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Design of $\text{Fe}^{\text{III}}\text{--Ln}^{\text{III}}$ binuclear complexes using compartmental ligands: synthesis, crystal structures, magnetic properties, and *ab initio* analysis[†]

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Three new binuclear $\text{Fe}^{\text{III}}\text{--Ln}^{\text{III}}$ complexes, with similar structures, have been synthesized using the end-off compartmental proligand, H_2valpn , resulting from the Schiff condensation between *o*-vanillin and 1,3-propylenediamine: $[\text{Fe}^{\text{III}}\text{Ln}^{\text{III}}(\text{valpn})(\text{hfac})_2(\text{OAc})\text{Cl}]$ ($\text{Ln} = \text{Gd}, \text{Tb}, \text{Dy}$; $\text{hfac}^- = \text{hexafluoroacetylacetone}$; $\text{AcO}^- = \text{acetate}$). The metal ions are triply bridged by two phenoxido oxygen atoms and by the acetato ligand (*syn-syn* bridging mode). Among all these compounds the $\text{Fe}^{\text{III}}\text{--Ln}^{\text{III}}$ exchange interaction was found to be ferromagnetic ($J_{\text{FeGd}} = +1.004(4) \text{ cm}^{-1}$; $\mathbf{H} = -\mathbf{JS}_{\text{Fe}}\mathbf{S}_{\text{Gd}}$). The $[\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}]$ derivative shows field-induced slow relaxation of the magnetization. The interpretation of the magnetic properties for the $[\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}]$, $[\text{Fe}^{\text{III}}\text{Tb}^{\text{III}}]$ and $[\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}]$ is done through *ab initio* calculations. It was concluded that the temperature behavior of the relaxation time in the $[\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}]$ derivative is due to dominant Orbach relaxation processes between states at the opposite sides and opposite heights of the blocking barrier. From this understanding, the relaxation mechanism for similar systems where a transition metal is in exchange coupling with a strong anisotropic lanthanide ion is elucidated, which could provide a strategy to synthesize other binuclear complexes with better magnetic properties.

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Introduction

Oligonuclear heterometallic complexes, combining 3d and 4f metal ions, are intensively investigated in molecular magnetism.¹ First of all, such compounds are useful models illuminating the factors that influence the nature and strength

of the 3d-4f magnetic couplings. Then, numerous 3d-4f complexes show Single Molecule Magnet (SMM) behavior and emphasize the role played by the magnetic anisotropy of both metal ions, as well as of the 3d-4f exchange interaction in the slow relaxation of the magnetization phenomena.² Theoretical and experimental studies indicate that the strength of the 3d-4f interaction has direct consequences on the quantum tunneling of magnetization (QTM).³ Concerning the $\text{Fe}^{\text{III}}\text{--Ln}^{\text{III}}$ pair, numerous clusters with various nuclearities have been reported. Most of these compounds have been obtained by reacting iron(III) and lanthanide(III) salts with aminoalcohols.⁴ Dodecanuclear complexes, $[\text{Fe}_6^{\text{III}}\text{Ln}_6^{\text{III}}]$, have been assembled by reacting pivalato iron(III) and lanthanide(III) precursors with phosphonic acids.⁵ Several $[\text{Ln}^{\text{III}}\text{Fe}_3^{\text{III}}]$ tetranuclear complexes with trichloroacetato bridging ligands have been reported by Powell *et al.*⁶ A high-nuclearity cluster, $[\text{Fe}_{12}^{\text{III}}\text{Sm}_{12}^{\text{III}}]$, the first Sm-based SMM, has been aggregated by benzoate ligands.⁷ Other $\text{Fe}^{\text{III}}\text{--Ln}^{\text{III}}$ oligonuclear complexes have been obtained using homo- and heteroleptic cyanido metalloligands, $([\text{Fe}^{\text{III}}(\text{CN})_6]^{3-}$ and $[\text{Fe}^{\text{III}}(\text{bipy})(\text{CN})_4]^-$).⁸ Tetranuclear cyanido-bridged $[\text{Fe}_2^{\text{III}}\text{Ln}_2^{\text{III}}]$ molecular squares have been assembled from $[\text{Fe}^{\text{III}}\{\text{HB}(\text{pz})_3\}(\text{CN})_3]^-$ and lanthanide ions carrying bidentate chelating ligands.⁹ By employing bis-calix[4]arene

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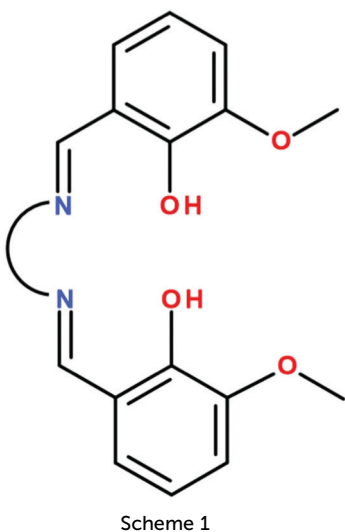
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Scheme 1

as a ligand, Brechin, Dalgarno *et al.* have obtained several heterometallic clusters, including a nonanuclear complex, $[\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}]^{10}$.

The number of binuclear $[\text{Fe}^{\text{III}}\text{Ln}^{\text{III}}]$ complexes is limited to a few examples: (i) cyanido-bridged complexes resulting by attaching a $[\text{Fe}^{\text{III}}(\text{CN})_6]^{3-}$ group to lanthanide ions coordinated by solvent molecules (H_2O , DMF, DMSO) or other organic ligands;¹¹ (ii) an $[\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}]$ complex showing SMM behavior which results from the self-assembly process involving $[\text{Fe}^{\text{III}}(\text{bpca})_2]^+$ as a metalloligand and dysprosium(III) nitrate ($\text{bpca} = \text{bis}(2\text{-pyridylcarbonyl})\text{amine}$);¹² (iii) a tris-phenoxido bridged binuclear $[\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}]$ complex synthesized from a bicompartimental ligand derived from *o*-vanillin and a tripodal amine.¹³ The Schiff-base bicompartimental ligands generated by *o*-vanillin, valXn^{2-} (Scheme 1), have been widely employed to design strictly binuclear 3d–4f complexes, the 3d metal ion being in almost all cases divalent and occupies the small inner compartment.^{14,15} Except the aforementioned $[\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}]$ complex,¹³ no other binuclear 3d–4f complexes with a trivalent 3d metal ion and bicompartimental ligands are known. The main reason is that, in the step-wise synthetic process, the $[\text{M}^{\text{III}}(\text{valXn})]^+$ intermediate bears a positive charge. Electrostatic repulsion between the metalloligand and the lanthanide cations hinders the formation of the heteronuclear binuclear complexes. Using valXn^{2-} ligands and Fe^{III} and Gd^{III} ions, it has been shown that, instead of binuclear complexes, tetranuclear $[\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}_2]$ species are assembled, the two $\{\text{Fe}^{\text{III}}(\text{valXn})\text{Ln}^{\text{III}}\}$ moieties being connected by an oxido bridge established between the two Fe^{III} ions.¹⁵ In these compounds, the weak $\text{Fe}^{\text{III}}\text{–Gd}^{\text{III}}$ exchange interaction is overwhelmed by the strong antiferromagnetic $\text{Fe}^{\text{III}}\text{–O–Fe}^{\text{III}}$ coupling. Finally, we recall that a non-compartmental ligand, obtained by reacting *o*-vanillin with *o*-amino-phenol, leads to trinuclear $[\text{Fe}^{\text{III}}_2\text{Ln}^{\text{III}}]$ complexes.¹⁶ Herein we report on a new general synthetic approach that allows an easy access to binuclear $\text{Fe}^{\text{III}}\text{–Ln}^{\text{III}}$ complexes.

Besides the new general synthetic approach, magnetic characterization of this first family of binuclear 3d–4f complexes

with a trivalent 3d metal ion and bicompartimental ligands is also investigated both in experiments and by using a state-of-the-art *ab initio* method. Interestingly, we found that exchange interactions between metal ions in these compounds are all ferromagnetic. Dynamic magnetic susceptibility measurements also revealed that among these compounds, only $[\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}]$ shows a field-induced slow relaxation of the magnetization. Based on *ab initio* calculations, physical insight into the electronic structure as well as relaxation mechanisms in this new family of $\text{Fe}^{\text{III}}\text{–Ln}^{\text{III}}$ complexes will be presented, which not only elucidates the magnetic properties in this new family of complexes but also may provide a recommendation to design other 3d–4f complexes with better magnetic properties in the future.

Theoretical investigation of polynuclear compounds is usually performed using phenomenological models based *e.g.* on the parameterization of the on-site magnetic anisotropy and zero-field splitting and of the inter-site magnetic interactions.^{17–20} Another phenomenological method involves the description of the zero-field splitting of the ground state spin state of the polynuclear compound (*e.g.* ZFS of the ground $S = 10$ in the case of Mn12). Quantum thermodynamical methods allowing the simulation of the magnetic properties of large spin clusters are also commonly used.^{21–25} The latter methods require experimental determination of the ground state spin state from low-temperature magnetic measurements. Broken-Symmetry DFT calculations are commonly used for the description of the inter-site magnetic exchange.^{26–28} Usually, it provides reasonable values for the magnetic interaction. This method was further expanded to extract the anisotropic magnetic exchange interactions. Fully *ab initio* computational methods are scarcely applied for the description of the electronic structure and properties of polynuclear compounds, because of the computational difficulties involved in their application: (a) large size of the active space; (b) a large number of roots required to be computed and (c) difficulties in molecular orbital optimizations for such states, which prevent the required level of numerical accuracy from being obtained ($<0.1 \text{ cm}^{-1}$). For these reasons, most used electron correlation methods are applied to binuclear compounds.²⁹ DMRG-based methods are used for describing large active spaces,^{30–33} but given the large number of roots required for the description of the spin-orbit coupling of lanthanides – their application for such compounds is yet limited.

To overcome the above computational difficulties, a semi-*ab initio* approach was proposed and successfully used for many cases.^{34–37} The proposed method involves (a) calculations of the on-site electronic structure and properties using high-level *ab initio* methods, (b) estimation of the inter-site magnetic exchange using the BS-DFT method,³⁸ and (c) evaluation of the energy spectra and properties of the entire polynuclear compound using the data obtained at (a) and (b) with a fine-tuning of the magnetic exchange in order to describe the available magnetization data. This method was successfully used for the description of the magnetism in a series of polynuclear compounds and offers the best balance between the computational cost and accuracy.

Experimental

Materials

All starting materials were reagent grade and used without purification. The syntheses of $[\text{Fe}^{\text{III}}(\text{valpn})(\text{H}_2\text{O})\text{Cl}]^{15b}$ and $[\text{Ln}^{\text{III}}(\text{hfac})_2(\text{Hhfac})(\text{OAc})(\text{H}_2\text{O})_2]^{39}$ ($\text{Ln} = \text{Gd}^{\text{III}}$, Tb^{III} , Dy^{III}) have been carried out as described in the respective papers.

Syntheses

$[\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}(\text{valpn})(\text{hfac})_2(\text{OAc})\text{Cl}]$ (1). $[\text{Gd}^{\text{III}}(\text{hfac})_2(\text{Hhfac})(\text{OAc})(\text{H}_2\text{O})_2]$ (97 mg, 0.111 mmol) was added under stirring to an acetonitrile solution (50 mL) of $[\text{Fe}^{\text{III}}(\text{valpn})(\text{H}_2\text{O})\text{Cl}]$ (50 mg, 0.111 mmol). The mixture is left under continuous stirring for approximately 30 minutes. The resulting dark violet solution was filtered and allowed to become concentrated by slow evaporation for several days in a hood at room temperature. Dark violet single crystals of compound **1** formed after several days and were collected and washed with a very small amount of acetonitrile. Yield: *ca.* 80%. Selected IR data (KBr pellets/cm⁻¹): 3474(w,br), 2913(w), 1656(s), 1627(m), 1558(m), 1470(m), 1254(s), 1215(s), 1114(s), 948(w), 798(w), 738(m), 659(m), 584(w). Elemental analysis: calcd for $\text{C}_{31}\text{H}_{25}\text{ClF}_{12}\text{FeGdN}_2\text{O}_{10}$: C, 35.05; H, 2.37; N, 2.63; Found: C, 35.50; H, 2.59; N, 2.44.

$[\text{Fe}^{\text{III}}\text{Tb}^{\text{III}}(\text{valpn})(\text{hfac})_2(\text{OAc})\text{Cl}]$ (2). The procedure is similar to that of compound **1**, using $[\text{Tb}^{\text{III}}(\text{hfac})_2(\text{Hhfac})(\text{OAc})(\text{H}_2\text{O})_2]$ (97 mg, 0.111 mmol). Dark violet single crystals of **2** formed after several days and were collected and washed with a very small amount of acetonitrile. Yield: *ca.* 80%. Selected IR data (KBr pellets/cm⁻¹): 3422(w,br), 2910(w), 1656(s), 1627(m), 1558(m), 1496(m), 1471(m), 1254(s), 1208(s), 1145(s), 949(w), 798(w), 739(m), 660(w), 584(w). Elemental analysis: calcd for $\text{C}_{31}\text{H}_{25}\text{ClF}_{12}\text{FeTbN}_2\text{O}_{10}$: C, 35.01; H, 2.36; N, 2.63; Found: C, 35.36; H, 2.62; N, 2.74.

$[\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}(\text{valpn})(\text{hfac})_2(\text{OAc})\text{Cl}]$ (3). The procedure is similar to that of compound **1**, using $[\text{Dy}^{\text{III}}(\text{hfac})_2(\text{Hhfac})(\text{OAc})(\text{H}_2\text{O})_2]$ (98 mg, 0.111 mmol). Dark violet single crystals of compound **3** formed after several days and were collected and washed with a

very small amount of acetonitrile. Yield: *ca.* 80%. Selected IR data (KBr pellets/cm⁻¹): 3431(w,br), 2910(w), 1657(s), 1628(m), 1559(m), 1471(m), 1255(s), 1208(s), 1145(s), 949(w), 856(w), 739(w), 660(m), 584(m). Elemental analysis: calcd for $\text{C}_{31}\text{H}_{25}\text{ClF}_{12}\text{FeDyN}_2\text{O}_{10}$: C, 34.88; H, 2.36; N, 2.62; Found: C, 35.23; H, 2.53; N, 2.81.

Physical measurements

IR spectra were recorded on a JASCO FTIR 4100 spectrometer in the 4000–400 cm⁻¹ range, with samples prepared as KBr pellets. The X-ray powder diffraction measurements were carried out on a Proto AXRD Benchtop using Cu-K α radiation with a wavelength of 1.54059 Å in the range 5–35° (2 θ). The absorption spectra were obtained on a JASCO V-670 spectrophotometer with solid sample accessory, in the 1800–250 nm range. Magnetic susceptibility measurements were performed on a Quantum Design SQUID MPMS-XL magnetometer at temperatures between 1.8 and 300 K. All data were corrected for the contribution of the sample holder and the diamagnetism of the samples estimated from Pascal's constants.

Crystal structure determination and refinement

X-Ray diffraction measurements for compounds **1**, **2** and **3** were performed on a STOE IPDS II diffractometer operating with Mo-K α ($\lambda = 0.71073$ Å) X-ray tube with graphite monochromator. The structures were solved by direct methods and refined by full-matrix least squares techniques based on F^2 . The non-H atoms were refined using anisotropic displacement parameters. Calculations were performed using the SHELX-2013 crystallographic software package. The structures were solved by direct methods using the SHELXS structure solution program. The H atoms attached to carbon were introduced in idealized positions using the riding model. A summary of the crystallographic data and the structure refinement for crystals **1–3** is given in Table 1. CCDC reference numbers: 1916564–1916566.†

Table 1 Crystallographic data, details of data collection and structure refinement parameters for **1**, **2** and **3**

| Compounds | 1 | 2 | 3 |
|-------------------------------------------------------|------------------------------------------------------------------------|------------------------------------------------------------------------|------------------------------------------------------------------------|
| Formula | $\text{C}_{31}\text{H}_{25}\text{ClF}_{12}\text{FeGdN}_2\text{O}_{10}$ | $\text{C}_{31}\text{H}_{25}\text{ClF}_{12}\text{FeTbN}_2\text{O}_{10}$ | $\text{C}_{31}\text{H}_{25}\text{ClF}_{12}\text{FeDyN}_2\text{O}_{10}$ |
| Molecular weight | 1062.08 | 1063.75 | 1067.33 |
| Temperature (K) | 293(2) | 293(2) | 293(2) |
| Crystal system | Monoclinic | Monoclinic | Monoclinic |
| Space group | $P2_1/a$ | $P2_1/a$ | $P2_1/a$ |
| a (Å) | 16.9739(2) | 16.9716(2) | 16.9946(2) |
| b (Å) | 12.6044(3) | 12.5632(3) | 12.5891(3) |
| c (Å) | 18.5271(4) | 18.5399(4) | 18.5684(4) |
| β (deg) | 98.272(5) | 98.459(5) | 98.404(5) |
| V (Å ³) | 3922.56(14) | 3910.03(14) | 3929.99(14) |
| Z | 4 | 4 | 4 |
| D_{calc} (g cm ⁻³) | 1.798 | 1.807 | 1.804 |
| F (000) | 2080 | 2084 | 2088 |
| Wavelength (Å) | 0.71073 | 0.71073 | 0.71073 |
| Final R_1^a , wR_2^b [$I > 2\sigma(I)$] | 0.0412, 0.0967 | 0.0420, 0.0924 | 0.0803, 0.1383 |
| R_1^a , wR_2^b (all data) | 0.0618, 0.1093 | 0.0624, 0.1138 | 0.1762, 0.1913 |
| Goodness-of-fit on F^2 | 1.023 | 1.111 | 1.130 |
| Largest peak in final difference (e Å ⁻³) | -0.837, 0.762 | -1.332, 1.069 | -0.982, 0.856 |

^a $R_1 = \sum |F_0| - |F_c| |/ \sum |F_0|$. ^b $wR_2 = [\sum w(F_0^2 - F_c^2)^2 / \sum w(F_0^2)^2]^{1/2}$; $w = 1/[\sigma^2(F_0^2) + (aP)^2 + bP]$ where $P = [\max(F_0^2, 0) + 2F_c^2]/3$.

Ab initio calculations

The *ab initio* calculations for each compound were carried out using CASSCF/RASSI-SO/SINGLE_ANISO approach implemented using the MOLCAS 8.2 program package.⁴⁰ The atomic coordinates for each compound were taken from structural data. The ANO-RCC-VDZP basis set was used for all atoms except C, F and H, for which the ANO-RCC-VDZ basis set was employed. The active space in the CASSCF calculations included 7 electrons in 7 orbitals for Gd^{III}, 8 electrons in 7 orbitals for Tb^{III}, 9 electrons in 7 orbitals for Dy^{III} and 5 electrons in 5 orbitals for Fe^{III} (4f orbitals of Gd^{III}, Tb^{III}, Dy^{III} and 3d orbital of Fe^{III}). On the basis of the resulting spin-orbit multiplets, *g*-factors and the orientation of the magnetic axes of doublet states at the metal sites, as well as the zero-field splitting tensors (ZFS parameters and the orientation of the anisotropy axes) of Gd^{III}, Tb^{III}, and Dy^{III} centres three complexes have been calculated *ab initio* within the CASSCF/RASSI/SINGLE_ANISO approach using the MOLCAS 8.2 package (see the ESI[†] for the details).^{34,40–45} Meanwhile, the dipolar coupling parameters were calculated directly from the *ab initio* results for the single-ion anisotropy of individual metal sites. Exchange coupling parameters were then determined by fitting the experimental magnetic susceptibility and magnetisation data using the POLY_ANISO module of the MOLCAS 8.2 package (see the ESI[†] for the details).^{34,40–45}

Results and discussion

The new compounds, $[\text{Fe}^{\text{III}}\text{Ln}^{\text{III}}(\text{valpn})(\text{hfac})_2(\text{OAc})\text{Cl}]$, have been obtained by reacting two neutral precursors: $[\text{Fe}^{\text{III}}(\text{valpn})(\text{H}_2\text{O})\text{Cl}]^{15b}$ and $[\text{Ln}^{\text{III}}(\text{hfac})_2(\text{Hhfac})(\text{OAc})(\text{H}_2\text{O})_2]$.³⁹ The hfac[–] ligands increase the Lewis acidity of the Ln^{III} ions, facilitating their interaction with the oxygen atoms from the open compartment of the iron(III) precursor. Moreover, the reaction between two neutral precursors will favor the assembly of the desired heterobinuclear complexes. For this study, we have chosen the following lanthanide ions: Gd^{III} 1, Tb^{III} 2, and Dy^{III} 3. This synthetic approach appears to be successful. It clearly indicates the principles to be followed to obtain strictly binuclear $[\text{Fe}^{\text{III}}\text{Ln}^{\text{III}}]$ complexes in a rational manner, paving the way for other syntheses. The diffuse reflectance spectra of the three compounds (Fig. 1) show the characteristic ligand-to-metal charge transfer band for Fe^{III}–phenoxido systems, which obscure the spin forbidden, and therefore weak, d–d bands of the $[\text{Fe}^{\text{III}}\text{N}_2\text{O}_3\text{Cl}]$ chromophore. (1: 554, 2: 555, 3: 535 nm). The purity of the crystalline phases was proved by powder X-ray diffraction (Fig. S1, ESI[†]).

Description of the structures

Compounds 1–3 are isostructural, therefore only the crystal structure of the $[\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}]$ derivative will be discussed here (Fig. 2). The metal ions are triply bridged by two phenoxido oxygen atoms and by the acetato ligand (*syn-syn* bridging mode). The Fe^{III} ion is hosted into the inner compartment, being coordinated by two nitrogen and two oxygen atoms from the organic ligand (Fe–N1 = 2.079(5); Fe–N2 = 2.091(5); Fe–O1 =

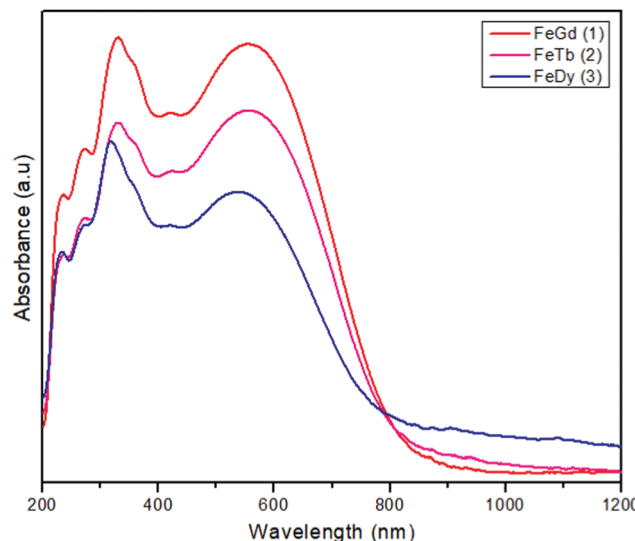


Fig. 1 Diffuse reflectance spectra for compounds 1 (red), 2 (violet) and 3 (blue).

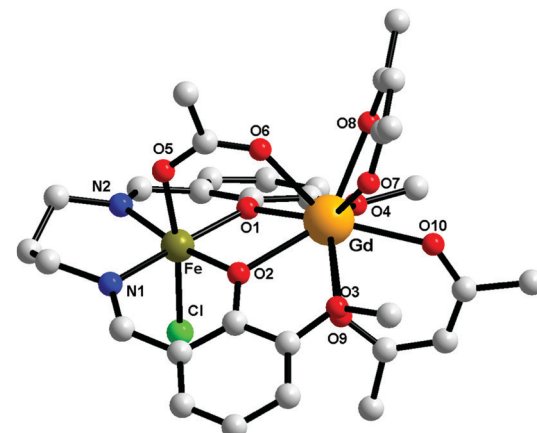


Fig. 2 Perspective view of complex 1. For clarity the hydrogen and fluorine atoms have been removed.

1.975(4); Fe–O2 = 1.978(3) Å); and by a chlorido ligand and an acetato oxygen atom into the apical positions (Fe–O5 = 2.050(4); Fe–Cl = 2.306(15) Å), with an octahedral geometry. The Ln^{III} ion shows a coordination number of 9, being coordinated only by oxygen atoms (4 from the two hfac[–] ligands, 4 from the open compartment of the Schiff base, and one from the bridging acetato ligand) with distances varying between 2.373(4) and 2.628(4) Å. The coordination geometry of Gd^{III} can be described best as a muffin (MFF-9)-like type with a CShM value of 0.585 (Table S1, ESI[†]).⁴⁶ The intramolecular Fe···Gd distance is 3.449 Å, while the shortest intermolecular Fe···Fe distance is 7.673 Å; the shortest intermolecular Gd···Gd distance is 10.296 Å and the shortest intermolecular Fe···Gd distance is 9.534 Å. The values of the two Gd–O(phenoxido)–Fe angles are Fe–O1–Gd = 103.41(14) and Fe–O2–Gd = 103.87(13)°. Selected bond distances and angles for the three compounds are collected in Table 2. We recall that the structural motif is

Table 2 Selected bond lengths (Å) and angles (°) for compounds **1**, **2**, and **3**

| Compounds | Distances (Å) | | Angles (°) | | | |
|-----------|---------------|------------|------------|------------|----------|------------|
| 1 | Gd-O10 | 2.362(4) | O10-Gd-O7 | 74.77(15) | O1-Fe-O5 | 92.37(14) |
| | Gd-O7 | 2.365(4) | O10-Gd-O6 | 141.49(14) | O2-Fe-N1 | 88.93(19) |
| | Gd-O6 | 2.365(4) | O7-Gd-O6 | 70.67(14) | O1-Fe-N1 | 172.40(18) |
| | Gd-O8 | 2.373(4) | O10-Gd-O8 | 80.72(14) | O5-Fe-N1 | 84.02(17) |
| | Gd-O1 | 2.390(3) | O7-Gd-O8 | 72.48(13) | O2-Fe-N2 | 171.62(17) |
| | Fe-O1 | 1.978(3) | O6-Gd-O8 | 73.22(13) | O1-Fe-N2 | 88.83(17) |
| | Fe-O5 | 2.050(4) | O10-Gd-O1 | 137.75(13) | O5-Fe-N2 | 86.71(16) |
| | Fe-N1 | 2.079(5) | O7-Gd-O1 | 143.26(13) | N1-Fe-N2 | 97.6(2) |
| | Fe-N2 | 2.091(5) | O6-Gd-O1 | 72.79(12) | O2-Fe-Cl | 95.34(11) |
| | Fe-Cl | 2.3062(15) | O8-Gd-O1 | 93.58(12) | O1-Fe-Cl | 94.32(11) |
| | Tb-O10 | 2.344(4) | O10-Tb-O7 | 74.43(16) | O1-Fe-O2 | 83.95(15) |
| | Tb-O7 | 2.350(4) | O10-Tb-O8 | 80.31(16) | O1-Fe-O5 | 92.54(15) |
| 2 | Tb-O8 | 2.352(4) | O7-Tb-O8 | 73.00(15) | O2-Fe-O5 | 88.59(16) |
| | Tb-O6 | 2.352(4) | O10-Tb-O6 | 141.12(15) | O1-Fe-N1 | 172.50(19) |
| | Tb-O1 | 2.381(4) | O7-Tb-O6 | 70.90(15) | O2-Fe-N1 | 89.2(2) |
| | Fe-O2 | 1.975(4) | O8-Tb-O6 | 73.27(15) | O5-Fe-N1 | 84.20(18) |
| | Fe-O5 | 2.055(4) | O10-Tb-O1 | 137.62(14) | O1-Fe-N2 | 88.97(18) |
| | Fe-N1 | 2.078(5) | O7-Tb-O1 | 143.83(14) | O2-Fe-N2 | 171.39(17) |
| | Fe-N2 | 2.081(5) | O8-Tb-O1 | 93.67(14) | O5-Fe-N2 | 86.82(17) |
| | Fe-Cl | 2.3052(16) | O6-Tb-O1 | 73.09(14) | N1-Fe-N2 | 97.6(2) |
| | Dy-O9 | 2.318(13) | O9-Dy-O6 | 141.4(4) | O2-Fe-N1 | 171.2(5) |
| | Dy-O6 | 2.336(11) | O9-Dy-O7 | 74.6(5) | O1-Fe-N1 | 88.5(5) |
| | Dy-O7 | 2.344(11) | O6-Dy-O7 | 70.7(4) | N2-Fe-N1 | 98.2(6) |
| | Dy-O8 | 2.348(11) | O9 Dy-O8 | 80.5(4) | O5-Fe-N1 | 84.6(5) |
| 3 | Dy-O2 | 2.355(10) | O6-Dy-O8 | 73.9(4) | O2-Fe-Cl | 95.1(3) |
| | Fe-O1 | 1.992(10) | O7-Dy-O8 | 73.1(4) | O1-Fe-Cl | 96.5(3) |
| | Fe-N2 | 2.059(13) | O9-Dy-O2 | 137.6(4) | N2-Fe-Cl | 90.1(4) |
| | Fe-O5 | 2.065(11) | O6-Dy-O2 | 73.1(3) | O5-Fe-Cl | 172.8(4) |
| | Fe-N1 | 2.081(14) | O7-Dy-O2 | 143.6(4) | N1-Fe-Cl | 89.7(4) |
| | Fe-Cl | 2.297(5) | O8-Dy-O2 | 93.2(4) | O2-Fe-Dy | 41.6(3) |

similar to the one described for $[M^{II}Ln^{III}(\text{valXn})(\text{hfac})_2(\text{OAc})]$ complexes,³⁹ but in our case an additional chlorido ligand is necessary in order to obtain a neutral species.

Magnetic properties

The new binuclear compounds are excellent candidates for magnetostructural correlations, which are scarce for the phenoxido-bridged Fe^{III} - Ln^{III} pairs. Let us start with compound **1**. The temperature dependence of $\chi_M T$ for this compound is shown in Fig. 3. As can be seen, the room temperature value of the $\chi_M T$ product ($12.26 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$) is close to the one of two magnetically independent Fe^{III} and Gd^{III} ions ($12.25 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$, considering $g_{\text{Fe}} = g_{\text{Gd}} = 2.00$). Below 100 K, $\chi_M T$ increases abruptly and reaches $20.05 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$ at 2 K, a value that corresponds to $S = 12/2$ for the ground state, resulting from the ferromagnetic interaction between Fe^{III} and Gd^{III} ions. The best fit to the data, using the Hamiltonian

$$H = -JS_{\text{Fe}}S_{\text{Gd}} + D_{\text{Fe}} \left[S_z^2 - \frac{1}{3}S_{\text{Fe}}(S_{\text{Fe}} + 1) \right] + E_{\text{Fe}} [S_x^2 - S_y^2] + g_{\text{Fe}}HS_{\text{Fe}} + g_{\text{Gd}}HS_{\text{Gd}} \quad (1)$$

leads to $J = +1.004(4) \text{ cm}^{-1}$ and $g_{\text{Fe}} = 1.979(3)$, $g_{\text{Gd}} = 2.0$ (fixed), $D_{\text{Fe}} = -0.51(8) |E_{\text{Fe}}| = 0.1(2) \text{ cm}^{-1}$. The value of the exchange parameter, J , is close to those found in trinuclear $[\text{Fe}^{III}\text{Gd}^{III}\text{Fe}^{III}]$ complexes, in which the metal ions are doubly bridged by phenoxido groups,¹⁶ but higher than the one found by Costes in his binuclear $[\text{Fe}^{III}\text{Gd}^{III}]$ complex with three phenoxido bridges ($J = +0.50 \text{ cm}^{-1}$),¹³ the intermolecular exchange interaction is assumed to be very weak because of the large intermolecular distances between the

magnetic centers (see above). The magnetization vs. field curves (inset to Fig. 3) can be well fitted using a Brillouin function for $S = 12/2$ and $g = 1.94$.

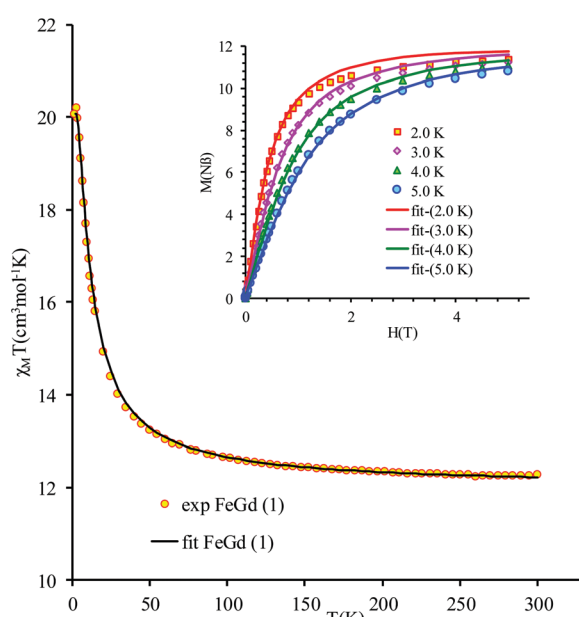


Fig. 3 $\chi_M T$ vs. T curve for compound **1**. Inset: Field dependence of the magnetization at various temperatures. The solid lines is the best simultaneous fit of magnetization and susceptibility data based on the anisotropic Hamiltonian (1).

Compounds **2** and **3** contain strongly anisotropic Tb^{III} and Dy^{III} ions, which have been chosen in order to induce the slow relaxation of the magnetization phenomenon. The $\chi_M T$ vs. T curves for **2** and **3** are shown Fig. 4. The corresponding room-temperature values 16.91 and $19.16 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$ are slightly higher than the sum of contributions of isolated magnetic ions in these complexes, 16.2 and $18.4 \text{ cm}^3 \text{ mol}^{-1} \text{ K}$, respectively (a straightforward calculation for $g_{\text{Fe}} = 2.00$, $S_{\text{Fe}} = 5/2$; $g_{\text{Tb}} = 3/2$, $J_{\text{Tb}} = 6$; $g_{\text{Dy}} = 4/3$, $J_{\text{Dy}} = 15/2$ can be done following ref. 47). Given small values of exchange parameters in these compounds (*vide infra*) and the fact that the magnetic susceptibility for individual magnetic centers in complexes are smaller than for corresponding isolated metal ions due to zero-field (crystal-field) splitting of their magnetic multiplets, the obtained higher values of room-temperature magnetic susceptibility suggest a rescaling down of the magnetic data for all complexes (see the ESI[†]).

The thermal dependences of $\chi_M T$ are similar: (i) a decrease from room temperature down to *ca.* 20 K , which is due to the depopulation of the Stark sublevels of the Ln^{III} ions; and (ii) an increase below 20 K , indicating the ferromagnetic coupling of the Fe^{III} and Ln^{III} ions. The M vs. H curves (insets to Fig. 4) confirm the expected magnetic anisotropy of these systems. Compounds **2** and **3** do not show frequency dependence of the magnetic susceptibility in zero static field, suggesting fast relaxation *via* QTM. The in-phase and out-of-phase magnetic susceptibilities of both compounds have been recorded at various static fields (0 – 3000 Oe), at 2 K (Fig. S2, ESI[†]), in order to find the optimal value of the field for the observation of slow relaxation of magnetization. For the terbium derivative, **2**, only weak frequency dependences of χ' and χ'' at 2500 Oe are observed (Fig. S3, ESI[†]). Conversely, the ac magnetic measurements indicate that the dysprosium derivative, **3**, (Fig. 5 and Fig. S4, Table S2, ESI[†]) shows slow relaxation of the magnetization under

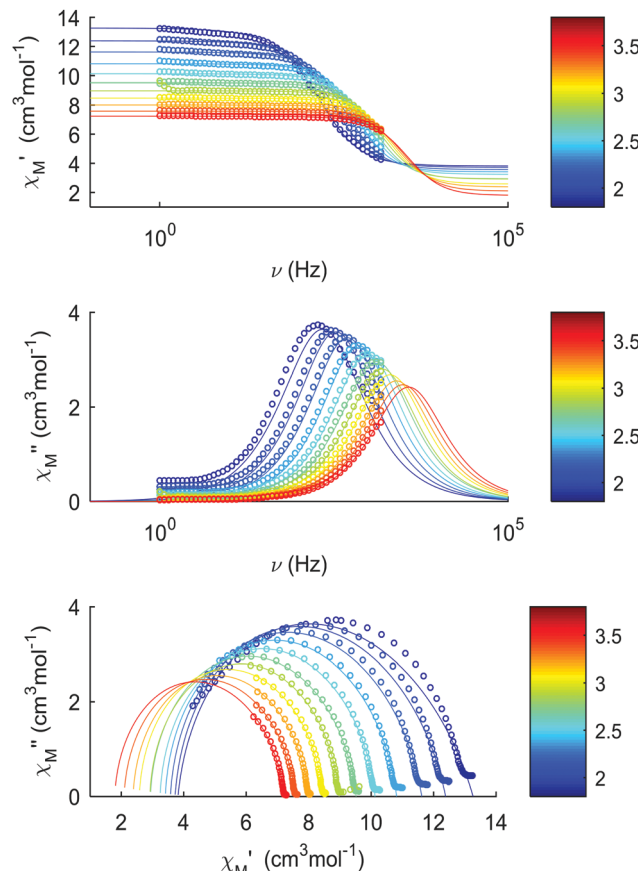


Fig. 5 Dynamic magnetic properties of compound **3**, recorded between 2 and 4 K : (a) in-phase component of the magnetic susceptibility, χ_M' , vs. frequency; (b) out-of-phase component of the magnetic susceptibility, χ_M'' , vs. frequency; (c) Cole–Cole plots.

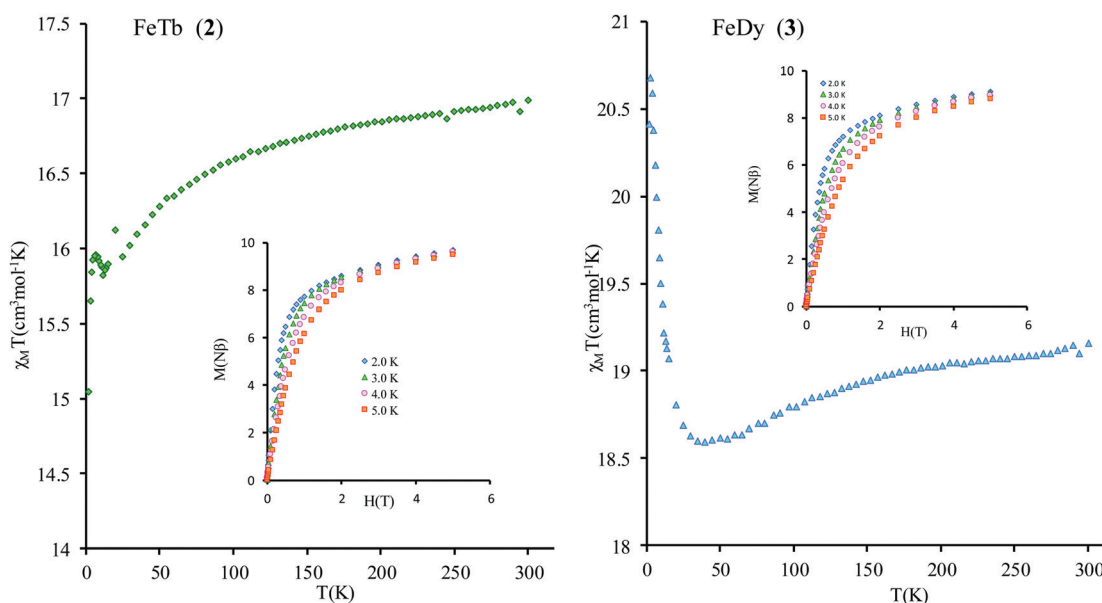


Fig. 4 $\chi_M T$ vs. T curve for compounds **2** (left) and **3** (right). Inset: M vs. H curves.

an applied static field (1000 Oe). Arrhenius plot of $\ln(\tau)$ vs. $1/T$ for compound **3** (Fig. 6) reveals that an approximately linear profile represents relatively well the relaxation curve, which indicates that the Orbach mechanism is dominant at an applied dc field. The corresponding effective blocking barrier to magnetization reversal from the fitting is around $U_{\text{eff}} \approx 9.72$ K if only the Orbach process is used and $U_{\text{eff}} \approx 8.39$ K if the Raman process is included. Detailed theoretical analyses of the magnetic relaxation behavior of all three complexes will be discussed in the following section.

Theoretical analysis

From *ab initio* results for **1**, which are listed in Tables S6–S9 and S20, S21 and the corresponding low-lying energy levels of each single metal ion plotted in Fig. S5–S10 (ESI†), as well as zero field splitting parameters D and E for Fe^{III} shown in Table 3, we see that the obtained ZFS splitting on Gd^{III} and Fe^{III} sites are relatively small, which is not surprising given that these ions are almost isotropic. In agreement with the fitted D and E parameters for Fe^{III} reported above, a negligible value for the

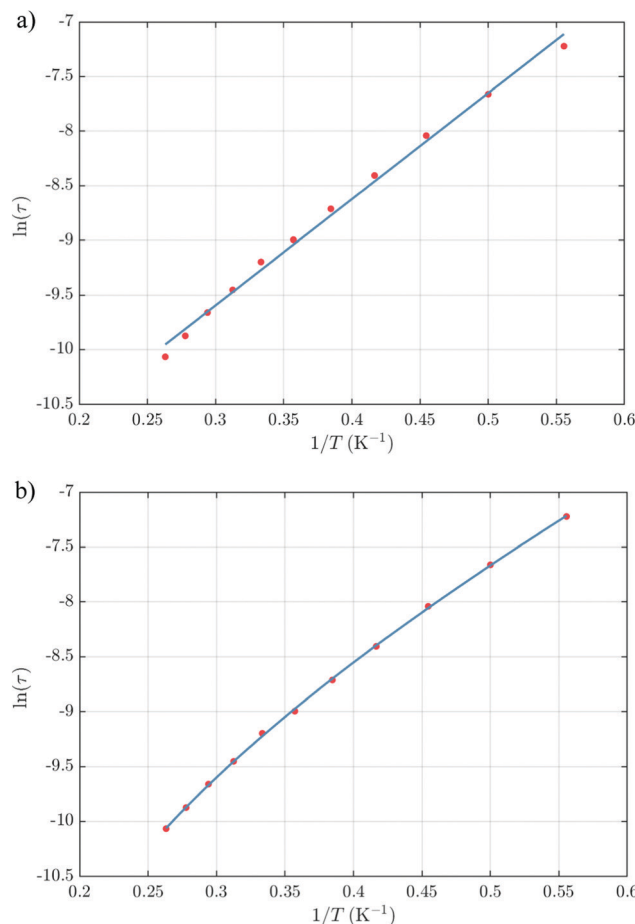


Fig. 6 Relaxation time of the magnetization $\ln(\tau)$ vs. T^{-1} plot for **3**. (a) Arrhenius fitting with $\ln(\tau) = \ln(\tau_0) + U_{\text{eff}}/T$ gives an effective blocking barrier of $U_{\text{eff}} \approx 9.72$ K and $\tau_0 \approx 3.69 \times 10^{-6}$ s if only the Orbach process is used. (b) Arrhenius fitting $\ln(\tau) = -\ln(\tau_0^{-1} \exp(-U_{\text{eff}}/T) + CT^n)$ gives an effective blocking barrier of $U_{\text{eff}} \approx 8.39$ K, $\tau_0 \approx 1.45 \times 10^{-5}$ s, $n = 4.16$, $C = 61.4 \text{ s}^{-1} \text{ K}^{-9}$ if the Raman process is included.

Table 3 The zero-field splitting parameters (cm^{-1}) at Fe^{III} in **1–3**

| Compound | D | E |
|--------------------------------------------------|--------|--------|
| $[\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}]$ | 0.2034 | 0.0109 |
| $[\text{Fe}^{\text{III}}\text{Tb}^{\text{III}}]$ | 0.2113 | 0.0103 |
| $[\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}]$ | 0.2129 | 0.0105 |

rhombic anisotropy parameter E is also observed. *Ab initio* calculations for **1** also reveal that the ZFS parameter D on Fe^{III} appears to be considerably smaller than the fitted one using the phenomenological model in eqn (1). The reason for this discrepancy is that in the latter no ZFS on Gd^{III} was taken into account, which was compensated by the artefactual increase of ZFS on Fe^{III} in order to fit the magnetic data. Another drawback of a pure phenomenological model is the opposite sign of D for Fe^{III} compared to the *ab initio* results. This is a known problem of a weak sensitivity of powder magnetic susceptibility to the sign of D , mentioned already by Kahn,⁴⁷ which makes it difficult to extract the sign of D from the fitting of magnetic data.

To elucidate the magnetic properties of the $[\text{Fe}^{\text{III}}\text{Ln}^{\text{III}}]$ series, we simulate the thermal dependences of $\chi_{\text{M}}T$ as well as the M vs. H curves in Fig. S10 and S11 (ESI†), for compound **1**. With an introduction of the intermolecular interaction $zJ = -0.0007 \text{ cm}^{-1}$, our *ab initio* results show that the simulated magnetic data for **1** are in good agreement with experiment. From the fitting, we also obtain the ferromagnetic exchange coupling $J = 0.898 \text{ cm}^{-1}$, which is consistent with the parameter used in the phenomenological model in eqn (1) but is slightly different with respect to the BS-DFT result ($J = 0.710 \text{ cm}^{-1}$).

Having all ZFS calculated, in Fig. 7, we show the low-lying spectrum of exchange/dipolar multiplets of the $[\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}]$ complex resulting from interaction between the ground multiplet $S = 5/2$ of the Fe^{III} site and $S = 7/2$ of the Gd^{III} site ($6 \times 8 = 48$

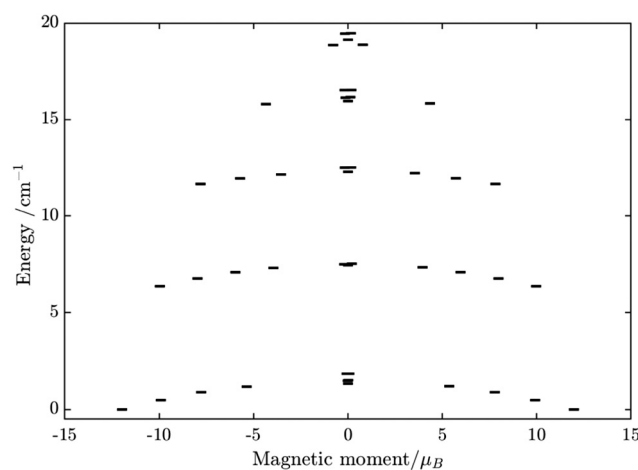


Fig. 7 Calculated low-lying spin-orbit manifolds of states arising from the magnetic interaction between the ground multiplet $S = 5/2$ of the Fe^{III} ion and the ground term ${}^8\text{S}$ of the Gd^{III} ion in the $[\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}]$ complex. The components of each doublet (black lines) are placed on the diagram according to their magnetic moment projection on the main magnetic axis of the ground doublet (with energy zero).

exchange states). These states subdivide into several groups of levels separated by the exchange interaction. The states in each such group correspond to a total spin $S_{\text{tot}} = S_{\text{Fe}} + S_{\text{Gd}}$ ($S_{\text{tot}} = 1, \dots, 6$) with total number of states being equal to $2S_{\text{tot}} + 1$ (Table S20, ESI[†]). In each group, these states basically split again into Ising doublets *via* a weaker ZFS interaction on Fe^{III} and Gd^{III} ions. The multiplet arising from the ground manifold of 13 states ($S_{\text{tot}} = 6$) spread over an energy less than 2 cm⁻¹; therefore, they are all populated at the investigated temperatures. Given that many of these doublets display a tunnelling gap as large as 10⁻² cm⁻¹ (Table S20, ESI[†]), no blocking of magnetization relaxation can be expected. This is in full agreement with the ac susceptibility measurements showing that **1** is not a SMM.

Substitution of the isotropic Gd^{III} ion for an anisotropic Tb^{III} ion with oblate-shape f-electron charge cloud⁴⁸ in complex **2** indeed increases the axiality of the crystal field on the lanthanide ion. This is clearly shown in Table S10 and Fig. S7 (ESI[†]) where the energy gap between the electronic ground and first excited doublet corresponding to the lanthanide center substantially changes from *ca.* 0.2 cm⁻¹ to *ca.* 40 cm⁻¹. This increase in the axiality effectively differentiates nature of the energy splitting in complex **2** from **1**. In particular, the energy diagram changes from subgroups characterized by the total spin $S_{\text{Fe}} + S_{\text{Gd}}$ in **1** to subgroup of pseudospin 1/2 (corresponding to each electronic doublet of Tb^{III} ion) in exchange coupling with S_{Fe} in **2** as seen in Fig. 8. It should also be mentioned that due to no change in the ligand, this substitution results in little effect on the Fe^{III} center (see Tables S12, S13 and Fig. S8, ESI[†]).

We further calculated the magnetic susceptibility and field-dependent magnetization for complex **2** and showed them in Fig. S12 and S13 (ESI[†]), respectively. An intermolecular antiferromagnetic exchange coupling, $zJ = -0.027$ cm⁻¹, was introduced to give a better fit to the $\chi_{\text{M}}T$ curve of **2** at low temperatures.^{34,41,42} It is noted that the ferromagnetic

exchange parameters $J = 0.310$ cm⁻¹ obtained from fitting of $\chi_{\text{M}}T$ is of the same order of magnitude with the one from BS-DFT calculation but *ca.* 3 times smaller (0.85 cm⁻¹ in Table S18, ESI[†]). This difference is understandable considering the low accuracy of the BS-DFT methodology for exchange parameter calculation.

Using the exchange parameter obtained from fitting, the low-lying manifolds of exchange/dipolar spin-orbit states of **2**, resulting from the magnetic interaction of the ground $S = 5/2$ of Fe^{III} centre with the ground non-Kramers doublet of the Tb^{III} ion (lower manifold of 12 states) and with the first excited doublet (higher manifold of 12 states) lying *ca.* 40 cm⁻¹ higher are plotted in Fig. 8. The multiplet states in each manifold group form Kramers doublets separated by the Fe^{III}-Tb^{III} exchange interaction and the ZFS interaction at the Fe^{III} site. At low temperature, the relaxation occurs mostly in the lowest multiplet of 12 states, which due to the axiality of Tb^{III} ions in **2**, can be considered in an effective manner by the exchange interaction between the ground doublet of the Tb^{III} ion (pseudospin 1/2) and the Fe^{III} ion (spin 5/2) in the crystal field.⁴⁹ However, as the ZFS of the Fe^{III} ion is weak (see Table S12, ESI[†]), the easy magnetic axis of the Tb^{III} ions ground doublet and the ZFS of Fe^{III} ion is also non-collinear (about 27.2°, Fig. 9),⁴⁹⁻⁵² and the exchange interaction between Tb^{III} and Fe^{III} ions (see Table S18, ESI[†]) is small, each eigenstate of the [Fe^{III}Tb^{III}] complex will accordingly be a strong mixture of different m_{Fe} states. This results in large transversal components of the g -tensor of doublets of the complex (see Table S23, ESI[†]) as well as large transition matrix elements between the eigenstates of **2**, which can be seen in Fig. 10 *via* the relative

transition conductance^{45,49-51} $p_{ij} = p_{ji} = \frac{(E_j - E_i)^3}{e^{E_j/kT} - e^{E_i/kT}} |\mu_{ij}|^2$ between localized states $|i\rangle$ and $|j\rangle$. Consequently, relaxation in

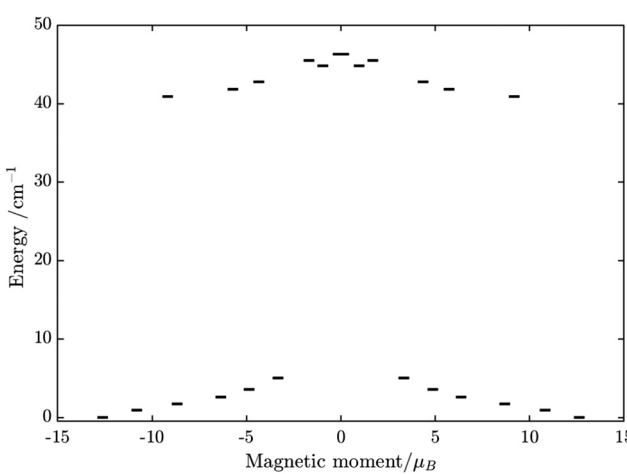


Fig. 8 Calculated low-lying spin-orbit manifolds of states arising from the magnetic interaction between the ground multiplet $S = 5/2$ of the Fe^{III} ion and the ground and first excited doublet of the Tb^{III} ion. The components of each doublet (black lines) are placed on the diagram according to their magnetic moment projection on the main magnetic axis of the ground doublet (with energy zero).

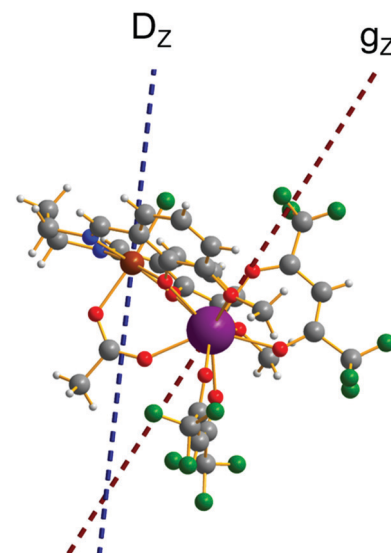


Fig. 9 Calculated molecular fragment of the [Fe^{III}Tb^{III}] complex (legend: Tb-indigo, Fe-brown, C-grey, N-blue, O-red, H-light grey). The dashed lines indicate the main anisotropy axis on Fe^{III} ion (blue) and the main magnetic axis on Tb^{III} ion (red).

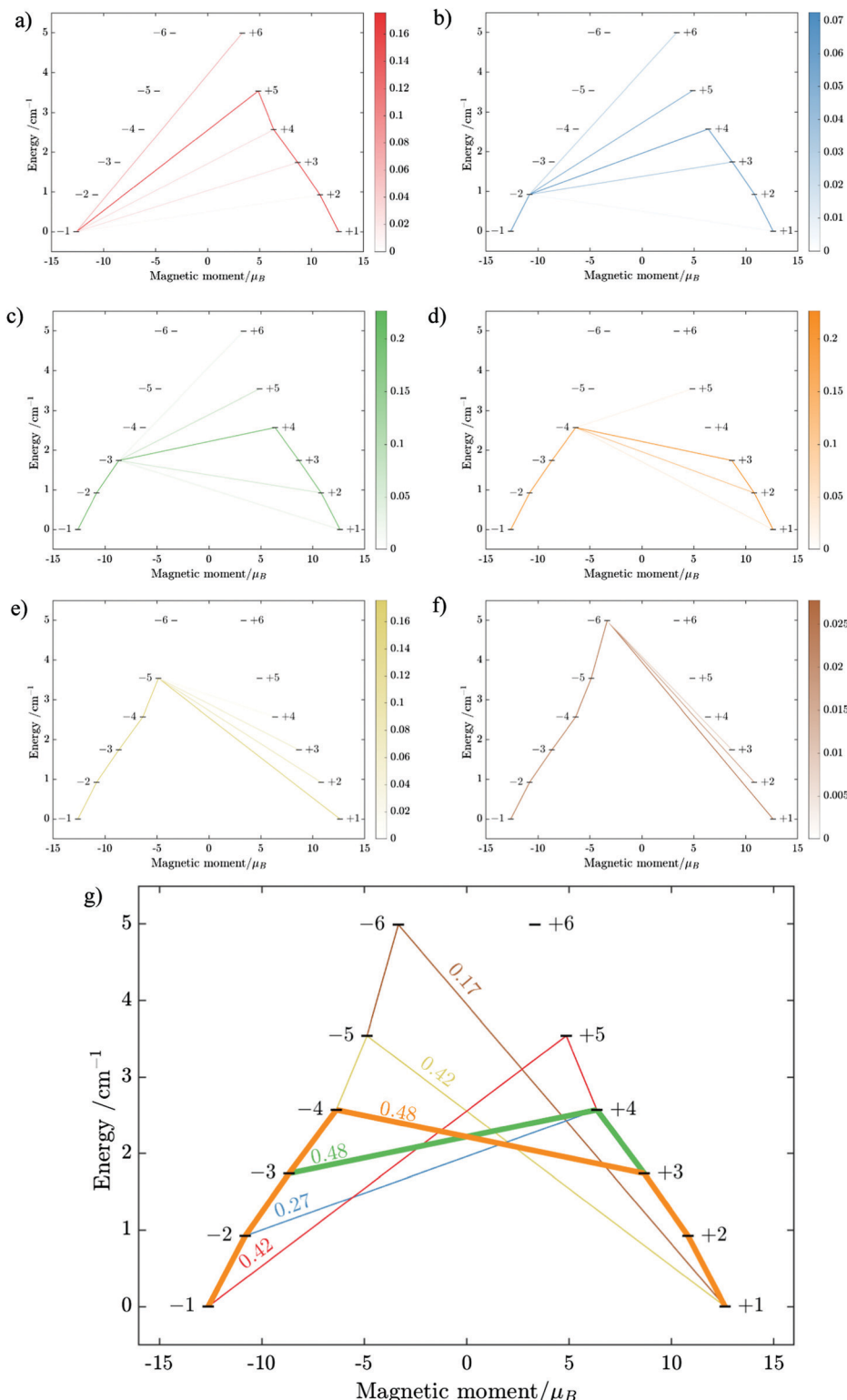


Fig. 10 Calculated relative transition conductance (non-horizontal lines) $p_{ij} = p_{ji} = \frac{(E_j - E_i)^3}{e^{E_j/kT} - e^{E_i/kT}} |\mu_{ij}|^2$ between any two localized states $|i\rangle$ and $|j\rangle$ of 6 lowest doublets involving in the magnetization relaxation at low temperature domain of $[\text{Fe}^{\text{III}}\text{Tb}^{\text{III}}]$ complex (2), assuming that the whole population is initially prepared on the left side of the potential well. Here $|\mu_{ij}|$ stands for average transition magnetic moment matrix element and taken from *ab initio* calculation, and $T = 2$ K is used for calculation of p_{ij} . Note that while relative transition conductive between states of different sides are shown with magnitude-dependent transparency, ones between states on the same side of the potential well are plotted without magnitude-dependent transparency. This is because population transition between states of the same side are much faster than ones between states of different sides, which is thus unimportant in determining the relaxation time of magnetization. Particularly, (a-f) p_{ij} for $i = -1 \dots -6$ respectively; (g) the largest p_{ij} for each i branch are plotted and compared to determined the dominant relaxation path (bold lines) from their corresponding p_{ij} . For $[\text{Fe}^{\text{III}}\text{Tb}^{\text{III}}]$ at low temperature, the dominant paths are from $|\pm 3\rangle \leftrightarrow |\mp 4\rangle$.

[Fe^{III}Tb^{III}], despite being slower than in [Fe^{III}Gd^{III}] due to the axiality of the Tb^{III} ion, is still overly fast. Specifically, at zero applied field, due to the large values of the transversal components of the *g*-tensor, even the internal field is able to open large tunnelling splitting gaps and accordingly accelerates the magnetization relaxation. At large applied field, although relaxation *via* QTM is suppressed, the large transition conductance, and accordingly large transition rates, between states with magnetic moments of opposite sign (see Fig. 10) are still sufficient to result in a fast relaxation of magnetization.⁵² This explains why some changes to the out-of-phase susceptibility (see Fig. S3, ESI[†]) can be observed at high magnetic field, but no slow relaxation is detected.

Similarly, the *ab initio* fragments calculations for complex 3 were performed. The spin-orbital energy levels and *g* tensors for each metal center in the crystal field are exhibited in Tables S14–S17, S24, S25 and Fig. S9, S10 (ESI[†]). As can be seen, while the ZFS splitting of Fe^{III} is similar to those in the above two complexes, the substitution from Tb^{III} to Dy^{III} further increases the axiality of the lanthanide center. In particular, the energy gap between the ground doublet and the first excited doublet of the Dy^{III} center in 3 is almost double of the Tb^{III} center in 2 (70 cm⁻¹ *vs.* 40 cm⁻¹). Other excited doublets of 3 also have a larger energy than corresponding ones in 2. Together with the fact that Dy^{III} is a Kramer ion, even without experiment, we can also guess that a slower relaxation than 2 in 3 is likely and may be observed.

Ab initio calculations of the magnetic susceptibility and field-dependent magnetization for the [Fe^{III}Dy^{III}] complex are also shown in Fig. S14 and S15 (ESI[†]), respectively. To have a good agreement with experiment, an intermolecular antiferromagnetic exchange coupling, $zJ = -0.023$ cm⁻¹ was introduced to describe the bending down of the $\chi_M T$ curve at low temperature.^{34,41,42}

The fitting process done using the POLY_ANISO package (MOLCAS 8.2) gives a value of $J = 0.452$ cm⁻¹, which is about 45% larger than J in 2. This relative difference of J between 2 and 3 is also verified by BS-DFT calculation (see Table S18, ESI[†]). It should also be noted that in complexes 1, 2, and 3, all *ab initio* calculations show that the exchange interaction between Ln^{III} and Fe^{III} ions is ferromagnetic, which is well supported by the BS-DFT calculations (see Table S18, ESI[†]). Meanwhile, the magnetic dipolar interaction (Table S19, ESI[†]) overall counteracts the ferromagnetic exchange interaction. However, in the [Fe^{III}Gd^{III}] compound, this effect is much weaker than in [Fe^{III}Dy^{III}], where it reduces the effect of ferromagnetic exchange by *ca.* 1/3. This is explained by a different orientation of the spins (magnetic moments) of Ln and Fe sites relatives to each other and relative to the Ln–Fe axis in the complexes in 1–3, with the easy magnetic axis of the Dy^{III} ion ground doublet and the ZFS of Fe^{III} also being non-collinear (at an angle of 44.4°, see Fig. 11).

Using fitting value of J , the energy spectrum of the complex is calculated and shown in Fig. 12 with the low-lying manifolds of exchange/dipolar spin–orbit states resulting from the magnetic interaction of the ground $S = 5/2$ of Fe^{III} centre with

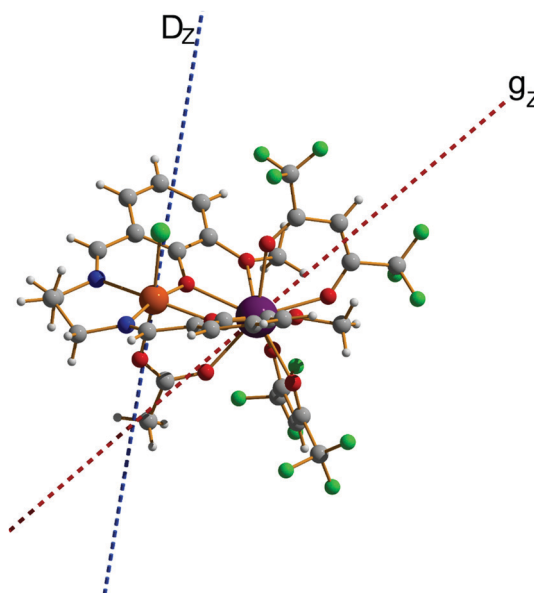


Fig. 11 Calculated molecular fragment of the [Fe^{III}Dy^{III}] complex (legend: Dy-indigo, Fe-brown, C-grey, N-blue, O-red, H-light grey). The dashed lines indicate the main anisotropy axis on Fe^{III} ion (blue) and the main magnetic axis on Dy^{III} ion (red).

the ground KD of the Dy^{III} ion (lower manifold of 12 states) and the first excited KD (higher manifold of 12 states) lying *ca.* 70 cm⁻¹ higher, which is larger than 2 due to Dy^{III} being more axial than the Tb^{III} ion (see Tables S10, S11 and S14, S15, ESI[†]). Relaxation in 3 also occurs mainly within the lowest exchange manifold (of 12 states) at low temperature. Due to *ca.* 45% stronger exchange interaction between Dy^{III} and Fe^{III} ions (see Table S18, ESI[†]), higher axiality and larger magnetic moment of the Dy^{III} ion, the mixture of different m_{Fe} in each eigenstate corresponding to the lower manifold of [Fe^{III}Dy^{III}] is

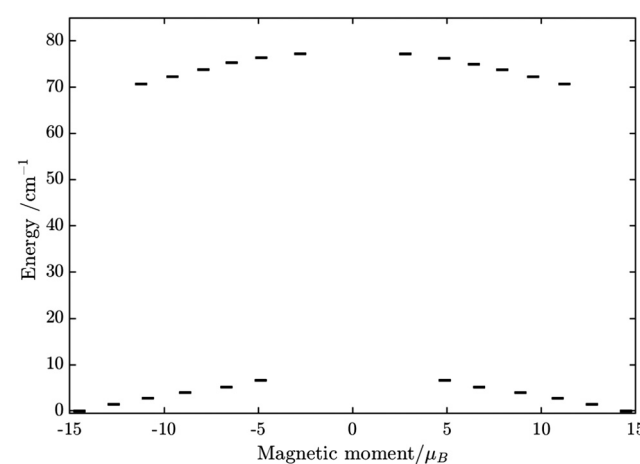


Fig. 12 Calculated low-lying spin–orbit manifolds of states arising from the magnetic interaction between the ground multiplet $S = 5/2$ of the Fe^{III} ion and the ground and first excited Kramers doublet of the Dy^{III} ion. The components of each doublet (black lines) are placed on the diagram according to their magnetic moment projection on the main magnetic axis of the ground doublet (with energy zero).

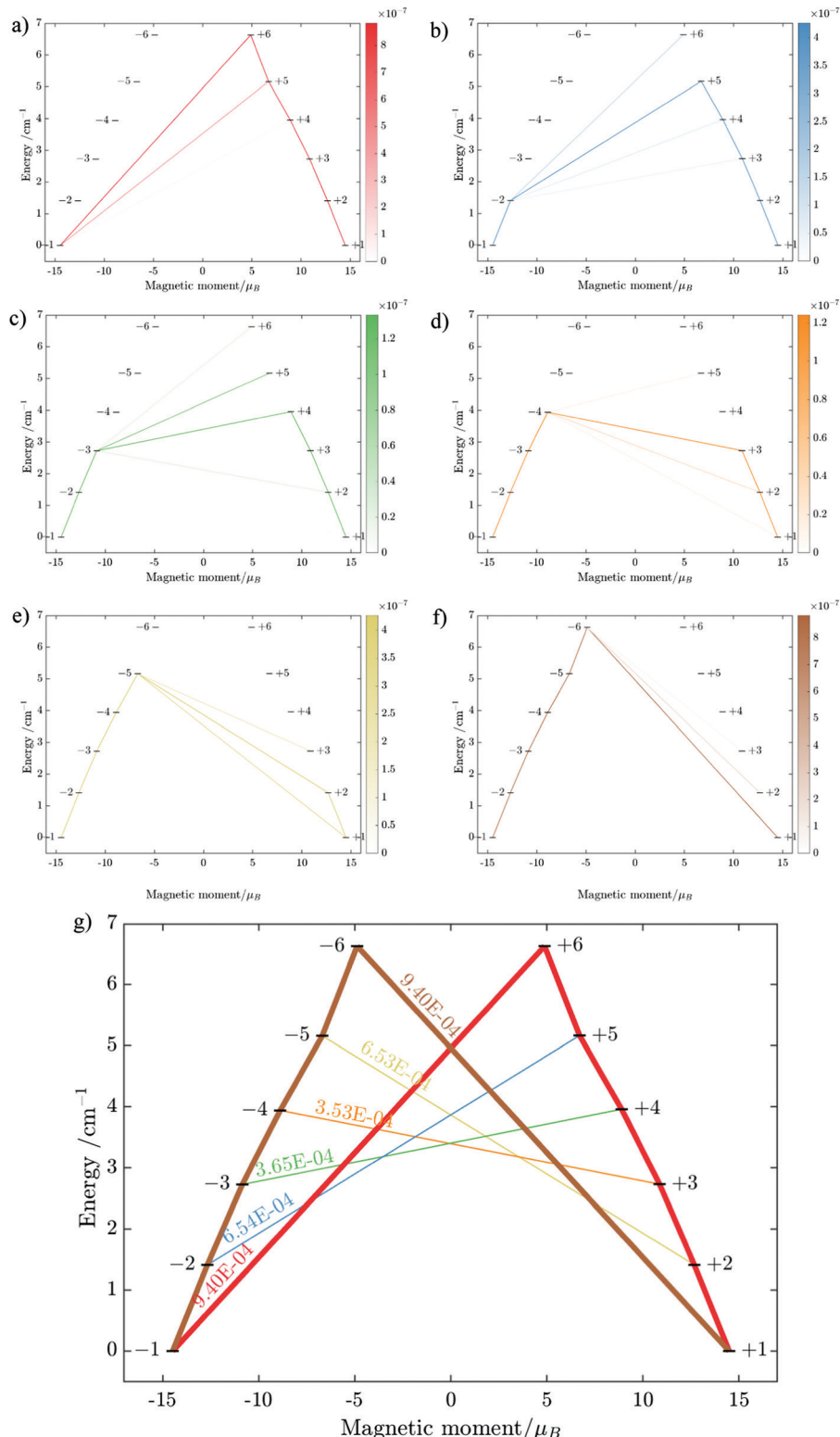


Fig. 13 Calculated relative transition conductance (non-horizontal lines) $p_{ij} = p_{ji} = \frac{(E_j - E_i)^3}{e^{E_j/kT} - e^{E_i/kT}} |\mu_{ij}|^2$ between any two localized states $|i\rangle$ and $|j\rangle$ of 6 lowest doublets involving in the magnetization relaxation at low temperature domain of $[\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}]$ complex (3), assuming that the whole population is initially prepared on the left side of the potential well. Here $|\mu_{ij}|$ stands for average transition magnetic moment matrix element and taken from *ab initio* calculation, and $T = 2$ K is used for calculation of p_{ij} . Note that while relative transition conductance between states of different sides are shown with magnitude-dependent transparency, ones between states on the same side of the potential well are plotted without magnitude-dependent transparency. This is because population transition between states of the same side are much faster than ones between states of different sides, which is thus unimportant in determining the relaxation time of magnetization. Particularly, (a-f) p_{ij} for $i = -1 \dots -6$ respectively; (g) the largest p_{ij} for each i branch are plotted and compared to the dominant relaxation path (bold lines) from their corresponding p_{ij} . For $[\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}]$ at low temperature, the dominant paths are from $|\pm 1\rangle \leftrightarrow |\mp 6\rangle$, which corresponds to an effective blocking barrier of around 9.54 K (6.63 cm^{-1}).

significantly lower.^{49,52} This leads to much smaller relative transition conductance, and accordingly much smaller transition rates between states having different signs of magnetic moments in 3 compared to 2, which can be seen from Fig. 10 and 13 *via* the relative transition conductance calculations. Indeed, these matrix elements for 3 are at least two orders of magnitude smaller than 2, which explains why under an applied field where QTM is suppressed, 3 relaxes much slower than 2 (see Fig. S2 and S3, ESI[†]) and is thus observable *via* the maximum in frequency-dependent out-of-phase susceptibility. However, at zero applied field, since the mixture of different m_{Fe} in eigenstates of 3 is still not sufficiently small due to the weak crystal field of the Fe^{III} ion, fairly decent tunnelling splitting gaps (see Table S24, ESI[†]) within (quasi-) doublets of 3 are still open, which speed up the relaxation *via* the QTM process between states with opposite signed magnetic moments^{49,52,55} (left and right states in Fig. 13).

From above, we have explained why the magnetization relaxation in 3 is slower than 2, which is slower than 1. A summary of the difference between these three binuclear complexes is also provided in Table 4. Since only slow relaxation is observed in 3, our next task is finding its dominant relaxation path and accordingly the effective blocking barrier. This can be done by first noticing that the exchange interaction splits the energy spectrum so that $| -m \rangle$ and $| +(7 - m) \rangle$ in Fig. 13 having approximately the same contribution of z-axis spin projection component of Fe^{III} spin (see Fig. 14 for a depiction of the splitting of the energy spectrum in 3 assuming that there is no involvement of the crystal field in the Fe^{III} ion). Accordingly, those transitions between $| -m \rangle$ and $| +(7 - m) \rangle$ are favoured as flipping one spin is much easier than two spins at the same time

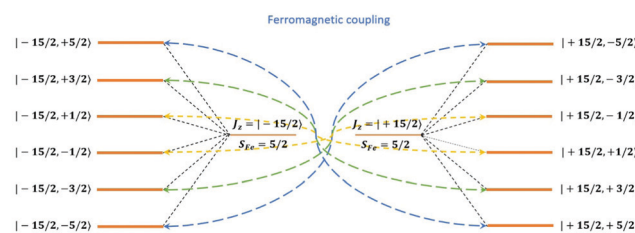


Fig. 14 Energy splitting of the lowest manifold arising from the exchange coupling between a strong anisotropic lanthanide ion (e.g., Dy^{III}) and a transition metal ion (e.g., Fe^{III}) in weak ZFS. Here orange lines are energy levels, colored lines with arrows indicates the most preferred relaxation channel for population at corresponding energy level. Lines with the same color are expected to have the same transition rate. Dominant path is the competition between different colored lines. Effective blocking barrier can thus range from the height of the fourth to the sixth energy levels depending on the specific system and temperature.

(coloured lines with arrow in Fig. 14). This is demonstrated in Fig. 13 for 3 where the relative transition conductance, and accordingly the transition rate, between states $| -m \rangle$ and state $| +(7 - m) \rangle$ is always fastest comparing to between $| -m \rangle$ and other $| +n \rangle$. Additionally, from Fig. 13g, the calculated data for the relative transition conductance also show that despite some contributions from other relaxation paths ($| \pm 2 \rangle$ to $| \mp 5 \rangle$ and $| \pm 3 \rangle$ to $| \mp 4 \rangle$), the dominant relaxation path is indeed from $| \pm 1 \rangle$ to $| \mp 6 \rangle$. This corresponds to the Orbach relaxation process with a blocking barrier of approximately 9.54 K (6.63 cm^{-1}), which is in excellent/good agreement with the fitted one 9.72 K from experimental data (see Fig. 6a) if only the Orbach process is taken into consideration, or 8.39 K (see Fig. 6b) if the Raman process is also included. This agreement validates our suggested mechanism for the relaxation in 3.

Table 4 Difference between three binuclear complexes with compartmental ligands **1**, **2**, and **3** from the theoretical investigations

| | $\text{Fe}^{\text{III}}\text{Gd}^{\text{III}}$ | $\text{Fe}^{\text{III}}\text{Tb}^{\text{III}}$ | $\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}$ |
|----------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------|
| ZFS on Fe^{III} | Small | Small | Small |
| ZFS on Ln^{III} | Small | Large | Largest |
| Order of interactions (Ex – Exchange, CF – Crystal field) | $H_{\text{EX}}^{\text{Gd-Fe}} > H_{\text{CF}}^{\text{Gd}}$ $H_{\text{EX}}^{\text{Gd-Fe}} > H_{\text{CF}}^{\text{Fe}}$ | $H_{\text{CF}}^{\text{Tb}} > H_{\text{EX}}^{\text{Tb-Fe}} > H_{\text{CF}}^{\text{Fe}}$ | $H_{\text{CF}}^{\text{Dy}} > H_{\text{EX}}^{\text{Tb-Fe}} > H_{\text{CF}}^{\text{Fe}}$ |
| Ground multiplet (relaxation at low T occurs here) | $S_{\text{total}} = 6$ 13 levels | $S_{\text{Tb}} = 1/2, S_{\text{Fe}} = 5/2, 12$ levels | $S_{\text{Dy}} = 1/2, S_{\text{Fe}} = 5/2, 12$ levels |
| Exchange constant $\text{Fe}^{\text{III}}\text{–Ln}^{\text{III}}$ (cm^{-1}) | 0.898 | 0.310 | 0.452 |
| Axiality of the ground multiplet | Low | High | Highest |
| Relaxation at zero field | Fast, not observable. | Fast, not observable. | Fast, not observable. |
| Main reason | Weak ZFS on Fe^{III} and isotropic Ln^{III} ion \rightarrow low axiality, large tunneling splittings, strong spin-phonon coupling. | Weak ZFS on Fe^{III} , weak $\text{Fe}^{\text{III}}\text{–Tb}^{\text{III}}$ exchange interaction and non-collinearity between easy magnetic axis of Tb^{III} ground doublet and ZFS on Fe^{III} \rightarrow large spin-phonon coupling and fast QTM | Weak ZFS on Fe^{III} \rightarrow fast QTM |
| Relaxation under an applied field | Fast, not observable. | Fast, not observable. | Slower, observable |
| Main reason | Weak ZFS on Fe^{III} and isotropic Ln^{III} ion \rightarrow low axiality, strong spin-phonon coupling. | Weak ZFS on Fe^{III} , weak $\text{Fe}^{\text{III}}\text{–Tb}^{\text{III}}$ exchange interaction and non-collinearity between easy magnetic axis of Tb^{III} ground doublet and ZFS on Fe^{III} \rightarrow strong spin-phonon coupling. | Stronger Fe^{III} exchange interaction, higher axiality \rightarrow weaker spin-phonon coupling. |
| U_{eff} | Not applicable | Not applicable | From lowest to highest energy level of the ground multiplet at low T |

From above analyses on the physics in complex series $[\text{Fe}^{\text{III}}\text{Ln}^{\text{III}}]$, a generalization for similar systems where a transition metal is in exchange coupling with the ground doublet of a strong anisotropic lanthanide ion can also be made. In particular, due to the exchange interaction, the energy spectrum of these complexes will be divided into multiple manifolds where each one effectively arises from exchange coupling between each doublet of the anisotropic lanthanide ion and the zero-field-split energy spectrum of the transition metal ion. Relaxation in these systems at low temperatures will then occur mainly in the lowest manifold of the $2 \times (2S_{\text{transition-metals-ion}} + 1)$ energy levels. In general, for these systems under an applied field, the dominant relaxation path, and accordingly the effective blocking barrier, will depend on the competition between relaxation channels connecting states with the same dominant z -component of the transition metal ion spin (see Fig. 14). On the other hand, tunnelling splitting in these systems, which plays an important role in relaxation at zero field, will depend on the mixing of different z -components of the transition metal ion spin in the localized states,⁵² hereinafter referred to as “purity” and thus depends strongly on the rhombic anisotropy parameter E of the crystal field of the transition metal ion, the magnitude of the exchange constant, the collinearity of the easy magnetic axis of the lanthanide ion's ground doublet and one of the ZFS of the transition metal ion, and to some degree the spin number of the lanthanide ion.^{50–57} Specifically, a weaker E , will not only decrease the tunnelling splitting gaps but also improve the “purity” of the eigenstates on the opposite sides of the potential wells and accordingly lower the transition rate between states with different dominant z -component contributions, *e.g.* $| -m \rangle$ and $| n \neq +(7 - m) \rangle$ in 3 (see Fig. 14). Meanwhile, a strong exchange interaction and collinearity of the axes of lanthanide ion's ground doublet and transition metal's ZFS, will further strengthen the axiality of the complex's eigenstates.^{50–54} All these effects are in a synergy to help reduce relaxation time. Thus, a system with a large J , a small E , and collinearity between easy magnetic axes of lanthanide's ion and ZFS of transition metal ion will highly likely show a slow relaxation of magnetization.

Conclusions

We presented a new synthetic approach allowing for an easy access to binuclear $\text{Fe}^{\text{III}}\text{--Ln}^{\text{III}}$ complexes. Applying this methodology, three new isostructural complexes, $[\text{Fe}^{\text{III}}\text{Ln}^{\text{III}}(\text{valpn})(\text{hfac})_2(\text{OAc})\text{Cl}]$ ($\text{Ln} = \text{Gd}, \text{Tb}, \text{Dy}$; $\text{hfac}^- = \text{hexafluoroacetylacetone}; \text{AcO}^- = \text{acetate}$), have been synthesized and thoroughly investigated. This strategy can be extended towards other 3d–4f complexes containing trivalent 3d metal ions. It was shown that only the third, $[\text{Fe}^{\text{III}}\text{Dy}^{\text{III}}]$ complex 3 exhibits SMM behavior within the investigated frequency domain of ac susceptibility, which was rationalized by *ab initio* analysis. In particular, *ab initio* analysis revealed that the temperature behavior of the relaxation time is due to the Orbach relaxation processes between the states on the opposite sides and opposite heights of the blocking barrier, which become

dominant in a sufficiently strong applied magnetic field ($H_{\text{dc}} = 800$ Oe), when the QTM is sufficiently suppressed. Our theoretical analyses not only elucidated the mechanism of the relaxation of magnetization in this series of binuclear complexes, but also shed light on the possible relaxation mechanism in similar systems where a transition metal is in exchange coupling with the ground doublet of a strong anisotropic lanthanide ion. From this understanding, similar systems with better magnetization relaxation may be developed in the future.

Conflicts of interest

There are no conflicts to declare.

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References

- 1 (a) M. Andruh, J.-P. Costes, C. Diaz and S. Gao, *Inorg. Chem.*, 2009, **48**, 3342; (b) M. Sakamoto, K. Manseki and H. Ōnse, *Coord. Chem. Rev.*, 2001, **219**–**221**, 379; (c) C. Benelli and D. Gatteschi, *Chem. Rev.*, 2002, **102**, 2369.
- 2 (a) R. Sessoli and A. K. Powell, *Coord. Chem. Rev.*, 2009, **293**, 2328; (b) K. Liu, W. Shi and P. Cheng, *Coord. Chem. Rev.*, 2015, **289**–**290**, 74; (c) H. L. C. Feltham and S. Booker, *Coord. Chem. Rev.*, 2014, **276**, 1; (d) L. Rosado Piquer and E. C. Sañudo, *Dalton Trans.*, 2015, **44**, 8771.
- 3 See, for example: (a) T. Gupta, M. F. Beg and G. Rajaraman, *Inorg. Chem.*, 2016, **55**, 11201; (b) L. Ungur, M. Thewissen, J.-P. Costes, W. Wernsdorfer and L. F. Chibotaru, *Inorg. Chem.*, 2013, **52**, 6328; (c) J.-W. Yang, Y.-M. Tian, J. Tao, P. Chen, H.-F. Li, Y.-Q. Zhang, P.-F. Yan and W.-B. Sun, *Inorg. Chem.*, 2018, **57**, 8065.
- 4 See, for example: (a) G. Peng, G. E. Kostakis, Y. Lan and A. K. Powell, *Dalton Trans.*, 2013, **42**, 46; (b) A. Baniodeh, I. J. Hewitt, V. Mereacre, Y. Lan, G. Novitchi, C. E. Anson and A. K. Powell, *Dalton Trans.*, 2011, **40**, 4080; (c) A. M. Ako, V. Mereacre, R. Clérac, I. J. Hewitt, Y. Lan, C. E. Anson and

A. K. Powell, *Dalton Trans.*, 2007, 5245; (d) J. W. Sharples and D. Collison, *Coord. Chem. Rev.*, 2014, **260**, 1.

5 E. Moreno Pineda, F. Tuna, Y.-Z. Zheng, S. J. Teat, R. E. P. Winpenny, J. Schnack and E. J. L. McInnes, *Inorg. Chem.*, 2014, **53**, 3032.

6 V. Mereacre, D. Prodius, C. Turta, S. Shova, G. Filoti, J. Bartolomé, R. Clérac, C. E. Anson and A. K. Powell, *Polyhedron*, 2009, **28**, 3017.

7 Y.-F. Zeng, G.-C. Xu, X. Hu, Z. Chen, X.-H. Bu, S. Gao and E. C. Sañudo, *Inorg. Chem.*, 2010, **49**, 9734.

8 (a) A. Figuerola, J. Ribas, M. Llunell, D. Cassanova, M. Maestro, S. Alvarez and C. Diaz, *Inorg. Chem.*, 2005, **44**, 6939; (b) B. Yan and Z. Chen, *Helv. Chim. Acta*, 2001, **84**, 817; (c) X.-J. Song, J.-J. Xu, Y. Chen, M. Muddassir, F. Cao, R.-M. Wei, Y. Song and X.-Z. You, *Polyhedron*, 2013, **66**, 212.

9 M.-G. Alexandru, D. Visinescu, S. Shova, W. X. C. Oliveira, F. Lloret and M. Julve, *Dalton Trans.*, 2018, **47**, 6005.

10 M. Coletta, R. McLellan, S. Sanz, K. J. Gagnon, S. J. Teat, E. K. Brechin and S. J. Dalgarno, *Chem. – Eur. J.*, 2017, **23**, 14073.

11 (a) A. Figuerola, C. Diaz, J. Ribas, V. Tangoulis, J. Granell, F. Lloret, J. Mahia and M. Maestro, *Inorg. Chem.*, 2003, **42**, 641; (b) W.-T. Chen, G.-C. Guo, M.-S. Wang, G. Xu, L.-Z. Cai, T. Akitsu, M. Akita-Tanaka, A. Matsushita and J.-S. Huang, *Inorg. Chem.*, 2007, **46**, 2105; (c) G. Li, T. Akitsu, O. Sato and Y. Einaga, *J. Am. Chem. Soc.*, 2003, **125**, 12396; (d) X.-R. Sun, Z.-D. Chen, F. Yan, S. Gao, K.-K. Cheung, C.-M. Che and X.-X. Zhang, *J. Clust. Sci.*, 2002, **13**, 103; (e) S. A. Stoian, C. Paraschiv, N. Kiritsakas, F. Lloret, E. Münck, E. L. Bominaar and M. Andruh, *Inorg. Chem.*, 2010, **49**, 3387.

12 M. Ferbinteanu, T. Kajiwara, K.-Y. Choi, H. Nojiri, A. Nakamoto, N. Kojima, F. Cimpoesu, Y. Fujimura, S. Takaishi and M. Yamashita, *J. Am. Chem. Soc.*, 2006, **128**, 9008.

13 J.-P. Costes, A. Dupuis and J.-P. Laurent, *Eur. J. Inorg. Chem.*, 1998, 1543.

14 M. Andruh, *Dalton Trans.*, 2015, **44**, 16633.

15 (a) J.-P. Costes, F. Dahan, F. Dumestre, J. M. Clemente-Juan, J. Garcia-Tojal and J.-P. Tuchagues, *Dalton Trans.*, 2003, 464; (b) M. Sarwar, A. M. Madalan, F. Lloret, M. Julve and M. Andruh, *Polyhedron*, 2011, **30**, 2414.

16 (a) I. Nemec, M. Machata, R. Herchel, R. Boča and Z. Trávníček, *Dalton Trans.*, 2012, **41**, 14603; (b) W.-W. Kuang, C.-Y. Shao and P.-P. Yang, *J. Coord. Chem.*, 2015, **68**, 1412.

17 S. Stoll and A. Schweiger, *J. Magn. Reson.*, 2006, **178**, 42.

18 S. Stoll and R. D. Britt, *Phys. Chem. Chem. Phys.*, 2009, **11**, 6614.

19 A. Bencini and D. Gatteschi, *EPR of exchange coupled systems: Springer - Verlag*, New York, 1990.

20 N. F. Chilton, R. P. Anderson, L. D. Turner, A. Soncini and K. S. Murray, *J. Comput. Chem.*, 2013, **34**, 1164.

21 J. Schnack, P. Hage and H. J. Schmidt, *J. Comput. Phys.*, 2008, **227**, 4512.

22 R. Schnalle and J. Schnack, *Int. Rev. Phys. Chem.*, 2010, **29**, 403.

23 M. Haertel, J. Richter, D. Ihle, J. Schnack and S.-L. Drechsler, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2011, **84**, 104411.

24 H. Veronika, K. Gieb, P. Müller, L. Ungur, L. F. Chibotaru, M. Heidemeier, E. Krickemeyer, A. Stammel, H. Böggel, C. Schröder, J. Schnack and T. Glaser, *Chem. Sci.*, 2012, **3**, 2868.

25 T. Glaser, V. Hoeke, K. Gieb, J. Schnack, C. Schröder and P. Müller, *Coord. Chem. Rev.*, 2015, **289**, 261.

26 L. Noddeman, D. A. Case and A. Aizman, *J. Am. Chem. Soc.*, 1988, **110**(4), 1001.

27 D. Gatteschi and K. Yamaguchi, *Opportunities for new physics in molecular magnetism. Molecular Magnetism: from Molecular Assemblies to the Devices*, Springer, Dordrecht, 1996, vol. 321, pp. 561–570.

28 M. Shoji, K. Koizumi, Y. Kitagawa, T. Kawakami, S. Yamanaka, M. Okumura and K. Yamaguchi, *Chem. Phys. Lett.*, 2006, **432**(1–3), 343.

29 I. Negodaev, C. de Graaf and R. Caballol, *Chem. Phys. Lett.*, 2008, **458**(4–6), 290.

30 S. Knecht, E. D. Hedegård, S. Keller, A. Kovyrshin, Y. Ma, A. Muolo, C. J. Stein and M. Reiher, *Chimia*, 2016, **70**, 24.

31 S. Battaglia, S. Keller and S. Knecht, *J. Chem. Theory Comput.*, 2018, **14**, 2353.

32 S. Wouters, T. Bogaerts, P. Van Der Voort, V. Van Speybroeck and D. Van Neck, *J. Chem. Phys.*, 2014, **140**, 241103.

33 S. Wouters, V. Van Speybroeck and D. Van Neck, *J. Chem. Phys.*, 2016, **145**, 054120.

34 L. F. Chibotaru, L. Ungur, C. Aronica, H. Elmoll, G. Pilet and D. Luneau, *J. Am. Chem. Soc.*, 2008, **130**, 12445.

35 L. F. Chibotaru, L. Ungur and A. Soncini, *Angew. Chem., Int. Ed.*, 2008, **120**, 4194.

36 S.-Y. Lin, W. Wernsdorfer, L. Ungur, A. K. Powell, Y.-N. Guo, J. Tang, L. Zhao, L. F. Chibotaru and H.-J. Zhang, *Angew. Chem., Int. Ed.*, 2012, **51**(51), 12767.

37 L. Ungur, S. K. Langley, T. N. Hooper, B. Moubaraki, E. K. Brechin, K. S. Murray and L. F. Chibotaru, *J. Am. Chem. Soc.*, 2012, **134**, 18554.

38 S. K. Langley, D. P. Wielechowski, V. Vieru, N. F. Chilton, B. Moubaraki, B. F. Abrahams, L. F. Chibotaru and K. S. Murray, *Angew. Chem., Int. Ed.*, 2013, **52**(46), 12014.

39 M. Towatari, K. Nishi, T. Fujinami, N. Matsumoto, Y. Sunatsuki, M. Kojima, N. Mochida, T. Ishida, N. Re and J. Mrozninski, *Inorg. Chem.*, 2013, **52**, 6160.

40 F. Aquilante, J. Autschbach, R. K. Carlson, L. F. Chibotaru, M. G. Delcey, L. De Vico, I. F. G. N. Ferre, L. M. Frutos, L. Gagliardi, M. Garavelli, A. Giussani, C. E. Hoyer, G. L. Manni, H. Lischka, D. Ma, P. A. Malmqvist, T. s. Muller, A. Nenov, M. Olivucci, T. B. Pedersen, D. Peng, F. Plasser, B. Pritchard, M. Reiher, I. Rivalta, I. Schapiro, J. Segarra-Mart, M. Stenrup, D. G. Truhlar, L. Ungur, A. Valentini, S. Vancoillie, V. Veryazov, V. P. Vysotskiy, O. Weingart, F. Zapata and R. Lindh, *J. Comp. Chem.*, 2016, **37**, 506.

41 L. Ungur and L. F. Chibotaru, <http://www.molcas.org/documentation/manual/>.

42 L. F. Chibotaru, L. Ungur and A. Soncini, *Angew. Chem., Int. Ed.*, 2008, **47**, 4126.

43 V. Vieru, L. Ungur and L. F. Chibotaru, *J. Phys. Chem. Lett.*, 2013, **4**, 3565.

44 A. Abragam and B. Bleaney, *Electron Paramagnetic Resonance of Transition Ions*, Clarendon Press, Oxford, 1970.

45 D. A. Garanin and E. M. Chudnovsky, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 1997, **56**, 11102.

46 (a) M. Llunell, D. Casanova, J. Girera, P. Alemany and S. Alvarez, *SHAPE, version 2.1*, Universitat de Barcelona, Barcelona, Spain; (b) S. Alvarez, P. Alemany, D. Casanova, J. Cirera, M. Llunell and D. Avnir, *Coord. Chem. Rev.*, 2005, **249**, 1693.

47 O. Kahn, *Molecular Magnetism*, Wiley-VCH, Dordrecht, 1993.

48 J. D. Rinehart and J. R. Long, *Chem. Sci.*, 2011, **2**, 2078.

49 L. F. Chibotaru, *Molecular Nanomagnets and Related Phenomena*, Springer Berlin Heidelberg, Berlin, Heidelberg, 2014, p. 185.

50 A. A. Patrascu, M. Briganti, S. Soriano, S. Calancea, R. A. Allão Cassaro, F. Totti, M. G. F. Vaz and M. Andruh, *Inorg. Chem.*, 2019, **58**, 13090.

51 A. A. Patrascu, S. Calancea, M. Briganti, S. Soriano, A. M. Madalan, R. A. A. Cassaro, A. Caneschi, F. Totti, M. G. F. Vaz and M. Andruh, *Chem. Commun.*, 2017, **53**, 6504.

52 D. Gatteschi and R. Sessoli, J. Villain, *Molecular Nanomagnets*, Oxford University Press, 2006.

53 V. Vieru, T. D. Pasatoiu, L. Ungur, E. Suturina, A. M. Madalan, C. Duhayon, J.-P. Sutter, M. Andruh and L. F. Chibotaru, *Inorg. Chem.*, 2016, **55**, 12158.

54 L. F. Chibotaru and L. Ungur, *J. Chem. Phys.*, 2012, **137**, 064112.

55 M. N. Leuenberger and D. Loss, *Phys. Rev. B: Condens. Matter Mater. Phys.*, 2000, **61**, 1286.

56 S.-G. Wu, Z.-Y. Ruan, G.-Z. Huang, J.-Y. Zheng, V. Vieru, G. Taran, J. Wang, Y.-C. Chen, J.-L. Liu, L. T. A. Ho, L. F. Chibotaru, W. Wernsdorfer, X.-M. Chen and M.-L. Tong, *Chem.*, 2021, **7**, 982.

57 M. N. Leuenberger and D. Loss, *Europhys. Lett.*, 1999, **46**, 692.