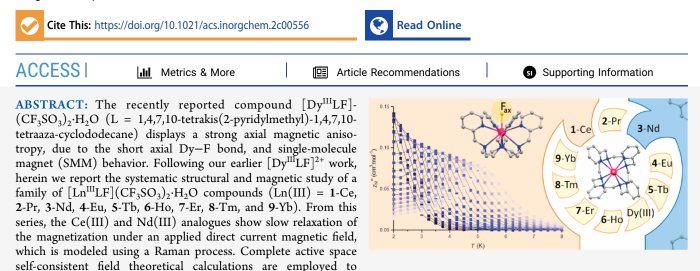
Inorganic Chemistry

Importance of an Axial Ln^{III}—F Bond across the Lanthanide Series and Single-Molecule Magnet Behavior in the Ce and Nd Analogues

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understand the relaxation pathways in 1-Ce and 3-Nd and also reveal a large tunnel splitting for 5-Tb. Additional computational studies on model compounds where we remove the axial F^- ligand, or replace F^- with I^- , highlight the importance of the F^- ligand in creating a strong axial crystal field for 1-Ce and 3-Nd and for promoting the SMM behavior. Importantly, this systematic study provides insight into the magnetic properties of these lighter lanthanide ions.

INTRODUCTION

Single-molecule magnets (SMMs) are molecular systems that have a bistable magnetic state and show magnetic hysteresis of molecular origin. They were studied for the first time in the 1990s after the magnetic characterization of $[Mn_{12}O_{12}(MeCO_2)_{16}(H_2O)_4] \cdot 2CH_3COOH \cdot 4H_2O_1^{-1}$ also known as $\{Mn_{12}\}^{2-4}$ Following studies on $\{Mn_{12}\}$, the main focus of the field was the study of molecular complexes containing 3d metals, such as $Mn_{,5,6}^{5,6}$ Fe,^{7,8} Ni,⁹ and Co.¹⁰ After 2003,¹¹ the interest shifted toward lanthanide ions because of their high intrinsic magnetic moments and strong spin–orbit coupling. Both properties are important in creating a magnetic easy axis that leads to improved SMM behavior.

Some of the great advances that have been reported, such as impressive magnetization reversal barriers (U_{eff}) and magnetic hysteresis at high temperatures,¹² have been focused around Dy(III), followed by Tb(III)/Tb(II) and Ho(III).^{13–17} While it is understandable to focus on Dy(III) due to its Kramers ion nature, large magnetic moment (J = 15/2), and high magnetic anisotropy, particularly useful properties when designing high-performance SMMs,^{18,19} there is still a shortage of studies on other lanthanide ions such as Ce(III) (²F_{5/2}) and Nd(III) (⁴I_{9/2}). Recent studies into the nature of the lanthanide relaxation,^{20,21} which is often found to dominate the spin dynamics in monometallic complexes of the early lantha-

nides.^{22,23} Indications for this include small values of the magnetization reversal barrier²⁴ that do not correspond to energy gaps between ground and excited m_J states and unreasonable values of the pre-exponential factor τ_0 when including an Orbach term.^{25,26} Understanding these relaxation processes is also a vibrant research area that is relevant to all spin-based systems.^{27,28}

Like Dy(III), both Ce(III) and Nd(III) have an oblate (equatorially expanded) electron distribution of the lowest J state. They are also both Kramers ions, which means their ground state is always doubly degenerate.¹⁸ This explains why one of the highest $U_{\rm eff}$ values recorded for Ce(III) and Nd(III) is for an aza-crown complex with nitrates in the axial positions.²⁹ In order to promote SMM behavior for such lanthanide ions, it is useful to design a coordination environment that creates strong axial magnetic anisotropy. If two suitable axial ligands are provided, such as in the case of pseudo- D_{5h} complexes of Dy(III),³⁰ one can obtain even zero-field single-ion magnets (SIMs) of Ce(III) and Nd(III), albeit

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with smaller energy barriers.^{31,32} Herein, we study a coordination environment that uses a fluoride ligand in the axial position to create a strong axial crystal field, following previous positive results with Dy(III) by us, using an octadentate N₈ ligand and Norel *et al.*, using a combination of a hexadentate N₆ ligand with either two pyridine or 1,4-dioxane ligands.^{33,34}

The ligand 1,4,7,10-tetrakis(2-pyridylmethyl)-1,4,7,10-tetraaza-cyclododecane, L, is a cyclen with four flexible "arms" containing pyridine and can encapsulate a metal through its eight nitrogen atoms (see Figure S1). The ligand L has been studied for its coordination with different metal ions,³⁵⁻ including lanthanides.^{42–47} With 3d metals, the ligand forms a pocket that fully encapsulates the metal. On the other hand, with lanthanides being larger cations, there is enough space in the first coordination sphere for another smaller ligand. This theory was tested by our group with the study of $[Dy^{III}LF]$ - $(CF_3SO_3)_2$ ·H₂O, which behaves as an SMM.³³ We showed that the fluoride ligand, due to its small size and negative charge, is an ideal candidate to create a strong axial magnetic anisotropy and generate the highest magnetization reversal barrier among a group of monodentate ligands (i.e., OH⁻, HCO₂⁻, ^tBuCO₂⁻, CF₃CO₂⁻, CH₃CH₂CH₂O⁻, and ^tBuCH₂CH₂O⁻) that were tested in silico.³³

Therefore, we decided to synthesize and study other analogues of this system with a terminal Ln(III)–F bond. Our synthetic strategy led to compounds with the formula [LnLF](CF₃SO₃)₂·H₂O: Ln(III) = Ce (1-Ce), Pr (2-Pr), Nd (3-Nd), Eu (4-Eu), Tb (5-Tb), Ho (6-Ho), Er (7-Er), Tm (8-Tm), and Yb (9-Yb) plus diluted compounds 10-Ce@La, 11-Nd@La, and 12-Tb@Y. We refer the interested reader to our earlier publication for full details on $[Dy^{III}LF](CF_3SO_3)_2$ ·H₂O.³³ Importantly, 1-Ce and 3-Nd show slow relaxation of the magnetization under an applied direct current (dc) field, expanding the limited family of reported Ce(III) and Nd(III) SMMs.

EXPERIMENTAL METHODS

All reagents were used as received without further purification. No safety hazards were encountered during the described experimental procedures.

1,4,7,10-Tetrakis(2-pyridylmethyl)-1,4,7,10-tetraaza-cyclododecane (L): 2-(chloromethyl) pyridine hydrochloride (2.51 g, 15.3 mmol) and cyclen (0.65 g, 3.87 mmol) were refluxed overnight in 60 mL of MeCN and excess Cs_2CO_3 (24.9 g, 76.4 mmol). The reflux gave a dark red solution that was filtered in vacuo and washed multiple times with CH_2Cl_2 . Rotary evaporation of the filtrate gave L in the form of yellow crystals (yield = 1.60 g, 78%). ¹H NMR in $CDCl_3$ at 298 K: δ 2.79 (16 H, s, NCH_2CH_2N), 3.66 (8 H, s, $NCH_2(C_5H_4N)$), 7.11 (4 H, ddd, J = 12.4 Hz, J = 7.8 Hz, J = 1.2 Hz, C_5H_4N), 7.74 (4 H, d, J = 7.8 Hz, C_5H_4N), 8.50 (4 H, dd, J = 4.8 Hz, J = 1.2 Hz, C_5H_4N), 1.67 (H₂O, s, solvent) 7.29 (CHCl₃, s, solvent) (see Figure S2).

For the synthesis of complexes 1-Ce to 12-Tb@Y, the following procedure is applicable: $Ln(CF_3SO_3)_3$ (0.08 mmol) was dissolved in 4 mL of MeOH along with L (47 mg, 0.08 mmol) and refluxed overnight. The solution was then dried in vacuo until an oil resulted, which was redissolved in 10 mL of MeCN and refluxed for a further 2 h. After drying the solution in vacuo again, the obtained oil was dissolved in 4 mL of CH₃Cl, stirred at high temperature, and dried in vacuo to obtain a white solid. The solid was dissolved in water with NH₄F (yield depending, 4 equiv) and stirred at high temperature for 10 min. After filtration and slow evaporation, colorless crystals formed in 2–3 days. For the 10% diluted analogues, 10-Ce@La to 12-Tb@Y, the same procedure is applicable by using a combination of salts,

0.072 mmol La(CF₃SO₃)₃ or Y(CF₃SO₃)₃ and 0.008 mmol Ln(CF₃SO₃)₃ (Ln(III) = Ce, Nd, and Tb) in the first step.

1-Ce [Ce(L)F](CF₃SO₃)₂·H₂O yield 53 mg. Elemental Anal. Calcd (found): C, 40.34 (40.30); H, 4.19 (4.05); N, 11.07 (10.94)%. **2**-Pr [Pr(L)F](CF₃SO₃)₂·H₂O yield 49 mg. Elemental Anal. Calcd

(found): C, 40.32 (40.43); H, 4.19 (4.15); N, 11.06 (11.03)%. **3**-Nd $[Nd(L)F](CF_3SO_3)_2$:H₂O yield 31 mg (56.3%). Elemental

Anal. Calcd (found): C, 40.19 (40.18); H, 4.17 (4.08); N, 11.03 (10.94)%.

4-Eu [Eu(L)F](CF₃SO₃)₂·H₂O yield 24 mg (44.4%). Elemental Anal. Calcd (found): C, 39.89 (39.90); H, 4.13 (4.03); N, 10.94 (10.88)%.

5-Tb [Tb(L)F](CF₃SO₃)₂·H₂O yield 14 mg (25.4%). Elemental Anal. Calcd (found): C, 39.62 (39.56); H, 4.11 (4.09); N, 10.87 (10.72)%.

6-Ho [Ho(L)F](CF₃SO₃)₂·H₂O yield 40 mg (30.5%). Elemental Anal. Calcd (found): C, 39.39 (39.61); H, 4.08 (4.07); N, 10.81 (10.89)%.

7-Er $[Er(L)F](CF_3SO_3)_2$ ·H₂O yield 20 mg (35.4%). Elemental Anal. Calcd (found): C, 39.30 (39.45); H, 4.07 (4.05); N, 10.78 (10.87)%.

8-Tm $[Tm(L)F](CF_3SO_3)_2$ ·H₂O yield 30 mg (22.7%). Elemental Anal. Calcd (found): C, 39.24 (39.48); H, 4.07 (4.02); N, 10.77 (10.9)%.

9-Yb [Yb(L)F](CF₃SO₃)₂·H₂O yield 7 mg (12.7%). Elemental Anal. Calcd (found): C, 39.08 (38.92); H, 4.05 (3.96); N, 10.72 (10.55)%.

10-Ce@La $[La_{0.9}Ce_{0.1}(L)F](CF_3SO_3)_2 \cdot H_2O$. Elemental Anal. Calcd (found): C, 41.14 (40.91); H, 4.06 (3.95); N, 11.29 (11.03)%.

11-Nd@La $[La_{0.9}Nd_{0.1}(L)F](CF_3SO_3)_2 \cdot 0.5H_2O$. Elemental Anal. Calcd (found): C, 40.77 (41.1); H, 4.13 (4.01); N, 11.19 (10.89)%.

12-Tb@Y $[Y_{0,9}Tb_{0,1}(L)F](CF_3SO_3)_2 \cdot 0.5H_2O$. Elemental Anal. Calcd (found): C, 43.11 (43.05); H, 4.31 (4.25); N, 11.83 (11.97)%.

THEORETICAL CALCULATIONS

In order to rationalize the magnetic properties observed through experiments, ab initio calculations were performed using the MOLCAS 8.2 suite.48-50 We used [ANO-RCC... 8s7p5d3f2g1h]⁵¹ for Ce, Nd, and Tb atoms; [ANO-RCC… 3s2p1d] for C, N, and F atoms; and [ANO-RCC…2s1p] for H atoms. Ce(III), Nd(III), and Tb(III) (f¹, f³, and f⁸) have a ${}^{2}F_{5/2}$, ${}^{4}I_{9/2}$, or ${}^{7}F_{6}$ ground state, respectively. Complete active space self-consistent field (CASSCF) calculations were carried out considering one electron in seven active orbitals [CAS-(1,7)] in 1-Ce, three electrons in seven active orbitals [CAS(3,7)] in 3-Nd, and eight electrons in seven active orbitals [CAS(1,7)] in 5-Tb. Further full CI method was employed to compute 7 doublets in 1-Ce, 35 quartets and 112 doublets in 3-Nd, and 7 heptets, 140 quintets, and 588 triplets in 5-Tb. All of these computed spin states are spin-free states. Afterward, using the restrictive active space spin-state interaction spin-orbit (RASSI-SO) program,⁵² 7 doublets were mixed in 1-Ce, 35 quartets and 112 doublets in 3-Nd, and 7 septets, 140 quintets, and 195 triplets in 5-Tb. Furthermore, these computed SO states were taken in the SINGLE ANISO code,⁴⁹ and g-tensors and other local magnetic properties were obtained. The model complexes were optimized using DFT calculations, employing the UB3LYP functional^{53,54} along with SDD^{55,56} for Y and 6-31G* for the other atoms, employing the G09 suite of programs.⁵⁷

RESULTS AND DISCUSSION

Description of the Crystal Structures. Single crystals were obtained for 1-Ce, 2-Pr, 3-Nd, 4-Eu, 5-Tb, 6-Ho, 7-Er, 8-Tm, 9-Yb, 10-Ce@La, 11-Nd@Y, and 12-Tb@Y, all of them are columnar prisms that gave good-quality single-crystal data

(see Tables S1–S6). The crystal system is orthorhombic for all analogues; however, the space group depends on the size of the lanthanide ion. The larger lanthanides (1-Ce to 3-Nd, 10-Ce@ La, and 11-Nd@La) crystallize in the centrosymmetric *Pccn* group, where the complexes are related by inversion, while the smaller ones (4-Eu to 9-Yb and 12-Tb@Y) crystallize in the enantiomorphic $P2_12_12$ group. The diluted samples were prepared with either La(III) or Y(III) to be consistent with the space group: La(III) crystallizes in the *Pccn* group, as do 1-Ce and 3-Nd, whereas Y(III) crystallizes in $P2_12_12$ as does 5-Tb.

In all cases, the asymmetric unit comprises a half-ligand L (which is completed by two-fold rotation) surrounding the metal, a fluoride ligand, the counterion $CF_3SO_3^-$, and a cocrystallized water molecule; the metal and fluoride and the cocrystallized water oxygen atom all lie on a 2-fold rotation axis with only 0.5 of each of these atoms in the asymmetric unit. All analogues pack in compact columns with the Ln(III)…Ln(III) intermolecular distances ranging from 7.95(9) Å (for 1-Ce) to 7.76(8) Å (for 9-Yb). Representative examples of the crystal packing in the *Pccn* and *P*2₁2₁2 groups can be found in Figure S3.

The lanthanide ion is coordinated with four nitrogen atoms of the aza-crown and encapsulated by four nitrogen atoms belonging to the pyridine group of the flexible arms of the ligand (see Figure 1). The Ln–N bond distances are longer for

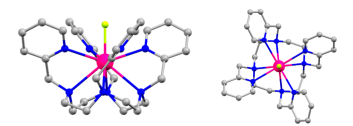


Figure 1. Structure of the cationic complex $[Nd^{III}LF]^{2+}$ in 3-Nd viewed along the *a*-axis (left) and *c*-axis (right). C, gray; N, blue; Nd, pink; F, lime green; and H omitted for clarity.

the nitrogen atoms belonging to the aza-crown than for the nitrogen atoms belonging to the pyridine groups, indicating that the lanthanide ion is not placed equidistantly within the $[N_8]$ cage, leaving space for further coordination with the electronegative anionic fluoride ligand. The Ln–F bond distance increases with the increasing ionic radius of the lanthanide ions, from 2.096(3) Å for 9-Yb to 2.206(3) Å for 1-Ce (see Tables 1 and S4–S6 for details). This distance is

considerably shorter than the distances for the Ln-N bonds, which range from 2.507(5) to 2.748(3) Å (see Tables 1 and S4–S6 for details). This $[N_8F]$ coordination environment is particularly promising to generate an axial crystal field for lanthanide ions with an oblate 4f-electron distribution, which is the case for the Kramers ions Ce(III), Nd(III), and Dy(III) and the non-Kramers ions Pr(III), Tb(III), and Ho(III).^{18,58}

All complexes were analyzed using SHAPE, ^{59–61} which compares the atomic coordinates to those of an ideal prism, with a value of 0 indicating a perfect match and higher values indicating higher distortion from the ideal geometry. ⁵⁹ For all complexes, the nine-coordinate environment around the lanthanide ion can be best described as a distorted capped square antiprism (see Figure S4 and Table S7), corresponding to a distorted $C_{4\nu}$ symmetry, which has not been reported for Ce(III) or Nd(III) SMMs before. The distortion is greater for the larger lanthanides, and it diminishes as the size of the metal ion decreases (Figure 2). For the larger lanthanide ions (1-Ce,

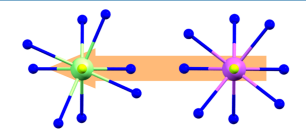


Figure 2. First coordination sphere of the lanthanide ions showing the increase in distortion from 9-Yb (pink) to 1-Ce (lime green).

2-Pr, and 3-Nd, see Table 1) which crystallize in the *Pccn* space group, the SHAPE values vary from 2.506 to 2.397. On the other hand, the analogues that crystallize in the $P2_12_12$ group (4-Eu, 5-Tb, 6-Ho, 7-Er, 8-Tm, and 9-Yb) have values far lower, ranging from 0.605 to 0.532 (see Table 1).

Phase purity was confirmed by powder X-ray diffraction (PXRD). Within the two different space groups, *Pccn* (see Figure S5) or $P2_12_12$ (see Figure S6), the different analogues have the same powder X-ray diffraction pattern.

Magnetic Properties. Magnetic susceptibility measurements in a dc field of 1000 Oe were recorded from 290 to 2 K for all complexes (Figure 3) except 4-Eu (where J = 0). The $\chi_{\rm M}T$ product of all compounds is constant at higher temperatures, and then $\chi_{\rm M}T$ decreases upon cooling due to the depopulation of the Stark levels (Figure 3). 5-Tb

Table 1. Selected	d Bond Lengths	and SHAPE Studi	ies for the [L	_n ^m LF] ²⁺ Comp	lexes
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	space group	Pseudo-symmetry	SHAPE studies	Ln–F bond length (Å)	Ln–N _{py} avg. bond length (Å)	Ln-N _{crown} avg. bond length (Å)	Ln…Ln closest distance (Å)
1-Ce	Pccn	capped square antiprism (C_{4v})	2.51	2.21	2.68	2.76	7.96
2 -Pr	Pccn		2.42	2.19	2.66	2.75	7.96
3-Nd	Pccn		2.40	2.19	2.64	2.73	7.96
4-Eu	$P2_{1}2_{1}2$		0.61	2.16	2.57	2.7	7.77
5-Tb	$P2_{1}2_{1}2$		0.57	2.14	2.55	2.69	7.76
Dy(III) 33	P2 ₁ 2 ₁ 2		0.58	2.12	2.53	2.68	7.76
6 -Ho	$P2_{1}2_{1}2$		0.51	2.13	2.52	2.67	7.76
7-Er	$P2_{1}2_{1}2$		0.54	2.13	2.52	2.67	7.76
8-Tm	$P2_{1}2_{1}2$		0.58	2.11	2.50	2.67	7.76
9 -Yb	$P2_{1}2_{1}2$		0.53	2.10	2.50	2.66	7.77

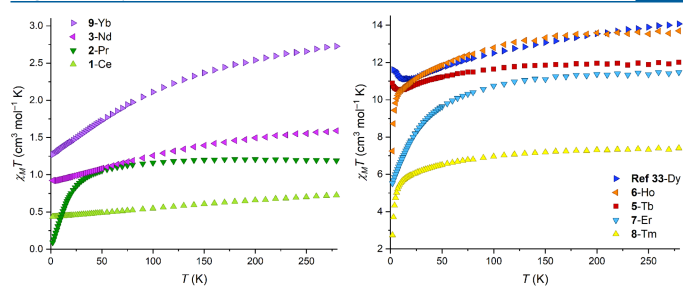


Figure 3. Temperature dependence of $\chi_M T$ from 290 to 2 K for all analogues measured, including the previously reported Dy analogue to allow comparison of the low-temperature region with 5-Tb.³³

Table 2. Theoretical and Experimental Values of the $\chi_M T$ Product (Given in cm³ mol⁻¹ K) and the Magnetization (Given in N β)

	1-Ce	2-Pr	3-Nd	5 -Tb	6 -Ho	7-Er	8-Tm	9 -Yb
$\chi_{ m M} T_{ m theo}$	0.80	1.60	1.64	11.81	14.06	11.48	7.15	2.57
$\chi_{\rm M} T_{\rm exp}$	0.72	1.20	1.59	12.01	13.70	11.47	6.97	2.67
$M_{ m sattheo}$	2.14	3.20	3.27	9.00	10.00	9.00	7.00	4.00
$M_{ m exp}$	1.04	0.51	1.46	5.13	4.92	5.00	3.62	1.94

experiences an increase in $\chi_M T$ from 10 K down to 2 K. The same behavior was seen for the Dy(III) analogue³³ of this family, and it was attributed to intermolecular interactions due to the short Dy…Dy distance, 7.7 Å.³³ The Tb…Tb distance in **5**-Tb is 7.76 Å, very similar to the case with Dy. In order to reduce the intermolecular interactions, the diluted analogue containing Y(III) and Tb(III) in a 9:1 ratio, **12**-Tb@Y, was synthesized. For this diluted sample, the $\chi_M T$ product is constant at higher temperatures and then decreases, consistent with a lack of intermolecular ferromagnetic interactions (see Figure S7). The higher magnetic moments of Dy(III) and Tb(III) over other lanthanides allow them to interact at shorter distances and could explain why ferromagnetic intermolecular interactions are only seen with these two ions in the series.⁶²

The $\chi_M T$ products at 290 K are in agreement with the theoretical values (see Table 2 and Figure 3). We note that 1-Ce, 2-Pr, 6-Ho, and 8-Tm have experimental values lower than the theoretical ones; this has been studied previously and could be related to ligand field effects.^{63,64} The temperature dependence of $\chi_M T$ was also calculated for 1-Ce, 3-Nd, and 5-Tb, which can be seen in Figure S8. Magnetization experiments were performed on all analogues with a variable dc field up to 7 T (Figure S9). Saturation was not reached for any of the analogues at 2 K, with the observed magnetization values being lower than the theoretical ones (M_{sat}), which is common when studying magnetically anisotropic complexes.^{65,66}

Under zero external dc field, no out-of-phase alternating current (ac) susceptibility signals were observed in any of the studied analogues, indicating no slow relaxation of the magnetization. However, when applying an optimum external dc field (Figure S10) to suppress quantum tunneling of the magnetization (QTM), 1-Ce and 3-Nd displayed out-of-phase ac susceptibility signals with fully formed peaks up to 5 K (see Figures S11-S14, 4, and 5).

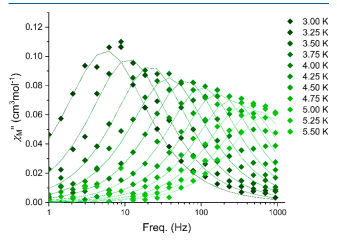


Figure 4. Out-of-phase ac magnetic susceptibility in an applied field $H_{dc} = 1200$ Oe for 1-Ce.

The effect of the applied dc field on the relaxation time was studied between 0 and 4000 Oe (see Figure S10). The optimum dc field that allows for the suppression of QTM was determined to be 1200 and 800 Oe for 1-Ce and 3-Nd, respectively.

For both 1-Ce and 3-Nd, it was possible to extract relaxation times from the Cole–Cole plots (see Figures S15 and S16).⁶⁷

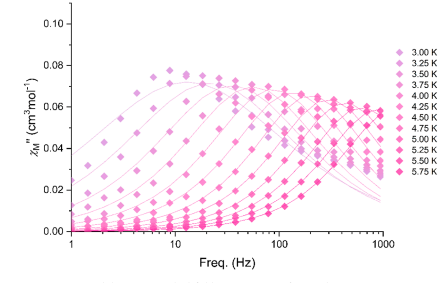


Figure 5. Out-of-phase ac magnetic susceptibility in an applied field H_{dc} = 800 Oe for 3-Nd.

The relaxation times for SMMs can be fitted using different relaxation pathways by using the following equation

$$\tau^{-1} = \tau_0^{-1} e^{-\frac{U_{\text{eff}}}{T}} + CT^n + AH^m T + \tau_{\text{QTM}}^{-1}$$
(1)

Equation 1 includes contributions from Orbach, where τ_0 is the pre-exponential factor, $U_{\rm eff}$ is the magnetization reversal barrier, and T is the absolute temperature; Raman, where C is a constant and n has values up to 9 for lanthanides; direct, where A is a constant, H is the magnetic field, and m is equal to 4 for Kramers ions and 2 for non-Kramers ions; and QTM, with τ_{QTM}^{-1} being the temperature-independent QTM parameter. Fitting the obtained relaxation data for 1-Ce and 3-Nd using eq 1 was unsuccessful. This could be due to the small temperature window in which the slow relaxation is observed, which makes separation of the different relaxation processes more difficult, and/or overparameterization. Considering this, the following simplifications were made. Due to the application of an external dc field during the ac measurements, QTM is considered to be suppressed, and therefore, the $au_{\rm QTM}$ ¹ term was not taken into account for both 1-Ce and 3-Nd.

First, we attempted to fit the field dependence of the relaxation times (see Figure S10) by using the equation

$$\tau^{-1} = AH^m T + \frac{B_1}{1 + B_2 H^2} \tag{2}$$

The direct terms are as defined above, and the second term represents the field dependence of QTM. Unfortunately, no satisfactory fit was obtained using this equation. We also tried to obtain the A parameter from fits that included the Orbach, Raman, and direct terms in eq 1: the A values obtained were negligible, and therefore, the direct relaxation process was not considered further. Next, attempts to fit the relaxation times by considering only the Orbach term and the Raman term in eq 1 yielded small values of $U_{\rm eff}$ of 37.5 and 73.7 K, for 1-Ce and 3-Nd, respectively, that are not consistent with the energies of the m_1 states obtained from the computational studies (vide infra). However, Raman relaxation is commonly observed in Ce(III) and Nd(III) SMMs, and we were able to fit the data using only a Raman process (see Figure 6).^{22,23} The best fits' values obtained were $C = 0.049(1) \text{ K}^{-n} \text{ s}^{-1}$, n = 6.56(1) for 1-Ce and $C = 0.038(15) \text{ K}^{-n} \text{ s}^{-1}$, n = 7.0(2) for 3-Nd, with fitting

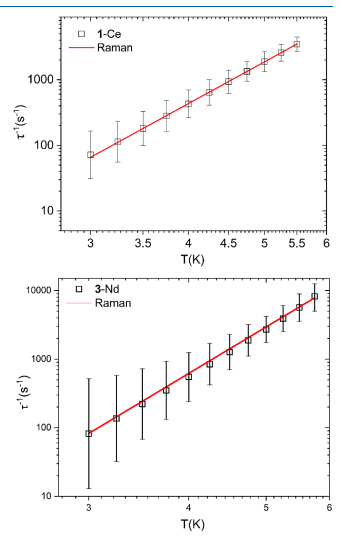


Figure 6. Temperature dependence of $1/\tau$ for 1-Ce (upper) and 3-Nd (lower). Solid red lines represent fits for Raman relaxation $\tau^{-1} = CT^n$ (see text for details). Black vertical bars are estimated standard deviations in the relaxation times derived from Debye fits according to ref 67.

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Table 3. Ce(III) Monometallic SMMs

Ce complexes ^a	Pseudo-symmetry	$H_{\rm dc}~({ m Oe})$	$U_{\rm eff}$ (K)	$ au_0 (s^{-1})$	$C (K^{-n} s^{-1})$	n	ref
[Ce(COT") ₂][Li(DME) ₃]	sandwich complex	400	30	1.20×10^{-6}			69
[CeCd ₃ (Hquinha) ₃ (<i>n</i> -Bu ₃ PO) ₂ I ₃] ₃ EtOH ₂ (H ₂ O)	D_{6h}	1500	27	8.20×10^{-7}			25
[Ce(NO ₃) ₃ (18-crown-6)]		1000	31.4	1.71×10^{-7}			29
			30.3	2.20×10^{-7}	0.1	5	
			25.6	9.00×10^{-7}	0.0016	9	
[Ce(NO ₃) ₃ (1,10-diaza-18-crown-6)]		1000	44	2.30×10^{-8}			29
			45	2.60×10^{-8}	0.52	5	
			23	6.00×10^{-6}	0.0022	9	
Ce(fdh) ₃ (bpy)		2000	33.3	1.80×10^{-7}	0.4	6	70
$[LCe(NO_3)_3]$	C_s	200			1.44	6.8	32
$[Ce(Cp^{ttt})_{2}\{(C_{6}F_{5}-\kappa^{1}-F)B(C_{6}F_{5})_{3}\}]$	sandwich complex	1000			0.0308	5.4	23
$[Ce(Cp^{ttt})_2(Cl)]$	sandwich complex	1000			0.00475	6.5	23
[Ce(18-crown-6) (Cl ₄ Cat) (NO ₃)]	D_{6h}	1500			1.22	5 (fixed)	22
[Ce(18-crown-6) (Br ₄ Cat) (NO ₃)]	D_{6h}	800			1.87	5 (fixed)	22

 a COT" = bis(trimethylsilyl)cyclooctatetraenyl dianion); DME = dimethyl ether; H₂quinha = quinaldic hydroxamic acid; fdh = 1,1,1-fluoro-5,5-dimethylhexa-2,4-dione; bpy = 2,2'-bipyridine; Cp^{ttt} = C₅H₂^tBu₃-1,2,4; X₄Cat = tetrahalocatecholate; and L = ^tBuPO(NHⁱPr)₂.

Table 4. Nd(III) Monometallic SMMs

Nd complexes ^a	Pseudo-symmetry	$H_{\rm dc}$ (Oe)	$U_{\rm eff}$ (K)	$ au_0~(\mathrm{s}^{-1})$	$C (K^{-n} s^{-1})$	п	ref
NdTp ₃	D_{3h}	100	4	4.20×10^{-5}			71
$[Nd(W_5O_{18})_2]^{9-}$	D_{4h}	1000	73.9	3.55×10^{-10}			68
$[L_1Nd(H_2O)_5][I]_3L_1(H_2O)$	D_{5h}	0	16.1	2.64×10^{-4}			31
	D_{5h}		24.7	5.03×10^{-6}			31
	D_{5h}	2000	39.2	8.98×10^{-7}			31
$[L_1Nd(H_2O)_5][I]_3L_1(H_2O)$	D_{5h}	2000				6.3	26
$\{[Nd((\mu_2L_2)_3(H_2O)_2] \cdot C_2H_3N\}_n$	$C_{2\nu}$ CHAIN	2000	27	4.10×10^{-7}			65
$[Nd(\mu_2-L_3) (L_3) (CH_3COO) (H_2O)_2]_n$	D _{3h} CHAIN	3500	29	3.10×10^{-7}			65
[NdCd ₃ (Hquinha) ₃ (<i>n</i> -Bu ₃ PO) ₂ I ₃]·3EtOH·2H ₂ O	D_{6h}	2500	22	3.90×10^{-7}			25
[Nd(NO ₃) ₃ (18-crown-6)]		1000	29.9	2.90×10^{-9}			29
			30.9	2.20×10^{-8}	4.1	5 (fixed)	
			33.4	1.69×10^{-9}	0.000025	9 (fixed)	
[Nd(NO ₃) ₃ (1,10-diaza-18-crown-6)]		1000	69	2.10×10^{-10}			29
			55	2.60×10^{-9}	0.05	5 (fixed)	
			73	1.40×10^{-10}	0.00107	9 (fixed)	
[Nd(CyPh ₂ PO) ₂ (H ₂ O) ₅]I3·2(CyPh ₂ PO)·3EtOH	D_{5h}	0				5.1	26
		2000				6.5	26
$(NH_2Me_2)_3 \{ [Nd(Mo_4O_{13})(DMF)_4]_3(BTC)_2 \} \cdot 8DMF$	D_{3h}	500	26.7	1.41×10^{-7}			72
first: only Orbach, second: Raman and Orbach		500	34.1	4.69×10^{-8}			
Nd(fdh) ₃ (bpy)		500	28.8	9.20×10^{-8}	0.93	6.6	70
$[Nd(Cp^{ttt})_{2}\{(C_{6}F_{5}-\kappa^{1}-F)B(C_{6}F_{5})_{3}\}]$	sandwich complex	1000			0.00117	6.3	23
$[Nd(Cp^{ttt})_2(Cl)]$	sandwich complex	1000	73.6	9.64×10^{-8}	0.0003	8.7	23

 ${}^{a}\text{Tp}^{-}$ = trispyrazolylborate; L₁ = ${}^{b}\text{BuPO}(\text{NH}{}^{i}\text{Pr})_2$; L₂ = 3,5-dinitrobenzoic acid; L₃ = 2,4-dinitrobenzoic acid; H₂quinha = quinaldic hydroxamic acid; CyPh₂PO = cyclohexyl(diphenyl)phosphine oxide; BTC = 1,3,5-benzenetricarboxylate; fdh = 1,1,1-fluoro-5,5-dimethylhexa-2,4-dione; bpy = 2,2'-bipyridine; Cp^{ttt} = C₅H₂'Bu₃-1,2,4; and COT'' = bis(trimethylsilyl)cyclooctatetraenyl dianion.

errors shown in parentheses. These values are in line with those observed previously for Ce(III) and Nd(III) SMMs. 26,29

In order to study the magnetism of these complexes further, the diluted analogues containing La(III), **10**-Ce@La, and **11**-Nd@La were synthesized. The dilution did not have a significant effect on the magnetic properties, with the out-of-phase peaks appearing at the same temperatures as in the non-diluted analogues (see Figure S17). In addition, the diluted analogue of Tb(III) with Y(III), **12**-Tb@Y, was also studied to see if it had improved magnetic properties since **5**-Tb showed no slow relaxation of the magnetization. However, only a negligible out-of-phase ac signal was observed under an applied dc field (see Figure S18).

For Ce(III) and Nd(III) single-ion magnets, the predominant magnetic relaxation process is usually Raman relaxation, but where the Orbach process is included in the analysis, magnetization reversal barriers up to 73 K for Nd(III) and 45 K for Ce(III) are reported.^{29,68} Tables 3 and 4 show a list of the single-ion magnets reported with Ce(III) and Nd(III). For both of these, the most common pseudo-symmetry falls within a dihedral group (D_{nh}) in contrast to 1-Ce and 3-Nd that are, to the best of our knowledge, the first Ce(III)/Nd(III) SMMs reported with $C_{4\nu}$ pseudo-symmetry.

Theoretical Studies. Among the reported compounds, 1-Ce, 3-Nd, and 5-Tb were analyzed using CASSCF, RASSI-SO, and SINGLE_ANISO calculations using the structures obtained from single-crystal X-ray diffraction as their inputs.

KD	energy (K)	composition $ m_{\rm J} $	g _{xx}	g_{yy}	g _{zz}	θ (deg)	
1	0.0	$0.86 \mid \pm 5/2 \rangle + 0.12 \mid \pm 3/2 \rangle$	1.235	1.335	3.295		
2	723.4	0.86 $\mid \pm 3/2 \rangle$ + 0.13 $\mid \pm 1/2 \rangle$	1.208	1.411	1.640	89.5	
3	1104.7	$0.99 \mid \pm 1/2 \rangle$	2.695	2.242	0.641	1.6	
^{<i>a</i>} The angle be	^a The angle between the ground-state g_{ax} and the respective excited-state g_{ax} axes is represented by θ .						

Table 5. Energies (K), m_J Composition of the Lowest Doublets, and g-Tensors of the Individual Lanthanide Magnetic Centers Associated with Each State for 1-Ce^{*a*}

For 1-Ce, the three lowest lying Kramers doublets (KDs) span the range 0–1105 K, and the ground state g values are g_{xx} = 1.235, $g_{yy} = 1.335$, and $g_{zz} = 3.295$. The composition of the ground state is predominantly $m_1 = \pm 5/2$, although there is mixing with the $m_{\rm I} = \pm 3/2$ state (see Table 5). This suggests a stronger axial contribution to the crystal field than the equatorial contribution, which is in line with other Ce(III) SIMs previously studied by some of us.^{32,51} Furthermore, the orientation of the ground state g_{zz} axis coincides with the Ce-F bond (see Figure S19). However, the transverse anisotropy present does lead to some mixing of the ground state with excited states (see Table 5). This leads to ground-state QTM, consistent with the absence of slow relaxation of the magnetization in the absence of an applied dc field. The first excited state lies at 723 K (see Figure 7 and Table 5); hence, if the QTM is even partially quenched by using an applied dc field, 1-Ce is likely to show slow relaxation of magnetization, as we observe experimentally.

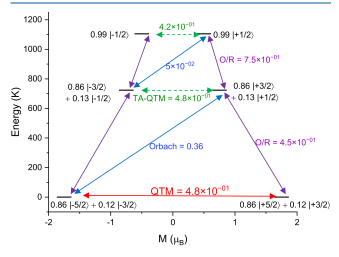


Figure 7. Energy-level distributions for 1-Ce with the indicated probability of the relaxation path: QTM (red arrow) from where the actual relaxation in zero field occurs; Orbach (blue arrow); Orbach/Raman (purple arrow); and TA-QTM (TA = thermally assisted; green dashed arrow). The numbers above each arrow represent the corresponding transverse matrix elements for the transition magnetic moments.

The computed LoProp⁷³ charge is -0.88 on the fluoride ion, whereas the combination of the nitrogen atoms in the equatorial plane yield total LoProp charges of -1.60 and -1.37 (considering two different sets of charges arising from the different nitrogen atoms, one for the nitrogen atoms in the aza-crown and another one for the nitrogen atoms in the pyridine groups) (see Table S8). Hence, there is a large contribution from the nitrogen atoms to the equatorial crystal field. It has been shown previously that, unlike Dy(III), Ce(III) ions are very sensitive to the equatorial ligand field,⁵¹ and this rationalizes our experimental observations. By considering the crystal field parameters (Table S9), it can be seen that although B_2^{0} is larger than the B_2^{q} non-axial components (where $q \neq 0$), which disfavors QTM, the B_4^0 parameter is smaller than some of the B_4^q non-axial components (where $q \neq$ 0), which favors QTM. Furthermore, other factors such as significant g_{xx}/g_{yy} values and a mixed ground KD also promote QTM (Table 5).

For 3-Nd, the computed ground-state g values are g_{xx} = 0.638, $g_{yy} = 0.707$, and $g_{zz} = 5.513$ (see Table 6). While the composition of the ground state is predominantly $m_1 = \pm 9/2$, there is mixing with the $m_{\rm I} = \pm 1/2$ state (see Table 6). The ground-state g_{zz} axis coincides with the Nd–F bond (Figure S19) as we saw for the Ce–F bond in 1-Ce. However, the first excited KD in 3-Nd lies at 223 K (see Table 6), which is far lower in energy than the first excited KD in 1-Ce. This is reflected in a smaller B_2^{0} parameter for 3-Nd than for 1-Ce (Table S9). Strong transverse anisotropy coupled with the mixing of the m₁ states facilitates ground-state QTM in 3-Nd, again suggesting that a zero-field SMM behavior should not be expected, as we observe experimentally. The relaxation mechanism calculated (see Figure 8) shows that if the ground-state QTM is quenched by a dc field, there is a possibility for slow relaxation of the magnetization, as we observe experimentally.^{26,65,71}

The analogue 5-Tb presents a different case to 1-Ce and 3-Nd, as Tb(III) is a non-Kramers ion. The *g* values of the ground state are calculated to be $g_{xx} = 0.000$, $g_{yy} = 0.000$, and $g_{zz} = 17.818$. The ground-state magnetic moment is aligned along the Tb–F bond, as in 1-Ce and 3-Nd (see Figure S19). However, a large tunnel splitting was calculated for 5-Tb, and this is due to the position of the nitrogen donors from the ligand L, which provide a significant equatorial ligand field that

Table 6. Energies (K), m_J Composition of Lowest Doublets, and g-Tensors of the Individual Lanthanide Magnetic Centers Associated with Each State for 3-Nd^{*a*}

KD	energy (K)	composition $ m_{\rm J} $	g_{xx}	g_{yy}	g_{zz}	θ (deg)
1	0.0	0.83 $\mid \pm 9/2 \rangle$ + 0.13 $\mid \pm 1/2 \rangle$	0.638	0.707	5.513	
2	223.3	0.57 $\mid \pm 5/2 \rangle$ + 0.41 $\mid \pm 3/2 \rangle$	3.521	2.911	1.175	1.2
3	337.8	0.88 $\mid \pm 7/2 \rangle$ + 0.07 $\mid \pm 1/2 \rangle$	0.439	0.722	4.252	0.04
4	503.9	$0.57 \mid \pm 3/2 \rangle + 0.41 \mid \pm 5/2 \rangle$	3.812	2.880	0.268	0.8

"The angle between the ground-state g_{zz} and the respective excited-state g_{zz} axes is represented by θ .

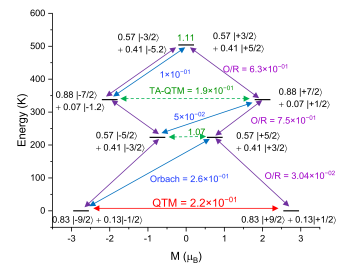


Figure 8. Energy-level distributions for 3-Nd with the indicated probability of the relaxation path: QTM (red arrow) from where the actual relaxation in zero field occurs; Orbach (blue arrow); Orbach/Raman (purple arrow); and TA-QTM (TA = thermally assisted; green dashed arrow). The numbers above each arrow represent the corresponding transverse matrix elements for the transition magnetic moments.

enhances the tunnel splitting (see Table S10). Hence, applying a magnetic field is probably not enough to quench QTM, which agrees with the lack of any significant slow magnetic relaxation for 5-Tb.

To understand further the magnetic properties of these complexes and the role of the fluoride ligand in promoting strong axiality, we have prepared a series of computational models 1-Ce(a), 3-Nd(a), and 5-Tb(a) where the axial F⁻ ligand is removed, so that each model has just a {LnN₈} coordination environment (see Figure S19).⁷⁴ The model structures were optimized using DFT calculations (see Experimental Methods). For the 1-Ce(a) and 3-Nd(a) models, the calculations reveal smaller g_{zz} values and larger g_{xx} and g_{yy} values, with the excited KDs closer in energy (see Table 7). This highlights the importance of the F⁻ ligand in creating a strong axial crystal field for 1-Ce and 3-Nd and the observed slow relaxation of the magnetization.

The ground m_J state is found to be $\pm 1/2$ for the model 1-Ce(a) and a mixture of $\pm 5/2$ and $\pm 3/2$ for the model 3-Nd(a) (see Figure 9). Hence, upon removal of the F⁻ ligand, the

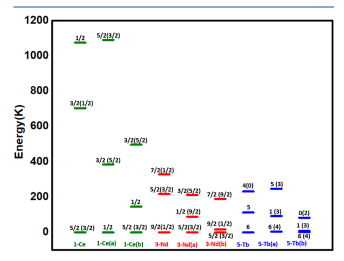


Figure 9. Comparative energies (in K) of the first three m_J states in 1-Ce, 1-Ce(a), and 1-Ce(b); 3-Nd, 3-Nd(a), and 3-Nd(b); and 5-Tb, 5-Tb(a), and 5-Tb(b). Models (a) have the axial F⁻ ligand removed and models (b) have the axial F⁻ replaced by I⁻. The major composition of the m_J states is shown, with the largest minor contribution given in brackets.

nature of the ground state changes, with the $[N_8]$ ligand stabilizing instead m_J states with stronger prolate 4f charge density for 1-Ce(a) and 3-Nd(a). For the model 5-Tb(a), the tunnel splitting is increased further after the removal of the F⁻ ligand (see Table S10). Furthermore, in 5-Tb(a), the direction of the ground-state g_{zz} axis passes through the direction that bisects the plane formed by the pyridine and aza-crown nitrogen atoms (see Figure S19). This shows that the F⁻ ligand is essential to align the g_{zz} axis along the pseudo- C_4 axis for 5-Tb. Furthermore, the ground-state to first-excited-state gap reduces upon removal of the F⁻ ligand, and the energy states are extremely mixed, although the ground m_J state is still predominantly ± 6 , which has an oblate charge density. We note that eight nitrogen donor atoms present in a pseudo- D_{4d} environment are known to stabilize the $m_J = \pm 6$ state, for

Table 7. Energies (K) and Composition of m_J States of Lowest Doublets and g-Tensors of the Individual Lanthanide Magnetic Centers Associated with Each State for Model Complexes 1-Ce(a), 1-Ce(b), 3-Nd(a), and 3-Nd(b)^a

	energy (K)	composition $ m_J $ states	g_{xx}	g_{yy}	g _{zz}	θ (deg)
1-Ce(a)	0.00	$0.99 \mid \pm 1/2 \rangle$	2.873	2.238	0.810	
	396.2	0.82 $\mid \pm 3/2 \rangle$ + 0.17 $\mid \pm 5/2 \rangle$	1.696	1.457	1.042	88.9
	1121.0	0.82 $\mid \pm 5/2 \rangle$ + 0.17 $\mid \pm 3/2 \rangle$	1.327	1.422	3.037	0.6
1-Ce(b)	0.0	$0.48 \mid \pm 5/2 \rangle + 0.40 \mid \pm 3/2 \rangle$	1.866	1.836	1.174	
	150.3	$0.99 \mid \pm 1/2 \rangle$	2.559	2.524	0.765	0.0
	511.2	$0.55 \mid \pm 3/2 \rangle + 0.44 \mid \pm 5/2 \rangle$	1.906	1.891	0.509	0.0
3-Nd(a)	0.0	$0.49 \mid \pm 5/2 \rangle + 0.48 \mid \pm 3/2 \rangle$	3.255	3.107	0.800	
	91.9	0.72 $\mid \pm 1/2 \rangle$ + 0.19 $\mid \pm 9/2 \rangle$ + 0.07 $\mid \pm 7/2 \rangle$	3.284	2.727	1.4103	0.3
	218.9	$0.50 \mid \pm 3/2 \rangle + 0.49 \mid \pm 5/2 \rangle$	3.598	2.956	0.764	0.2
	329.8	0.89 $\mid \pm 7/2 \rangle$ + 0.09 $\mid \pm 9/2 \rangle$	0.997	1.375	4.046	1.0
3 -Nd(b)	0.0	$0.55 \mid \pm 5/2 \rangle + 0.45 \mid \pm 3/2 \rangle$	3.298	3.167	1.046	
	17.4	0.49 $\mid \pm 9/2 \rangle$ + 0.46 $\mid \pm 1/2 \rangle$ + 0.07 $\mid \pm 7/2 \rangle$	2.081	2.191	3.181	0.0
	193.8	0.85 l± 7/2⟩ 0.14 l± 9/2⟩	1.475	1.520	3.516	0.0
	243.7	0.55 l± 3/2⟩ 0.49 l± 5/2⟩	3.349	3.346	0.456	0.0

^{*a*}The angle between ground-state g_{zz} and the respective excited-state g_{zz} axes is represented by θ .

example, in $[Tb(pc)_2]^-$, which has a sandwich-like ligand arrangement.

We also have modeled another set of molecules, 1-Ce(b), 3-Nd(b), and 5-Tb(b), where the axial F^- ligand is replaced by an I⁻ ligand. The model structures were optimized using DFT calculations (see Experimental Methods). The optimized Ln-I distance is 3.3 Å, compared to the Ln-F distance of 2.2 Å (see Table 1). The orientation of the g_{zz} axis is along the pseudo- C_4 axis in 1-Ce(b) and 3-Nd(b) as in 1-Ce and 3-Nd, that is, along the Ln-I bond (Figure S19). However, for these models, there is a reduction in the axial crystal field: the energy gap between the first KD and the second KD, which was 723 K in 1-Ce is lowered to 150 K for 1-Ce(b), while the 223 K gap in 3-Nd decreases to 17 K in 3-Nd(b) (see Table 7 and Figure 9). Again, this highlights the importance of the F⁻ ligand in creating a strong axial crystal field for 1-Ce and 3-Nd. In model 5-Tb(b), there is also a significant tunnel splitting (see Table S10); however, the first exited state is very low in energy (7.1/8.6 K). In this model, the g_{zz} axis lies in the plane between the pyridine and aza-crown nitrogens, as it did in model 5-Tb(a), reflecting the fact that the I⁻ ligand does not provide a sufficient axial crystal field to offset the eight nitrogen atoms in a D_{4d} environment. Here, we can draw parallels to earlier work on Na[Tb^{III}(DOTA)(H₂O)]·4H₂O (H₄DOTA is 1,4,7,10tetraazacyclododecane-1,4,7,10-tetraacetic acid), which has a weak axial H₂O ligand, where the easy axis is found perpendicular to the Ln-H₂O bond.⁷⁵

CONCLUSIONS

In summary, we report the structural and magnetic study of a family of lanthanide compounds featuring an axial Ln–F bond, including eight new analogues (1-Ce, 2-Pr, 3-Nd, 5-Tb, 6-Ho, 7-Er, 8-Tm, and 9-Yb). From these, the 1-Ce and 3-Nd analogues show slow relaxation of the magnetization under an applied dc field of 1200 and 800 Oe, respectively, which is modeled using a Raman process. The strong axial magnetic anisotropy generated by the fluoride ligand helps promote the SMM behavior in the oblate lanthanides Ce(III) and Nd(III), both Kramers ions, and the relaxation pathways have been elucidated by performing ab initio calculations. The Tb(III) complex does not show any significant slow relaxation of the magnetization, even when diluted with Y. We have shown that this can be attributed to a large tunnel splitting in the ground state and the non-Kramers nature of the ion. Furthermore, the analysis of 1-Ce(a), 3-Nd(a), and 5-Tb(a) model complexes, where the axial fluoride ligand is removed to study the effect of the [N₈] coordination cage, and 1-Ce(b), 3-Nd(b), and 5-Tb(b), where the F^- is replaced by a I^- , show that the crystal field splitting is dramatically reduced. This highlights the importance of the F⁻ ligand in creating a strong axial crystal field for 1-Ce and 3-Nd and for promoting the SMM behavior.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.inorgchem.2c00556.

Ligand ¹H NMR; crystallographic tables; crystal packing diagrams; PXRD patterns; further magnetic data; and further computational studies (PDF)

Accession Codes

CCDC 2117093-2117101 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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Notes

The authors declare no competing financial interest.

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