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Chiral or Luminescent Lanthanide Single-Molecule Magnets Involving Bridging Redox Active Triad Ligand

Bertrand Lefeuvre, Jessica Flores Gonzalez, Carlo Andrea Mattei, Vincent Dorcet, Olivier Cador and Fabrice Pointillart *

ISCR (Institut des Sciences Chimiques de Rennes)—UMR-CNRS 6226, Université de Rennes, 35000 Rennes, France; bertrand.lefeuvre@univ-rennes1.fr (B.L.); jessica.flores-gonzales@univ-rennes1.fr (J.F.G.); carlo-andrea.mattei@univ-rennes1.fr (C.A.M.); vincent.dorcet@univ-rennes1.fr (V.D.); olivier.cador@univ-rennes1.fr (O.C.)

* Correspondence: fabrice.pointillart@univ-rennes1.fr; Tel.: +33-(0)2-23-23-67-52; Fax: +33-(0)2-23-23-68-40

Abstract: The reactions between the bis(1,10-phenanthro[5,6-b])tetrathiafulvalene triad (L) and the metallo-precursors Yb(hfac)$_3$(H$_2$O)$_2$ (hfac$^-$ = 1,1,1,5,5,5-hexafluoroacetylactonato anion) and Dy(facam)$_2$ (facam$^-$ = 3-trifluoro-acetyl(-)-camphorato anion) lead to the formation of two dinuclear complexes of formula [Yb$_2$(hfac)$_6$(L)]·2(C$_7$H$_6$O) and [Dy$_2$((+)-facam)$_6$(L)]·2(C$_6$H$_{14}$) (2). The X-ray structures reveal that the L triad bridges two terminal Yb(hfac)$_3$ or Dy(facam)$_2$ units. (1)-2(C$_7$H$_6$O) behaved as a near infrared Yb$^{III}$ centered emitter and a field-induced Single-Molecule Magnet (SMM) while (2)-2(C$_6$H$_{14}$) displayed SMM behavior in both zero- and in-dc field. The magnetization mainly relays through a Raman process for both complexes under an optimal applied magnetic field.

Keywords: tetrathiafulvalene; triads; lanthanides; chirality; luminescence; single-molecule magnet

1. Introduction

Lanthanide ions are playing a crucial role in the design of Single-Molecule Magnets (SMM) thanks to their strong magnetic anisotropy and high magnetic moment [1–10]. The ideal adequacy between the nature of the used lanthanide and crystal field environment allowed chemists to elaborate SMMs displaying magnetic memory at a temperature up to 80 K [11,12]. Thus, such molecular objects are back in the race for potential applications in data storage as well as quantum computing [13,14]. Lanthanide elements are well-known to be highly luminescent [15,16] pushing up both chemist and physicist communities for having an interest in the design of luminescent SMMs for which magneto-optical correlation could be performed [17–20]. In a general manner, the addition of one or more physical properties to the SMM behavior paves the way to the design of multi-properties SMM as well as the study of a possible synergy between the different properties. Moreover, when chirality is added to the (luminescent) SMM behavior [21–28], chiral SMM [29], ferroelectric SMM [30] as well as chiral luminescent SMM [31–36] and magneto-chiral SMM [37,38] can be obtained. To these three properties, the introduction of redox-active ligands is of interest because it could permit both optical and magnetic properties depending on the oxidation state of the ligand [39–46]. One of the most used redox-active ligands is made from the tetrathiafulvalene (TTF) core because of its two mono radical and dicationic stable states which are easily accessible by chemical ways and because it can be easily decorated with one to four substituted groups able to coordinate transition metal and lanthanide ions [18,47–49]. Among the plethora of chemical substituents [50,51], the 1,10-phenanthroline (phen) is a remarkable choice to design donor-acceptor dyads (called also push-pull ligands) [52–55] or acceptor-donor-acceptor triads [56,57]. Once associated with metallic precursors, auspicious optical and magnetic properties are observed highlighted by
the recent design of a redox-active chiral luminescent SMM displaying circularly polarized luminescence [58].

In the last example, the bis(1,10-phenanthro[5,6-b]tetrafluorovalene triad (L) was employed. The following lines propose to select this triad to bridge two Yb(hfac)$_3$·(H$_2$O)$_2$ (hfac$^{-}$ = 1,1,1,5,5,5-hexafluoroacetylacetonato anion) and two Dy(facam)$_3$ (facam = 3-trifluoro-acetyl-(+)-camphorato anion). Thus the two dinuclear complexes of formula [Yb$_2$(hfac)$_6$(L)]·2(C$_7$H$_{16}$) ((1)·2(C$_7$H$_{16}$)) and [Dy$_2$((+)facam)$_6$(L)]·2(C$_6$H$_{14}$) ((2)·2(C$_6$H$_{14}$)) have been obtained. Their single crystal structure, magnetic and optical properties have been investigated.

2. Results and Discussion

2.1. Crystal Structure Description of [Yb$_2$(hfac)$_6$(L)]·2(C$_7$H$_{16}$) ((1)·2(C$_7$H$_{16}$))

(1)·2(C$_7$H$_{16}$) crystallizes in the P-1 (N°2) triclinic space group. The X-ray crystallographic data for compound (1)·2(C$_7$H$_{16}$) are given in Table S1. The asymmetric unit is composed of two half dinuclear complexes of formula [Yb$_2$(hfac)$_6$(L)] and two n-heptane molecules of crystallization. Thus, two crystallographically independent Yb$^{III}$ centers are present in the molecular structure (Figure S1). At this point, it is worth noticing that the use of n-heptane instead of n-hexane induces the reduction of crystallographically independent metal centers from six to two [59]. All the Yb$^{III}$ centers are linked to three hfac$^{-}$ anions and one L ligand giving an N2O6 surrounding with a D$_{2d}$ symmetry (triangular dodecahedron) for Yb$_1$ and D$_{4d}$ symmetry (square antiprism) for Yb$_2$ (Figure 1, Table S2). The distortion from ideal symmetry can be explained by the Yb-O bond lengths (2.285 Å) shorter than the Yb-N ones (2.481 Å) (Table S3) and it is visualized by continuous shape measures performed with SHAPE 2.1 (Table S2) [60]. The two 1,10-phenanthroline coordination sites are occupied by Yb(hfac)$_3$ units and L plays the role of bridge between two Yb(hfac)$_3$ units (Figure 1). The bridging triads L are planar and the average C=C central distances (1.340 Å) agree with the neutral form of L [61,62]. Finally, the crystal packing highlighted the one-dimensional stacking of dinuclear complexes through π-π interactions (Figure 2). No significant S···S short contacts are present since the shortest S···S distance has been identified equally to 3.955 Å. The shortest intramolecular and intermolecular Yb-Yb distances are 17.742 Å and 9.489 Å respectively.

Figure 1. Coordination reactions between L and Yb(hfac)$_3$·(H$_2$O)$_2$ and Dy(facam)$_3$ as well as molecular structures of [Yb$_2$(hfac)$_6$(L)]·2(C$_7$H$_{16}$) ((1)·2(C$_7$H$_{16}$)) and [Dy$_2$((+)facam)$_6$(L)]·2(C$_6$H$_{14}$) ((2)·2(C$_6$H$_{14}$)).
Figure 2. Crystal packing of (1)-2(C$_7$H$_{16}$) highlighting the 1D stacking of L (spacefill representation). The Yb(hfac)$_3$ units are represented by balls and sticks.

2.2. Crystal Structure Description of [Dy$_2$((+)(facam)$_6$(L))·2(C$_6$H$_{14}$) ((2)-2(C$_6$H$_{14}$))

(2)-2(C$_6$H$_{14}$) crystallizes in the C2 (N°5) monoclinic chiral space group. The X-ray crystallographic data for compound (2)-2(C$_6$H$_{14}$) are given in Table S1. This compound has been found isostructural to the Yb$^{III}$ analog recently published by some of us [57]. The asymmetric unit consists of one dinuclear complex of formula [Dy$_2$(facam)$_6$(L)] and two n-hexane molecules of crystallization. Thus, two crystallographically independent Dy$^{III}$ centers are present in the molecular structure (Figure S1). Both Dy$^{III}$ centers are linked to three facam$^-$ anions and one L ligand giving an N2O6 surrounding with a D$_{4d}$ symmetry (square antiprism, Table S2) (Figure 1). The Dy-O bond lengths (2.285 Å) are shorter than the Dy-N ones (2.481 Å) (Table S3). One could remark that three short Dy-O and three longer Dy-O bond lengths for each Dy$^{III}$ center are observed due to the dissymmetric nature of the facam$^-$ ancillary ligand. The short Dy-O distance involves the oxygen atom of the carbonyl group linked to the CF$_3$ moiety (Table S3). The two Dy(facam)$_3$ units are bridged by a triad L which is not rigorously planar (Figure 1). The average C=C central distances (1.331 Å) agree with the neutral form of L [61,62].

Finally, the crystal packing did not show any significant π–π stacking or S···S short contact (Figure 3) on contrary to what is observed in the X-ray structure of the achiral analog [59]. The higher steric hindrance of the facam$^-$ anions compared to hfac$^-$ could explain this difference. Nevertheless, short S2···O3 contacts (3.115 and 3.183 Å) between the TTF core and one facam$^-$ ligand of the neighboring molecule ensure the coherence of the crystal packing. The shortest intramolecular and intermolecular Dy-Dy distances are 17.867 Å and 10.211 Å, respectively.

Figure 3. Crystal packing of (2)-2(C$_6$H$_{14}$) with the triads L and Dy(facam)$_3$ units drawn respectively in spacefill and ball and sticks representation.
2.3. Electrochemical Properties

The two complexes 1 and 2 displayed two monoelectronic oxidations respectively determined at 0.87 V/1.22 V and 0.83 V/1.17 V (vs. SCE) (Figures S3 and S4, Table S4) and attributed to the formation of the radical cation and dication species of the TTF core. These potential values are in line with those observed for their analogs previously published [58,59]. Cyclic voltammetry demonstrated the reversible redox-activity of the two triad-based complexes.

2.4. Photophysical Properties

Irradiation of the HOMO → LUMO Intra-Ligand Charge Transfer (ILCT) of L (Figures S5 and S6) [57] at 22,220 cm⁻¹ (450 nm) in complex 1 induced a Near InfraRed (NIR) emission centered at 9900 cm⁻¹ in the solid-state at 10 K (Figure 4). Such NIR signal is attributed to the YbIII centered 2F5/2 → 2F7/2 emission. The TTF-centered dyads or triads have been demonstrated to be efficient chromophores for the sensitization of NIR lanthanide luminescence through the antenna process [18,58]. Such efficiency for 1 is proven by the absence of residual ligand-centered emission which attests to a rather efficient energy transfer from the triad L to the YbIII ion. The low-temperature measurement allowed us to observe a well-resolved emission spectrum with several emission contributions in the energy range of 10,300–9700 cm⁻¹. At least eight contributions could be identified with maxima at 10,235, 10,204, 9975, 9940, 9753, 9711, 9671, and 9629 cm⁻¹ which is more than the four expected sub-states due to the splitting of the 2F7/2 ground state under the crystal field effect [63,64]. Moreover, one could remark that the energy difference between two consecutive transitions is about 37(5) cm⁻¹ leading to the suspicion that the four additional transitions may originate from the second crystal field sublevel of the excited 2F5/2 state as proposed on the right part of Figure 4 [65,66]. These so-called “hot bands” usually disappear upon the lowering of the temperature that depopulates upper excited states [28,67–69]. Consequently, these “hot bands” as well as other contributions of weak intensities that appeared in the emission spectrum may be also attributed to the vibronic coupling between electronic transitions and low energy vibrational modes as recently suggested for such YbIII ion in molecular systems [38,70].

![Figure 4](image-url)
Similar light excitation into the ILCT at 22,220 cm$^{-1}$ (Figure S4) for 2 cannot give the visible Dy$^{III}$ centered emission because of the too low-energy absorption band of L.

2.5. Magnetic Properties

The thermal dependence of the magnetic susceptibility ($\chi_M$) and the field dependence of the magnetization (M) are given for two Yb$^{III}$ and Dy$^{III}$ centers for 1 (Figure S7) and 2 (Figure S8), respectively. The room temperature values of the $\chi_M$T product are 4.97 cm$^3$ K mol$^{-1}$ for 1 and 27.47 cm$^3$ K mol$^{-1}$ for 2 in agreement with the expected values for two Yb$^{III}$ ions (5.14 cm$^3$ K mol$^{-1}$, $^2F_{7/2}$ ground state multiplet) and two Dy$^{III}$ ions (28.56 cm$^3$ K mol$^{-1}$, $^6H_{15/2}$ ground state multiplet) [71]. Upon cooling, $\chi_M$T decreases monotonically down to 2.34 cm$^3$ K mol$^{-1}$ for 1 and 20.27 cm$^3$ K mol$^{-1}$ at 2 K. The decrease of the $\chi_M$T products can be mainly attributed to the thermal depopulation of the $M_J$ state even if the intermolecular dipolar interaction effect cannot be ruled out in case of 2. Both compounds exhibited classical magnetization for Yb$^{III}$ and Dy$^{III}$ ions in N2O6 environment with experimental values of 3.26 N$\beta$ for 1 and 9.58 N$\beta$ for 2 for the saturation values sign of significant magnetic anisotropy.

1 does not show any out-of-phase component of the ac susceptibility in zero external dc fields (Figure S9). Fast relaxation through Quantum Tunneling of the Magnetization (QTM) could be the origin of the lack of SMM behavior in zero fields. Nevertheless, it is well known that QTM can be canceled by applying an external dc field. Thus, the field dependence of the ac magnetic susceptibility was performed (Figure S9). The application of a moderate external dc field (0–3000 Oe) induces an out-of-phase signal in the frequency window (40–10,000 Hz) that does not significantly shift with the applied field but grows in amplitude (Figure S9). The optimum field value was determined from the $\tau$ vs. $H$ plot as the value at which the best compromise between slow relaxing fraction, $\tau$ value and maximum amplitude are reached (Figure 5a). The $\tau$ vs. $H$ curve was fitted with the equation 1 for the 0–3000 Oe field range (Figure 5a) [45,72].

$$\tau^{-1} = \frac{B_1}{1 + B_2H^2} + 2B_3H^m + B_4$$

Figure 5. (a) Field dependence of the relaxation time at 2 K in the field range of 0–3000 Oe for 1 (size of the black dot is proportional to the slow relaxing fraction) with the best-fitted curve (full black line) with the combination Orbach + Raman (dashed brown line), QTM (dashed orange line) and direct (dashed blue line) processes. (b) frequency dependence of $\chi_M''$ for 1 at 1000 Oe in the temperature range 2–10 K; (c) temperature dependence of the relaxation time for 1 at 1000 Oe in the temperature range 2–4.25 K with the best-fitted curve (full black line) with Raman process only. Error lines are calculated using the log-normal distribution model at the 1$\sigma$ level [73].
The three terms correspond respectively to the QTM, Direct, and field-independent magnetic relaxation processes (Raman and Orbach). The best fit was obtained for $B_1 = 2.53 \times 10^6$ s$^{-1}$; $B_2 = 0.11$ Oe$^{-2}$; $B_3 = 1.83(19) \times 10^{-11}$ s$^{-1}$ K$^{-1}$ Oe$^{-4}$ (m fixed to 4) and $B_4 = 8804(114)$ s$^{-1}$ leading at 1000 Oe to $\tau^{-1}$ (QTM) = 23.0 s$^{-1}$ (dashed orange line on Figure 5a), $\tau^{-1}$ (Direct) = 36.6 s$^{-1}$ (dashed blue line on Figure 5a) and $\tau^{-1}$ (Raman + Orbach) = 8804 s$^{-1}$ (dashed brown line on Figure 5a). Thus, at 1000 Oe field value, both QTM and Direct processes could be discarded while the thermal variation of the magnetic relaxation times could be fitted by using a combination of Orbach and Raman processes.

At such a field, the variation of the ac susceptibility (Figure 5b and Figure S10) with the oscillation frequency of the magnetic field can be quantitatively analyzed in the framework of the extended Debye model (Figure S11). The slowly relaxing fraction of the magnetic susceptibility represents almost the entire sample (90–95%) which means that the vast majority of the magnetic moments are involved in the relaxation process and can be visualized from the Cole–Cole plot (Figure S12). The $\alpha$ parameter ranges from 0.01 and 0.09 between 2 and 4.25 K (Table S6) which is in agreement with a single relaxation time. From these indications, one could conclude that both crystallographically independent Yb$^{III}$ displayed indistinguishable ac magnetic dynamics. The high-temperature part of the thermal variation of the relaxation time (Figure 5c) can be fitted leading to an Orbach process ($\tau^{-1} = \tau_0^{-1} \exp(\Delta/T)$ characterized by $\tau_0 = 7(2) \times 10^{-7}$ s and $\Delta = 12(1)$ K (Figure S13). The value of the effective energy barrier is much smaller than the energy gap given by the luminescence spectrum of $1$ at 10 K (264 cm$^{-1}$ (380 K)) which is a strong indication of the involvement of the under-energy barrier process in magnetic relaxation. Attempt to fit the log($\tau$) vs. $T$ plot with combinations of Orbach + Raman process ($\tau^{-1} = \tau_0^{-1} \exp(\Delta/T) + C T^n$) led to the following parameters $\tau_0 = 2.97(78) \times 10^{-7}$ s and $\Delta = 21(2)$ K and $C = 1957(220)$ s$^{-1}$ K$^{-n}$ with $n = 2.21(16)$ (Figure S14). The Orbach process appeared too slow to contribute to the magnetic relaxation in agreement with the discrepancy between the effective energy barrier and emission spectrum. As the ac magnetic susceptibility measurements have been performed under an applied dc field, the Direct process should contribute to the magnetic relaxation nevertheless the best fit of the log($\tau$) vs. $T$ plot with combinations of Raman + Direct processes ($\tau^{-1} = C T^n + A H^4 T$) gave the following parameters $C = 369(47)$ s$^{-1}$ K$^{-n}$ with $n = 3.53(8)$ and $A = 2.44(18) \times 10^{-9}$ s$^{-1}$ Oe$^{-4}$ K$^{-1}$ (Figure S15) demonstrating that the Direct process could be involved in the magnetic relaxation at the lowest temperature (2 K). Consequently, the most reasonable fit of the thermal dependence of the relaxation time was obtained by using a Raman process only ($\tau^{-1} = C T^n$) with the parameters $C = 1279(70)$ s$^{-1}$ K$^{-n}$ with $n = 2.75(5)$ (Figure 5c). The value of $n$ is initially predicted equal to 9 for Kramers ions [74] when the Raman process operating through acoustic phonons (lattice vibrations) but can be found between 2 and 7 [75–77] because of the presence of optical phonons (molecular vibrations) [12,14].

2 shows a frequency dependence of the ac susceptibility in zero external dc fields with a non-zero out-of-phase component ($\chi_M'$) (Figure 6a). The variation of the ac susceptibility (Figure 6a and Figure S16) with the oscillation frequency of the magnetic field can be quantitatively analyzed in the framework of the extended Debye model (Figure S17). The normalized Cole–Cole plot (Figure S18) allowed us to determine that the non-relaxing fraction is about 10% of the sample meaning that the measured out-of-phase component ($\chi_M'$) represents almost the entire compound. The $\alpha$ parameter ranges from 0.18 to 0.42 between 2 and 14 K (Table S7) which is in agreement with a wide range of relaxation time due to the involvement of several relaxation processes at zero applied field. From the thermal dependence of the relaxation time (Figure 6d), one could remark that the magnetic performances of 2 are limited due to a thermally independent behavior of log($\tau$) vs. $T$ attributed to the QTM. The latter could be canceled by applying an external dc field. Thus field dependence of the magnetic susceptibility was performed (Figure S19). The application of a moderate external dc field (0–3000 Oe) induces a shift to lower frequency of the out-of-phase signal from 1000 Hz at 0 Oe to about 0.2 Hz for $H > 600$ Oe (Figure S19). Significant dipolar intermolecular interactions are present as double contributions at 200 Oe.
was observed for the out-of-phase signal. The optimum field value of 1200 Oe was determined from the τ vs. H plot curve which was fitted with the equation for the 0–3000 Oe field range (Figure 6b). The best fit was obtained for $B_1 = 6043.6 \, s^{-1}$, $B_2 = 2.23 \times 10^{-2} \, Oe^{-2}$, $B_3 = 6.77 \times 10^{-14} \, s^{-1} \, K^{-1} \, Oe^{-4}$ (m fixed to 4) and $B_4 = 0.75 \, s^{-1}$ leading at 1200 Oe to $\tau^{-1}$ (QTM) = 0.19 $s^{-1}$ (dashed orange line on Figure 5a), $\tau^{-1}$ (Direct) = 0.28 $s^{-1}$ (dashed blue line on Figure 5a) and $\tau^{-1}$ (Raman + Orbach) = 0.75 $s^{-1}$ (dashed brown line on Figure 5a).

Thus, at 1200 Oe field value, both QTM and Direct processes might be discarded while the thermal variation of the magnetic relaxation times could be fitted by using a combination of Orbach and Raman processes. At such a field, the variation of the ac susceptibility (Figure 6c and Figure S20) with the oscillation frequency of the magnetic field can be quantitatively analyzed in the framework of the extended Debye model (Figure S21). One more time the slowly relaxing fraction can be determined from the normalized Cole–Cole (about 90%) (Figure S22) and the α parameter at 1200 Oe are close to the values found at 0 Oe (from 0.10 to 0.44) (Table S9). One could notice that applying a dc magnetic field should not modify the thermally activated relaxation processes such as Orbach ($\tau^{-1} = \tau_0^{-1} \exp(\Delta/T)$) and Raman ($\tau^{-1} = CT^n$) processes. Consequently, Orbach and Raman parameters for thermal variation of the relaxation times at 0 Oe and 1200 Oe should be shared. The effective energy barrier was first determined from the fit of the high-temperature region of log($\tau$) vs. T plot of $2$ at 1200 Oe using an Orbach process only (Figure S23). The best fit was obtained for $\tau_0 = 3.2(7) \times 10^{-6} \, s$ and $\Delta = 53(2) \, K$ parameters which are used to simultaneously fit the log($\tau$) vs. T plots at 0 Oe and 1200 Oe with shared Raman parameters and additional QTM process in zero fields. It is worth noticing that this effective energy barrier is close to the value of 55.1 K found in a similar dinuclear complex involving the simple 2,2'-bipyrimidine bridging ligand instead of $L$ [78]. The best simultaneous fit was obtained for $\tau_0 = 3.2 \times 10^{-6} \, s$ (fixed) and $\Delta = 53 \, K$ (fixed), $C = 0.10(2) \, s^{-1} \, K^{-n}$ with $n = 3.94(15)$ and $\tau_{TI} = 1.30(8) \times 10^{-4} \, s$ (Figure S24). One more time the Raman contribution at a given temperature (5 K) are of the same order of magnitude in the present and published examples. From the previous fit, it could be concluded that the Orbach process only weakly contributes in the temperature range of 2–14 K. Thus a simultaneous fit could be performed by using a combination of Raman and QTM ($H = 0$ Oe) and Raman only ($H = 1200$ Oe) processes for which the Raman parameters are shared. The best fit is depicted in Figure 6d with the following parameters $C = 0.049(5) \, s^{-1} \, K^{-n}$ with $n = 4.6(6)$ and $\tau_{TI} = 1.30(5) \times 10^{-4} \, s$. Importantly, $\tau_{TI}$ fitted from thermal variation matches the zero-field limit of Equation (1) ($B_1^{-1}$) obtained from field variations.

The Raman process operates through both acoustic phonons (lattice vibrations) and optical phonons (molecular vibrations) since the n value is weaker than the one expected ($n = 9$) for a pure acoustic phonon-assisted Raman relaxation.

One could remark that changing the ancillary hfac$^-$ with the facam$^-$ ligand enhanced the magnetic performances for both Dy$^{III}$ and Yb$^{III}$. Indeed, at 2 K, $1$ displayed a $\chi_M''$ peak centered at 1500 Hz while $[\text{Yb(facam)}_3(L)]_2$ displayed a $\chi_M''$ peak centered at 100 Hz [58]. Similarly, $2$ displayed a $\chi_M''$ peak centered at 1000 Hz ($H = 0$ Oe) and 0.2 Hz ($H = 1200$ Oe) while $[\text{Dy(hfac)}_3(L)]_2 \cdot \text{CH}_2\text{Cl}_2 \cdot \text{C}_6\text{H}_{14}$ displayed no SMM behavior at 0 Oe and a $\chi_M''$ peak centered at 2 Hz at 1000 Oe [59].
3. Experimental Section

3.1. Synthesis of \([\text{Yb(hfac)}_3(H_2O)](\text{L})\cdot 2(\text{C}_6\text{H}_{14})\) (1)

The precursor \(\text{Yb(hfac)}_3(H_2O)_2\) (hfac\(^{-}\) = 1,1,1,5,5,5-hexafluoroacetylacetonate) \([79]\), \(\text{Dy}((+)\text{facam})_3\) (facam\(^{-}\) = 3-trifluoro-acetyl-(+)-camphorato anion) \([58]\) and the bis(1,10-phenantro[5,6-b])tetrafluorvalene ligand \([57]\) were synthesized by following previously reported methods. All other reagents were purchased from Aldrich Co., Ltd. (St. Louis, MO, USA) used without further purification. All solid-state characterization studies (elementary analysis, IR, photo-physical and magnetic susceptibility measurements) were performed on dried samples and are considered without solvent of crystallization.

3.2. Synthesis of Complex \([(\text{Dy(hfac)}_3(\text{L}))\cdot 2\text{C}_6\text{H}_{14}h]\) 1

11 mg of \(\text{L}\) (0.022 mmol) were added to 20 mL of 1,2-dichloroethane. The suspension was heated to reflux and then a solution of 10 mL of 1,2-dichloroethane containing 36.1 mg of \(\text{Yb(hfac)}_3(H_2O)_2\) (0.044 mmol) was added. After 6 h of reflux, the 1,2-dichloroethane
was eliminated under vacuum and the residue was dissolved in CH2Cl2. n-heptane was layered on the CH2Cl2 solution of 1 in the dark to give red single crystals suitable for X-ray diffraction study. Yield 37.5 mg (73%) for 1. I.R. bands (KBr): 2970, 1650, 1564, 1538, 1498, 1465, 1255, 1218, 1147, 1098, 801, 662, and 587 cm\(^{-1}\). Crystallographic data are summarized in Table S1. Complete crystal with a full-matrix least-squares method on \(F^2\) was heated to reflux and then a solution of 10 mL of 1,2-dichloroethane containing 39.6 mg was eliminated under vacuum and the residue was dissolved in CH2Cl2· n-heptane was layered on the CH2Cl2 solution of 2 in the dark to give red single crystals. Yield 41.3 mg (81%) for 2. I.R. bands (KBr): 2969, 1649, 1564, 1537, 1501, 1464, 1260, 1216, 1147, 1100, 801, 660, and 590 cm\(^{-1}\). Anal. Calcd (%) for C\(_{60}\)H\(_{18}\)Yb\(_6\)F\(_{108}\)N\(_{12}\)O\(_{36}\)S\(_{12}\) (1): C 33.64, H 1.51, N 2.49; found: C 33.71, H 1.66 N, 2.37.

3.3. Synthesis of [Dy\((+\text{facam})_6\)(L)]·2(C\(_6\)H\(_{14}\)) ((2·2-C\(_6\)H\(_{14}\))

11 mg of L (0.022 mmol) were added to 20 mL of 1,2-dichloroethane. The suspension was heated to reflux and then a solution of 10 mL of 1,2-dichloroethane containing 39.6 mg of Dy\((+\text{facam})_3\) (0.044 mmol) was added. After 6 h of reflux, the 1,2-dichloroethane was eliminated under vacuum and the residue was dissolved in CH2Cl2· n-heptane was layered on the CH2Cl2 solution of 2 in the dark to give red single crystals. Yield 41.3 mg (81%) for 2. I.R. bands (KBr): 2969, 1649, 1564, 1537, 1501, 1464, 1260, 1216, 1147, 1100, 801, 660, and 590 cm\(^{-1}\). Anal. Calcd (%) for 2 C\(_{98}\)H\(_{96}\)Dy\(_2\)F\(_{18}\)N\(_4\)O\(_{12}\)S\(_4\): C 50.76, H 4.14, N 2.42; found: C 51.07, H 4.19 N, 2.34.

3.4. Crystallography

Single crystal of (1·2-(C\(_7\)H\(_{18}\)) and (2·2-(C\(_6\)H\(_{14}\)) were mounted on a D8 VENTURE Bruker-AXS diffractometer for data collection (MoK\(_\alpha\) radiation source, \(\lambda = 0.71073 \text{ Å}\)), from the Centre de Diffractométrie X (CDIFX), Université de Rennes 1, France. The structure was solved with a direct method using the SHELXT program [80] and refined with a full-matrix least-squares method on \(F^2\) using the SHELXL-14/7 program [81]. A SQUEEZE procedure of PLATON [82] was performed as the structures for (1·2-(C\(_7\)H\(_{18}\)) and (2·2-(C\(_6\)H\(_{14}\)) contain large solvent-accessible voids in which residual peaks of diffraction were observed. Crystallographic data are summarized in Table S1. Complete crystal structure results as a CIF file (CCDC 2,086,542 for (1·2-(C\(_7\)H\(_{18}\)) and CCDC 2,086,543 for (2·2-(C\(_6\)H\(_{14}\))) including bond lengths, angles, and atomic coordinates are deposited as Supplementary Materials.

3.5. Physical Measurements

The elementary analyses of the compounds were performed at the Centre Régional de Mesures Physiques de l’Ouest, Rennes.

Cyclic voltammetry was carried out in dried and degassed CH2Cl2 solution, containing 0.1M N(C\(_4\)H\(_4\))PF\(_6\) as supporting electrolyte. Voltammograms were recorded at 100 mVs\(^{-1}\) at a platinum disk electrode. The potentials were measured vs. a saturated calomel electrode (SCE).

Absorption spectra were recorded on a JASCO V-650 spectrophotometer in diluted solution, using spectrophotometric grade solvents.

Solid-state emission and excitation spectra were measured using a Horiba Jobin Yvon Fluorolog-3\(^\circ\) spectrofluorimeter, equipped with a three slit double grating excitation and emission monochromator with the dispersion of 2.1 nm mm\(^{-1}\) (1200 grooves mm\(^{-1}\)). The steady-state luminescence was excited by unpolarized light from a 450 W xenon CW lamp and detected at an angle of 90° by a liquid nitrogen-cooled InGAs detector. Spectra were reference corrected for both the excitation intensity variation and the emission spectral response. Solid sample was placed in a 0.5 mm diameter quartz tube that was set into an Oxford Instrument cryostat (Optistat CF2) insert directly in the sample chamber of the spectrofluorimeter. The interference signals due to scattered excitation light were suppressed by a 780 nm high pass filter placed at the entry of the emission monochromator.

The dc magnetic susceptibility measurements were performed on a solid polycrystalline sample with a Quantum Design MPMS-XL SQUID magnetometer between 2 and 300 K in the applied magnetic field of 0.2 T for temperatures of 2–20 K, 1 T for temperatures of 20–300 K for 1 while 0.02 T for temperatures of 2–20 K, 0.2 T for temperatures of 20–80 K and 1 T for temperatures of 80–300 K were used for 2. The ac magnetic susceptibility measurements were performed using a Quantum Design MPMS-XL SQUID for frequen-
cies between 1 and 1000 Hz and Quantum Design PPMS magnetometers for frequencies between 50 and 10,000 Hz. These measurements were all corrected for the diamagnetic contribution as calculated with Pascal’s constants.

4. Conclusions and Outlook

Two dinuclear complexes of formula \([\text{Yb}_2(hfac)_6(L)]\cdot 2(\text{C}_7\text{H}_{16})\) and \([\text{Dy}_2((+)\text{facam})_6(L)]\cdot 2(\text{C}_7\text{H}_{14})\) were obtained using the bridging bis(1,10-phenantro[5,6-b])tetraathiafulvalene triad and their structural properties determined by X-ray diffraction on single crystal. Both compounds displayed field-induced Single-Molecule Magnet behavior with the magnetization mainly relaxing through a Raman process. The \(\text{Dy}^{III}\) derivative displayed also a slow magnetic relaxation of its magnetization in zero applied dc field with a Quantum Tunneling of the Magnetization identified at low temperature. Light irradiation of the visible HUMO→LUMO intra-ligand charge transfer of the \(\text{Yb}^{III}\) derivative induced Near-Infrared emission of the metal center. The energy gap between the two lowest energy emissive contributions (260 cm\(^{-1}\)) confirmed the non-significant Orbach process in the magnetic relaxation. In both compounds, the TTF-based bridging triad is in its neutral form nevertheless its redox-activity opens the route to redox switching of the optical and magnetic properties.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/inorganics9070050/s1: CIF and CIF-checked file; Figure S1: ORTEP view of the asymmetric unit for (1) 2(C7H16). Thermal ellipsoids are drawn at 30% probability. Hydrogen atoms and solvent molecules of crystallization are omitted for clarity; Figure S2: ORTEP view of the asymmetric unit for (2) 2(C6H14). Thermal ellipsoids are drawn at 30% probability. Hydrogen atoms and solvent molecules of crystallization are omitted for clarity; Figure S3: Cyclic voltammograms of 1 in CH3Cl2 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 S4: Cyclic voltammery of 2 in CH3Cl2 at a scan rate of 100 mVs\(^{-1}\). The potentials were measured versus a saturated calomel electrode (SCE) with Pt wire as the counter electrodes; Figure S5: Experimental UV-visible absorption spectrum at room temperature of 1 in CH3Cl2 solution (C = 4 × 10\(^{-5}\) mol L\(^{-1}\)); Figure S6: Experimental UV-visible absorption spectrum at room temperature of 2 in CH3Cl2 solution (C = 4 × 10\(^{-5}\) mol L\(^{-1}\)); Figure S7: Thermal dependence of the \(\chi_M T\) product between 2 and 300 K for 1. In the inset, the field dependence of the magnetization at 2 K for 1; Figure S8: Thermal dependence of the \(\chi_M T\) product between 2 and 300 K for 2. In the inset, the field dependence of the magnetization at 2 K for 2; Figure S9: In-phase (left) and out-of-phase (right) components of the ac magnetic susceptibility for 1 at 2 K under a DC magnetic field from 0 to 3000 Oe; Figure S10: Frequency dependence of the in-phase component of the magnetic susceptibility under an applied magnetic field of 1000 Oe between 2 and 10 K for 1; Figure S11: Frequency dependence of the in-phase (\(\chi_{M'}\)) and out-of-phase (\(\chi_{M''}\)) components of the ac susceptibility measured on powder at 2 K and 1000 Oe with the best-fitted curves (red lines) for 1; Figure S12: Normalized Cole–Cole plot for 1 at several temperatures between 2 and 4.25 K under an applied magnetic field of 1000 Oe. Black lines are the best-fitted curves; Figure S13: Temperature dependence of the relaxation time for 1 at 1000 Oe in the temperature range 2–4.25 K. Green dashed lines corresponds to the thermally activated contribution (Orbach process with the parameters given in the main text); Figure S14: Temperature dependence of the relaxation time for 1 at 1000 Oe in the temperature range of 2–4.25 K with the best-fitted curve (full black line) with the combination Orbach + Raman processes. The Orbach and Raman contributions to the relaxation time are respectively represented in dashed green line and dashed red line (the parameters are given in the main text); Figure S15: Temperature dependence of the relaxation time for 1 at 1000 Oe in the temperature range of 2–4.25 K with the best-fitted curve (full black line) with the combination Raman + Direct processes. The Raman and Direct contributions to the relaxation time are respectively represented in dashed red line and dashed blue line (the parameters are given in the main text); Figure S16: Frequency dependence of the in-phase component of the magnetic susceptibility under a zero applied magnetic field between 2 and 15 K for 2; Figure S17: Frequency dependence of the in-phase (\(\chi_{M'}\)) and out-of-phase (\(\chi_{M''}\)) components of the ac susceptibility measured on powder at 2 K in zero applied dc field with the best-fitted curves (red lines) for 2; Figure S18: Normalized Cole–Cole plot for 2 at several temperatures between 2 and 15 K under a zero applied magnetic field.
Black lines are the best-fitted curves; Figure S19: In-phase (left) and out-of-phase (right) components of the ac magnetic susceptibility for 2 at 2 K under a DC magnetic field from 0 to 2400 Oe; Figure S20: Frequency dependence of the in-phase component of the magnetic susceptibility under a 1200 Oe applied magnetic field between 2 and 13 K for 2; Figure S21: Frequency dependence of the in-phase ($\chi_M'$) and out-of-phase ($\chi_M''$) components of the ac susceptibility measured on powder at 2 K and 1200 Oe with the best-fitted curves (red lines) for 2; Figure S22: Normalized Cole–Cole plot for 2 at several temperatures between 2 and 13 K under a 1200 Oe applied magnetic field. Black lines are the best-fitted curves; Figure S23: Temperature dependence of the relaxation time for 2 at 1200 Oe in the temperature range 2–13 K. Green line corresponds to the thermally activated contribution (Orbach process with the parameters given in the main text). Figure S24: Temperature dependence of the relaxation time for 2 at 0 Oe (full gray circles) and 1200 Oe (open gray circles) in the temperature range of 2–15 K with the best-fitted curve (full black line) with the combination Orbach + Raman + QTM (for H = 0 Oe) processes. The Orbach, Raman, and QTM contributions to the relaxation time are respectively represented in the dashed green line, dashed red line, and dashed orange line (the parameters are given in the main text). Table S1: X-ray crystallographic data for (1)-2(C7H16) and (2)-2(C6H14); Table S2: SHAPE analysis of the coordination polyhedra around the lanthanide in the polymeric compounds; Table S3: Selected bond lengths for (1)-2(C7H16) and (2)-2(C6H14) in Å; Table S4: Oxidation potentials of the complexes 1 and 2 (V vs. SCE, nBu4NPF6, 0.1 M in CH2Cl2 at 100 mV.s⁻¹); Table S5: Best fitted parameters ($\chi_T$, $\chi_S$, $\tau$ and $\alpha$) with the extended Debye model for compound 1 at 2 K in the magnetic field range 200–3000 Oe; Table S6: Best fitted parameters ($\chi_T$, $\chi_S$, $\tau$ and $\alpha$) with the extended Debye model for compound 1 at 1000 Oe in the temperature range 2–4.25 K; Table S7: Best fitted parameters ($\chi_T$, $\chi_S$, $\tau$ and $\alpha$) with the extended Debye model for compound 2 at 0 Oe in the temperature range 2–14 K; Table S8: Best fitted parameters ($\chi_T$, $\chi_S$, $\tau$ and $\alpha$) with the extended Debye model for compound 2 at 2 K in the magnetic field range 200–2400 Oe; Table S9: Best fitted parameters ($\chi_T$, $\chi_S$, $\tau$ and $\alpha$) with the extended Debye model for compound 2 at 1200 Oe in the temperature range 2–13 K.

Author Contributions: B.L. made the synthesis, J.F.G., C.A.M., and O.C. performed the magnetic measurements and interpreted them, V.D. performed the single-crystal X-ray diffraction study, F.P. elaborated and supervised the project, O.C. and F.P. participated in the writing process of the manuscript. All authors have read and agreed to the published version of the manuscript.

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Abbreviations
The following abbreviations are used in this manuscript

- **SMM**: Single-Molecule Magnet
- **QTM**: Quantum Tunneling of the Magnetization
- **Hfac**⁻: 1,1,5,5,5-hexafluoroacetylacetonato anion
- **Facam**⁻: 3-trifluoro-acetyl-(+)-camphorato anion
- **Phen**: 1,10-phenantroline
- **TTF**: Tetrathiafulvalene
- **HOMO**: Highest Occupied Molecular Orbital
- **LUMO**: Lowest Unoccupied Molecular Orbital
- **ILCT**: Intra-Ligand Charge Transfer
- **CH2Cl2**: Dichloromethane
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