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Redox-Active Dysprosium Single-Molecule Magnet: Spectro-Electrochemistry and Theoretical Investigations

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Abstract: The mononuclear single-molecule magnet (SMM) [Dy(tta)3(L)]·C6H14 (1) (where tta = 2-thienoyletrifluoroacetate and L = 4,5-bis(propylthio)tetrathiafulvalene-2-(2-pyridyl)benzimidazole-methyl-2-pyridine) was studied by spectro-electrochemistry. The resulting electronic spectra of the three oxidation states 1, 1+, and 12+ were rationalized by time-dependent density functional theory (TD-DFT) calculations starting from the DFT optimized structures. The modulation of the magnetic anisotropy of the DyIII center upon oxidation was also inspected at the Complete Active Space Self-Consistent Field (CASSCF) level of calculation.

Keywords: dysprosium; tetrathiafulvalene; single-molecule magnet; spectro-electrochemistry; TD-DFT calculations; ab initio calculations

1. Introduction

Tetrathiafulvalene (TTF) is a well-known organic moiety successfully used in the elaboration of molecular conductors [1–7] and molecular switches [8–11] as well as solar-energy systems [12,13] due to its ability to be a reversible, strong electron-donating species. To reach the design of multifunctional materials, the intense effort by chemists to combine the electronic conductivity with other physical properties is the focus of current research [14–17]. For instance, during the last decade, the “through bond” strategy, consisting in the functionalization of the TTF core with organic groups suitable for a coordination reaction with metal precursors [18–24], has been used to combine lanthanide magnetism and the attractive reversible redox properties of TTF derivatives [25–28]. Indeed, lanthanide ions possess a strong anisotropy and a large magnetic moment which make them good candidates for obtaining single-molecule magnets (SMMs) [29–37]. The motivation to design these new molecular objects is driven by their potential abilities to pave the way for a new generation of quantum devices, such as high-density data storage, quantum computing, and spintronics [38–43]. Thus, the combination of the lanthanides and TTF-based ligands led to exciting multiple properties SMMs [44–54]. Nevertheless, in these SMMs, TTF-based ligands are in their neutral state, and the examples of lanthanide coordination complexes involving radical cation TTF species are still very scarce [55,56]. In fact, there are only two examples to date of the direct comparison between two systems in which the TTF-based ligand is in its neutral and radical cation oxidation states to evaluate the effect of the oxidation on the magnetic properties [57,58].

In the present article, we focus our attention on a mononuclear complex of formula [Dy(tta)3(L)]·C6H14 (1) (where tta = 2-thienoyletrifluoroacetate and L = 4,5-bis(propylthio)tetrathiafulvalene-2-(2-pyridyl)benzimidazole-methyl-2-pyridine) that displays SMM behavior with
hysteresis at low temperature [59]. During the investigation of 1, its stability in CH$_2$Cl$_2$ solution was demonstrated by the study of the electronic spectrum and by measuring the hysteresis loop in frozen solution. Complex 1 was also at the origin of the highlight of isotopic effects on the relaxation time of the magnetization [60–62]. In spite of the plethora of studies performed on this system, no study was realized to probe the physical properties as well as the magnetic anisotropy after oxidation of the L ligand. Thus, to go one step further, we studied, herein, the electronic spectra of the first and second oxidation states of 1 (i.e., 1$^{+}$ and 1$^{2+}$, respectively) by spectro-electrochemistry followed by their rationalization upon TD-DFT (time-dependent density functional theory) calculations on the DFT optimized complexes. Finally, the influence of the oxidation on the magnetic properties was evaluated using Complete Active Space Self-Consistent Field/Restricted Active Space State-Interaction - Spin Orbit (CASSCF/RASSI-SO) calculations.

2. Results and Discussion

2.1. X-ray Structures of [Dy(tta)$_3$(L)] and DFT Optimized Structures of [Dy(tta)$_3$(L)] (1$_{opt}$), [Dy(tta)$_3$(L$^{+}$)] (1$_{opt}^{+}$) and [Dy(tta)$_3$(L$^{2+}$)] (1$_{opt}^{2+}$)

The X-ray structure of 1 was refined and described in a previous work [60]. Briefly, the experimental molecular structure was used as starting point for the DFT optimized structures of [Dy(tta)$_3$(L)] (1$_{opt}$), [Dy(tta)$_3$(L$^{+}$)] (1$_{opt}^{+}$) and [Dy(tta)$_3$(L$^{2+}$)] (1$_{opt}^{2+}$) (Figure S2). 1 consists in a mononuclear complex of Dy$^{III}$ (Figure 1 and Figure S3). The Dy$^{III}$ ion lies in an N$_2$O$_6$ coordination sphere with a square antiprism symmetry (SAPR-8 at 0.538 using SHAPE program [63]). The six oxygen atoms and two nitrogen atoms belong to three hfac$^-$ anions and the L ligand, respectively. The average Dy-N and Dy-O distances were 2.530 Å and 2.333 Å, respectively. The central C=C bond length (1.347 Å) and the non-planar conformation of the TTF core attest the neutrality of L.

![Figure 1. X-ray structure of 1 (yellow) and optimized structures for the different oxidation states 1$_{opt}$ (red), 1$_{opt}^{+}$ (blue) and 1$_{opt}^{2+}$ (green).](image)

The DFT optimization of the structure of 1 led to a complete boat conformation for the TTF core (Figure 1 and Figure S2). While in the crystal structure, the π-bonding between TTFs of neighboring molecules maintained the structure flat, in the absence of such interaction, the preferable minima for the ligand was a boated one. The Dy-N (2.551 Å) and central C=C (1.352 Å) distances were found slightly longer than for the X-ray structure (Dy-N = 2.530(5) Å and C=C = 1.347(9) Å) while Dy-O (2.322 Å) was slightly shorter (Dy-O = 2.333(4) Å). The symmetry of the Dy$^{III}$ surrounding remained D$_{4d}$ with a very similar distortion compare to the ideal square antiprism (0.526). The crystal packing of 1 was driven by the formation of head-to-tail dimers of complexes. One can remark that the bent ligand character is more pronounced in 1$_{opt}$ than for 1. It could be due to the fact that the flat ligand allows an optimal π-π interaction in the head-to-tail dimers of complexes while for the optimized
structure, the neighboring molecules are not taken into account in the calculations. Thus, the ligand is free to adopt such pronounced boat conformation in $1_{\text{opt}}$.

After the mono- (Figure S4) and di-oxidation (Figure S5) of the TTF core, the optimized structures revealed an elongation of the central C=C bond (1.391 Å for $1^{+}_{\text{opt}}$ and 1.419 Å for $1^{2+}_{\text{opt}}$) due to the loss of electrons. The latter led to an increase of the aromaticity of $L$ due to the removal of electrons in the $\pi$ C=C and $\pi^*$ C=S molecular orbitals (MOs) (see $1_{\text{opt}}$, $1^{+}_{\text{opt}}$, and $1^{2+}_{\text{opt}}$), and thus the TTF moiety became increasingly planar (Figure 1). The two previous observations are in agreement with the well-known consequences of a TTF core centered oxidation [64–66].

The decrease of the electron density on the TTF core affected the electron density of the benzoimidazole-pyridine moiety, too, and consequently the Dy-N (2.562 Å for $1^{+}_{\text{opt}}$ and 2.318 Å for $1^{2+}_{\text{opt}}$) and Dy-O (2.569 Å for $1^{+}_{\text{opt}}$ and 2.315 Å for $1^{2+}_{\text{opt}}$) distances were slightly longer and shorter than for the neutral species, respectively. The symmetry of the Dy$^{III}$ surrounding remained D$_{4d}$ with a slightly higher distortion compared to the ideal square antiprism (0.600 for $1^{+}_{\text{opt}}$ and 0.590 for $1^{2+}_{\text{opt}}$).

The X-ray structure of $1$ and the three optimized structures demonstrated the integrity of the coordination complexes for all the oxidation states of the TTF core.

2.2. Electrochemical Properties

The redox properties of the ligand $L$ and complex $1$ were investigated by cyclic voltammetry. The cyclic voltammograms for $L$ and $1$ showed two mono-electronic oxidations at 0.504 V (reversible) and 0.912 V (quasi-reversible) for $L$ and 0.519 V (reversible) and 0.941 V (quasi-reversible) for $1$. They correspond successively to the formation of the TTF fragment radical cation and dication (Figures S6 and S7). Oxidation potentials were shifted to higher positive values relative to the parent TTF due to partial electron transfer from the donor (TTF) to the benzoimidazole-pyridine acceptor moiety. More interesting is the slight shifts to higher oxidation potentials from $L$ to $1$ which illustrate the weak decrease of electron density on the TTF core after coordination. The electrochemical properties attest to the redox-activity of $L$ and the corresponding complex. Such redox properties are required to perform spectro-electrochemistry and give crucial information on the potential values which must be applied (see next section).

2.3. Photo-Physical and Spectro-Electrochemical Properties

The electronic properties of the complex were investigated in its neutral state ($1$), in its first ($1^{+}$) and second ($1^{2+}$) oxidized states during electrochemical oxidation in the 9000 to 27,000 cm$^{-1}$ energy range.

The electronic spectrum of $1$ in this energy range was mainly composed of one intense absorption band centered at 23200 cm$^{-1}$ (red lines in Figure 2 and red decomposition in Figure 3a) which has been identified as intra-ligand charge transfer (ILCT) from the TTF fragment to both bzip and 2-thienyltrifluoroacetate (tt$^-$) units (Table 1) [59]. The modification of the electronic spectrum was further studied during electrochemical oxidation in an optically transparent thin-layer electrochemical (OTTLE) cell. The applied potential of the device was firstly set between the first and second oxidation potentials of $1$ to study the radical cation form $1^{+}$. Under an applied potential of 0–0.7 V (Figure 2a), several new absorption bands appeared (Figure 2b).
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Figure 3. (a) Experimental UV-visible absorption spectrum of 1 in the visible-NIR range during its first oxidation in 0.2 M n-Bu4NPF6/CH2Cl2. (b) Absorption spectra and spectroscopic changes for 1 in the visible-NIR range during its first oxidation in 0.2 M n-Bu4NPF6/CH2Cl2. (c) Absorption spectra and spectroscopic changes for 1 in the visible-NIR range during its second oxidation in 0.2 M n-Bu4NPF6/CH2Cl2. (d) Absorption spectra for 1 (red curve), 1+ (blue curve), and 12+ (green curve).

Figure 3. (a) Experimental UV-visible absorption spectrum of 1 in CH2Cl2 solution (open grey circles), respective Gaussian decompositions (dashed red lines), and best fit (full black line, R² = 0.9980). (b) Experimental UV-visible absorption spectrum of 1+ in CH2Cl2 solution (open grey circles), respective Gaussian decompositions (dashed red lines), and best fit (full black line, R² = 0.9974). (c) Experimental UV-visible absorption spectrum of 12+ in CH2Cl2 solution (open grey circles), respective Gaussian decompositions (dashed red lines), and best fit (full black line, R² = 0.9965). (d) Experimental positions and intensities of the Gaussian decompositions for 1 (red sticks), 1+ (blue sticks), and 12+ (green sticks).
Both experimental (Figures 2d and 3a,b) and calculated (Figure 3) absorption spectra are in perfect agreement. The three lowest-energy bands are identified as intra-donor (ID) excitations involving π−−π* transitions. The HOMO−LUMO excitations (HOMO→LUMO, Table 1) and to the HOMO−LUMO ILCT transitions, and the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO), respectively. Therefore, ILCT stands for intra-ligand charge transfer. The theoretical values are evaluated at the PCM(CH₂Cl₂)-PBE0/SVP level of approximation.

| Exp. Energy (cm⁻¹) | Theo Energy (cm⁻¹) | Osc. | Type | Assignment | Transition |
|--------------------|--------------------|------|------|------------|------------|
| 1                  |                    |      |      |            |            |
| 23,200             | 21,044             | 0.23 | ILCT | π₁TTF→π₁*_{bzip} | H→L (91%) |
| 21,356             |                    | 0.04 | ID   | π₁TTF→π₁*_{u} | H→L + 1 (72%) |
| 12,300             | 12,475             | 0.35 | ID ILCT | π₁TTF→π₁*_{TTF} | H→L-2 (75%) |
| 15,800             | 18,491             | 0.02 | ID   | π₁TTF→π₁*_{TTF} | H→L-3 (75%) |
| 18,900             | 19,844             | 0.02 | ID   | π₁TTF→π₁*_{TTF} | H→L-2 (30%) |
| 21,500             | 22,325             | 0.20 | ILCT | π₁TTF→π₁*_{bzip} | H⁺→L (65%) |
| 22,900             | 23,823             | 0.06 | ID   | π₁TTF→π₁*_{TTF} | H⁺→L + 6α (37%) |
| 24,200             | 24,449             | 0.22 | IA ID ILCT | π₁_{bzip}→π₁*_{bzip} | H⁺→L+1α (15%) |
| 25,400             | 25,732             | 0.12 | ID   | π₁TTF→π₁*_{TTF} | H⁺→L+6α (21%) |
| 12,000 *           | 12,475             | 0.35 | ID ILCT | π₁TTF→π₁*_{TTF} | H⁻→L-2 (30%) |
| 15,400             | 13,620             | 0.66 | ID ILCT | π₁L→π₁*TF | H⁻→L-2 (69%) |
| 14,100             | 14,085             | 0.19 | ILCT | π₁L→π₁*TF | H⁻→L (65%) |
| 14,900             | 14,689             | 0.11 | ILCT | π₁L→π₁*TF | H⁻→L (69%) |
| 16,200             | 15,063             | 0.05 | ILCT | π₁L→π₁*TF | H⁻→L (65%) |
| 18,600 *           | 19,844             | 0.02 | ID   | π₁TTF→π₁*_{TTF} | H⁻→L-2 (49%) |
| 21,500 *           | 22,325 23,823      | 0.20 | ILCT ID | π₁TTF→π₁*_{bzip} | H⁻→L + 1α (15%) |
| 23,500             | 23,769             | 0.02 | ID   | π₁TTF→π₁*_{TTF} | H⁻→L (49%) |
| 24,700 *           | 24,449 25,732      | 0.220 | IA ID ILCT | π₁_{bzip}→π₁*_{bzip} | H⁻→L + 1α (15%) |

* These absorption contributions come from the 1⁺⁺.

The resulting electronic spectrum of 1⁺⁺ is depicted in Figure 3b, and it was decomposed in seven contributions. The three lowest-energy bands are identified as intra-donor (ID) excitations involving the radical cation TTF⁺⁺ core. At this point, it is worth noting that the lowest-energy band centered at 12,300 cm⁻¹ (calculated at 12,475 cm⁻¹, Figure 4) is also composed of an intra-ligand charge transfer (ILCT) excitation (HOMO-2β→LUMOβ, Figure 5 and Table 1) starting from the tta⁻⁻ ancillary ligand to the TTF⁺⁺. Such ILCT excitations are allowed because of (i) the strong decrease of the donating character of the radical-cation TTF⁺⁺ core compared to the neutral TTF core [67,68] and (ii) the stronger accepting character of the tta⁻⁻ anions than bzip moiety as attested by the nature of the ILCT in 1 and as already observed in other TTF-based systems involving tta⁻⁻ anions [59]. Then four new additional absorption bands appeared after oxidation of the complex at 21,500, 22,900, 24,200, and 25,400 cm⁻¹ (Figures 2–4) which are attributed to π→π* intra-TTF⁺⁺ (ID) excitations (HOMOα→LUMO + 5/6 + 6α, Figure 5 and Table 1) and to the HOMOα→LUMOα ILCT. The HOMOα and LUMOα are respectively centered TTF⁺⁺ and bzip orbitals (Figure 5), and consequently the HOMOα→LUMOα ILCT of 1⁺⁺ (21500 cm⁻¹, Table 1) is analogous to the HOMO→LUMO ILCT of 1 (23,200 cm⁻¹, Table 1). Both experimental (Figures 2d and 3a,b) and calculated (Figure 3) absorption spectra are in perfect agreement.
agreement. The reversibility of the first electrochemical oxidation of 1 was checked showing an almost complete reversible process (Figure S8).

*Figure 4.* Theoretical absorption spectra for 1_{opt} (red curve), 1^{+}_{opt} (blue curve), and 1^{2+}_{opt} (green curve); the bars represent the mean contribution of the absorption spectra that are listed in Table 1.

*Figure 5.* Molecular Orbital diagram of 1_{opt}, 1^{+}_{opt}, and 1^{2+}_{opt}. Energy levels of the centered TTF donor, bzip acceptor, and tta⁻ anion are represented in orange, blue, and green, respectively. Full vertical black lines represent the electrons of the occupied orbitals.

Spectroscopic changes under an applied potential of 0.75 to 1.10 V are depicted in Figure 2a,c. Comparison of the experimental (Figure 3c) and calculated absorption spectra (Figure 4) leads to the conclusion that even at 1.10 V a mixture of the 1⁺ (blue decompositions) and 1^{2+} (green decompositions) species is observed. Unfortunately, in our experimental conditions, the application of higher potentials led to irreversible decomposition of the species. Nevertheless, the absorption contributions attributing to 1^{2+} could be determined. Thus, in the energy range of 9000 to 27,000 cm⁻¹, five decompositions have been identified (Figure 3c,d). The four absorption bands centered at 13,400, 14,100, 14,900, and 16,200 cm⁻¹ and calculated at 13620, 14085, 14689, and 15063 cm⁻¹ (Table 1) are attributed to ILCT between the highest occupied molecular orbital (HOMO)-4 (fully delocalized orbital on the whole ligand) and the lowest unoccupied molecular orbital (LUMO) (centered on the dicationic TTF^{2+}.
core) and ILCT between tta\textsuperscript{−} centered orbital and the LUMO. At higher energy, an ID HOMO-20 → LUMO (23,500 cm\textsuperscript{−1}) excitation was also identified. The electronic spectrum of 1\textsuperscript{2+} was composed of excitations involving the LUMO as the accepting orbital because of the drastic decrease of the donating character of the dicationic TTF\textsuperscript{2+} core compared to the neutral TTF and radical cationic TTF\textsuperscript{+} cores. In the fully oxidized 1\textsuperscript{2+} compound, the tta\textsuperscript{−} ancillary ligands can be seen as the best accepting groups of the system. Finally, the HOMO-4 for 1\textsuperscript{2+} is the equivalent of the HOMO for 1 and 1\textsuperscript{+}.

Interestingly, the delocalization of these highest-energy TTF-centered orbitals is proportional to the aromaticity of the ligand. Thus, the HOMO is totally centered on the TTF core (Figure 5), which adopts a boat conformation (Figure 1), in 1; the HOMO is mainly centered on the TTF\textsuperscript{+} core, which is almost planar (Figure 1), in 1\textsuperscript{+} and the HOMO is totally delocalized on the whole L ligand, which is totally flat (Figure 1), in 1\textsuperscript{2+}.

The second oxidation of 1 is badly reversible (Figure S9) in agreement with the electrochemical properties, which show only a quasi-reversible second oxidation at 0.94 V (Figure S7).

2.4. Magnetic Properties

SA-CASSCF/RASSI-SO calculations were performed for each oxidation state of 1 to tentatively evaluate the potential impact of the oxidation of the TTF core on the magnetic anisotropy of the Dy\textsuperscript{III} center. Calculations demonstrated that the axial character of the magnetic anisotropy tensor of the almost pure ground Kramers doublet $M_J = \pm 15/2$ is conserved after the TTF-centered oxidation (Table S1). The calculated $g_{zz}$ components are equal to 19.52 for 1, 19.56 for 1\textsuperscript{opt}, 19.57 for 1\textsuperscript{+ opt}, and 19.45 for 1\textsuperscript{2+ opt} (Table S1). Whatever the oxidation state, the main component of the magnetic anisotropy ($g_{zz}$) remains oriented perpendicular to the plane formed by the nitrogen atoms of the benzo-imidazole-2-pyridine moiety as expected for an oblate ion (Figure S10). Finally, the energy splitting of the $6\text{H}_{15/2}$ ground state is almost not affected by the oxidation process (Figure S11). In other words, the oxidation of the TTF core has presumably no influence on the magnetic anisotropy from a molecular perspective. However, from a computational perspective, these calculations cannot completely exclude an influence of the oxidation on the magnetism. Taking the 1\textsuperscript{+ opt} as an example, to affirm this, it would be necessary to perform a calculation on an active space that includes at least the metal ion, the electron on the TTF, and all its correlated orbitals. However, such calculations demand is, for now, prohibitive. In any case, the observations made from our state-of-the-art calculations are in agreement with what has been observed in previously studied systems highlighting that to obtain a redox-magnetic switching, the organic radical should be directly connected to the lanthanide ion [69–71].

3. Materials and Methods

3.1. Synthesis. General Procedures and Materials

All solvents were dried using standard procedures. 4,5-bis(propylthio)-tetrathiafulvalene-2-(2-pyridyl)benzimidazole-methyl-2-pyridine ligand (L) [72], Dy(tta)\textsubscript{3}·2H\textsubscript{2}O [73] precursor and [Dy(tta)\textsubscript{3}(L)]·C\textsubscript{6}H\textsubscript{14} (1) [59] compound were prepared according to the literature procedure. All the reagents were purchased from Aldrich and Alfa Aesar Co. Ltd. and were used without further purification.

3.2. Physical Measurements

Cyclic voltammetry was carried out in CH\textsubscript{2}Cl\textsubscript{2} solution, containing 0.1 M (C\textsubscript{4}H\textsubscript{9})\textsubscript{4}NPF\textsubscript{6} as supporting electrolyte. Voltammograms were recorded at 100 mV s\textsuperscript{−1} at a platinum disk electrode. The potentials were measured versus a saturated calomel electrode (SCE). Optical spectra were measured using the KBr disk method on Perkin-Elmer 1600 Series FT-IR (resolution 4 cm\textsuperscript{−1}) for infrared (IR). Absorption spectra were recorded on a Varian Cary 5000 UV-Visible-NIR spectrometer. Electrochemical studies were carried out under argon using an Eco Chemie Autolab PGSTAT
Wavefunction-based calculations were carried out on the experimental molecular structure of \(1\), with the SVP basis sets [78]. The first 50 monoelectronic excitations were calculated. A Pt mesh was used as the working electrode, a Pt wire was used as the counter electrode, and an Ag wire was used as a pseudo reference electrode. The electrodes were arranged in the cell such that the Pt mesh was in the optical path of the quartz cell. The anhydrous freeze–pump–thaw degassed sample-electrolyte solution (0.2 M n-Bu4NPF6) was cannula-transferred under argon into the cell previously thoroughly deoxygenated.

3.3. Computational Details

DFT geometry optimizations and TD-DFT excitation energy calculations of the \(Y^{III}\) analogs of the corresponding complexes \(1_{opt}\), \(1^{+}_{opt}\), and \(1^{2+}_{opt}\) were carried out with the Gaussian 09 (revision A.02) package [74] employing the PBE0 hybrid functional [75,76]. The “Stuttgart/Dresden” basis sets and effective core potentials were used to describe the yttrium atom [77], whereas all other atoms were described with the SVP basis sets [78]. The first 50 mono-electronic excitations were calculated. In all steps, a modeling of bulk solvent effects (solvent = dichloromethane) was included through the Polarisable Continuum Model (PCM) [79], using a linear-response non-equilibrium approach for the TD-DFT step [80,81].

Molecular orbitals were sketched using the Gabetor graphical interface [82]. Wavefunction-based calculations were carried out on the experimental molecular structure of \(1\) and on the DFT optimized structures \(1_{opt}\), \(1^{+}_{opt}\), and \(1^{2+}_{opt}\) by using the SA-CASSCF/RASSI-SO approach, as implemented in the MOLCAS quantum chemistry package (versions 8.0) [83]. In this approach, the relativistic effects are treated in two steps based on the Douglas–Kroll Hamiltonian. First, the scalar relativistic effects were included in the basis-set generation and were used to determine the spin-free wavefunctions and energies in the complete active space self-consistent field (CASSCF) method [84]. Next, spin-orbit coupling was added within the restricted-active-space-state-interaction (RASSI-SO) method, which uses the spin-free wavefunctions as basis states [85,86]. The resulting wavefunctions and energies are used to compute the magnetic properties and g-tensors of the lowest states from the energy spectrum by using the pseudo-spin \(S = \frac{3}{2}\) formalism in the SINGLE-ANISO routine [87,88]. Cholesky decomposition of the bielectronic integrals was employed to save disk space and speed-up the calculations [89]. In all calculations, the active space of the self-consistent-field (CASSCF) method consisted of the nine 4f electrons of the Dy\(^{III}\) ion spanning the seven 4f orbitals, i.e., \(\text{CAS}(9,7)\text{SCF}\). State-averaged (SA-)CASSCF calculations were performed for all of the sextets (21 roots), all of the quadruplets (224 roots), and 300 out of the 490 doublets (due to hardware limitations) of the Dy\(^{III}\) ion. Twenty-one sextets, 128 quadruplets, and 107 doublets were mixed through spin–orbit coupling in RASSI-SO.

All atoms were described using the Relativistic (semi-)Core Correlation - Atomic Natural Orbital (ANO-RCC) basis sets [90–92]. The following contractions were used: [3s2p1d] for the C and F atoms, [4s3p1d] for the S atoms, and [2s1p] for the H atoms.

4. Conclusions

To conclude, we have shown by spectro-electrochemistry that it is possible to form reversibly the radical cation and the dicationic species of a well-known redox-active SMM. The electronic spectra of the three compounds \(1\), \(1^{+}\), and \(1^{2+}\) were experimentally decomposed and rationalized by TD-DFT calculations. Upon oxidation, the donating character of the TTF core decreased, playing the double role of electron donor and acceptor in \(1^{+}\), until becoming an electron-acceptor in \(1^{2+}\). Consequently, the lowest-energy electronic transitions were identified as TTF → bzip ILCT for \(1\), TTF → TTF ID for \(1^{+}\), and tta → TTF ILCT for \(1^{2+}\). Finally, ab initio calculations demonstrated that the magnetic anisotropy of the Dy\(^{III}\) is kept under the oxidation process in at the molecular level paving the route to design conducting SMM with these systems. Efforts to obtain single crystals of the oxidized systems and measurements of electronic conductivity are still in progress in our laboratory.
**Supplementary Materials:** The following are available online at [http://www.mdpi.com/2312-7481/5/3/46/s1](http://www.mdpi.com/2312-7481/5/3/46/s1), Figure S1. X-ray structure of 1 and optimized structures for the different oxidation states (I_{opt}, 1^{+}_{opt}, and 1^{2+}_{opt}); Figure S2. Experimental X-ray structure of 1. n-hexane molecule of crystallization and hydrogen atoms are omitted for clarity; Figure S3. Optimized structure of 1; Figure S4. Optimized structure of 1^{+}. Figure S5. Optimized structure of 1^{2+}; Figure S6. Cyclic voltammograms of 1 (gray line) and 1 (black line) in CH_{2}Cl_{2} at a scan rate of 100 mV s^{-1}. The potentials were measured vs. a saturated calomel electrode (SCE) with Pt wires as working and counter electrodes; Figure S7. Cyclic voltammograms of 1 (black line, ten first cycles) in CH_{2}Cl_{2} at a scan rate of 100 mV s^{-1}. The potentials were measured vs. a saturated calomel electrode (SCE) with Pt wires as working and counter electrodes; Figure S8. Initial absorption spectra of 1 (red line), after oxidation under a 0.7 V applied current 1^{+} (blue line), and after switching off the applied current for 20 min (dashed red line). The full red line is recovered after 35 minutes; Figure S9. Initial absorption spectra of 1 (red line), after oxidation under a 1.1 V applied current 1^{2+} (green line) and after switching off the applied current for 20 min (dashed red line); Figure S10. Representation of the calculated local ground state magnetic anisotropy axes for the complexes 1, I_{opt}, 1^{+}_{opt}, and 1^{2+}_{opt}; Figure S11. Energy splitting of the 6H_{15/2} ground state for the complexes 1, I_{opt}, 1^{+}_{opt}, and 1^{2+}_{opt}; Table S1. Main components of the magnetic anisotropy g-tensor (g_{XX}, g_{YY}, and g_{ZZ}) and wavefunction composition of the ground state for the complexes 1, I_{opt}, 1^{+}_{opt}, and 1^{2+}_{opt}.

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**Abbreviations**

The following abbreviations are used in this manuscript:

- SMM: Single Molecule Magnet
- CH_2Cl_2: dichloromethane
- tta^-: 2-thenoyltrifluoroacetate
- TTF: Tetrathiafulvalene
- HOMO: Highest Occupied Molecular Orbital
- LUMO: Lowest Unoccupied Molecular Orbital
- ILCT: Intra-Ligand Charge Transfer
- DFT: Density Functional Theory
- TD-DFT: Time-Dependent Density Functional Theory
- CASSCF: Complete Active Space Self-Consistent Field
- RASSI-SO: Restricted Active Space State Interaction—Spin-Orbit

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