is attractive and significantly larger in the π -ecl configuration even if in this configuration the

^b This accounts for the dispersion recovered by Grimme's D2 semiempirical approach.

^cThe values reported in this entry are just the sum of the values of "DFT dispersion" and "Grimme dispersion".

^b This accounts for the dispersion that the PBE functional is able to recover by itself.

polarized charges of the same sign are on top of each other. The repulsion energy between radicals, in turn, is much larger in the π -ecl dimers than in the π -slip dimers. Therefore, the π -ecl dimers of LT are more stable than the π -slip dimers of HT because the larger attractive components of the interaction energy (of which, the polarization and dispersion are the dominant ones) compensate the increase in repulsion energy in going from π -ecl(LT) to π -slip(HT). On the other hand, the values gathered in Table 10 demonstrate that the weakening of the lateral S···S bonds in going from HT to LT is due to a weakening of all the attractive components of the interaction energy, especially the dispersion and polarization energies.

Conclusions

On the basis of the computational study herein presented, we have identified the intermolecular interactions that drive the crystal packing of the two polymorphs of TTTA and the intermolecular interactions that play a prime role in establishing the different stability of the polymorphs of this prototypical example of a bistable material based on dithiazolyl radicals. The formation of π -stacks of radicals in the two polymorphs is driven by strong π - π interactions, which are the strongest of all intermolecular interactions between TTTA radicals. Among the lateral interactions that connect the π -stacks together, the stronger ones are those mediated by sixcenter S···N bridges, followed by those mediated by four-center S···N bridges.

The difference in stability between polymorphs is primarily governed by two types of intermolecular interactions: the π - π intermolecular interactions along the π -stacks and the S···S lateral contacts between π -stacks. The π - π interactions result in a substantial stabilization of the LT phase (relative to the HT phase) because the binding between radicals in the eclipsed π -dimers of the LT phase is stronger than in the slipped π -dimers of the HT phase. The stronger binding of the eclipsed π -dimers mainly originates in larger values of the polarization and dispersion components of the energy interaction. The interstack S···S contacts, in turn, result in a destabilization of the LT phase (relative to the HT phase) because some of the lateral S···S contacts of LT have a significantly weaker interaction than the lateral S···S contacts of HT. The interplay between the effects of the π - π and the S···S

interactions (in which the effect of the π - π interactions is the dominant one) in defining the difference in energy between polymorphs is tuned to a small but non-negligible extent by other types of interstack interactions, including those interactions mediated by six- and four-center S···N bridges.

The results herein presented are not only relevant for the protypical TTTA compound, but also for other members of the family of dithiazolyl-based switchable materials because many of the supramolecular motifs identified as being important for driving the crystal packing and/or determining the difference in stability between polymorphs of TTTA are common to other members of this family. 12,14-16,23 Therefore, our results provide valuable information for the rationalization of the crystal packing and spin transition temperatures of DTA-based switchable crystals, as well as for the design of new materials of this type.

Acknowledgements

FM, MD, JJN and JRA acknowledge financial support from MINECO through Grant CTQ2017-87773-P/AEI/FEDER, UE and Spanish Structures Excellence María de Maeztu program through grant MDM-2017–0767. FM, MD, JJN and JRA are also thankful to the Catalan DURSI (2017SGR348 grant) and to BSC and CSUC for the allocation of computer time. TF acknowledges funding from the European Union's Horizon 2020 research and innovation programme under the Marie Curie Skłodowska-Curie grant agreement no. 642294.

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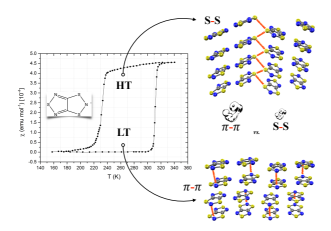
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A computational study of the intermolecular interactions existing in the two polymorphs of the bistable neutral radical 1,3,5-trithia-2,4,6-triazapentalenyl (TTTA) has shown that the energy difference between its two phases is mainly controlled by an interplay between π - π interactions and S···S interactions.