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Slow magnetic relaxation in Ni–Ln (Ln = Ce, Gd, Dy) dinuclear complexes

METAMORPHOSIS: The preparation of a gadolinium(III) complex with an unsymmetrical coordination environment yields a product with unexpected single-molecule-magnet behavior. Related Ce(III) and Dy(III) compounds have also been prepared and their magnetism has been characterised.



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## Introduction

Since the identification of the first single-molecule magnet (SMM)  ${Mn_{12}}$  in 1993,<sup>1</sup> research on this phenomenon has continued apace and has also been extended to lanthanide based materials.<sup>2</sup> As early as 2004, heteronuclear 3d-4f complexes were observed to display slow magnetic relaxation.<sup>3</sup> The advantage of transition metal and lanthanide complexes lies in a fusion of the properties of both. In fact, the magnetic study of complexes comprising transition metals and lanthanides dates back to 1985, when Gatteschi et al. observed unexpected ferromagnetic coupling between copper $\left(\mathrm{II}\right)$  and gadolinium $(m)$  ions.<sup>4</sup>

More recent research has yielded a few examples of a significant reduction in quantum tunnelling magnetization

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# Slow magnetic relaxation in Ni–Ln (Ln = Ce, Gd, Dy) dinuclear complexes†

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Three new isomorphous complexes  $[Ni(O-van-en)LnCl<sub>3</sub>(H<sub>2</sub>O)] [H<sub>2</sub>(O-van-en) = N,N'-ethylene-bis(3-1)$ methoxysalicylaldiminate; Ln = Ce  $(1)$ , Gd  $(2)$ , Dy  $(3)$ ] were prepared by a stepwise reaction using mild conditions and were structurally characterised as dinuclear molecules in which Ni and Ln are coordinated by the compartmental Schiff base ligand (o-van-en)<sup>=</sup> and doubly bridged by O atoms. While the nickel(III) centre is diamagnetic within the  $N_2O_2$  square-planar coordination of the Schiff base ligand, the lanthanide atoms are octa-coordinated to give an  $\{LnCl<sub>3</sub>O<sub>5</sub>\}$  chromophore with a fac-arrangement of the chlorido ligands. AC magnetic measurements revealed that all three complexes, including the nominally isotropic Gd(III) system, show field induced slow magnetic relaxation with two or three relaxation channels: at  $T =$ 1.9 K the low-frequency relaxation time is  $\tau_{LF}(1) = 0.060$  s at  $B_{DC} = 0.5$  T,  $\tau_{LF}(2) = 0.37$  s at  $B_{DC} = 0.3$  T, and  $\tau_{LF}(\mathbf{3}) = 1.29$  s at  $B_{DC} = 0.15$  T. PAPER<br>
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(QTM) due to non-negligible magnetic exchange interactions between lanthanide and transition metal ions. $5$  A recent study on 3d–4f complexes with diamagnetic 3d metal ions such as zinc(II) or cobalt(III) showed an enhancement of the  $U_{\text{eff}}$  barrier in comparison to their mononuclear lanthanide analogues;<sup>6</sup> it was suggested that the presence of a diamagnetic 3d cation near the lanthanide central atom, with a monatomic O bridge, induces a large charge polarisation on the bridging oxygen atom that favours an increase in the  $U_{\text{eff}}$  barrier. This observation suggests a new strategy in designing 3d–4f complexes with diamagnetic 3d ions.<sup>5,7-10</sup>

Regarding the 4f elements, heavy lanthanide ions were widely used for the preparation of SMM materials due to their large angular momentum  $J$  and large magnetic anisotropy.<sup>11-13</sup> However, a recent study of light lanthanide ions revealed surprising magnetic behaviour for cerium $(m)$ .<sup>14</sup> Despite the small magnetic moment, which arises from its small angular momentum  $J$  and small Landé  $g$  factor, the spin–orbit coupling is strong enough to create significant magnetic anisotropy and thus an energy barrier between the two orientations of the magnetization. The first cerium $(m)$  based SMM was reported in 2013<sup>15</sup> and later other cerium( $\text{m}$ ) complexes were reported to be SMMs. $16-19$ 

Among the lanthanide ions, the case of gadolinium $(m)$ complexes represents another breakthrough in the theory of slow relaxation of magnetization. Despite its essentially isotropic ground state with  $L = 0$ , several materials have been reported for which magnetic anisotropy around the gadoli $nium(m)$  centre was induced by local coordination, magnetic exchange coupling or electron density donation.<sup>20-29</sup> These



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materials containing gadolinium $(m)$  may be considered as members of a new class of SMMs with the highest known spin state  $(4f^7)$  for a single ion.

Some considerable effort has been extended to produce strictly dinuclear 3d-4f molecular complexes<sup>30-32</sup> to serve as building blocks for the further preparation and study of polynuclear clusters $33-36$  and polymers. $37-39$  A related fact in the context of the present work is that salen-type Schiff bases originating from o-vanillin and related aldehydes have been used in coordination chemistry due to their ability to form complexes with a rich variety of geometries and structural dimensions.<sup>40</sup>

In what follows we describe the preparation, chemical and structural characterization and magnetic properties of three isomorphous dinuclear  $[Ni(o\nuan-en)Ln(H<sub>2</sub>O)Cl<sub>3</sub>]$  complexes [Ln = Ce (1), Gd (2), Dy (3); H<sub>2</sub>L = H<sub>2</sub>(o-van-en) = N,N'-ethylenebis(3-methoxysalicylaldiminate)]. The choice of Ce(III),  $Dy(m)$ and Gd(III) was designed to expose the main differences in the magnetic behaviours of a light lanthanide ion, a heavy lanthanide ion and the lanthanide ion with a spherical orbital ground state, respectively.

## Results and discussion

#### Synthesis and identification

The three complexes  $[Ni(o\text{-}van\text{-}en)LnCl<sub>3</sub>(H<sub>2</sub>O)]$   $[Ln = Ce (1),$ Gd (2), Dy (3)] were first synthesized in microcrystalline form using mild solution techniques in a modification of the method of Costes and co-workers.<sup>41</sup> The synthetic procedure is depicted in Scheme 1. The products were characterized using chemical analysis and IR spectroscopy; the observed absorption bands are listed in the ESI.† The imine group absorption,  $\nu$ (C=N), observed slightly above 1600 cm<sup>−1</sup>, can be considered as a characteristic band. In the IR spectrum of the ligand, this vibration was observed at 1631 cm−<sup>1</sup> which agrees with the reported value of 1628 cm−<sup>1</sup> reported for the same Schiff base,<sup>42</sup> and is similar to the value of 1625 cm<sup>-1</sup> for a related Schiff base L  $[L = N, N'-bis([E]-3-)$ -phenylallylidene)-3-methyl-1,2phenylenediamine].<sup>43</sup> The ligand bonded in the complex [CdLX<sub>2</sub>] (X = Cl, Br, I, SCN) exhibits a shift of 1–7 cm<sup>-1</sup> to lower wavenumbers. In the spectra of the complexes 1, 2 and 3 the vibration  $\nu$ (C=N) was observed at 1622–1623 cm<sup>-1</sup>.

Recrystallization of these microcrystalline products or reaction by diffusion techniques yielded single crystals of 1–3. The phase identities of the microcrystalline products with the corresponding single-crystal samples were established by LeBail refinement using the powder diffraction patterns for all three samples (see Fig. S1†).

#### Crystal structures

Compounds 1–3 are isomorphous, so we will limit our description and discussion to complex 1 with the corresponding geometric parameters of 2 and 3 given in parentheses. We note that all three crystal structures were studied at room temperature. The crystal structure of 1 consists of neutral heterodi-



Scheme 1 Schematic representation of the synthetic route for preparation of 1, 2 and 3.

nuclear molecules of  $[Ni(o\nuan-en)Ce(H_2O)Cl_3]$  (Fig. 1). The  $Ni(n)$  atom is coordinated in square-planar form by the  $N_2O_2$ donor set of the potentially ditopic ligand  $(o$ -van-en)<sup>2-</sup>. The  $Ce(m)$  central atom is octa-coordinated by four oxygen atoms from the  $(o$ -van-en)<sup>2-</sup> ligand, one aqua and three chlorido ligands, yielding an  ${LnCl<sub>3</sub>O<sub>5</sub>}$  chromophore (Fig. 1). Calculations using the program  $SHAPE^{44}$  indicate that the coordination polyhedron is close to a bicapped (O1 and O4) trigonal prism [Fig. S2;† the same holds for the Dy $(m)$  and  $Gd(m)$  complexes]. It is significant (vide infra) that the three chlorido ligands form a single face of the polyhedron.

Analogous  $[Ni(o\text{-}van\text{-}en)Ln(NO_3)_3]\cdot (CH_3)_2CO$  heterodinuclear complexes with auxiliary nitrato ligands and contain-



Fig. 1 Molecular structure of  $[Ni(o-van-en)Ce(H<sub>2</sub>O)Cl<sub>3</sub>]$ ; thermal ellipsoids are drawn with 50% probability.

ing various lanthanide atoms have been reported; $45,46$  but to our knowledge no analogous complexes with other 3d elements with auxiliary chlorido ligands have been studied. On the other hand, eight analogous heterodinuclear complexes containing chlorido ligands were reported with modified Schiff base ligands. $34,47-51$  An interesting example is the complex  $[NiCl_2(o\text{-}van-bu)YbCl(H_2O)_3]$  with a hexacoordinated Ni(II) central atom (donor set  $Cl_2O_2N_2$ ).<sup>48</sup>

The observed Ni–N and Ni–O as well as the Ce–O bond distances are similar to those reported in the analogous complexes  $[Ni_2(o\text{-}van\text{-}en)_2Ce(NO_3)_2](NO_3)$ ,<sup>52</sup> ( $[Cu(o\text{-}van\text{-}en)Gd(NO_3)_3]$ . 0.25H<sub>2</sub>O,<sup>53</sup> [Co(*o-van-en*)(CH<sub>3</sub>COO)<sub>2</sub>Dy(NO<sub>3</sub>)<sub>2</sub>]<sup>54</sup>) (Table 1). The Ln–Cl bond distances in 1 are in the range of  $2.717(2)$ –2.802 $(2)$ Å [2.630(3)–2.714(3) Å for 2 and 2.6005(10)–2.6955(9) Å for 3], which can be compared to the average Ce–Cl distance of 2.7467 Å (2.6829 Å for Gd–Cl and 2.6357 Å for Dy–Cl) found in the CSD.<sup>55</sup> The CSD further reveals that in all Ni–Ln (Ln = Ce, Gd, Dy) with Ln–Ni distances shorter than 5 Å the two metal atoms are bridged by monatomic O bridges, and the mean values of the Ni–Ln distances are 3.58 Å, 4.53 and 4.45 Å for Ce–Ni, Gd–Ni and Dy–Ni, respectively. In our compounds 1–3 the observed Ln–Ni distances are significantly shorter than the corresponding calculated mean values (Table 1). The remaining geometric parameters in all three complexes exhibit expected values. **Datton Terractions**<br>
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Although molecular compounds with Ln–Cl bonds are known, these represent only about 6% of all molecular lanthanide complexes.55 Those in which the lanthanide is bonded to three terminal chlorido ligands are just around 1%. For comparison, the number of molecular lanthanide compounds with three coordinated  $NO<sub>3</sub><sup>-</sup>$  groups is more than five times that with three Cl<sup>−</sup> ligands. Nevertheless, given the large number of molecular Ln compounds that have been characterized, there are some 400 compounds in the latter class –  $Ln-Cl<sub>3</sub>(terminal)$ . The only eight heterodinuclear Tr–Ln complexes with Schiff bases similar to the one in compounds 1–3 and with chlorido ligands bonded to the lanthanide metal that have been structurally characterized have only one or two chlorido ligands bonded to the lanthanide, $34,47-51$  not three as in compounds 1–3. Also, two related compounds with three Cl<sup>−</sup> ligands bonded to the same lanthanide metal have been found, but with higher nuclearity, *i.e.*,  $Tr<sub>2</sub>Ln<sub>2</sub>$  compounds in which some of the chlorido ligands act as bridges between the lanthanide centers.47,56

There are four dinuclear molecules per unit cell of 1 (and likewise for 2 and 3). The packing of the complex molecules is governed by medium-strength hydrogen bonds of the O–H⋯Cl type yielding a chain-like supramolecular structure running along the *a* axis with additional, weaker C<sub>ar</sub>-H…Cl, N=C-H…Cl type hydrogen bonds and weak  $\pi-\pi$  stacking interactions of the aromatic rings of the neighbouring chains forming supramolecular layers in the  $ab$  plane (Fig. S3 and Table S2 $\dagger$ ). There are only close contacts of the C–H⋯Cl type between planes, involving methyl and methylene groups of the ligand.

#### DC magnetic data

In compounds  $1-3$  the central Ni( $\pi$ ) atom is in a square planar environment formed by four oxygen donor atoms. Thus, it is magnetically silent, except for some temperature-independent magnetism arising from the presence of low-lying excited states. The lanthanide  $Ln(m)$  centres bear orbital and spin angular momentum and the ground multiplet is  ${}^{2}F_{5/2}$  for Ce(m),  ${}^{8}S_{7/2}$  for Gd(m) and  ${}^{6}H_{15/2}$  for Dy(m), with the magnetogyric ratios  $g_I = 6/7$ , 2 and 4/3, respectively. Thus the magnetization per formula unit should saturate to  $M_1 = M_{\text{mol}}/(N_A \mu_B)$  =  $g_T J = 6/7 \times 5/2 = 15/7$ ,  $2 \times 7/2 = 7$ , and  $4/3 \times 15/2 = 10$ . The hightemperature limit of the effective magnetic moment, based upon the magnetic susceptibility, is  $\mu_{eff}/\mu_B = g_I [J(J + 1)]^{1/2}$  $(6/7)(5/2 \times 7/2)^{1/2} = 2.54$ ,  $2 \times (7/2 \times 9/2)^{1/2} = 7.94$ , and  $(4/3)$  $(15/2 \times 17/2)^{1/2} = 10.6$ .

The DC magnetic data for 1 are shown in Fig. 2. Clearly visible linear dependence of the effective magnetic moment between  $T = 9-300$  K indicates some temperature-independent paramagnetism as expected for the  $Ni(II)$  centre in a square planar environment:  $\chi_{\text{TIP}} = 6 \times 10^{-9} \text{ m}^3 \text{ mol}^{-1}$ . With the above correction the room-temperature effective magnetic moment (the high-temperature limit) adopts a value of  $\mu_{\text{eff}} = 2.36 \mu_{\text{B}}$ , that is not far from the single ion value (2.54). The magnetization data at  $T = 2.0$  and  $B = 7$  T,  $M_1 = 1.0$ , is much lower relative to isolated multiplet  ${}^{2}F_{5/2}$  for Ce(III) (2.1).

Owing to the effect of the crystal field, the six-fold degenerate ground atomic multiplet  ${}^{2}F_{5/2}$   $|R'_{3}:J;M_{J}\rangle = |5/2;\pm 1/2,$ 



Table 1 Selected geometric parameters in 1, 2 and 3 with their s.u.'s



Fig. 2 DC magnetic data for 1. Solid lines – fitted. Colored circles – TIP corrected data.

 $\pm 3/2, \pm 5/2$ ) is split into three crystal-field multiplets (Kramers doublets) within the respective double group:  $|R'_{3}: J; M_{J}\rangle \stackrel{\text{crystal field}}{\longrightarrow} |G': I'; \gamma'\rangle$ . The magnetic data was fitted assuming that only the lowest Kramers doublet is thermally populated. Then the effective Hamiltonian in use is

$$
\hat{H}_a^{\text{eff}} = \mu_B B (g_z^{\text{eff}} \hat{f}_z^{\text{eff}} \cos \theta_a + g_{xy}^{\text{eff}} \hat{f}_x^{\text{eff}} \sin \theta_a)
$$
(1)

with  $f^{\text{eff}} = 1/2$ ; *a* denotes grids distributed uniformly over the meridian. To this end, the optimization routine converged to the following set of magnetic parameters:  $g_{xy}^{\text{eff}}$  = 0.80(19),  $g_{z}^{\text{eff}}$  = 4.50(1), and the molecular-field correction (operative at low temperature)  $zj/hc = -0.27$  cm<sup>-1</sup>. Large  $g_z$  and small  $g_{xy}$  are typical values for the anisotropic lanthanide ions.

Ab initio calculations for 1 gave the values  $g(0.035, 0.251, 0.035)$ 3.880} that support the magnetic data analysis. The first excited Kramers doublet lies 300  $cm^{-1}$  above the ground one and its thermal population at  $T = 300$  K is only 9%.

The DC magnetic data for 2 confirms the behaviour of the  ${}^{8}S_{7/2}$  ground term (Fig. 3): the effective magnetic moment is almost constant over the whole temperature range with  $\mu_{\text{eff}}$  = 7.8–7.9 $\mu_{\rm B}$ ; the magnetization per formula unit at  $B = 7.0$  T and  $T = 2.0$  K saturates to  $M_1 = 6.6$ . This data is consistent with an isotropic g-factor slightly below 2.0. The data fitting to the Curie law and the Brillouin function gave  $g = 1.952(6)$ . A superposition of the magnetization data vs.  $BT^{-1}$  confirms an invisible magnetic anisotropy of 2.

The DC magnetic data for 3 is shown in Fig. 4. The effective magnetic moment displays the high-temperature (HT) limit of  $\mu_{\text{eff}}$  = 12.15 $\mu_{\text{B}}$  that decreases only slightly down to T ~ 60 K; then it decreases more rapidly to  $11.1\mu_B$  at the lowest temperature of the measurements. The observed HT limit is somewhat higher than that expected for a single  $Dy(m)$  ion (10.6). This might be due to the presence of low-lying excited states of the cooperating  $Ni(n)$  centre. More surprising is the saturation value of the magnetization  $M_1(J) = 7.9$  at  $T = 2.0$  and  $B = 7.0$  T, which differs substantially from the single-ion expectation  $(M_1 = 10)$ . This can arise in a partial quenching of the orbital angular momentum of the  $Dy(m)$  centre by the asymmetric ligand field of the  ${DyO_4O'Cl_3}$  chromophore. Another reason could be the zero-field splitting of the ground crystal-field multiplet inherent to an anisotropic ion such as  $Dy(m)$ . In such a



Fig. 3 DC magnetic data for 2. Solid lines – fitted. The state of temperatures.



Fig. 4 DC magnetic data for 3. Dashed lines are visual guides.

case, the higher terms of the series of the crystal field potential will also contribute

$$
\hat{V} = \sum_{k=0,2,4,6} \sum_{m=0}^{+\hat{k}} B_k^m \cdot \hat{O}_k^m(\hat{J}_z, \hat{J}_\pm)
$$
 (2)

where  $\hat{O}_{k}^{m}(\hat{J}_{z}\hat{J}_{\pm})$  are the appropriate equivalent operators. The sixteen members of the  ${}^{6}H_{15/2}$  multiplet are then split into eight Kramers doublets.

#### Field scan of the AC susceptibility

The AC susceptibility data was acquired at the field amplitude of  $B_{AC}$  = 0.38 mT and depending upon the external magnetic field  $B_{\text{DC}}$ , temperature, and the frequency f of the oscillating field.

The first scan for a variable magnetic field, fixed low temperature  $T = 2.0$  K, and a set of four representative frequencies  $f = 1.1$ , 11, 111, and 1116 Hz is shown in Fig. 5. It is seen that the out-of-phase susceptibility  $\chi''$  is zero at  $B_{\text{DC}} = 0$ , which is traditionally explained as an effect of the fast magnetic tunnelling. With the applied external field, the out-of-phase susceptibility rises to a maximum and then attenuates at higher fields. The position of the maximum slightly depends upon the frequency of the oscillating field. The subsequent data measurements were done at the external field for which the out-ofphase susceptibility component displays a maximum:  $B_{\text{DC}}$  = 0.1 or 0.5, 0.3, and 0.15 T for 1 through 3.

#### Temperature scan of the AC susceptibility

Temperature evolution of the AC susceptibility components for a set of frequencies varying between  $f = 0.1$  and 1500 Hz at a selected external field  $B_{\text{DC}}$  is shown in Fig. S4 (see ESI†). It can be concluded that the samples 1 through 3 show field induced slow magnetic relaxation typical for single ion magnets (SMM). On heating, the in-phase susceptibility components for various frequencies tend to merge at the critical temperature  $(ca. 10, >7,$ and 7 K); at the same time the out-of-phase component vanishes since then the system behaves as an ordinary paramagnet.

#### Frequency scan of the AC susceptibility for 3, Ni–Dy complex

The same data sets have been rearranged to the frequency dependence of the AC susceptibility components for a set of



Fig. 5 AC magnetic data for 1, 2, and 3. Field scans for a set of frequencies at low temperature.

For 3, it is evident (Fig. 6) that the out-of-phase component consists of three contributions: low-frequency (LF), intermediate-frequency (IF), and the high-frequency (HF). For the lowesttemperature measurements  $(T = 1.9 \text{ K})$  the LF component dominates with the peak-position at  $f_{\text{max}}$  ~ 0.16 Hz. Then the corresponding relaxation time is estimated to be as long as



Fig. 6 AC magnetic data for 3. Solid lines – fitted by the extended three-set Debye model.

 $\tau(LF) = 1/(2\pi f_{\text{max}}) \sim 0.995$  s. Both, in-phase and out-of-phase susceptibilities were fitted simultaneously by employing the three-set Debye model, as explained in the ESI.† Using the resulting relaxation parameters (thermal susceptibilities  $\chi_{Tk}$ , distribution parameters  $\alpha_k$ , and relaxation times  $\tau_k$ ,  $k = 1-3$ ), the convoluted susceptibility components are drawn as solid lines in Fig. 6. It is evident that these lines pass through the experimental points and this is also supported by the statistical analysis in the ESI.†

At low temperature the HF component recovers only a tail; however, with increasing temperature the LF component decays progressively in favor of the HF one; the IF component balances in-between. At temperature  $T > 5$  K, the IF component cannot be fitted satisfactorily and then only a two-set Debye model is applicable. At the lowest measurement temperature,  $T = 1.9$  K, the mole fractions of the individual relaxation species are  $x(LF) = 0.49$ ,  $x(F) = 0.28$ , and  $x(HF) = 0.23$ , respectively. At the same time, the three relaxation times are  $\tau(LF)$  = 1.29 s,  $\tau(\text{IF}) = 74$  ms, and  $\tau(\text{HF}) = 964$  μs. In contrast, at  $T =$ 6.3 K,  $x(LF) = 0.13$  and  $x(HF) = 0.87$  hold true.

In Fig. 7, the Argand diagram is shown; it is a convolution of three distorted semicircles (arcs) of different height, position and width. Also an Arrhenius-like plot  $\ln \tau$  vs.  $T^{-1}$  is shown on the right for the individual relaxation processes. The three highest-temperature points for the HF relaxation channel have been fitted by a straight line in order to get parameters for the Orbach relaxation process  $\tau = \tau_0 \exp(U_{\text{eff}}/k_{\text{B}}T)$ :  $U_{\text{eff}}/k_{\text{B}} =$ 65 K and  $\tau_0 = 1.9 \times 10^{-10}$  s.

The temperature evolution of the HF relaxation time can be represented by various graphs as shown in Fig. S5 (see ESI†). The dependence  $\ln \tau$  vs.  $\ln T$  at its two edges can be recovered by straight lines:  $\ln \tau = -8.81 + 11.3(\ln T)$  at the highest temperatures, and  $\ln \tau = -D_{\tau}$  at the lowest temperatures. Therefore, the whole dataset has been fitted via the equation

$$
\tau^{-1} = CT^n + D_\tau \tag{3}
$$

where the first term refers to the Raman process and the second one the quantum tunnelling process. The optimization routine gave  $n = 10.8(3)$ ,  $C = 3.6(16) \times 10^{-4} \text{ K}^{-n} \text{ s}^{-1}$ , and



Fig. 7 Argand diagram (left) and the Arrhenius-like plot (right) for 3. Straight line – high-temperature limit of the Arrhenius equation referring to the Orbach process. Solid curve – fitted using Raman and quantum tunneling processes.

 $D_{\tau}$  = 1.0(1) × 10<sup>3</sup> s<sup>-1</sup>. This means that the high-temperature limit of the high-frequency relaxation process can be equally reproduced by the Raman process itself. A value of  $n = 12$  has been found for a  $Co(n)$  compound.<sup>57</sup>

#### Frequency scan of the AC susceptibility for 1, Ni–Ce complex

The frequency dependence of the AC susceptibility components for 1 is shown in Fig. 8. For  $T = 1.9$  K the well-developed peak of  $\gamma''$  at  $f \sim 10$  Hz implies the relaxation time  $\tau(LF) \sim$ 16 ms. This is much shorter relative to 3. However, the peak is slightly asymmetric with an arm, and its reliable fit requires a two-set Debye model where the HF component is also considered. On heating the peak is shifted to higher frequencies but for  $T > 4$  K the two-set model becomes unstable; then the single-set model has been applied for higher temperatures. The resulting relaxation parameters are listed in the ESI† along with statistical analysis.

The Argand diagram and the Arrhenius-like plot for 1 are displayed in Fig. 9. In this case the LF relaxation time was fitted *via* eqn (3) giving  $n = 6.59(7)$ ,  $C = 174(22) \times 10^{-4} \text{ K}^{-n} \text{ s}^{-1}$ , and  $D_{\tau} = 51(1) \text{ s}^{-1}$ . At the same time the parameters of the Orbach relaxation process are  $U_{\text{eff}}/k_{\text{B}} = 43 \text{ K}$  and  $\tau_0 = 3.1 \times 10^{-7} \text{ s}$ .

As expected, the external magnetic field has a visible effect on the relaxation behaviour of 1 (see ESI†): the relaxation time



Fig. 8 AC magnetic data for 1. Solid lines – fitted by the extended twoset Debye model.



Fig. 9 Argand diagram (left) and the Arrhenius-like plot (right) for 1. Full circles – single-set Debye model; solid curve – fitted with Raman & tunneling process; dashed – three-point linear extrapolation for the Orbach process. Fig. 11 Argand diagram (left) and the Arrhenius-like plot (right) for 2.

at  $T = 1.9$  K and  $B_{\text{DC}} = 0.5$  T becomes  $\tau(LF) = 60.3(5)$  ms as compared to 18.7(11) ms at  $B_{\text{DC}} = 0.1$  T. A successful data fitting requires the two-set Debye model for  $T > 3.5$  K. SIM behaviour of the Ce( $\text{III}$ ) ions has been reviewed recently.<sup>14</sup> For instance, AC susceptibility studies of single-molecule magnet behaviour in polynuclear assembly of  $Ce(m)$  with polymolybdates reveal a two-channel slow relaxation with the relaxation times  $\tau = 6009$ and 217 μs at  $T = 1.9$  K and  $B_{\text{DC}} = 0.02$  T.<sup>18</sup> There is a sizable field influence since at  $B_{\text{DC}}$  = 0.14 T the relaxation times merge to  $τ = 605$  μs.

#### Frequency scan of the AC susceptibility for 2, Ni–Gd complex

The frequency dependence of the AC susceptibility components for the Gd compound 2 is shown in Fig. 10. In this case the two-set Debye model reproduces the experimental data with well separated branches. The existence of slow relaxation is unexpected in the case of the isotropic gadolinium centre. Even more surprising is the Arrhenius-like plot drawn in Fig. 11 where the HF relaxation time is only slightly temperature dependent between 1.9 and 6.5 K. Moreover, the HF branch shows a "strange" behaviour (Fig. S6†): on cooling the relaxation time passes through a maximum and then it is shortened:  $\tau(HF) = 223 \text{ }\mu\text{s}$  at  $T = 3.9 \text{ as compared to } \tau(HF) = 120 \text{ }\mu\text{s}$ at  $T = 1.9$  K. Paper<br>  $D_x = 1.0(1) \times 10^3 \text{ s}^3$ . This means that the high-temperature at  $T = 1.9$  K access Common Maximum article is like the state of the st

The strange behaviour has been observed recently in  $Cu(II)$ , Ni( $\pi$ ), Co( $\pi$ ), and Mn( $\pi$ ) complexes showing distorted octa-



Fig. 10 AC magnetic data for 2. Solid lines – fitted by the extended two-set Debye model.



hedral geometry.<sup>58–61</sup> It was possible to reproduce such data by using a phenomenological equation

$$
\tau^{-1} = CT^n + E_\tau T^{-k} \tag{4}
$$

that contains the conventional Raman term (parameters C and  $n > 0$ ), along with the "strange" term (parameters  $E<sub>\tau</sub>$  and  $k > 0$ ). Linear fits for the truncated data allowed bracketing starting values for data fitting *via* eqn (4); the final parameters are:  $n =$ 1.18(26),  $C = 4.7(26) \times 10^2 \text{ K}^{-n} \text{ s}^{-1}$ ,  $E_\tau = 2.4(3) \times 10^4 \text{ K}^k \text{ s}^{-1}$ , and  $k = 1.7(2)$ . Two slow relaxation processes have also been described for several  $Gd(m)$  compounds.<sup>20,22,23,28,29</sup> For instance, for a  $Gd(m)EDTA$  chelate compound,<sup>29</sup> the values of  $U_{\text{eff}}$  = 6.1 K ( $\tau$  = 4 × 10<sup>-2</sup> s) and  $U_{\text{eff}}$  = 84 K ( $\tau$  = 8 × 10<sup>-7</sup> s) were reported  $(B_{\text{DC}} = 0.45 \text{ T}).$ 

The geometry and the electron density of the coordination sites has been proved to exert a very important influence on the SMM behaviour of lanthanide ions. There are few studies of the influence of ions and since the number of chlorido derivatives is smaller than for instance, nitrato, the studies reported about the influence of the chlorido ligands in the SMM behaviour are very rare and it may require more research.

The magnetic study on the  $\left[ \text{Ln}_2(\text{mq})_4 \text{X}_6 \right]$ (EtOH)<sub>2</sub> (Fig. 12, left) (mq = 8-hydroxy-2-methylquinoline) compounds which have been prepared for Ln = Dy, with  $X = NO_3^-$  or Cl<sup>−</sup> shows that for these compounds the energy barrier is higher when  $X = Cl^{-} (NO_3^-: \Delta/k_B = 40.0 \text{ K and } Cl^{-}: \Delta/k_B = 102.4 \text{ K}).^{62} \text{ We}$ note that for Ln = Gd,  $X = NO_3^-$ , Cl<sup>-</sup>, no slow relaxation was observed. However, a recent study on  $[Ln(tpm)X_3]\cdot yMeCN^{63}$ (tpm = tris(3,5-dimethyl-pyrazolyl)methane) shows that at zero DC field, no significant out-of-phase susceptibility was detected for Ln = Dy or Er, none for  $X = NO_3^-$  nor  $X = Cl^-$ , but under applied fields, it was observed for  $Ln = Dy$  and  $NO_3^-$  but not for Cl−, while for Ln = Er, the situation was the opposite, indicating that  $NO<sub>3</sub><sup>-</sup>$  in this case is a more suitable anion for obtaining SMM with Dy while Cl<sup>−</sup> is a better anion for Er. No such comparative information for  $Ln = Gd$  has been reported, to our knowledge.

All three of our compounds, including the  $Gd(m)$  one, with three Cl<sup>−</sup> ligands and under applied external field, show slow relaxation. We note that the location of the three coordinated chlorido atoms in our compounds (Fig. 12, right) is different

**D** CI3

Fig. 12 Left: [Ln<sub>2</sub>(mq)<sub>4</sub>Cl<sub>6</sub>] showing the disposition of the Cl<sup>−</sup> ligands. Right: Molecular drawing of compounds 1-3 showing the "fac" disposition of the three clorido ligands.

than in the above-mentioned compounds  $\left[ \text{Ln}_2(\text{mq})_4\text{Cl}_6 \right]$ . From Fig. 12, it can be seen that in compounds 1–3 the chlorido ligands are in what can be called fac-configuration – that is, concentrated in the same part of the coordination sphere, while the other compounds present a more symmetrical distribution; that has to affect the electron distribution in the coordination sphere.

### **Conclusions**

Using mild chemical conditions a new kind of 3d–4f dinuclear NiLn Schiff base compound,  $[Ni(o\nuan-en)LnCl<sub>3</sub>(H<sub>2</sub>O)]$  (Ln = Ce, Gd, Dy) was prepared and structurally characterized. In all three complexes the square four-coordinated nickel $(n)$  atom and octacoordinated Ln( $\text{m}$ ) atom (donor set O<sub>5</sub>Cl<sub>3</sub>) are located in the neighbouring compartments provided by the Schiff base ligand and linked by two monatomic O bridges. The coordination of the Ln atom is highly unsymmetrical with all chlorido ligands situated in a "fac" fashion. AC susceptibility measurements reveal that all three complexes show field supported slow magnetic relaxation. This result in the case of the  $Gd(m)$  compound suggests a need for further detailed magnetic studies, in the field of SMM, of complexes with the chlorido ligands concentrated in a face of the coordination polyhedron. AC magnetic measurements revealed that all three complexes, including the nominally isotropic  $Gd(m)$  system, show field induced slow magnetic relaxation with two or three relaxation channels: at  $T = 1.9$  K the low-frequency relaxation time is  $\tau_{LF}(1) = 0.060$  s at  $B_{DC} = 0.5$  T,  $\tau_{LF}(2) = 0.37$  s at  $B_{DC} = 0.3$  T, and  $\tau_{LF}(3) = 1.29$  s at  $B_{DC} = 0.15$  T. **Duton Transactors**<br>
Such a possible to reproduce such data by  $\pi$  that in the such situation comparison in the subsequent state of the duties<br>  $\tau^2 = C\tau^2 = B, \tau^{-4}$  (4) Equation for 8 PM. The intermediate and in composit

### **Experimental**

#### Syntheses

Ethylenediamine, o-vanillin, NiCO<sub>3</sub>, GdCl<sub>3</sub>·6H<sub>2</sub>O, CeCl<sub>3</sub>·7H<sub>2</sub>O,  $DyCl<sub>3</sub>·6H<sub>2</sub>O$ , ethanol and diethyl ether, all of analytical grade, were purchased from commercial sources and used as received.

Synthesis of the ligand  $H_2$ (*o-van-en*). For the synthesis of  $H<sub>2</sub>(o-van-en)$  a modification of the previously reported method $^{35}$  was used. Ethylenediamine  $(0.1 \text{ cm}^3, 1.5 \text{ mmol})$  was added to 50 cm<sup>3</sup> of an ethanol solution of  $o$ -vanillin (0.4542 g, 3 mmol). The mixture was placed in a boiling flask and refluxed for 5 hours. The final solution was filtered and left for crystallization. After a few hours, dark yellow crystals of the product appeared. The crude product was filtered and washed with 2 ml of diethylether. Yield: 90%. Purity and identity of the product were checked by CHN analysis, IR, UV-Vis, <sup>1</sup>H and <sup>13</sup>C NMR spectroscopies. The corresponding analytical data for  $H_2$ (o-van-en) as well as for complexes 1, 2 and 3 are listed in ESI.†

Synthesis of  $\left[\text{Ni}(o\text{-}van\text{-}en)\text{Ce}(H_2O)Cl_3\right](1)$ . Solid  $H_2(o\text{-}van\text{-}en)$  $(2 \text{ g}, 6 \text{ mmol})$  and nickel(II) carbonate  $(0.7 \text{ g}, 6 \text{ mmol})$  were added to 100  $\text{cm}^3$  of water and heated in an open beaker. After one hour of reaction the resulting brown microcrystalline

product (crude nickel complex) was separated by filtration, washed with 2 ml of ethanol and dried in air. The dry product was used in the next step of the synthesis without further purification. Solid CeCl<sub>3</sub>·7H<sub>2</sub>O (0.3 g, 0.8 mmol) was dissolved in 10 cm<sup>3</sup> of ethanol and the resulting clear solution was added to the brown ethanol solution of  $[Ni(\rho\nu an-en)]$  (50 cm<sup>3</sup> of ethanol). After 15 min of stirring, the orange microcrystalline complex  $[Ni(o\text{-}van\text{-}en)Ce(H_2O)Cl_3]$  (1) was separated by filtration and washed three times with 2 ml of ethanol. Single crystals suitable for X-ray analysis were obtained by recrystallization of the microcrystalline complex using room temperature diffusion of a MeOH solution into iPrOH.

Synthesis of  $\left[\text{Ni}(o\text{-}van\text{-}en)\text{Gd}(\text{H}_2\text{O})\text{Cl}_3\right]$  (2). The same procedure was used as for the  $Ce(m)$  complex with the modification that instead of CeCl<sub>3</sub>·7H<sub>2</sub>O, GdCl<sub>3</sub>·6H<sub>2</sub>O (0.3 g, 0.8 mmol) was used. Single crystals of  $\text{[Ni}(o\text{-}van\text{-}en)\text{Gd}(\text{H}_2\text{O})$  $Cl<sub>3</sub>$ ] (2) were obtained by crystallization using diffusion of a MeOH solution of GdCl3 into an iPrOH solution of [Ni(o-vanen)] at room temperature.

Synthesis of  $[Ni(o\text{-}van\text{-}en)Dy(H_2O)Cl_3]$  (3). For the synthesis of the complex  $[Ni(o\text{-}van\text{-}en)Dy(H_2O)Cl_3]$  the same procedure was used as for the  $Ce(m)$  complex with the modification that instead of CeCl<sub>3</sub>·7H<sub>2</sub>O, DyCl<sub>3</sub>·6H<sub>2</sub>O (0.3 g, 0.8 mmol) was used. Crystals of  $[Ni(o\text{-}van\text{-}en)Dy(H_2O)Cl_3]$  (3) were obtained by recrystallization using diffusion of a methanol solution of the microcrystalline product into iPrOH with a temperature gradient of from 70 °C at the bottom of the beaker to room temperature at the top of the solution.

#### Physical measurements

Elemental analyses (C, H, N) were performed on a PerkinElmer 2400 Series II CHNS/O analyser. Infrared spectra were recorded on a PerkinElmer Spectrum 100 CsI DTGS FTIR Spectrometer with UATR 1 bounce-KRS-5 in the range of 4000–300  $\text{cm}^{-1}$ . Powder X-Ray data were collected using a RIGAKU D/max 2500 diffractometer equipped with a copper rotating anode, operating at 40 kV and 80 mA, with a graphite monochromator. Measurements were made for  $2\theta$  from  $2.5^{\circ}$  to  $40^{\circ}$  in steps of 0.02° with a rate of 1 s per step. The magnetic susceptibility (determined as  $\chi = M/H$ ) was taken at  $B = 0.1$  T using a SQUID magnetometer (Quantum Design, MPMS-XL7) in the temperature range from 1.9 to 300 K. Raw susceptibility data was corrected for the diamagnetic contribution. The magnetization data was acquired at low temperature  $T = 2.0$  and 4.6 K until  $B_{\rm DC}$  = 7.0 T. The AC susceptibility data was taken at the applied external fields using the amplitude of the oscillating field  $B_{AC}$  = 0.38 mT for a set of temperatures and for 22 frequencies between  $f = 0.1$  to 1500 Hz.

#### Crystallography

X-ray experiments were carried out on a four-circle κ-axis Xcalibur2 diffractometer equipped with a Sapphire3 CCD detector (Oxford Diffraction). The CrysAlis software package [Oxford Diffraction 2006 CrysAlisPro CCD and CrysAlisPro RED] was used for data collection and reduction. Crystals of compound 1 were found to exhibit multiple domains. For the

crystal used for data collection, two domains were indexed and data integration was conducted simultaneously for both. The structures for  $1$ , 2 and 3 were solved using SHELXT.<sup>64</sup> Refinement based on intensities was performed using SHELXL-2014/7.<sup>65</sup> All non-hydrogen atoms were refined anisotropically. Hydrogen atoms were located in difference Fourier maps. For the final refinement, H atoms bound to C were placed at calculated positions and refined as riders with  $U_{\text{iso}}(H) = 1.2U_{\text{eq}}(C)$ for non-methyl H and with  $U_{\text{iso}}(H) = 1.5U_{\text{eq}}(C)$  for methyl H. Using the program CALC-OH the positions of the water hydrogen atoms for the O1 aqua ligand were found; their isotropic thermal parameters were tied with the parent atom  $[U_{\text{iso}}(H) = 1.2U_{\text{eq}}(O)]^{66}$  For compound 1 the final refinement was conducted using data from both domains. The structure figures were drawn using Diamond.<sup>67</sup> Crystal data and final parameters for the structure refinements are summarized in Table S1† and possible hydrogen bonds are given in Table S2.† Paper **Contents** (cross article complex) was separated by dittribution, crystal used for data collection, two contrasts were independent on a set of the synchronic synchronic synchronic synchronic interest are a set of a

#### Ab initio calculations

Ab initio calculations were performed with ORCA 4.1.2 computational package using the experimental geometry of complex under study.<sup>68</sup> The relativistic effects were included in the calculations with second-order Douglas-Kroll-Hess (DKH)<sup>69</sup> procedure together with the scalar relativistic contracted version of def2-SV(P) basis function for all elements, except Ce atom. For Ce, the SARC2-DKH-QZVP basis function was used.<sup>70</sup> The calculations were based on state average complete active space self-consistent field (SA-CASSCF) wave functions. The active space of the CASSCF calculations comprised of one electron in seven metal-based f-orbitals. The state averaged approach was used, in which 7 doublet states were equally weighted. The spin–orbit effects have been included through quasi-degenerate perturbation theory in which an approximation to the Breit–Pauli form of the spin–orbit coupling operator (SOMF) was utilized.<sup>71</sup>

## Conflicts of interest

There are no conflicts to declare.

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