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The effect of the second coordination sphere on the magnetism of  $[Ln(NO_3)_3(H_2O)_3] \cdot (18$ -crown-6) (Ln = Dy and Er)†

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The objective of this work was the exploration of the effect of the second coordination sphere on the magnetic properties of  $[Ln(NO_3)_3(H_2O)_3] \cdot (18C6)$  (Ln = Dy (1) and Er (2)) compounds comprising cocrystallized 18-crown-6 ethers. Both compounds were identified as field-induced single molecule magnets (SMMs) with estimated magnetization reversal barriers  $U_{\rm eff}=66-71~{\rm K}$  for 1 and  $U_{\rm eff}=21-24~{\rm K}$ for 2. Theoretical calculations with the B3LYP functional revealed substantial change and redistribution of the electrostatic potential upon accounting for the second coordination sphere represented by two 18C6 molecules, which resulted in the change of the crystal-field around metal atoms. As a result, the multireference CASSCF calculations exposed significant impact of the second coordination sphere on the energy splitting of the respective  $^6H_{15/2}$  (Dy<sup>III</sup>) and  $^4I_{15/2}$  (Er<sup>III</sup>) ground states, the magnetization reversal barrier and the magnetic anisotropy parameters. Moreover, the calculated magnetization reversal barriers,  $U_{\text{calc.}} = 57 \text{ K}$  for 1 and  $U_{\text{calc.}} = 16 \text{ K}$  for 2, are in good agreement with the experimental values accentuating the importance of the second coordination sphere on the magnetic properties of SMMs.

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# Introduction

The lanthanide coordination compounds are of great interest to many magnetochemists around the world since Ishikawa's seminal report about the phthalocyanine complexes [Bu<sub>4</sub>N]  $[Ln(Pc)_2]$  (Ln = Tb and Dy, Pc = phthalocyanine, Bu<sub>4</sub>N<sup>+</sup> = tetrabutylammonium cation) behaving as single-molecule magnets (SMMs).1 SMMs are molecular nanomagnets exhibiting slow relaxation of the magnetization and magnetic bistability of molecular origin.2,3 SMM based materials can be potentially utilized in devices with high storage capacity, in sensors, quantum computing, spintronics, etc.4,5 The key concept of SMMs is the existence of the energy barrier acting against the reversal of the magnetization and so far many lanthanide-based SMMs have been reported.6 In the case of dysprosium(III) and erbium(III) ions, this barrier is defined by the ligand-field induced splitting of the ground state  ${}^{6}H_{15/2}$  and  $^4I_{15/2}$ , respectively, into the respective Kramers sublevels of the Jmanifold. Thus, the coordination number and the shape of the coordination polyhedron are crucial properties defining the

energy barrier properties,7 hence SMMs performance. In last

years, the enormous progress was achieved regarding the

maximising the effective energy barrier ( $U_{\text{eff}}$ ) in Dy(III) SMMs, e.g.  $U_{\text{eff}} = 1025 \text{ K}$  was reported for [Dy(bbpen)Br] (H<sub>2</sub>bbpen = N,N'-bis(2-hydroxybenzyl)-N,N'-bis(2-methylpyridyl)ethylenediamine),  $U_{\text{eff}} = 1815 \text{ K}$  was found in  $[Dy(OtBu)_2(py)_5][BPh_4]$ , (py = pyridine)9 and the record-holding SMM is [Dy(Cp<sup>ttt</sup>)<sub>2</sub>]  $[B(C_6F_5)_4]$  (Cp<sup>ttt</sup> = 1,2,4-tri(tert-butyl)cyclopentadienide) with  $U_{\rm eff} = 1837$  K and magnetic hysteresis present up to 60 K (ref.  $(Cp^{iPr5}$  $[(Cp^{iPr5})Dy(Cp*)]^+$ propylcyclopentadienyl, Cp\* = pentamethylcyclopentadienyl) which has 80 K highest magnetic blocking temperature. 11 In addition, even though erbium based compounds exhibit slow relaxation of magnetization less frequently and with lower  $U_{\text{eff}}$ barriers than their dysprosium analogues continuous research progresses steadily to greater values of  $U_{\rm eff}$ . This can be demonstrated on several examples featuring predominantly organometallic ligands: e.g.  $[(Cp^*)Er(COT)]$   $(Cp^* = pentam$ ethylcyclopentadienide; COT = cyclooctatetraenide dianion) with hysteresis loop from 1.8 to 5 K and two magnetic relaxation barriers 197 K and 323 K,12 [(Dsp)Er(COT)] (Dsp = 3,4dimethyl-2,5-bis(trimethylsilyl) phospholyl) shows slow relaxation with the zero applied magnetic field and barrier 358 K,13 lastly boron containing six membered borabenzene ligand in  $[(C_5H_5BH)Er(COT)]$  with  $U_{eff} = 371$  K and  $[(C_5H_5BCH_3)Er(COT)]$ which for ErIII complexes holds currently highest energy barrier of 421 K.14

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However, the SMMs properties are not solely defined by the effective energy barrier due to the fact that the slow relaxation magnetization can be governed by various mechanisms - onephonon direct and two-phonon Orbach and Raman processes, 15 and additionally also by the quantum tunnelling of the magnetization. However, the identification and comprehension of the relaxation processes is not an easy task and a better insight into these mechanisms is crucial for the further development of the single-molecule magnets. 16,17

One of broadly used ligands in search of tuning magnetic behaviour of the lanthanide-based compounds appears to be 18crown-6 (18C6). This hexadentate ether serves frequently in formation of half-sandwich or sandwich complexes.<sup>18</sup> Nevertheless, only in a few works this ligand directly bonds to lanthanide atom and at the same time magnetic properties were studied. Gadolinium based complex  $[{Gd(18C6)(\mu\text{-OH})(CH_3CN)}_2]$  $[Ni(dmit)_2]_2$  (dmit<sup>2-</sup> = 2-thioxo-1,3-dithiole-4,5-dithiolate) with the nickel encompassing complex counter anion is reported to have antiferromagnetic exchange between gadolinium atoms.19 Likewise magnetic properties of synthesized thiocyanate containing [Eu(18C6)(NCS)<sub>3</sub>] and [Tb(18C6)(NCS)<sub>3</sub>] compounds were studied. However, only Tb(III) complex was subjected to analysis of frequency dependent behaviour in AC magnetic susceptibility measurement, but no slow relaxation of magnetization was detected.<sup>20</sup> Simple motif  $[Ln(NO_3)_3(18C6)]$  (Ln = Ce<sup>III</sup>, Pr<sup>III</sup>, Nd<sup>III</sup>) was explored relatively recently in 2016 probing magnetic properties of nitrato ligands with several lanthanides, where under applied external magnetic field of 1000 Oe slow relaxation of magnetization was observed for Ce(III) and Nd(III) complexes.21 The same year were published two new structures of dysprosium containing coordinated 18C6 ligand, namely [Dy(18C6)(NO<sub>3</sub>)<sub>2</sub>]BPh<sub>4</sub> and [Dy(18C6)(NO<sub>3</sub>)<sub>2</sub>]ClO<sub>4</sub>, in which diverse coordination of nitrato ligands induced differences in the slow relaxation of the magnetization demonstrated by  $U_{\rm eff} = 43$  K and  $U_{\rm eff} = 63$  K, respectively.<sup>22</sup> Currently, the newest research of magnetic properties in 18C6based coordination compounds with lanthanides includes four new compounds of general formula [Ln(H<sub>2</sub>O)<sub>3</sub>(18C6)](ClO<sub>4</sub>)<sub>3</sub> (Ln = Tb<sup>III</sup>, Dy<sup>III</sup>, Er<sup>III</sup>, Yb<sup>III</sup>) and two of them were identified as fieldinduced SMMs.23

All of the previously mentioned compounds contained 18C6 bonded directly to the central lanthanide atom. However, several works also include the ether only as co-crystalized molecules. Zhao-Yang et al. explored the impact of chlorido and nitrato anions in two glycol coordinated dysprosium complexes where the purpose of crown-ether utilization was mainly to further separation of metal atoms and to promote crystallization. The prepared complexes [Dy(H2TEG)A3] (18C6) (H<sub>2</sub>TEG = triethylene glycol; A = Cl<sup>-</sup>, NO<sub>3</sub><sup>-</sup>) exhibit field induced two step magnetic relaxation having 28 K and 54 K energy barriers.24 Interestingly, the very simple compound Dy(NO<sub>3</sub>)<sub>3</sub>·6H<sub>2</sub>O or more precisely [Dy(NO<sub>3</sub>)<sub>3</sub>(H<sub>2</sub>O)<sub>4</sub>]·2H<sub>2</sub>O was reported recently as field-induced SMM.25 The fact that second coordination sphere and its impact on magnetic behaviour of SMMs remains partly overlooked with the exception of work by Petit et al. on Co(II) complex,<sup>26</sup> inspired us to study the influence of the second coordination sphere on the electronic structure and magnetic properties of Kramers ions based Dy(III) and Er(III)

complexes [Ln(NO<sub>3</sub>)<sub>3</sub>(H<sub>2</sub>O)<sub>3</sub>]·(18C6) for which we report the static and dynamic magnetic properties and theoretical calculations based on CASSCF multireference method.

# Results and discussion

### Synthesis and characterization

The infrared spectroscopy measurements confirmed the presence of the characteristic vibration bands associated to bonds in nitrato (NO<sub>3</sub>), water (OH) and crown ether (CH). The complete crystallographic information and structural description can be acquired from original literature.27 This work focuses predominantly on magnetic properties arising from covalent and noncovalent interactions in the studied compounds and thus the basic characterization of the crystal structures will be briefly described. The phase purity of the prepared compounds 1 and 2 was confirmed by powder X-ray diffraction (Fig. S1, ESI†). Both compounds of general formula  $[Ln(NO_3)_3(H_2O)_3]\cdot (18C6)$ , Ln = Dy (1), Er (2), are isostructural and the complex molecules are composed of three water and three nitrate molecules coordinated to the metal atom by the oxygen donor atoms forming thus polyhedron with coordination number nine and geometry closest to muffin (Table S1, ESI†) as evaluated by the Shape algorithm.28

All three nitrato ligands coordinate metal atoms in bidentate fashion with the Ln–O bond lengths ranging from 2.38 to 2.43 Å (in 1 and 2). The metal-ligand bond lengths are slightly shorter in the case of the aqua ligands: 2.36-2.39 Å (1) and 2.34-2.39 Å (2). The crystal structures of 1 and 2 are composed of the [Ln(NO<sub>3</sub>)<sub>3</sub>(H<sub>2</sub>O)<sub>3</sub>] complex molecules assembled into 1D chains by O-H···O hydrogen bonding between the aqua ligands and intercalated 18C6 molecules. Two of three coordinated aqua ligands are symmetry related by the mirror plane and form a pair of rather strong O-H···O hydrogen bonds (d(O···O)) =2.82-2.87 Å in 1 and 2 solely with one 18C6 molecule (Fig. 1). The third aqua ligand act as a bridging moiety, which interconnects two 18C6 molecules via two mirror related hydrogen bonds 2.832(11) for 1 and 2.814(11) Å for 2 (Fig. 1). Further stabilization is provided by plethora of weaker interactions

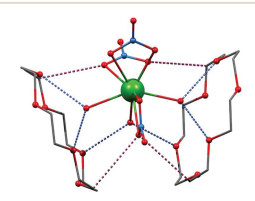


Fig. 1 The molecular structure of 1 showing the second coordination sphere formed by the net of O-H···O hydrogen bonds between O-H atoms of coordinated water and oxygen of crown ether (blue dashed lines) and C-H···O (nitrato) contacts (violet dashed line).

including non-covalent contacts between the aliphatic part (CH<sub>2</sub> donor groups) of 18C6 and nitrate oxygen atoms forming weak C-H···O (nitrato) hydrogen bonds (d(C···O) = 3.33-3.61 Åin 1 and 2) - Fig. 1 and S2.† The above-mentioned main supramolecular chain motif is extended to 2D layer by noncovalent O···O interaction between the nitrates of two adjacent complex molecules. This interaction is rather close with  $d(O \cdot \cdot \cdot O) = 2.957(16)$  in 1 and 2.986(16) Å in 2.

# Magnetic properties

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The temperature and field dependent static magnetic data were acquired on polycrystalline samples of 1 and 2 as depicted in Fig. 2. The value of  $\mu_{\rm eff}/\mu_{\rm B}$  at room temperature is 10.3 for 1 and 9.2 for 2, which are values close to the theoretically expected values of 10.65 (Dy<sup>III</sup>,  ${}^{6}H_{15/2}$ ) and 9.58 (Er<sup>III</sup>,  ${}^{4}I_{15/2}$ ). The  $\mu_{\rm eff}/\mu_{\rm B}$  is gradually decreasing upon cooling the samples down to 1.9 K both for 1 and 2, which is the most likely due to depopulation of ligand field multiplets arising from ground atomic terms effected by spin-orbit coupling and ligand field of the chromophores. There are no maxima on susceptibility, which eliminates existence of substantial intermolecular contacts providing magnetic exchange interactions of the antiferromagnetic nature.

The reciprocal susceptibilities were analysed with the Curie-Weiss law in the temperature range of 1.9-300 K for 1 and in the temperature range of 50-300 K for 2 (Fig. S3, ESI†), which resulted in  $C = 1.648 \times 10^{-4} \text{ m}^3 \text{ mol}^{-1} \text{ K}$ ,  $\Theta = -2.2 \text{ K}$  and g = -2.2 K1.28 for 1 and  $C = 1.405 \times 10^{-4} \,\mathrm{m}^3 \,\mathrm{mol}^{-1} \,\mathrm{K}$ ,  $\Theta = -18.8 \,\mathrm{K}$  and g

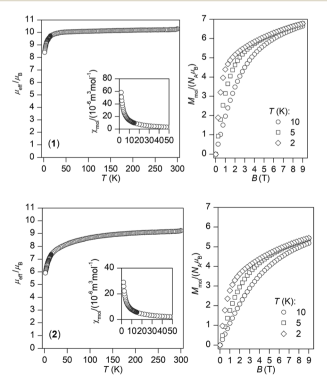


Fig. 2 Magnetic data for complexes 1 and 2. Temperature dependence of the effective magnetic moment (left) and the isothermal molar magnetizations measured at various temperatures (right)

= 1.18 for 2. All the Weiss constants are of negative values and gfactors are close to theoretical Landé g-factors, i.e. 1.33, and 1.20 for DyIII and ErIII, respectively. The isothermal magnetization data,  $M_{\rm mol}/N_{\rm A}\mu_{\rm B}$ , measured at T=2 K saturate to 6.8 for 1 and 5.4 for 2 and these values are well below theoretically predicted values based on I and Landé g-factors, which are 10.0 for DyIII and 9.0 for ErIII. This points out to large magnetic anisotropy of these complexes.

The AC susceptibility measurements were made firstly in zero and nonzero static magnetic fields as depicted in Fig. S4 (ESI).† None of these compounds showed nonzero out-of-phase signal of AC susceptibility at zero static magnetic field, but evidently, small magnetic field is efficient to observe slow relaxation of magnetization and suppress tunnelling effect. Therefore, the temperature dependence of AC susceptibility was measured at  $B_{\rm DC}=0.1~{\rm T}$  for frequencies 1–1500 Hz as depicted in Fig. 3 and 4. Evidently, the relaxation of the compound 1 is rather complicated, exhibiting three relaxation pathways - there are three maxima in the frequency-dependent plot of  $\chi''$  (Fig. 3 bottom). Usually, the existence of the multiple relaxation processes is assigned to the intermolecular interactions in solid state. However, the shortest distance between Dy atoms is quite large and equals 7.927 Å, thus is seems that such interactions should be negligible. On contrary, it was recently demonstrated that multiple relaxation processes can show up even in well isolated single-ion metal complexes due to the existence of the multilevel energy spectrum, thus they are of the intramolecular origin.<sup>29</sup> The relaxation pattern in 2 is much simpler, exhibiting just one dominant relaxation process, but without well-defined maxima (Fig. 4). Therefore, the simplified model was utilized to analyze AC susceptibility data for both compounds, which is derived under the assumption that the adiabatic susceptibility is usually approaching zero in the single-molecule magnets, thus it holds  $\chi_S \to 0$ ,

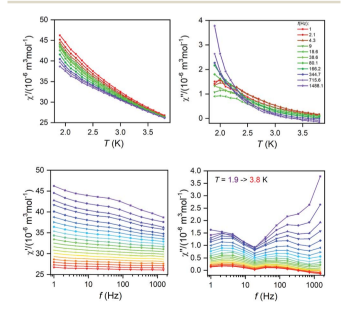


Fig. 3 Temperature and frequency dependent in-phase  $\chi'$  and out-ofphase  $\chi''$  molar susceptibilities for 1 at the applied external field  $B_{DC} =$ 0.1 T. Lines serve as guide to the eye

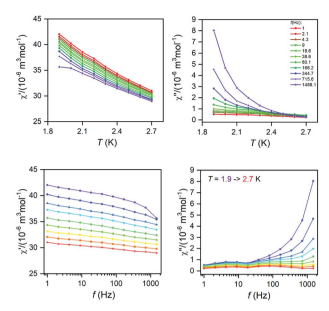


Fig. 4 Temperature and frequency dependent in-phase  $\chi'$  and out-of-phase  $\chi''$  molar susceptibilities for **2** at the applied external field  $B_{DC}=0.1$  T. Lines serve as quide to the eye.

$$\chi' = \frac{\chi_{\rm T} - \chi_{\rm S}}{1 + \omega^2 \tau^2} + \chi_{\rm S} \tag{1}$$

$$\chi'' = \varpi \tau \frac{\chi_T - \chi_S}{1 + \omega^2 \tau^2} \tag{2}$$

can be approximated as

$$\chi''/\chi' \cong \varpi\tau = 2\pi f\tau \tag{3}$$

Such approximation was already used to analyze magnetic data for other SMM compounds<sup>30</sup> and parameters of Orbach relaxation processes are derived from linear regression analyses using following equation

$$\ln(\chi''/\chi') = \ln(2\pi f \tau_0) + U_{\text{eff}}/kT \tag{4}$$

The analyses of ac susceptibility data for highest frequencies resulted in the magnetic moment reversal barrier parameters  $U_{\rm eff}=66$ –71 K for 1 (Fig. S5, ESI†) and  $U_{\rm eff}=21$ –24 K for 2 (Fig. S6, ESI†).

### Theoretical calculations

With the aim to explore the impact of the second coordination sphere on the electronic structure and magnetic properties of 1 and 2, the post-Hartree–Fock multireference calculations were done utilizing Complete Active Space Self-Consistent Field method (CASSCF). Thus, MOLCAS 8.0 computational package was employed for CASSCF calculations with respective relativistic ANO-RCC basis set and Douglas–Kroll–Hess (DKH) approximation accounting for the relativistic effects. First, the CASSCF calculations were done for the complexes  $[Ln(NO_3)_3(H_2O)_3]$  (Ln = Dy and Er), which resulted in the

energies of the ligand-field multiplets showed in Fig. 5, where a scheme of the magnetization blocking barrier is depicted.

The respective effective g-factors of the Kramers doublets are listed in Tables S2 and S3.† There is large axial anisotropy of the lowest Kramers doublet for both complexes  $(g_z \gg g_x, g_y)$  and the orientation of the respective easy axis of the magnetization is showed in Fig. 5 and 6. The energy of the first excited Kramers doublet is  $\varepsilon_1 = 143$  K for Dy<sup>III</sup> and  $\varepsilon_1 = 42$  K for Er<sup>III</sup> complex and the easy axis of these excited states is also depicted in Fig. 5 and 6. Secondly, the similar calculations were done for the complexes  $[Ln(NO_3)_3(H_2O)_3] \cdot (18C6)_2$  (Ln = Dy and Er), in which two non-coordinated 18-crown-6 molecules bound [Ln(NO<sub>3</sub>)<sub>3</sub>(H<sub>2</sub>O)<sub>3</sub>] through the non-covalent interactions form the second coordination sphere. Surprisingly, the incorporating the second coordination sphere caused deep impact on the energy level splitting of  ${}^6H_{15/2}$  (Dy<sup>III</sup>) and  ${}^4I_{15/2}$  (Er<sup>III</sup>) ground states as showed in Fig. 5 and 6. The energy of the first excited state was lowered for both compounds as  $\varepsilon_1 = 57$  K for Dy<sup>III</sup> and  $\varepsilon_1 = 16 \text{ K for Er}^{\text{III}}$  complex. Next, the ground state Kramers doublet properties were also significantly modified, the rhombic g-factors,  $g_x$  and  $g_y$  increased for Dy<sup>III</sup> complex (Table S4†), which resulted in the larger probabilities of the quantum tunnelling (Fig. 5).

On contrary, the probability of the quantum tunnelling decreases in  ${\rm Er^{III}}$  complex (Fig. 6) due to smaller rhombic g-

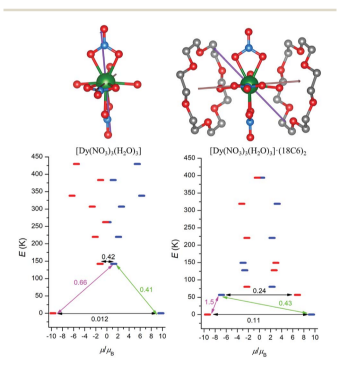


Fig. 5 Top: The molecular structures of  $[Dy(NO_3)_3(H_2O)_3]$  and  $[Dy(NO_3)_3(H_2O)_3] \cdot (18C6)_2$  used for CASSCF calculations overlaid with the easy axes of the ground and first excited Kramers doublets coloured with violet and brown colour, respectively. Bottom: *ab initio* computed magnetization blocking barrier for both molecular structures. The thick blue/red bars indicate the Kramer's doublets (KDs) as a function of magnetic moment. Green lines indicate the magnetization reversal mechanism. The magenta lines show the possible pathway of the Orbach process. The black lines represent the presence of QTM/TA-QTM between the connecting pairs.

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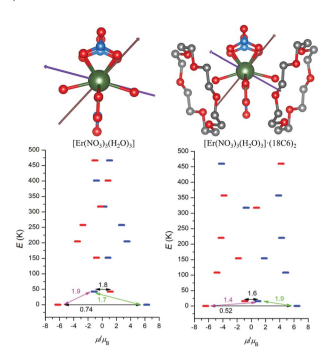


Fig. 6 Top: The molecular structures of  $[Er(NO_3)_3(H_2O)_3]$  and  $[Er(NO_3)_3(H_2O)_3] \cdot (18C6)_2$  used for CASSCF calculations overlaid with the easy axes of the ground and first excited Kramers doublets coloured with violet and brown colour, respectively. Bottom: *ab initio* computed magnetization blocking barrier for both molecular structures. The thick blue/red bars indicate the Kramer's doublets (KDs) as a function of magnetic moment. Green lines indicate the magnetization reversal mechanism. The magenta lines show the possible pathway of the Orbach process. The black lines represent the presence of QTM/TA-QTM between the connecting pairs.

factors  $g_x$  and  $g_y$  (Table S5†). To summarize, these quantum calculations agree with the experimental observations: (i) both compounds 1 and 2 are field-induced SMMs, which is consistent with calculated quantum tunnelling probabilities, (ii) the derived effective reversal barriers  $U_{\text{eff}} = 66-71 \text{ K}$  for 1 and  $U_{\text{eff}} =$ 21–24 K for 2 are rather close to calculated ones,  $U_{\rm calc.} = 57$  K for 1 and  $U_{\rm calc.} = 16$  K for 2. To better understand the substantial effect of the second coordination sphere on the calculated energy levels and magnetic anisotropy, the additional DFT calculations were done. The B3LYP functional was used to calculate the electrostatic potential retaining the molecular geometry for the ligand environment of the previous calculations  $[Ln(NO_3)_3(H_2O)_3]$  and  $[Ln(NO_3)_3(H_2O)_3] \cdot (18C6)_2$  (Ln = Dy and Er) and removing the central Ln atoms. Then, the electrostatic potential was calculated in a sphere of 1 Å radius located at the lanthanide atom Ln position. The color mapped plots are shown in Fig. 7 and S7 (ESI)† and it is evident that incorporating 18C6 molecules into the calculations results in less negative electrostatic potential, which rationalize the results of CASSCF calculations in such a way, that weakening of the crystal-field potential around central lanthanide atoms leads to the smaller energy splitting. Thus, the non-covalent contacts of the 18C6 molecules in the second coordination sphere are sufficient to induce significant changes of the electrostatic potential, hence the crystal/ligand-field resulting in the variation of the multiplets energy levels.

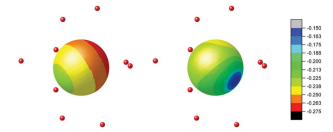


Fig. 7 The electrostatic potentials of the ligands projected in a sphere of 1 Å radius located at the dysprosium position in molecular systems  $[Dy(NO_3)_3(H_2O)_3]$  (left) and  $[Dy(NO_3)_3(H_2O)_3] \cdot (18C6)_2$  (right) obtained by removing the dysprosium atom. The color scale of the electrostatic potential is shown on the right. The donor oxygen atoms from nitrato and aqua ligands are shown as red balls.

# **Experimental section**

All the materials were purchased in Sigma-Aldrich and used without any purification. Basic chemical composition was determined by Flash 2000 (Thermo Scientific) analyser. IR measurements were performed on FT-IR spectrometer Thermo Nicolet Nexus 670, magnetic measurements on PPMS Dynacool (Quantum Design) and SQUID magnetometer XL-7 (Quantum Design). Structure was confirmed by X-ray diffractometer MiniFlex600 (Rigaku). The procedure for synthesis of  $[M(NO_3)_3(H_2O)_3]\cdot(18C6)$  was performed similarly as described in literature.  $^{27}$ 

## Characterization

Lanthanide salt of  $Dy(NO_3)_3 \cdot 6H_2O$  (0.329 mmol) or  $Er(NO_3)_3 \cdot 5H_2O$  (0.271 mmol) was dissolved in mixture of acetonitrile and methanol 3 : 1. To this solution a stoichiometric amount of solid 18C6 was added which resulted in forming precipitate that redissolved quickly. The mixture was left stirring for approximately 15 minutes, filtrated and left to evaporate slowly. After a few days, bulky crystals were collected, washed with methanol and dried under vacuum in desiccator containing silica gel.

[Dy(NO<sub>3</sub>)<sub>3</sub>(H<sub>2</sub>O)<sub>3</sub>]·(18C6) 1a. Colourless plates, isolated 117 mg, 53%. EA(calc.) for  $C_{12}H_{30}DyN_3O_{18}$  (FW = 666,88): C, 21.61; H, 4.53; N, 6.30. Found: C, 21.56; H, 4.64; N, 6.60. FT-IR (mid ATR,  $\nu$ /cm<sup>-1</sup>): 3345  $\nu$ (H<sub>2</sub>O); 3232  $\nu$ (H<sub>2</sub>O); 2907  $\nu$ (CH); 2873  $\nu$ (CH); 2845  $\nu$ (CH); 1632  $\nu$ (H<sub>2</sub>O); 1495 (NO<sub>3</sub>); 1271 (NO<sub>3</sub>).

[Er(NO<sub>3</sub>)<sub>3</sub>(H<sub>2</sub>O)<sub>3</sub>]·(18C6) 1b. Pink plates, isolated 101 mg, 55%. EA(calc.) for  $C_{12}H_{30}ErN_3O_{18}$  (FW = 671,64): C, 21.46; H, 4.50; N, 6.26. Found: C, 21.52; H, 4.72; N, 5.84. FT-IR (mid ATR,  $\nu/cm^{-1}$ ): 3334  $\nu(H_2O)$ ; 3233  $\nu(H_2O)$ ; 2907.2  $\nu(CH)$ ; 2873.6  $\nu(CH)$ ; 2845  $\nu(CH)$ ; 1633  $\nu(H_2O)$ ; 1495 (NO<sub>3</sub>); 1273 (NO<sub>3</sub>).

### Theoretical calculations

The respective CASSCF calculations were performed with the MOLCAS 8.0 program package³¹ on the  $[Ln(NO_3)_3(H_2O)_3]$  and  $[Ln(NO_3)_3(H_2O)_3]\cdot(18C6)_2$  (Ln = Dy and Er) molecular systems using atomic coordinates derived from the experimental X-ray structures (CSD codes LEYHUS and LEYJII). The active space was defined by seven f-orbitals as CAS(9,7) for 1 and CAS(11,7)

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for 2. The RASSCF method was employed in CASSCF calculations<sup>32</sup> with the following numbers of multiplets: 21 sextets, 224 quartets and 490 doublets for DyIII, 35 quartets and 112 doublets for ErIII. The spin-orbit coupling based on atomic mean field approximation (AMFI)33 was considered using RASSI-SO with the following numbers of multiplets: 21 sextets, 128 quartets and 130 doublets for DyIII, 35 quartets and 112 doublets for ErIII. The relativistic effects were treated with the Douglas-Kroll Hamiltonian.34 The following basis sets were utilized: Ln.ANO-RCC-VQZP (Ln = Dy and Er), O.ANO-RCC-VDZP, N.ANO-RCC-VDZP, C.ANO-RCC-VDZ and H.ANO-RCC-VDZ.35 Then, SINGLE\_ANISO module36 was used to calculate the effective spin Hamiltonian parameters for Kramers doublets of the ground states and the respective parameters of the blocking energy barrier. The calculated easy axes were visualized with VESTA 3 program.37

The DFT calculations were done with ORCA 4.0 software<sup>38</sup> utilizing B3LYP functional39 together with relativistic basis sets DKH-def2-TZVP for N and O atoms, and DKH-def2-SVP for C and H atoms. 40 Increased integration grids (Grid6 in ORCA convention) and tight SCF convergence criteria were used in all calculations.

# Conclusions

The magnetic properties of the compounds [Ln(NO<sub>3</sub>)<sub>3</sub>(H<sub>2</sub>O)<sub>3</sub>]·(18C6) (Ln = Dy (1)) and Er (2)) were studied both experimentally and theoretically. The AC susceptibility confirmed the onset of slow relaxation of the magnetization typical for SMMs under applied static magnetic field. The  $Dy^{III}$  compound 1 evidenced three relaxation pathways in contrast to ErIII compound 2 for which only one relaxation pathway was found. The analysis of the experimental AC magnetic data revealed  $U_{\rm eff} = 66$ -71 K for 1 and  $U_{\rm eff} = 21$ –24 K for 2. The subsequent CASSCF calculations were done both for the complexes with the primary coordination sphere composed of aqua and nitrato ligands [Ln(NO<sub>3</sub>)<sub>3</sub>(H<sub>2</sub>O)<sub>3</sub>], and for complexes comprising also the secondary coordination sphere of two 18-crown-6 molecules [Ln(NO<sub>3</sub>)<sub>3</sub>(H<sub>2</sub>O)<sub>3</sub>]·(18C6)<sub>2</sub>. It was found that secondary coordination sphere significantly influenced the calculated energy splitting, hence the magnetization reversal barrier and magnetic anisotropy of the resulting Kramers doublets both for 1 and 2. Moreover, the calculated magnetization reversal barriers,  $U_{\text{calc.}} = 57 \text{ K}$  for 1 and  $U_{\text{calc.}} = 16 \text{ K}$  for 2, after incorporating the second coordination sphere are in good agreement with the experimental ones emphasizing the importance of non-covalent interactions induced by 18-crown-6 cocrystallized molecules on nitrato and aqua ligands. This was also supported by DFT calculations of the electrostatic potentials around metal atoms, which rationalized the CASSCF results.

# Conflicts of interest

There are no conflicts to declare.

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# Notes and references

- 1 N. Ishikawa, M. Sugita, T. Ishikawa, S. Koshihara and Y. Kaizu, J. Am. Chem. Soc., 2003, 125, 8694-8695.
- 2 (a) R. Sessoli, L. Hui, A. R. Schake, S. Wang, J. B. Vincent, K. Folting, D. Gatteschi and G. Christou, J. Am. Chem. Soc., 1993, 115, 1804-1816; (b) R. Sessoli, D. Gatteschi, A. Caneschi and M. A. Novak, *Nature*, 1993, 365, 141–143.
- Gatteschi, R. Sessoli and J. Villain, Molecular Nanomagnets; Oxford University Press, Oxford, 2006.
- 4 (a) Molecular Magnets Physics and Applications, ed. J. Bartolomé, F. Luis and J. F.Fernández, Springer-Verlag, Berlin, Heidelberg, 2014; (b) B. Warner, F. E. Hallak, H. Prüser, J. Sharp, M. Persson, A. J. Fisher and C. F. Hirjibehedin, *Nat. Nanotechnol.*, 2015, **10**, 259–263; (c) A. Ardavan, O. Rival, J. J. L. Morton, S. J. Blundell, A. M. Tyryshkin, G. A. Timco and R. E. P. Winpenny, Phys. Rev. Lett., 2007, 98, 057201.
- 5 M. N. Leuenberger and D. Loss, *Nature*, 2001, 410, 789-793; L. Bogani and W. Wernsdorfer, Nat. Mater., 2008, 7, 179-186.
- 6 (a) D. N. Woodruff, R. E. P. Winpenny and R. A. Layfield, Chem. Rev., 2013, 113, 5110-5148; (b) P. Zhang, Y.-N. Guo and J. Tang, Coord. Chem. Rev., 2013, 257, 1728-1763; (c) F. Pointillart, O. Cador, B. Le Guennic and L. Ouahab, Coord. Chem. Rev., 2017, 346, 150-175; (d) A. Dey, P. Kalita and V. Chandrasekhar, ACS Omega, 2018, 3, 9462-9475.
- 7 (a) J. Long, F. Habib, P.-H. Lin, I. Korobkov, G. Enright, Ungur, W. Wernsdorfer, L. F. Chibotaru and M. Murugesu, J. Am. Chem. Soc., 2011, 133, 5319-5328; (b) J. D. Rinehart and J. R. Long, *Chem. Sci.*, 2011, 2, 2078–2085.
- 8 J. Liu, Y.-C. Chen, J.-L. Liu, V. Vieru, L. Ungur, J.-H. Jia, L. F. Chibotaru, Y. Lan, W. Wernsdorfer, S. Gao, X.-M. Chen and M.-L. Tong, J. Am. Chem. Soc., 2016, 138, 5441-5450.
- 9 Y.-S. Ding, N. F. Chilton, R. E. P. Winpenny and Y.-Z. Zheng, Angew. Chem., Int. Ed., 2016, 55, 16071-16074.
- 10 (a) F. Guo, B. M. Day, Y. Chen, M. Tong, A. Mansikkamäki and R. A. Layfield, Angew. Chem., Int. Ed., 2017, 56, 11445-11449; (b) C. A. P. Goodwin, F. Ortu, D. Reta, N. F. Chilton and D. P. Mills, Nature, 2017, 548, 439-442.
- 11 F.-S. Guo, B. M. Day, Y.-C. Chen, M.-L. Tong, A. Mansikkamäki and R. A. Layfield, Science, 2018, eaav0652.
- 12 S.-D. Jiang, B.-W. Wang, H.-L. Sun, Z.-M. Wang and S. Gao, J. Am. Chem. Soc., 2011, 133, 4730-4733.
- 13 S.-M. Chen, J. Xiong, Y.-Q. Zhang, Q. Yuan, B.-W. Wang and S. Gao, Chem. Sci., 2018, 9, 7540-7545.
- 14 Y.-S. Meng, C.-H. Wang, Y.-Q. Zhang, X.-B. Leng, B.-W. Wang, Y.-F. Chen and S. Gao, Inorg. Chem. Front., 2016, 3, 828-835.
- 15 R. L. Carlin, *Magnetochemistry*, Springer-Verlag, Berlin, 1986.
- 16 L. Escalera-Moreno, J. J. Baldoví, A. Gaita-Ariño and E. Coronado, Chem. Sci., 2018, 9, 3265-3275.
- 17 Y.-S. Ding, K.-X. Yu, D. Reta, F. Ortu, R. E. P. Winpenny, Y.-Z. Zheng and N. F. Chilton, Nat. Commun., 2018, 9, 3134.

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18 (a) C. Pettinari, F. Marchetti and A. Drozdov, in Comprehensive Coordination Chemistry II, 2003, vol. I, pp. 211-246; (b) F. T. Edelmann, Coord. Chem. Rev., 2014, 261,

- 73-155; (c) M. J. Hardie and C. L. Raston, J. Chem. Soc., Dalton Trans., 2000, 15, 2483-2492.
- 19 S. Nishihara, T. Akutagawa, T. Hasegawa and T. Nakamura, Inorg. Chem., 2003, 42, 2480-2482.
- 20 S. Petrosyants, Z. Dobrokhotova, A. Ilyukhin, N. Efimov, Y. Mikhlina and V. Novotortsev, Inorg. Chim. Acta, 2015, 434, 41-50.
- 21 H. Wada, S. Ooka, T. Yamamura and T. Kajiwara, Inorg. Chem., 2016, 56, 147-155.
- 22 Y.-S. Ding, T. Han, Y.-Q. Hu, M. Xu, S. Yang and Y.-Z. Zheng, Inorg. Chem. Front., 2016, 3, 798-807.
- 23 L. Maxwell, M. Amoza and E. Ruiz, Inorg. Chem., 2018, 57, 13225-13234.
- 24 G. Peng, Y.-Y. Zhang and Z.-Y. Li, Inorg. Chem. Commun., 2017, 77, 40-43.
- 25 E. Mamontova, J. Long, R. Ferreira, A. Botas, D. Luneau, Y. Guari, L. Carlos and J. Larionova, Magnetochemistry, 2016, 2, 41.
- 26 S. Petit, G. Pilet, D. Luneau, L. F. Chibotaru and L. Ungur, Dalton Trans., 2007, 4582-4588.
- 27 R. D. Rogers and A. N. Rollins, J. Chem. Crystallogr., 1994, 24, 321-329.
- 28 (a) M. Pinsky and D. Avnir, Inorg. Chem., 1998, 37, 5575-5582; (b) A. Ruiz-Martínez, D. Casanova and S. Alvarez, Chem. - Eur. J., 2008, 14, 1291-1303; (c) A. Ruiz-Martínez, D. Casanova and S. Alvarez, Dalton Trans., 2008, 19, 2583-2591.
- 29 L. T. A. Ho and L. F. Chibotaru, Phys. Rev. B, 2016, 94, 1-5. 30 (a) J. Bartolomé, G. FIloti, V. Kuncser, G. Schinteie, V. Mereacre, C. E. Anson, A. K. Powell, D. Prodius and C. Turta, Phys. Rev. B: Condens. Matter, 2009, 80, 014430; (b) R. Ishikawa, R. Miyamoto, H. Nojiri, B. K. Breedlove and M. Yamashita, Inorg. Chem., 2013, 52, 8300-8302; (c) I. Nemec, R. Marx, R. Herchel, P. Neugebauer, J. van Slageren and Z. Travnicek, Dalton Trans., 2015, 44, 15014-15021.
- 31 (a) F. Aquilante, L. De Vico, N. Ferré, G. Ghigo, P. Å. Malmqvist, P. Neogrády, T. B. Pedersen, M. Pitoňák,

- M. Reiher, B. O. Roos, L. Serrano-Andrés, M. Urban, V. Veryazov and R. Lindh, J. Comput. Chem., 2010, 31, 224-247; (b) J. A. Duncan, J. Am. Chem. Soc., 2009, 131, 2416; (c) G. Karlström, R. Lindh, P. Å. Malmqvist, B. O. Roos, Ryde, V. Veryazov, P.-O. Widmark, M. Cossi, B. Schimmelpfennig, P. Neogrády and L. Seijo, Comput. Mater. Sci., 2003, 28, 222-239; (d) V. Veryazov, P.-O. Widmark, L. Serrano-Andres, R. Lindh and B. O. Roos, Int. J. Quantum Chem., 2004, 100, 626-635.
- 32 P. Å. Malmqvist, B. O. Roos and B. Schimmelpfennig, Chem. Phys. Lett., 2002, 357, 230-240.
- 33 (a) B. A. Hess, C. M. Marian, U. Wahlgren and O. Gropen, Chem. Phys. Lett., 1996, 251, 365-371; B. Schimmelpfennig, AMFI, an atomic mean-field spinorbit integral program, Stockholm University, 1996.
- 34 (a) N. Douglas and N. M. Kroll, Ann. Phys., 1974, 82, 89-155; (b) B. A. Hess, Phys. Rev. A, 1986, 33, 3742-3748.
- 35 (a) B. O. Roos, R. Lindh, P. A. Malmqvist, V. Veryazov and P.-O. Widmark, J. Phys. Chem. A, 2008, 112, 11431-11435; (b) B. O. Roos, R. Lindh, P. A. Malmqvist, V. Veryazov and P.-O. Widmark, Chem. Phys. Lett., 2005, 409, 295-299.
- 36 (a) L. F. Chibotaru, L. Ungur and A. Soncini, Angew. Chem., Int. Ed., 2008, 47, 4126-4129; (b) L. F. Chibotaru, L. Ungur, C. Aronica, H. Elmoll, G. Pillet and D. Luneau, J. Am. Chem. Soc., 2008, 130, 12445-12455; (c) L. F. Chibotaru and L. Ungur, J. Chem. Phys., 2012, 137, 064112-064122; (d) L. Ungur, M. Thewissen, J.-P. Costes, W. Wernsdorfer and L. F. Chibotaru, Inorg. Chem., 2013, 52, 6328-6337.
- 37 K. Momma and F. Izumi, J. Appl. Crystallogr., 2011, 44, 1272-127639.
- 38 (a) F. Neese, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2012, 2, 73-78; (b) F. Neese, Wiley Interdiscip. Rev.: Comput. Mol. Sci., 2018, 8, e1327.
- 39 (a) A. D. Becke, Phys. Rev. A: At., Mol., Opt. Phys., 1988, 38, 3098-3100; (b) C. Lee, W. Yang and R. G. Parr, Phys. Rev. B: Condens. Matter Mater. Phys., 1988, 37, 785-789; (c) P. J. Stephens, F. J. Devlin, C. F. Chabalowski and M. J. Frisch, J. Phys. Chem., 1994, 98, 11623-11627.
- 40 F. Weigend and R. Ahlrichs, Phys. Chem. Chem. Phys., 2005, 7, 3297-3305.