

Tetrathiafulvalene-Based Magnets of Lanthanides

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Introduction 29

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Magnets have been discovered about four millenniums ago in ancient Greece. Nowadays, in 2015, permanent magnets market is valued to 15 billion € and is expected to grow according to the demands for medical and industrial devices. In this market, the segment occupied by the badly named rare-earth¹-based magnets continues to expand owing to superior properties (such as saturation magnetization).

In the 1990s, a new class of magnets emerged in the scientific community with the discovery of the single-molecule magnets (SMMs) [1]. In these magnets, the magnetic memory is stored by the magnetic moment of a single molecule constituted of 12 manganese ions. This scientific finding reduced the size of a storage unit (byte) to nanometer. At the same time, the storage capacity of hard disks based on molecules would increase drastically. The drawback is the operating temperature range, below liquid helium (-269° C). In the last three decades, the quest for better SMMs never really stopped. In 2003, Ishikawa et al. [2] discovered that a single lanthanide ion ($Ln = Tb^{III}$) embedded in a double-decker complex behaved as a SMM. To date, the lanthanide series is the most productive SMMs line in Mendeleev's periodic table with a recent tremendous record of closure of the magnetic hysteresis loop at 60 K [3, 4], close to liquid nitrogen. Tetrathiafulvalene (TTF) and its analogues are well known in the field of molec-

ular materials to produce organic metals, semiconductors, and superconductors [5, 6]. The functionalization of the electron donor TTF core by an acceptor moiety contributed to the development of functional materials such as switches, sensors, photovoltaic cells, and nonlinear optical systems [7–9]. It was then logical to adapt the acceptor moiety to coordinate transition metals for (1) elaboration of multifunctional materials with both paramagnetism and electrical conductivity [10, 11] and (2) the synthesis of polynuclear transition metal complexes exhibiting SMM properties embedded in a conducting material [12-16]. One must admit that all tentative proposals were not very successful except Oshio's work [17] which shows SMM behavior but without conductivity. The first TTF-Ln system was reported by Faulkner et al. [18] with the assembly in

solution of tetrathiafulvalene carboxylic acid and ytterbium. In 2003 [19], the first structurally characterized TTF-Ln system in the solid state was reported. There was no chemical bond between the lanthanide ion and the TTF moiety, so the chemical approach is the so-called through space. We have published almost 10 years ago the first isolated TTF-Ln coordination complex [20] based on the spin-only Gd(III) ion, which, of course, does not behave as a SMM. Since 2009, several groups [21-24], including us [25], have paid much attention to TTF-Ln systems with the objectives to combine conductivity (electrical transport), magnetism (magnetic memory), and luminescence (light emitter) in a single chemical object. In this frame, the TTF moiety turns out to be a powerful sensitizer of the NIR luminescence of

¹Most of the rare earths are not rare: cerium is more abundant than copper on earth, and thulium (the most rare) is more abundant than silver.

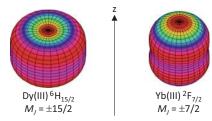
lanthanides [26]. Nevertheless and despite colossal efforts of the scientific commu- 69 nity, electrical conductivity has never been observed in TTF-Ln systems. One could 70 say that the necessary oxidation (partial) of the TTF moiety to promote electronic 71 transport does not preserve the chemical integrity of the complex (dissociation) 72 except when both TTF core and coordinating moiety are not fused [27]. Despite 73 several attempts, there is no SMM based on TTF and strongly anisotropic 3-D 74 transition metals such as Co(II) and Ni(II) [12, 13, 28, 29]. But TTF performed in 75 the field of SMMs with plethora of mononuclear and polynuclear complexes which 76 possess a magnetic memory in the absence and, in less extent, in the presence of an 77 external constant magnetic field. Since highly anisotropic magnetic moments are 78 necessary, Dy(III) and Tb(III) are the ideal candidates [30-37]. These two ions 79 represent almost 99% of reported Ln-based SMMs [38] with a preponderant role 80 played by Dy(III)-based systems.

Preamble 2 82

Naturally, researchers have at first focused their attention on the synthesis of mononuclear TTF-Dy complexes. Like most of the time, the first attempt was not very 84 successful [39]. The reaction of two equivalents of tetrathiafulvalene-amido-2-pyri- 85 dine-N-oxide with [Dy(hfac)₃]·2H₂O (hfac⁻: 1,1,1,5,5,5-hexafluoroacetylacetonate 86 anion) precursor produced mononuclear species in which Dy(III) is surrounded by 87 eight oxygen atoms: two from pyridine-N-oxide moieties and six from three 88 bidentates hfac⁻. This fully oxygenated environment adopts a coordination polyhe- 89 dron close to square antiprism (SAP) D_{4d} with CShM = 0.528 [40]. The material does not show any out-of-phase component of the AC susceptibility in zero external dc 91 field down to 2 K below oscillating field frequencies of 1 kHz. In other words, it is not 92 a magnet. Probably, the charge distribution around the Dy(III) center does not match 93 the axiality required by the simple but chemically implementable precepts exposed by 94 Rinehart et al. [41]. Performant magnets are obtained when the largest M_J states are 95 stabilized for a given multiplet ground state ($M_J = \pm 15/2$ for Dy(III)). The analysis of the electron density distribution provides a simple tool to anticipate what might be the 97 ground state in a given environment [42]. The charge density distribution of the Ising 98 component (the largest M_I values) of the multiplet ground state of the oblate Dy(III) is 99 represented on Fig. 1. The electrons are principally located in a plane (xy), so the 100 disposition of negatively charged ligands in the z direction will stabilize this Kramers 101 state. It is the opposite for the prolate Yb(III) for which the negative charges must lie in 102 the xy plane. This textbook analysis guided the synthesis of the most efficient SMM 103 reported so far [3, 4]. Additionally, advanced quantum calculations have demon- 104 strated that such approach might provide high-temperature SMMs (if chemically 105 accessible) [43, 44].

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Fig. 1 Angular dependences of the total 4f charge density for largest M_J states of the multiplet ground state for Dy(III) and Yb(III)



of 3 Mononuclear TTF-Dy(III) SMMs: The N₂O₆ Saga

We have sought, at first, for neutral complexes to crystallize with bidentate TTF-based ligands in order to (1) stabilize the complexes and (2) minimize the 109 degrees of freedom of the coordination sphere. The use of three negatively charged 110 acetylacetonate (-1) ancillary ligands counterbalances the charge +3 of the lantha-111 nide ions and insures complex neutrality. To complete the coordination sphere, TTF-based ligands with nitrogen-coordinating sites have been designed to 113 desymmetrize the ligand field (preamble). The first two complexes synthesized with 114 this approach are $[Dy(hfac)_3L^1]$ and $[Dy(hfac)_3L^2]$ (with $L^1 = 2-\{4,5-[4,5-bis]\}$ 115 (propylthio)-tetrathiafulvalenyl]-1H-benzimidazol-2-yl}pyridine and $L^2 = 2-\{1-1\}$ 116 methylpyridyl-4,5-[4,5-bis(propylthio)tetra-thiafulvalenyl]-1H-benzimidazol-2-yl} 117 pyridine) [45] (Fig. 2). Except the obvious chemical difference at N3 (alkylation), the 118 analysis of the crystallographic structure revealed the presence of intermolecular 119 hydrogen bond at N3 site in [Dy(hfac)₃L¹]. Such intermolecular connection does 120 not exist in [Dy(hfac)₃L²], and it has a dramatic consequence on the coordination 121 polyhedra. Indeed, the polyhedron is strongly distorted in $[Dy(hfac)_3L^1]$ with respect 122 to the one in [Dy(hfac)₃L²] (Fig. 2). As a consequence, the magnetic properties 123 124 of these two, apparently similar, complexes differ. In zero external dc field, [Dy (hfac)₃L²] behaves as a SMM with the apparition of a frequency-dependent out-of-125 phase component of the ac susceptibility. Such signal is absent for [Dy(hfac)₃L¹] 126 which is not a SMM, as far as the crystalline condensed phase is concerned. The hydrogen-bonding network plays a crucial role in the modification of this behavior. Magnetic measurements in solution reveal the true nature of the complexes. Of 129 course, one must be sure that the complexes are stable in solution. They both behave 130 the same, as molecular magnets. This proves two important characteristics: (1) the behavior of SMM can be preserved in solution. This is an important issue since one 132 may say that the observed behavior is truly of molecular origin and then the molecular 133 magnet can be manipulated. (2) The destruction of the intermolecular network by 134 dissolution restores the molecular property. The absence of SMM behavior in crys-135 talline condensed phase must be taken carefully, and the impact of the crystal packing 136 must be analyzed prior any hasty conclusion. 137 138

The ground multiplet ground state ${}^{6}H_{15/2}$ of Dy(III) splits under the effect of crystal field in several sublevel characterized by pure M_{J} levels or a mixture of M_{J} levels depending on the symmetry of the ligand field. In this frame, the effective 1/2

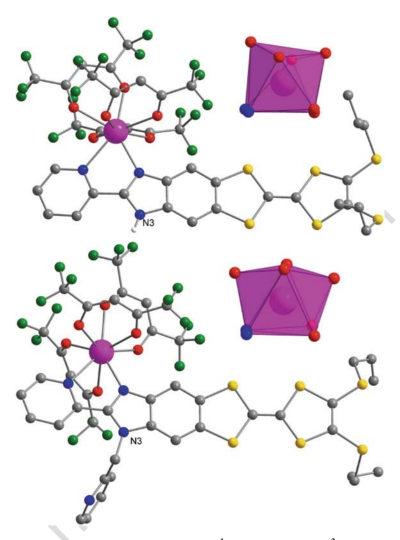


Fig. 2 Representation of the complexes $[Dy(hfac)_3L^1]$ (top) and $[Dy(hfac)_3L^2]$ (bottom) with the alkylated nitrogen atom N3. Coordination polyhedra are also represented

is often used to describe these Kramers sublevels. Then, the Kramers ground state 141 possesses an effective spin 1/2 with an effective g-tensor in the reference frame of 142 the complex. It is not difficult to show that for the two Ising components $M_J = \pm 15/2$ 143 of the $^6H_{15/2}$ multiplet the g-tensor is characterized by $g_x = g_y = 0$ and $g_z = 20$ with 144 z the axis of projection. Experimentally, the orientation and the amplitude of the g- 145 tensor are available under certain conditions: the complex must crystallize in the 146 triclinic system with only one Dy(III) crystallographic site. Measurements of the 147 magnetization on an oriented single crystal in three perpendicular planes as a 148 function of the angle (θ) between the magnetic field (H) with the axes of the single 149

150 crystal allow the determination without ambiguity of the g-tensor. The magnetic susceptibility in a plane can be fitted with the following equation:

$$\chi_{\rm M}T = \frac{MT}{H} = \chi_{\alpha\alpha}T\cos^2\theta + \chi_{\beta\beta}T\sin^2\theta + 2\chi_{\alpha\beta}T\sin\theta\cos\theta$$

where α and β are the directions X, Y, and Z of the crystal reference frame in a 152 circular permutation (Fig. 3) and $\chi_{\rm M}$ and T are, respectively, the molar magnetic 153 susceptibility and the temperature expressed in Kelvin. 154

In the effective spin-1/2 frame, the principal values of g-tensor are $g_z = 14.22$,

 $g_y = 3.96$, and $g_x = 9.43$. These values are far away from those expected for a purely 156 axial system and explain why [Dy(hfac)₃L¹] is not a SMM in the condensed crystalline phase. The orientation of the g_z is represented on Fig. 4 with a black arrow. In the present case, in order to simulate physical properties such as magnetism, the neighboring molecules need to be explicitly integrated. The calculated orientation of the most magnetic axis is also represented on Fig. 4. Clearly, calculations with N-H bond fail to reproduce the orientation of the magnetic poles as well as the standards $\chi_M T$ vs. T and M vs. H plots (Fig. 4 top right, M the magnetization expressed in Bohr magneton per mole) [46]. One can notice however that in this chemical configuration, the magnetic moment of a Dy(III) ion resides in a direction passing through the most negatively charged direction (two hfac anions) and perpendicular to the plane defined with the less electronegative nitrogen atoms from imidazole and pyridine moieties, in agreement with basic electrostatic considerations. The hydrogen atom must be positioned between two heteroatoms: the nitrogen N3 of imidazole of one complex and one oxygen atom of one hfac moiety of a neighboring complex. In other words, in the crystal, the N-H bond disappeared with a hydrogen atom, in average, localized at an intermediate position between the two atoms. Nevertheless, the calculated orientation is still at 30° of the experimental one and that dynamical effects should also probably be included to properly account for the experience.

Fig. 3 Angular dependence of $\chi_M T$ measured at 2 K for $[Dy(hfac)_3L^1]$ with a 1 kOe magnetic field in three perpendicular planes (XY, ZY, and XZ). In inset a schematic representation of a single crystal of [Dy $(hfac)_3L^1$] with crystallographic axes in the frame of the single crystal

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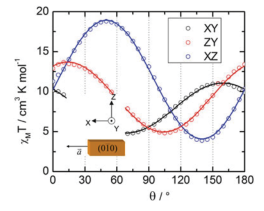
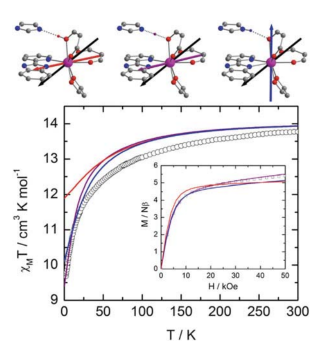


Fig. 4 (Top) orientations of the experimental z magnetic axis represented in black for [Dy(hfac)₃L¹] with the calculated orientation for different positions of the hydrogen atom (O-H in red, N-H in blue, and middle in purple). (Bottom) thermal variations of the experimental (white symbols) and calculated (full-colored lines) $\chi_{\rm M}T$ within the inset of the experimental and calculated field variation of the magnetization at 2 K



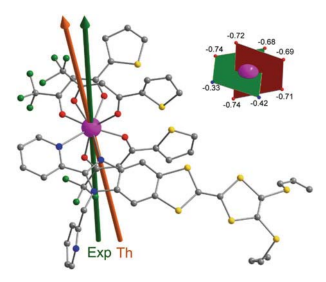
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At this stage L^2 seems to be a good starting point, but the use of hfac produces a 176 SMM with relatively low efficiency. Indeed, the energy necessary to reverse the 177 magnetic moment is less than 20 K (the barrier) and a temperature-independent 178 relaxation process (quantum tunneling of the magnetization) of the order of 100 µs. 179 In order to improve the SMM in keeping this topology, one possibility is to play with 180 the ancillary ligand. Thiophene groups are less electroattractive than CF₃, and then 181 the substitution of one CF₃ by one thiophene should increase the negative charges on 182 coordinated oxygen atoms.

The magnetic properties of $[Dy(tta)_3L^2]$ (tta⁻: 2-thenoyltrifluoroacetonate) have 184 been studied in the crystalline condensed phase and in frozen solution [47]. Qualita- 185 tively, the magnetism corresponds to $[Dy(hfac)_3L^2]$: it is a SMM in solid state and in 186 solution. Quantitatively, the energy barrier has been multiplied by a factor two and validates our approach. Ab initio calculations showed that the negative charges carried by oxygen atoms are larger, in amplitude, than for $[Dy(hfac)_3L^2]$ (-0.71 vs. -0.68 in average) according to electrostatic considerations [46, 47]. Interestingly, 190 SHAPE analyses [40] on $[Dy(tta)_3L^2]$ and $[Dy(hfac)_3L^2]$ reveal nearly the same 191 distortions. In both structures, Dy(III) resides in SAP environment with 192 CShM = 0.538 and 0.597, respectively. Angular-resolved magnetometry measurements show that the anisotropy axis (the easy magnetization axis) is parallel to the 194 most negatively charged direction (Fig. 5). This experimental finding is supported by 195 ab initio calculations with a gap between the calculated and the experimental easy 196 axis of only 7.6°. Furthermore, the calculated g_z (19.5) is very close to the Ising limit. 197 The examination of the temperature dependence of the relaxation time of the 198

Fig. 5 Representation of the molecule $[Dy(tta)_3L^2]$ avec and the calculated (orange) and experimental (green) magnetic axes. Insert: scheme of the coordination sphere of Dy (III) ion with the calculated charges of the coordinated atoms

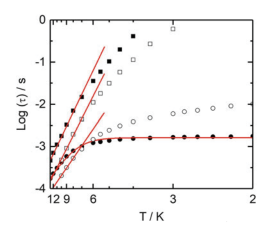
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magnetic moment reveals the presence of a thermally activated regime at a high temperature, while the system enters a thermally independent regime below 6 K (Fig. 6). It can be reproduced with a combination of the Orbach [48] (over the barrier) and tunneling (through the barrier) processes: $\tau^{-1} = \tau_0^{-1} \exp(\Delta/T) + \tau_{\rm TI}^{-1}$ with Δ =42 K, $\tau_0 = 8 \times 10^{-6}$ s, and $\tau_{\rm TI} = 1.62 \times 10^{-3}$ s. Dy(III) is a Kramers ion, 201 202 203 and the magnetic moment should not be able to tunnel through the barrier: the two 204 Ising components cannot be mixed by modulation of the crystal field. The applica-205 tion of a moderate external dc field (1 kOe) destroys this relaxation path, and the 206 system falls in a pure thermally activated regime with nearly the same activation 207 energy. Additional perturbations must affect the Kramers ground state in zero 208 external dc field to allow the system to oscillate between the "up" and "down" 209 states. In the condensed crystalline state, the molecules are closely packed, and 210 interactions of dipolar origin may propagate through space. This is especially true when someone deals with heavy lanthanides which possess the largest magnetic 212 moments of the periodic table. Transverse component of this internal field can mix 213 the Kramers doublets and facilitate the tunneling. The dilution of the complex in a 214 diamagnetic medium (at low concentration) minimizes this internal field and is supposed to suppress the tunneling. However, one can see on Fig. 6 that dissolution 216 in dichloromethane is not enough: leveling of the relaxation time still persists at low 217 temperature. Compared to the application of an external dc field of 1 kOe, which 218 completely lifts the degeneracy of the ground state and destroys the tunneling effect (Fig. 6), in solution, the relaxation time remains rapid (about 100 times faster). The 220 consequence is that, even in solution, the hysteresis loop remains closed in zero field 221 while it is opened up in field (butterfly-shaped hysteresis) at any temperatures above 222 500 mK [47]. 223

Then, if the closure of the hysteresis loop at the origin does not arise from 225 intermolecular considerations, it might come from inside the complex. Dysprosium

Fig. 6 Log-scale representation of the thermal variation of the relaxation time of [Dy(tta)₃L²] in solid state (full symbols) and in solution (empty symbols), in zero external dc field (circles) and under 1 kOe external dc field (squares). Solid red lines correspond to the best fitted curves with a modified Arrhenius law at zero field and a Arrhenius law in field



is one of the elements in the periodic which consists of different and stable isotopes 226 (161 Dy, 162 Dy, 163 Dy, and 164 Dy) in quasi-equivalent natural abundance. Two of 227 them, with an even mass number, possess a nuclear spin (I=0), and the two others 228 have a nuclear spin I=5/2. We have then decided to study the influence of this 229 nuclear spin on the relaxation of the electronic magnetic moment [49] coupled with 230 magnetic dilution. Hyperfine interactions and dilution are known to affect the 231 relaxation of the magnetic moments [50–52]. Clearly, metal-centered isotopic 232 enrichment modifies the relaxation rate in the quantum regime (Fig. 7).

Below 6 K when the system enters in the quantum regime, the relaxation time of the magnetic moment is ten times slower for the isotopically enriched complex [164 235 Dy(tta) $_3$ L 2] (I=0) than for the isotopically enriched complex [161 Dy(tta) $_3$ L 2] (I=5/236 2). This is true in zero external dc field in the condensed crystalline phase but also 237 when the enriched complex is diluted in a diamagnetic isomorphous crystalline 238 matrix ([164 Dy $_{0.04}$ Y $_{0.96}$ (tta) $_3$ L 2] $_6$ C $_6$ H $_{14}$ vs. [161 Dy $_{0.03}$ Y $_{0.97}$ (tta) $_3$ L 2] $_6$ C $_6$ H $_{14}$). The dramatic difference is that the dilution in a diamagnetic medium of these isotopically 240 enriched complexes slows the relaxation enough to observe the opening of the 241 hysteresis loop at the origin for the 164 Dy derivative and not for the 161 Dy derivative. 242 At this stage we proved that isotopes chemistry drives some electronic properties. 243 Nevertheless, one must keep in mind that the hysteresis (the memory) in the absence 244 of an external stimulus shows up only when the molecules are far away from each 245 other. Thus, the deposition of juxtaposed SMMs on surfaces can lead to unexpected 246 results because they will behave, in a certain manner, collectively and not individually (Fig. 8).

In addition, we recently investigated the magnetism of the last two stable enriched 249 complexes [162 Dy(tta) $_3$ L 2] and [163 Dy(tta) $_3$ L 2] [53]. [162 Dy(tta) $_3$ L 2] is the copy 250 paste of [164 Dy(tta) $_3$ L 2] because the nuclear spin of Dy(III) is zero in both cases. 251 The nuclear spins of 163 Dy and 161 Dy are indeed equal (I=5/2), but the 252 hyperfine coupling constant A_{HF} differs [54, 55] and then the relaxation rate 253 affected. This is perceptible in condensed crystalline phase with [163 Dy(tta) $_3$ L 2] 254

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Fig. 7 Frequency dependences of $\chi_{M}^{\ \prime\prime}$ of [164 Dy(tta) $_{3}$ L 2] and [161 Dy (tta) $_{3}$ L 2] in zero field in the temperature range 2–14 K

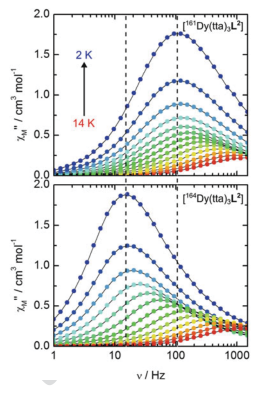
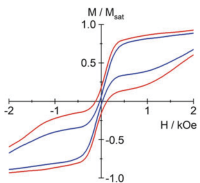


Fig. 8 Normalized magnetic hysteresis loops measured at 460 mK for [164 Dy_{0.04}Y_{0.96}(tta)₃L 2]•C₆H₁₄ (red line) and [161 Dy_{0.03}Y_{0.97}(tta)₃L 2]•C₆H₁₄ (blue line)



slightly faster than [161 Dy(tta) $_3$ L 2], but at this stage, the relaxation is essentially driven by intermolecular interactions. Once diluted, [161 Dy $_{0.05}$ Y $_{0.95}$ (tta) $_3$ L 2] $_6$ C $_6$ H $_{14}$ is about ten times slower than [163 Dy $_{0.05}$ Y $_{0.95}$ (tta) $_3$ L 2] $_6$ C $_6$ H $_{14}$. Sign and/or amplitude of the hyperfine coupling seem to play also a fundamental role on the relaxation in quantum regime.

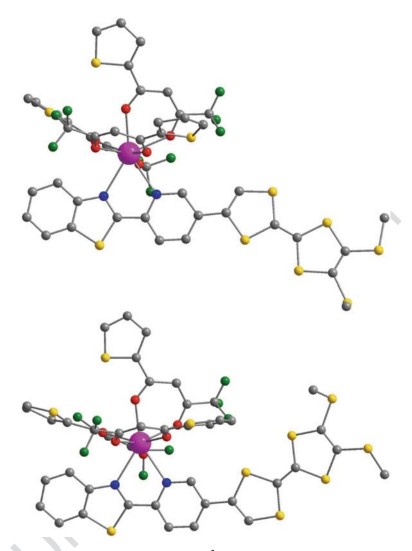
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4 A Journey in TTF-Ln SMMs

There are additional TTF-Dy(III) $[N_2O_6]$ mononuclear complexes which behave as 261 SMMs in the literature. Also, higher coordination number, typically N_3O_6 , has been 262 envisaged with less success. Environment such as O_8 for which the charge distribution is more symmetric has also been investigated. Some of these complexes are 264 stricto sensu (chemically) mononuclear complexes, but some of them are polynuclear complexes but in which the distances between metallic centers are so large that 266 we can consider they are mononuclear from a magnetic point of view. In other 267 words, there are no interactions between those centers. Such complexes might be 268 described as an assembly of mononuclear SMMs.

4.1 Nitrogen-Based Donor-Acceptor Type Dyads

The first example we want to introduce is based on a similar TTF ligand with a 271 benzothiazole group. The ligand L^3 ($L^3 = 4-[6-(1.3-benzothiazol-2-y])$ pyridin-3-272 yl]-4',5'-bis(methyl-thio)tetrathiafulvene) is similar to L^1 with however no possible 273 intermolecular hydrogen bond [56]. Such benzothiazole group might also be of 274 interest to realize photoswitchable conductors and photoelectric conversion mate- 275 rials [57]. Reaction of L³ with [Ln(tta)₃]·2H₂O gives [Ln(tta)₃L³] complex which 276 crystallize in the P-1 triclinic space group for light elements and in the P2₁/a 277 monoclinic space group for heavy elements. For intermediate Dy(III), the two 278 polymorphs can be obtained. Like in the previous section, Dy(III) ion is in a N₂O₆ 279 environment (Fig. 9) with a square antiprismatic idealized coordination polyhedron. 280 The complexes behave as SMMs with slightly different energy barriers (57 K 281 vs. 42 K). The coordination polyhedron is slightly more distorted in the triclinic 282 phase than in the monoclinic one with however the highest energy barrier. One 283 should then conclude that there is no direct correlation between symmetry and 284 activation energy. Additional factors, which are not that clear and yet to be identi- 285 fied, influence the energy splitting diagram. Quantum chemistry calculations qual- 286 itatively reproduce this experimental fact: the first excited state is located higher in 287 energy in the monoclinic phase than in the triclinic. These barriers are also similar to 288 the one found for $[Dy(tta)_3L^2]$ (due the similarities between the two environments). Here again, the calculated and the experimental orientation of the easy magnetic axis 290 are in very good agreement with less than 10° of mismatch and an orientation in the 291 most negatively charged direction. The dilution+enrichment protocol has been 292 applied to both polymorphs, but only nuclear spin-free isotopes were employed to 293 slow down the relaxation as much as we could. The same receipts give the same 294 results: the hysteresis loops open in zero field for the enriched and diluted 295 complexes.



 $\textbf{Fig. 9} \ \ \text{Representation of the complex } [\text{Dy}(\text{tta})_3L^3] \ \text{in the triclinic (top) and monoclinic (bottom)} \\ \text{forms}$

All the TTF-based ligand envisaged so far could be coordinated only by one metal center, so we imagined and designed new TTF-based ligand to incorporate, in a bridging ligand, different coordination sites. The ideas behind were to (1) select different metals in incorporating different chelating sites (e.g., tris-) and (2) to accommodate the same metal in different environments to tune its magnetic properties (Fig. 10) [58].

When reacted with $[Dy(hfac)_3] \cdot 2H_2O$, L^4 coordinates from both bischelating sites to form $[Dy_2(hfac)_6L^4]$ neutral complex. Each Dy(III) ion is surrounded by six

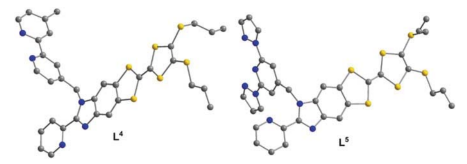


Fig. 10 Representations of L⁴ and L⁵ ligands

oxygen and two nitrogen atoms and lies in almost the same N_2O_6 D_{4d} environment 305 and is separated by more than 10 Å [58]. From a magnetic point of view, these two 306 sites should behave the same, and this is what is observed. Both behave as SMMs in 307 zero external dc field, and the extended Debye analysis [59] reveals only one 308 relaxation process. Seventy percent of the magnetic moments relax at the same 309 frequency. The relaxation is however too fast in zero field to be quantitatively 310 analyzed. In applying an external field to suppress the thermally independent regime, 311 only one relaxation time that includes both sites is identified. At the optimum field, 312 i.e., the external field for which the relaxation is the slowest (800 Oe), the 313 thermal variation of the relaxation time does not follow the Arrhenius law expected 314 for the Orbach process [60] but can be easily reproduced with a Raman process 315 ($\tau = CT_{-}^n$, $C = 4.8 \times 10^{-3}$, and n = 6.26). This tends to prove that the relaxation 316 does not occur through the first excited state, at least, as long as the in-field relaxation 317 is concerned.

The reaction of $[Dy(hfac)_3]\cdot 2H_2O$ with L^5 leads to dinuclear species 319 $([Dy_2(hfac)_6L^5])$ with two different coordination polyhedra [61], N_2O_6 320 $(CShM_{SAPR-8}\ (D_{4d})=0.435)$ and $N_3O_6\ (CShM_{TCTPR-9}\ (D_{3h})=0.586)$. The N_2O_6 321 sites behave in a standard way (SMM in zero external field), while the 322 nonacoordinated site does not show any out-of-phase signal in zero external field. 323 The application of a moderate external dc field slows down the relaxation with the 324 emergence of two identifiable processes that can be safely attributed to the two 325 different sites. An extended Debye model featuring two relaxation times has been 326 employed to treat the ac data. Interestingly, the analyses reveal the ratio of the 327 magnetic susceptibility which relaxes at the two relaxation time to be close to 50:50 328 in agreement with the chemical structure. Furthermore, the energy barrier for the 329 octacoordinated site is in good agreement with its mononuclear equivalent 330 [45]. From a chemical point of view, it was interesting to look for site selectivity 331 with regard to different $Dy(\beta-diketonate)_3$ precursors. A 1:1 ratio of $[Dy(hfac)_3]\cdot 332$ $2H_2O$ and $[Dy(tta)_3]\cdot 2H_2O$ was reacted with L^5 , and it forms dinuclear species 333

[Dy₂(hfac)₃(tta)₃L⁵]. The crystal structure of the complex reveals ligand exchange, with the N₂O₆ site made of two tta⁻ and one hfac⁻ ligands and the N₃O₆ site made of one tta and two hfac ligands. We think this ligand exchange occurs as a consequence of a subtle balance between the size of the metallic precursors and the 337 coordination which leads to a minimum steric hindrance. The magnetic behaviors 338 of these two metallic sites are qualitatively identical to [Dy₂(hfac)₆L⁵] with the N₂O₆ 339 site being a SMM in zero field while the N₃O₆ site being a SMM only in field [61]. 340 At this stage, it appears that N₂O₆ coordination polyhedron around Dy(III) 341 systematically produces SMMs in zero field. Liu et al. [62] developed a TTF-fused 342 donor-acceptor system based on dipyrido[3,2-a:2',3'-c]phenazine (dppz) which can be reacted with Dy(β-diketonate)₃ precursors ([Dy(hfac)₃]·2H₂O and [Dy(tta)₃]· 2H₂O) [63]. Two mononuclear units are obtained with Dy(III) in the standard 345 N₂O₆ SAP coordination polyhedron made of three β-diketonate ligands and one 346 dipyridyl moiety (CShM_{SAPR-8} (D_{4d}) = 0.724 for the hfac⁻ derivative and 347 $CShM_{SAPR-8}$ (D_{4d}) = 0.507 for the tta⁻ derivative) from the ligand L⁶ (L⁶ = TTFfused dipyrido[3,2-a:2',3'-c]phenazine). Both compounds behave as SMM in zero 349 field, but they are "faster," and they relax at higher frequencies, than previous N_2O_6 350 systems with however the tta derivative slower than the hfac. Typically, at 2 K, the maximum on the $\chi_{\rm M}^{\prime\prime}$ vs. ν curves shows up at 700 Hz for [Dy(tta)₃L⁶], while it is located above 1,500 Hz for $[Dy(hfac)_3L^6]$. To compare, for $[Dy(hfac)_3L^2]$, the 353 maximum was at 970 Hz and at 56 Hz for [Dy(tta)₃L²] in the same sample environment. Such comparison, if natural, is however dangerous since in the low-temperature regime, where thermally independent processes take over all the others, the relaxation in the condensed crystalline phase is governed by the combi-357 nation of magnetic intermolecular interactions (of dipolar origin) and hyperfine 358 coupling. One can say that this N₂O₆ topology (three bischelating oxygenate ligand 360 and one bischelating nitrogenated ligand) provides efficient magnets that qualitatively behave the same and quantitatively almost the same. The difference resides in 361 the electron withdrawing or donating ability of the chemical groups on the 362 β-diketonate ligands.

364 5 Oxygen-Based TTF-Based Ligands

The strong oxophilic characters of lanthanide authorize the synthesis of fully oxygenated coordination polyhedron around metal centers. However, such environment does not, a priori, create the expected dissymmetry of charges to produce SMMs in the specific case of Dy(III) at least as far as O_8 environments are concerned. The reaction of $[Dy(hfac)_3]\cdot 2H_2O$ with 4,4',7,7'-tetra-*tert*-butyl-2,2'-bi-1,3-benzo-dithiole-5,5',6,6'-tetrone ligand [64] (\mathbf{L}^7) forms a dinuclear complex $[Dy_2(hfac)_6(H_2O)_2\mathbf{L}^7]$ [65]. In this complex, two Dy(III) ions, related by an inversion center, in O_9 coordination polyhedron, are linked by an acceptor-donor-acceptor triad (Fig. 11). Eight of the nine are coming from bischelating ligands (three hfac and one quinone), and the last one is coming from a water molecule. The intramolecular

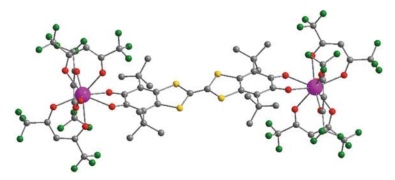


Fig. 11 View of the dinuclear compound [Dy₂(hfac)₆(H₂O)₂L⁷]

Dy-Dy distance is close to 17.5 Å, so there is no superexchange interaction between 375 the two metals.

In this environment, Dy behaves as a SMM in zero external field. Interestingly, 377 the equivalent complex obtained from the reaction of L^7 with $[Dy(tta)_3] \cdot 2H_2O$ does 378 not produce SMM. The steric hindrance of tta avoids water molecule to coordinate 379 Dy(III), and therefore in this dinuclear complex, the coordination polyhedron is only 380 made of eight oxygen atoms which is probably less suitable to promote axial 381 anisotropy. This simple analysis is counterbalanced by other investigations [66] 382 which clearly demonstrate that O₈ environment could perfectly produce SMMs 383 with Dy(III) ions. Probably, the primary approach which consists of considering 384 only the point charge model might be oversimplified. Some authors pointed out with 385 a deeper analysis that dipole and quadrupole moments in the electrostatic potential 386 expansion play a significant role on the magnetic anisotropy [67, 68].

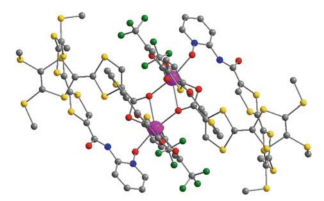
To conclude this section, we would like to briefly discuss one peculiar 388 system. Two different TTF-based ligands are used to produce a dinuclear Yb 389 (III)-based complex $[Yb(tta)_2L^8L^9]_2$ [69]. The redox active ligand 4,5-bis 390 (thiomethyl)-4'-carboxylictetrathiafulvalene (L⁸) is bridging two Yb(III) ions 391 through $\mu_2(\eta_1,\eta_2)$ oxygen atoms, and 4,5-bis(thiomethyl)-4'-ortho-pyridyl-N- 392 oxide-carbamoyl-tetrathiafulvalene (L⁹) is terminal (Fig. 12). The coordination 393 sphere around each Yb(III) is made of eight oxygen atoms, and the two ions are 394 separated by only 3.89 Å.

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The static magnetic properties reveal what could be analyzed as a sign of 396 ferromagnetic interactions between the two ${}^{2}F_{7/2}$ multiplet ground states. Indeed, 397 on cooling from room temperature, $\chi_{\rm M}T$ decreases continuously in agreement with 398 the thermal depopulation of M_J states, passes through a broad minimum at 9 K, and 399 then slightly increases on cooling further down to the lowest temperature. We have 400 attempted to fit the $\chi_M T$ vs. T plot taking into account the crystal field effects by the 401 extended Stevens operators technique [48] and the interaction between magnetic 402 moments. The Hamiltonian to consider is the following:

Fig. 12 View of the dinuclear compound [Yb $(tta)_2L^8L^9$]₂



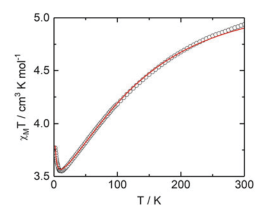
$$\widehat{H} = \sum_{i=1}^{2} \left(B_{2}^{0} \widehat{\boldsymbol{O}_{2}^{\theta} i} + B_{2}^{2} \widehat{\boldsymbol{O}_{2}^{2} i} + B_{4}^{0} \widehat{\boldsymbol{O}_{4}^{\theta} i} + B_{4}^{2} \widehat{\boldsymbol{O}_{4}^{2} i} + B_{4}^{4} \widehat{\boldsymbol{O}_{4}^{\theta} i} + B_{6}^{0} \widehat{\boldsymbol{O}_{6}^{\theta} i} + B_{6}^{2} \widehat{\boldsymbol{O}_{6}^{2} i} + B_{6}^{4} \widehat{\boldsymbol{O}_{6}^{\theta} i} + B_{6}^{6} \widehat{\boldsymbol{O}_{6}^{2} i} \right) + \beta \left(g_{J} \widehat{\boldsymbol{J}}_{1} + g_{J} \widehat{\boldsymbol{J}}_{2} \right) \cdot \widehat{\boldsymbol{H}} - J \widehat{\boldsymbol{J}}_{1} \cdot \widehat{\boldsymbol{J}}_{2}$$

$$(1)$$

The first line corresponds to the crystal field effect at the two lanthanide sites with $\widehat{O_k^q}$ the operator equivalents which can be expressed as polynomials of the total angular momentum matrices $(J^2, J_z, J_+, \text{ and } J_-)$ associated with the ${}^2F_{7/2}$ multiplet ground state. The second line corresponds to the Zeeman effects on J_i which are coupled through J. A homemade program has been developed to fit the magnetic data. Surprisingly a fairly good agreement (Fig. 13) is obtained without any interaction $(J=0 \text{ cm}^{-1})$, so one can consider this dimer as two isolated Yb(III) centers. The wave function analysis reveals that the Kramers ground state is the pure $M_J=\pm 7/2$ component, separated only by 2.6 cm $^{-1}$ from the first excited state $M_J=\pm 1/2$. The stabilization of the largest M_J component can be viewed in the frame of point charge model. The carboxylate group acts as a tweezer which projects the -1 charge in a plane containing two hfac $^-$ ligands (Fig. 12). Then, ligand charges are condensed in a plane around this prolate ion (Fig. 1) and stabilize the largest M_J doublet state.

As a consequence of this Ising-type anisotropy, the complex behaves as a SMM with however a small energy barrier (\sim 21 K) determined from the temperature dependence of the relaxation time. This value can be compared with estimated gap between the ground and the first excited states (see above, \sim 4 K) from DC magnetic measurements. To support this interpretation, luminescence provides a unique tool to probe energy levels. The low-temperature (77 K) excitation of the sample at 20,000 cm⁻¹ sensitizes the $^2F_{5/2} \rightarrow ^2F_{7/2}$ transitions in the range of 10,400–9,400 cm⁻¹. The excitation corresponds to LLCT (ligand-to-ligand charge transfer) and highlights the role of antenna played by the redox-active TTF ligands. We must mention that Dy(III) luminescence cannot be probed with TTF-based ligand since the emission lines fall in the absorption bands of the ligands. The

Fig. 13 Temperature dependences of $\chi_{\rm M}T$ (open circles) with the best fitted curve (red line)



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emission profile can be deconvoluted in four transitions (9,703, 9,936, 10,197, and 429 10,213 cm⁻¹). The gap between the two most energetic transitions (23 K) gives the 430 exact gap between the Kramers doublet ground state and the first excited state. It is 431 almost in perfect agreement with magnetism.

6 Polynuclear TTF-Dy(III) SMMs

One question emerges from the previous paragraph. Does the nonexistence of 434 interactions between Yb(III) in Yb derivative [Yb(tta)₂L⁸L⁹]₂ can be generalized 435 to other lanthanide? The synthesis of $[\mathrm{Dy}(\mathrm{tta})_2\mathbf{L^8L^9}]_2$, isostructural of Yb derivative, 436 provides an answer [70]. On cooling, $\chi_{\mathrm{M}}T$ vs. T plot passes through a broad 437 minimum at 16 K that cannot be reproduced without taking into account interactions. 438 The best fits are obtained with a ferromagnetic interaction $J = 2.98 \times 10^{-3} \text{ cm}^{-1}$. 439 This interaction is very weak with respect to the energy engaged in crystal field 440 splitting (hundreds of wave numbers). In this frame the Kramers ground state for 441 each Dy(III) ion corresponds to more than 99% of the $M_J=\pm 15/2$. In this effective 442 spin-1/2 model, one expects g=20 in one direction and 0 in a perpendicular plane. 443 Then, the coupling of dipolar origin between the two magnetic moment can be easily 444 calculated from the Hamiltonian $\hat{H} = -J_{dd}\hat{\sigma_l} \cdot \hat{\sigma_2}$ where $\hat{\sigma_i}$ are the operators 445 associated with the effective 1/2 and $J_{dd} = -(\mu_0 g^2 \beta^2 / 4\pi h c r^3)(1 - 3\cos^2\theta)$ cm⁻¹ 446 (r is the distance between the metal centers, and θ is the angle between the anisotropy 447 axis and the r vector). Such interaction can be ferro- or antiferromagnetic depending 448 on θ . The amplitude can also be rather large at short distances, with $r=4~\text{Å}~J_{dd}$ 449 varies from -2 to +5 cm⁻¹, to compare with J=0.67 cm⁻¹ estimated from DC 450 measurements in the same spin-1/2 model ($2.98 \times 10^{-3} \times 15 \times 15$). Transferred to 451 Yb(III) system, the interaction of dipolar origin is more than six times stronger in the 452 very hypothetical case of identical θ . Taking into account the interaction between the 453 two Ising centers, we can describe the ground state with two components: $|\uparrow\uparrow\rangle$ and 454

 $|\downarrow\downarrow\rangle$ separated by J/2 (0.335 cm⁻¹) from the excited state described by $|\uparrow\downarrow\rangle$ and $|\downarrow\uparrow\rangle$. The complex is a SMM. However, the thermal variation of the relaxation time at low 456 temperature is very different from what we are used to observe on mononuclear 457 complexes. Indeed, there is no leveling of the relaxation time on cooling in zero 458 459 external dc field down to 2 K: τ increases continuously on cooling. This is certainly the consequence of the thermal population of the four levels. Remarkably, within an 460 external DC field, τ does not greatly vary. This is a consequence of the condensation 461 of the four states on few tenths of wave numbers. 462

The implication of magnetic interactions on slow relaxation dynamics in dimers is confirmed by other investigations on TTF-based Dy(III) dinuclear complexes. The 464 reaction of tetrathiafulvalene-3-pyridine-N-oxide ligand (L¹⁰) with [Dy(tta)₃]·2H₂O gives the centrosymmetric complex [Dy(tta)₃L¹⁰]₂ [39]. The $\chi_{\rm M}T$ vs. T plot shows strong antiferromagnetic interactions between $^{6}{\rm H}_{15/2}$ multiplets. In the effective spin-1/2 model, the interaction is estimated at -2.3 cm^{-1} with a g value (19.2) close to the Ising limit (20). The nonmagnetic ground state is then described by $|\uparrow\downarrow\rangle$ and $|\downarrow\uparrow\rangle$, with the first excited state $(|\uparrow\uparrow\rangle)$ and $|\downarrow\downarrow\rangle$) at 1.3 cm⁻¹. Despite the nonmagnetic nature of the ground state, [Dy(tta)₃L¹⁰]₂ behaves as a SMM (Fig. 14). The thermal variation of the relaxation time τ does not follow a simple mathematical law since various energy levels are involved at temperatures as low as 2 K. The application of an external dc field corroborates the interpretations based on dc measurements. The field behavior of τ does reflect the low-level energy diagram. At low field and temperatures below 8 K, τ decreases with the field (Fig. 15) with a clear dip at 1.6 kOe. At such temperatures, the first magnetically active $(1\uparrow\uparrow)$ and $|\downarrow\downarrow\rangle$) excited states are thermally populated. It must be pointed out that the transition between these two states necessitates to flip simultaneously both magnetic moments, so the transition probability is very small and the relaxation time long. On increasing the magnetic field, there is a crossing between levels (Fig. 15), and, at the intersection, transition between two states involves "only" to flip one magnetic moment, and the relaxation time shortens. The minimum of τ should occur at a field which can be related to the interaction between Dy(III). With J = -2.3 cm⁻¹, the crossing should

occur at 1.3 kOe which relatively close to the measured value. This in-field behavior

Fig. 14 Temperature dependences of χ_{M}' and χ_{M}'' measured at 1 Hz (black), 10 Hz (light gray), 100 Hz (mid gray), and 1,000 Hz (dark gray) for [Dy(tta)3 L^{10}]₂ in the absence of an external dc field. Full symbols correspond to $\chi_{\rm M}$ and empty symbols to χ_{M}'

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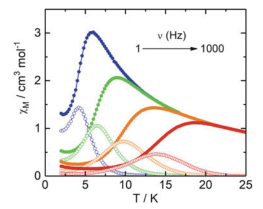
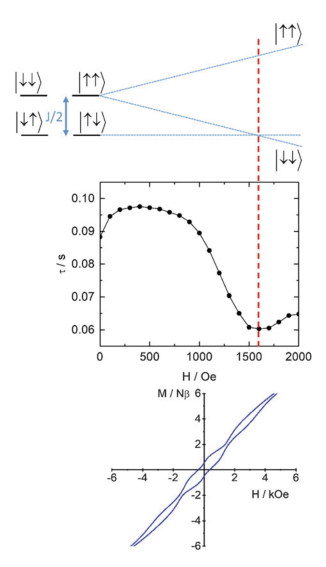


Fig. 15 (Top) field dependence of the relaxation time τ at 5 K with the magnetic field evolution of the energy levels in an Ising pattern for [Dy(tta)₃L¹⁰]₂. (Bottom) hysteresis loop for [Dy(tta)₃L¹⁰]₂ measured at 1.5 K at a sweep rate of 66 Oe s⁻¹



has also a consequence on the magnetic hysteresis. At 1.5 K, the magnetic hysteresis 486 loop measured at 66 Oe s⁻¹ differs significantly from those of mononuclear species 487 (Fig. 15) [71]. The butterfly transforms into a double butterfly. The neck at 1.3 kOe 488 traduces the acceleration of the relaxation at crossing field. In addition, the loop is 489 opened at the origin. One may say that this nonmagnetic object possesses a magnetic 490 memory anyway.

 $[Dy(hfac)_2(SO_3CF_3)\textbf{L}^{11^{\bullet}+}]_2 \text{ is another example of dinuclear TTF-Ln-based complexes} \quad [27] \quad \text{with} \quad 4,5\text{-bis}(3\text{-pyridyl-N-oxidemethylthio})-4',5'\text{-methyldithio-detathiafulvene ligand } (\textbf{L}^{11}). \quad \text{This complex has some common points with } [Dy 494 (tta)_3\textbf{L}^{10}]_2 \text{: the coordination polyhedron is made of eight oxygen atoms, and 495}$

pyridine N-oxide bridges two Dy(III) ions. However, in this system, one monoanionic β-diketonate moiety has been substituted by one monoanionic sulfo-497 nate. One oxygen atom from pyridine N-oxide group completes the coordination 498 sphere. The ligand L¹¹ has been oxidized during galvanostatic. The TTF core is 499 almost planar in agreement with its radical cationic form L¹¹*-. Two non-coordinated sulfonate anions balance the positive charge of the complex. In 501 the crystal lattice, the TTF cores are dimerized with short intermolecular S···S 502 contacts (~3.35 Å), and then the radicals are magnetically inactive (strongly antifer-504 romagnetically coupled). This is confirmed by the very weak EPR signal centered at g~2.007 measured at 77 K. The electrical resistivity measured at room temperature 505 on single crystals corresponds to an insulator. The analysis of the static magnetic 506 properties reveals a weak antiferromagnetic coupling $(J = -3 \times 10^{-3} \text{ cm}^{-1})$ according to Eq. 1) and a Kramers ground state mainly constituted of $M_I = \pm 13/$ 2. The $\chi_{M}^{"}$ vs. ν curves at zero field does not pass through the characteristic 509 maximum at low temperature, so even if it is clear that [Dy(hfac)₂(SO₃CF₃)L^{11•+}]₂ 510 behaves as a SMM, it is less efficient than the previous example probably because the environment around the lanthanide is chemically different. 512

The last two examples we would like to tackle are polynuclear complexes 513 which feature more than two metal centers. The rational design of a lanthanide-514 based complex featuring different lanthanide motifs can be safely envisaged in combining $[Dy(tta)_3L^{10}]_2$ and $[Dy(tta)_3L^2]$. To do so a TTF-based ligand has been designed to feature a bridging site of pyridine N-oxide type and a 517 bischelating nitrogen-based site [72]. The ligand 2-{1-methylpyridine-N-oxide-4,5-[4,5-bis(pro-pylthio)tetrathiafulvalenyl]-1H-benzimidazol-2-yl $pyridine (L^{12})$ was then treated with two equivalents of [Dy(tta)₃]·2H₂O to give the complex 520 $[Dy_4(tta)_{12}(L^{12})_2]$ (Fig. 16). In the complex, the reader will recognize one moiety 521 similar to $[Dy(tta)_3L^{10}]_2$ and two moieties similar to $[Dy(tta)_3L^2]$. The distance between these moieties (metal-metal) is above 10 Å, so there is no direct interaction between these three distinct SMMs. The magnetism of this object 524 should coincide with the superposition of two different SMMs. In zero external field, the $\chi_{\rm M}$ vs. ν curves at various temperatures between 2 and 525 526

In zero external field, the $\chi_{\rm M}^n$ vs. ν curves at various temperatures between 2 and 11 K clearly show two well-separated relaxations which can be confronted to the measurements on the isolated species. The presence of a slow and a fast process at low and high frequencies, respectively, matches almost perfectly with the isolated species. The low-frequency side corresponds to the dinuclear part and the high frequency to the mononuclear. It is also possible to analyze quantitatively the thermal and the in-field behaviors with a combination of two extended Debye models. At this stage our synthetic approach allowed us to conceive a complex which contains two different SMMs which act differently in the temperature and time scales. This rational design is very promising to elaborate multifunctional complexes (Fig. 17).

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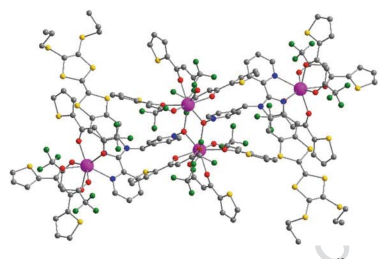
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The last example we would like to comment concerns the polymeric species. [Yb 538 (hfac)₃]·2H₂O was reacted with the disodium salt of $\mathbf{L^{13}}$ with $\mathbf{H_2L^{13}}=4,5$ -bis 539 (carboxylic)-4',5'-methyldithiotetrathiafulvene in dimethylformamide (DMF) to 540 produce $\{[YbL^{13}(H_2O)_3(DMF)]\cdot(HL^{13})\cdot(H_2O)\}_n$ [73]. The monodimensional



 $\textbf{Fig. 16} \quad \text{Single-crystal X-ray structure of the tetranuclear complex } [Dy_4(tta)_{12}(\textbf{L^{12}})_2]$

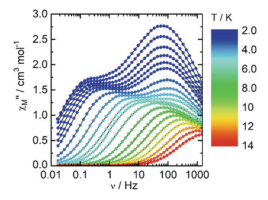
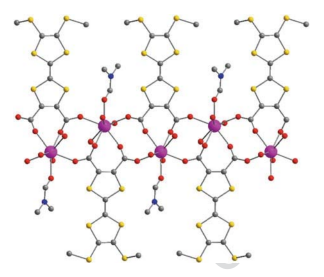


Fig. 17 Temperature and frequency dependence of the out-of-phase component of the ac susceptibility $[\mathrm{Dy_4(tta)_{12}L^{12}_2}]$ measured in zero external dc field

polymer consists of chain of Yb(III) bridged by carboxylate anions in a $\mu_2(\eta_1,\eta_1)$ 541 mode (Fig. 18). Coordination compounds of lanthanide ions with TTF-based ligands 542 are not so common and are essentially 0D. This is the first example of coordination 543 polymer of lanthanide with TTF-based ligand. The coordination polyhedron is made 544 of eight oxygen atoms in a D_{4d} environment (CShM_{SAPR-8} = 0.454). Our efforts to 545 produce the dysprosium derivative were unsuccessful. It must be mentioned that no 546 hfac⁻ anions are present in the structure and that the polymer cannot be obtained 547 from nitrate or halogenate salts of Yb(III). The system crystallizes in the triclinic 548 space group $P\bar{1}$ with one Yb(III) site, so the g-tensor can be extracted from single-549 crystal rotating magnetometry. In the effective spin-1/25 model, $g_x = 3.24$, 550 $g_y = 1.53$, and $g_z = 4.25$. These values are far away from the Ising limit for which 551

Fig. 18 Representation of the one-dimensional structure. The counterion (HL¹³)⁻ has not been represented



552 $g_x = g_y = 0$ and $g_z = 8.00$. As a result, the polymer does not behave as a SMM in 553 zero external dc field. Only when an external dc field is applied that χ_M " shows up. g_z 554 orientation almost coincides with the fourfold axis of the square antiprism. We 555 tentatively tried to reproduce the static magnetic properties using the ab initio 556 CASSCF/PT2/SI-SO approach. Unfortunately, all tentative efforts failed to properly 557 reproduce the magnetic susceptibility and magnetization curves. This underlines the 558 difficulties already observed in the literature to efficiently model both the wave 559 function and the energy of the low-lying multiplets of Yb(III) complexes.

560 7 Conclusions

In this chapter we wish to have convinced the reader that TTF-based ligands can be employed to produce SMMs. We have focused the first part of this chapter on strategies to enhance the magnetic performance of Dy-based SMM in a N₂O₆ environment. A simple molecular engineering consisting in the modulation of the electron withdrawing strength of the β-diketonate ancillary ligand highlighted the importance of the electron charge density carried by the first neighboring atom to control the energy crystal splitting and well isolate the ground multiplet state. The 567 canceling of the intermolecular (hydrogen bond and dipolar) interactions, thanks to 568 magnetic dilutions and spin-free isotopic enrichment, showed their efficiency to decrease the quantum tunneling of the magnetization and therefore optimize the magnetic properties of the SMM. Then the influence of the nature of the coordination sphere was studied by the analysis of mononuclear SMM library in which the Dy(III) is in N₂O₆, N₃O₆, O₈, and O₉ environments. When the Dy(III) is placed in a N₂O₆ environment, the Ising character of the magnetic anisotropy is enhanced compared to the N₃O₆ environment. In a general manner, SMM behavior is detected when the 575 negative charge is localized along an axis and in a plan, respectively, for the Dy(III) 576 and Yb(III) ions [41]. In a second part, we have increased the nuclearity of the 577 complexes. The role of the intramolecular magnetic interactions on the slow mag- 578 netic relaxation has been demonstrated.

Finally it is worth to notice that a large panel of theoretical and experimental tools 580 is available and can be used to reach a high level of understanding of the lanthanide 581 SMM magnetic properties, i.e., experimental measurements of the angular dependence of the magnetization, correlation between magnetism and experimental luminescence, crystal-field determination by Stevens method, and ab initio calculations. 584

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Tetrathiafulvalene-Based Magnets of Lanthanides

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