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Axial Ligand Field in D_{4d} Coordination Symmetry: Magnetic Relaxation of Dy SMMs Perturbed by Counter-anions

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ABSTRACT: A series of mononuclear Dy^{III} complexes with the general formula $[DyLz_2(salicylaldehyde)_2] \cdot X \cdot \text{solvent}$ ($Lz = 6\text{-Pyridin-2-yl-[1,3,5]triazine-2,4-diamine}$; $X = OH^-$ (**1-OH**), Cl^- (**2-Cl**), Br^- (**3-Br**)) have been synthesized using mixed salicylaldehyde/pyridin-triazine ligands and discriminative counter-anions. The Dy^{III} ion in these three complexes resides in a similar D_{4d} coordination geometry with counter-anions perturbing the coordination environment and bond lengths and angles in the lattice. Magneto-structural studies reveal that the asymmetric distribution of salicylaldehyde/pyridin-triazine ligands and the presence of discriminative counter-anion result in the coexistence of large anisotropy and quantum tunneling of magnetization. The magnetic anisotropy is dominated by the axial ligand field with short $Dy-O_{sali}$ distances and large $\angle O_{sali}-Dy-O_{sali}$ angles, while the quantum tunneling relaxation is probably dictated by the $\pi-\pi$ stacking of the Lz ligands which induces an axial constriction of the coordinating plane. *Ab initio* calculations substantiate the diversity of the magnetic behaviors in these complexes and highlight the importance of axial ligand field with short $Dy-O_{sali}$ distances, large $\angle O_{sali}-Dy-O_{sali}$ angles and less ligand stacking in these pseudo- D_{4d} symmetrical single-molecule magnets.

INTRODUCTION

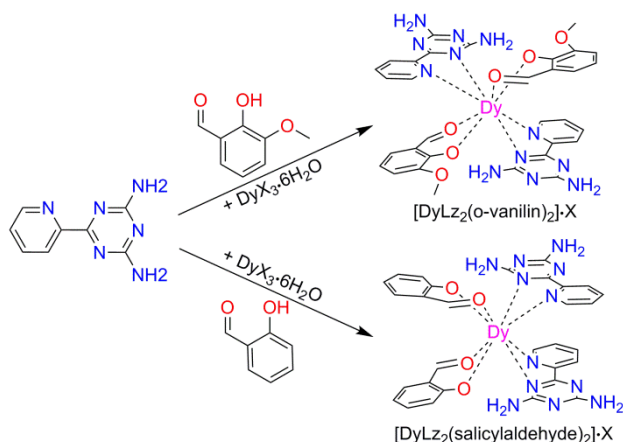
Since the discovery of single-molecule magnetic behavior in the celebrated molecule of $(Bu_4N)[Tb(Pc)_2]$ ($H_2Pc = \text{phthalocyanine}$) in 2003,¹ the research on lanthanide single-molecule magnets (SMMs) has been well developed and hundreds of lanthanide compounds with diverse structural topologies have been reported in the past decade.²⁻⁸ Compared with transition elements, lanthanide ions provide larger magnetic moments and more significant magnetic anisotropy profiting from the strong coupling between the spin and orbit angular moments and the weak crystal-field splitting effects,⁹⁻¹² thus making them ideal candidates for the construction of SMMs. However the relatively weak magnetic coupling between lanthanide ions in multinuclear compounds limits the performance of robust SMM behavior, except for low-symmetry dysprosium compounds, such as Dy_5 ,¹³ Dy_4K_2 ,¹⁴ Dy_3 ¹⁵ and Dy_2 ¹⁶⁻¹⁹ complexes. According to the crystal field approach toward 4f SMMs,²⁰⁻²² the splitting of the $^{2S+1}L_J$ multiplets varies with the coordination symmetry of the lanthanide ion and determines the magnetic anisotropy and magnetic relaxation behavior.²³⁻²⁸ Theoretically speaking, lanthanide SMMs with high symmetry, such as C_3 , S_8 , D_{4d} , D_{5h} and D_{6d} , are inclined to show very high-

energy barriers and less quantum tunneling of magnetization (QTM).²⁹⁻³³ Indeed, a series of Dy^{III} , Tb^{III} and Er^{III} based mononuclear SMMs with landmark significance have been reported showing the above symmetries and robust anisotropy barriers and blocking temperatures.^{1, 34-36}

On the other hand, a simple but amenable electrostatic model based on basic overall shape of free-ion electron density has been widely used recently to understand and predict the magnetic anisotropy of lanthanide SMMs.³⁷⁻³⁹ For instance, the Kramers ion Er^{III} ($^4I_{15/2}$) and Dy^{III} ($^6H_{15/2}$) with prolate and oblate shaped electron density would provide large magnetic anisotropy in highly equatorial and strongly axial ligand field, respectively.⁴⁰⁻⁴⁴ In particular, the later oblate shaped Dy^{III} ion in near-perfect pentagonal bipyramidal symmetry with weak equatorial pyridine donors and strong *tert*-butoxide donors was reported recently showing record energy barrier of 1815 K.³⁴ The anionic character of coordinate donors and the steric tunability of counter-anion may be exploited to stabilize the 4f electronic density.^{37, 45, 46} Therefore, rational optimization of the ligand field around a lanthanide ion is the key step to obtain high performance SMMs.^{22, 47}

On the basis of above theoretical and conceptual aspects, we intend to design SMMs based on Dy^{III} ion with

high coordination number and strong axial ligand field, such as eight coordinate Dy^{III} in D_{4d} symmetry, which are stable in both solid-state and frozen solution.^{48–56} Recently, we obtained a series of mononuclear dysprosium complexes with a Dy^{III} ion located in approximate D_{4d} coordination geometry and two *o*-vanilin in *trans* configuration (Scheme 1), showing remarkable relaxation barrier and blocking temperature.⁵⁷ In the present work, we replaced the *o*-vanilin with salicylaldehyde and used halogen as counter anions, and obtained a series of mononuclear dysprosium complexes of formula $[\text{DyLz}_2(\text{salicylaldehyde})_2]\cdot\text{X}\cdot\text{solvent}$ (Lz = 6-Pyridin-2-yl-[1,3,5]triazine-2,4-diamine; $\text{X} = \text{OH}^-$ (**1-OH**), Cl^- (**2-Cl**), Br^- (**3-Br**)). In these complexes, the magnetic relaxation barrier varies largely according to the change of counter anions. Magneto-structural studies and *ab initio* calculations indicate that the magnetic anisotropy is dominated by the short $\text{Dy-O}_{\text{sali}}$ distances and large $\angle\text{O}_{\text{sali}}\text{-Dy-O}_{\text{sali}}$ angles in the pseudo- D_{4d} geometry.



Scheme 1 Preparation of the complexes $[\text{DyLz}_2(\text{o-vanilin})_2]\cdot\text{X}$ and $[\text{DyLz}_2(\text{salicylaldehyde})_2]\cdot\text{X}$.

EXPERIMENTAL

General Synthetic Considerations. All chemicals and solvents were commercially obtained and used as received without any further purification. FTIR spectra were measured using a Nicolet 6700 Flex FTIR spectrometer equipped with smart iTR™ attenuated total reflectance (ATR) sampling accessory in the range from 500 to 4000 cm^{-1} . Elemental analysis for C, H and N were carried out on a Perkin-Elmer 2400 analyzer. Ligand Lz (2,4-diamino-6-pyridyl-1,3,5-triazines) was prepared according to a previously published method under ambient conditions.⁵⁸ All doped samples were analyzed using Inductively Coupled Plasma Optical Emission Spectrometry (ICP OES). Powder X-ray diffraction measurements were recorded on Bruker D8 advance X-Ray diffractometer using $\text{Cu-K}\alpha$ radiation.

Synthesis of 1-OH. $\text{Dy}(\text{SO}_3\text{CF}_3)_3\cdot 6\text{H}_2\text{O}$ (0.2 mmol) was added to a solution of Lz (0.2 mmol) and salicylaldehyde (0.2 mmol) in 15 mL mixture of methanol and dichloromethane (1:2, v/v), and then triethylamine (0.4 mmol) was added. The resultant solution was stirred for 3 h and subsequently filtered. The filtrate was exposed to air to allow the slow evaporation of the solvent. Yellow crystals of **1-OH** suitable for X-ray diffraction analysis were collected after 1 week. Yield in ~ 70%. Selected IR (cm^{-1}) for **1-OH**: 3336 (m), 3145 (b), 1619 (s), 1535 (s), 1440 (m), 1397 (w), 1322 (s), 1244 (s), 1147 (s), 1025 (m), 900 (s), 792 (m), 760 (m), 637 (m). Anal. Calcd. for $[\text{DyLz}_2(\text{salicylaldehyde})_2]\cdot\text{OH}\cdot 2\text{CH}_3\text{OH}$ ($\text{C}_{32}\text{H}_{35}\text{DyN}_{12}\text{O}_7$, MW = 862.22): C, 44.58%; H, 4.09%; N, 19.49%. Found: C, 44.36%; H, 3.98%; N, 19.58%. Doping of complex **1-OH** was performed by adding $\text{Dy}(\text{SO}_3\text{CF}_3)_3\cdot 6\text{H}_2\text{O}$ and $\text{Y}(\text{SO}_3\text{CF}_3)_3\cdot 6\text{H}_2\text{O}$ together (with the ratio of 1:20) in the synthesis process. ICP OES analysis found $[\text{Dy}_{0.067}\text{Y}_{0.933}\text{Lz}_2(\text{salicylaldehyde})_2]\cdot\text{OH}\cdot 2\text{CH}_3\text{OH}$.

Syntheses of complexes 2-Cl and 3-Br. Complexes **2-Cl** and **3-Br** were synthesized using a similar procedure as for **1-OH** with the replacement of $\text{Dy}(\text{SO}_3\text{CF}_3)_3\cdot 6\text{H}_2\text{O}$ by $\text{DyCl}_3\cdot 6\text{H}_2\text{O}$ and $\text{DyBr}_3\cdot 6\text{H}_2\text{O}$, respectively. Selected IR (cm^{-1}) for **2-Cl**: 3273 (m), 3130 (b), 1617 (s), 1580 (w), 1561 (w), 1536 (s), 1510 (m), 1437 (m), 1392 (m), 1325 (s), 1257 (w), 1184 (m), 1147 (s), 1014 (m), 900 (s), 791 (m), 769 (m), 625 (m), 589 (w). Anal. Calcd. for $[\text{DyLz}_2(\text{salicylaldehyde})_2]\cdot\text{Cl}\cdot\text{H}_2\text{O}$ ($\text{C}_{31}\text{H}_{30}\text{ClDyN}_{12}\text{O}_5$, MW = 848.62): C, 43.88%; H, 3.56%; N, 19.81%. Found: C, 43.81%; H, 3.68%; N, 20.04%. Selected IR (cm^{-1}) for **3-Br**: 3296 (m), 3133 (b), 1618 (s), 1535 (s), 1512 (m), 1437 (m), 1392 (m), 1325 (s), 1196 (m), 1149 (s), 1014 (m), 900 (s), 791 (m), 767 (m), 589 (m). Anal. Calcd. for $[\text{DyLz}_2(\text{salicylaldehyde})_2]\cdot\text{Br}\cdot\text{H}_2\text{O}\cdot\text{CH}_3\text{OH}$ ($\text{C}_{31}\text{H}_{32}\text{BrDyN}_{12}\text{O}_6$, MW = 911.10): C, 40.87%; H, 3.54%; N, 18.45%. Found: C, 40.72%; H, 3.68%; N, 18.57%. Doping of complex **2-Cl** was performed by adding $\text{DyCl}_3\cdot 6\text{H}_2\text{O}$ and $\text{YCl}_3\cdot 6\text{H}_2\text{O}$ together (with the ratio of 1:20) in the synthesis process. ICP OES analysis found $[\text{Dy}_{0.058}\text{Y}_{0.942}\text{Lz}_2(\text{salicylaldehyde})_2]\cdot\text{Cl}\cdot\text{H}_2\text{O}$.

Crystallography. Single-crystal X-ray data of the titled complexes were collected on a Bruker Apex II CCD diffractometer equipped with graphite-monochromatized $\text{Mo-K}\alpha$ radiation ($\lambda = 0.71073$ Å) at 293(2) K. The structures were solved by direct methods and refined by full-matrix least-squares methods on F^2 using SHELXTL-2014.^{59, 60} All non-hydrogen atoms in the whole structure were refined with anisotropic displacement parameters. Hydrogen atoms were introduced in calculated positions and refined with fixed geometry with respect to their carrier atoms. Crystallographic data are listed in Table S1. CCDC 1513743–1513745 contain the supplementary crystallographic data for this paper. These data can be obtained free of charge from the Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.

Magnetic Measurements. Magnetic susceptibility measurements were recorded on a Quantum Design

MPMS-XL7 SQUID magnetometer equipped with a 7 T magnet. Direct current (dc) magnetic susceptibility measurements were performed on polycrystalline samples of **1·OH**, **2·Cl** and **3·Br** in the temperature range 2–300 K, in an applied field of 1000 Oe. The variable-temperature magnetization was measured in the temperature range of 1.9–300 K. The dynamics of the magnetization were investigated from the ac susceptibility measurements in the zero static fields and a 3.0 Oe ac oscillating field. Diamagnetic corrections were made with the Pascal’s constants⁶¹ for all the constituent atoms as well as the contributions of the sample holder.

Table 1. Selected bond distances (Å) and angles (°) for complexes **1·OH**, **2·Cl** and **3·Br**.

	1·OH		2·Cl	3·Br
Dy1-O1	2.365(5)	Dy1-O1	2.252(6)	2.237(7)
Dy1-O1' ^{a)}	2.365(5)	Dy1-O2	2.356(7)	2.352(7)
Dy1-O2	2.203(5)	Dy1-O3	2.242(6)	2.241(7)
Dy1-O2'	2.203(5)	Dy1-O4	2.369(6)	2.366(7)
Dy1-N1	2.509(6)	Dy1-N1	2.513(8)	2.503(8)
Dy1-N1'	2.509(6)	Dy1-N6	2.593(7)	2.572(7)
Dy1-N6	2.593(5)	Dy1-N7	2.519(7)	2.522(8)
Dy1-N6'	2.593(5)	Dy1-N12	2.591(7)	2.588(7)
O2-Dy1-O2'	149.018	O1-Dy1-O3	142.855	142.958
Dy...Dy	7.682	Dy...Dy	8.093	8.080
$\pi\cdots\pi$ / Å ^{b)}	3.826	$\pi\cdots\pi$ / Å	3.596	3.590
ϕ / ° ^{c)}	58.646	ϕ / °	46.946	46.942
θ / ° ^{d)}	6.841	θ / °	2.525	2.593
d_{pp} / Å ^{e)}	2.639	d_{pp} / Å	2.613	2.619

^{a)} Symmetry transformation: 1 -x, -y+3/2, z. ^{b)} Distance between the centers of the triazine groups. ^{c)} Angle defined as the rotation angle between the two Lz ligands. ^{d)} Angle between the upper and lower planes containing the coordinated atoms. ^{e)} The sum distance from the center Dy^{III} ion to two coordination planes.

Ab initio Calculations. Wavefunction-based calculations were carried out on the X-ray structures of **1·OH**, **2·Cl** and **3·Br** using the SA-CASSCF/RASSI-SO approach as implemented in MOLCAS 8.0,⁶² in which the relativistic effects are treated by means of the Douglas-Kroll Hamiltonian in a two steps scheme. First, the scalar terms are included in the basis-set generation and are used to determine the spin-free states in the Complete Active Space Self-Consistent Field (CASSCF) method.⁶³ Next, spin-orbit coupling is added within the Restricted Active Space State-Interaction (RASSI-SO) method,^{64, 65} in which the spin-free states serve as basis states. The resulting energies and wave functions are finally used to compute the magnetic properties (*i.e.* magnetization and magnetic susceptibility curves, anisotropy tensors of the low-energy states of the system, as well as the associated wave functions in term of M_J eigenstates) using the pseudo-spin $S = 1/2$ approximation as defined in the SINGLE_ANISO routine.^{66, 67} Cholesky decomposition is used when computing bielectronic integrals.⁶⁸ The active space of the CASSCF calculations consisted of the nine 4f electrons of the Dy^{III} ion spanning the seven 4f orbitals, *i.e.* CAS(9,7)SCF. The State-Averaged CASSCF calculations were performed for all the 21 sextet roots, all the 224 quadruplet roots and 300 out of 490 doublet roots. In the RASSI-SO calculation, the 21 sextet roots were allowed to mix through spin-orbit coupling with the first 128 quadruplet roots and the first 107 doublet roots. All atoms were described by ANO-type basis sets from the ANO-RCC library of MOLCAS.⁶⁹⁻⁷¹ The following contractions were used: [8s7p4d3f2g1h] for the Dy^{III} ion, [4s3p2d] for the four N and the four O atoms of the first coordination sphere, [3s2p1d] for the remaining N atoms and all C atoms, and [2s] for the H atoms.

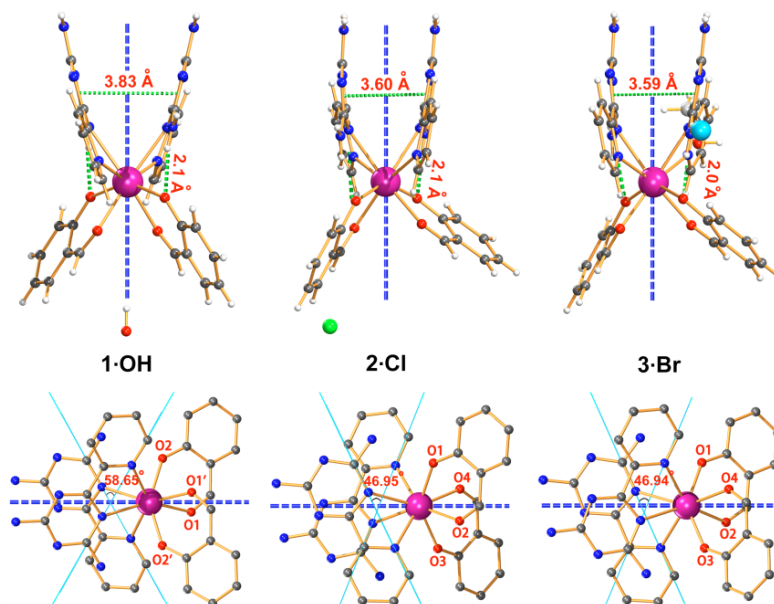


Figure 1. Side (top) and top (bottom) views of complexes **1·OH**, **2·Cl** and **3·Br** with parameter labels. The dashed blue lines represent the location of the pseudo- C_2 molecular axes. The crossed solid azure lines represent the orientations of the two Lz ligands with labeled rotation angle (ϕ). The pink, blue, dark grey, red, green and azure spheres represent Dy, N, C, O, Cl and Br atoms, respectively; solvents (top) and/or hydrogen atoms (bottom) have been omitted for clarity.

RESULTS AND DISCUSSION

Structural Descriptions. Compounds **1·OH**, **2·Cl** and **3·Br** were isolated from the reaction of $DyX_3 \cdot 6H_2O$ ($X = CF_3SO_3^-$ for **1·OH**, Cl^- for **2·Cl**, Br^- for **3·Br**) with Lz and salicylaldehyde in the presence of triethylamine in 15 mL mixture of methanol and dichloromethane (1:2, v/v) after several days (Scheme 1). Details for the structure solution and refinement are summarized in Table S1. Powder XRD analyses are given in Figure S2. Selected bond distances and angles are given in Table 1. Single-crystal X-ray diffraction analyses (Figure 1, Table S1) show that complex **1·OH** crystallizes in the tetragonal $I4_1/a$ space group with the asymmetric unit containing half of the complex (one ligand Lz and one ligand salicylaldehyde and one half Dy^{III} ion). Complexes **2·Cl** and **3·Br** crystallize in the triclinic $P-1$ space group with the asymmetric unit containing two ligands Lz, two salicylaldehyde and one Dy^{III} ion. The metallic core arrangements of **1·OH**, **2·Cl** and **3·Br** are identical, each displaying a sandwich-like structure in which the ligands Lz and salicylaldehyde construct two coordinate planes with the Dy^{III} ion sandwiched between them. The Dy^{III} ion is eight coordinated and resides in a square-antiprismatic environment where the planes are constructed by hydrogen bonding-associated Lz and salicylaldehyde ligands through the hydroxy oxygen of the salicylaldehyde and one hydrogen atom of the Lz ligand (Figure 1). Each complex is further charge balanced by counter ions in the lattice. Thus the only chemical differences found between the three complexes are the counter ions (OH^- for **1·OH**, Cl^- for **2·Cl** and Br^- for **3·Br**). Because of the different counter ions in **1·OH**, **2·Cl** and **3·Br**, the complexes display distinct properties that distinguish each other.

Structural Comparisons. As shown in Figure 1 and Table 1, the variation of counter ion in the lattice results in the modification of the coordination environment as well as the arrangement of ligands. In all three complexes, the Lz and salicylaldehyde ligands are in *cis* configuration relative to each other with the average Dy-N distance of 2.55 Å (for all three complexes) and Dy-O distances of 2.28 Å (**1·OH**) and 2.30 Å (**2·Cl** and **3·Br**). The anionic donor O atoms are far closer to the Dy^{III} ion than the neutral N atoms, highlighted by the Dy- O_{sali} distances, $\angle O_{sali}-Dy-O_{sali}$ angles (here the O_{sali} represents the hydroxy oxygen of the salicylaldehyde) and $\pi \cdots \pi$ stacking effects of ligands Lz, creating a predominantly axial distribution of the electronic structure. A closer look at the structure reveals that the Dy- O_{sali} distances and $\angle O_{sali}-Dy-O_{sali}$ angles are dominated by the distance and angle between the upper and lower coordinate planes (θ) as well as the rotation angle between them (ϕ). In complex **1·OH**, a relatively

large θ angle (6.84°) with respect to **2·Cl** (2.53°) and **3·Br** (2.59°) indicates a constriction of the coordinate planes at the O_{sali} atoms side (Figure S1), resulting in a short Dy- O_{sali} distance of 2.20 Å (2.25 and 2.24 Å for **2·Cl**, 2.24 Å for **3·Br**, Tables 1 and S3). The $\angle O_{sali}-Dy-O_{sali}$ angle of 149.02° for **1·OH** is larger than 142.86° for **2·Cl** and 142.96° for **3·Br** (140.46° for perfect D_{4d} geometry), which is also consistent with the large ϕ angle of 58.65° for **1·OH** (46.95° , 46.94° for **2·Cl** and **3·Br**, respectively). The $\pi \cdots \pi$ stacking effects are also influenced by the rotation angles ϕ of the Lz ligands. When the ϕ angles increase, the $\pi \cdots \pi$ distances get larger, therefore the $\pi \cdots \pi$ distance of 3.826 Å for **1·OH** is larger than the values of 3.596 Å, 3.590 Å for **2·Cl** and **3·Br** (Table 1). The SHAPE software⁷² was used to quantify the coordination geometry of the Dy^{III} center and gave the continuous shape measure deviations (CSMD) of 1.327, 0.643 and 0.725 for **1·OH**, **2·Cl** and **3·Br** (Table S2, D_{4d} coordination symmetry), respectively. This demonstrates that the coordination polyhedron in **1·OH** is more distorted than in **2·Cl** and **3·Br**. Moreover, we investigated the closest intermolecular Dy $\cdots$ Dy distances to check their effects on the dynamical magnetic properties. In the case of complex **1·OH**, the relatively short Dy $\cdots$ Dy distance (7.68 Å) does not necessarily preclude weak dipolar interactions since the later are also governed by the angles between the magnetic moments (*vide infra*). The Dy $\cdots$ Dy distances in **2·Cl** (8.09 Å) and in **3·Br** (8.08 Å) are a little longer, which also suggests the presence of potential weak intermolecular magnetic interactions.

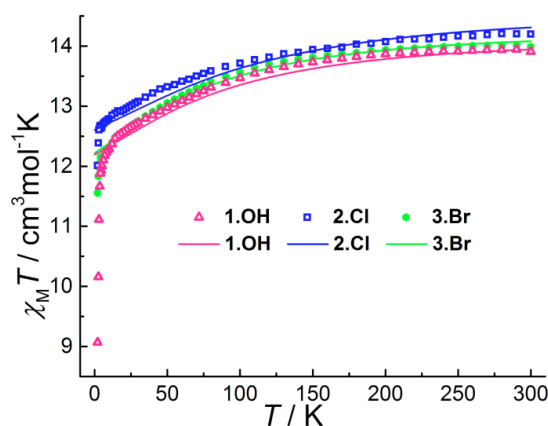


Figure 2. Temperature dependence of $\chi_M T$ product at 1 kOe for **1·OH** (red), **2·Cl** (blue) and **3·Br** (green). The solid lines correspond to *ab initio* calculations.

Magnetic properties. In order to probe the magnetic properties, direct current (dc) magnetic susceptibility measurements were collected on polycrystalline samples under an applied field of 1 kOe in the temperature range 2–300 K. The $\chi_M T$ (χ_M molar magnetic susceptibility) vs. T plots for **1-OH**, **2-Cl** and **3-Br** (Figure 2), reveal the room temperature $\chi_M T$ values of 13.91, 14.18 and 13.91 cm³·K·mol⁻¹, respectively, which are in good agreement with the value

expected for the free-ion approximation of Dy^{III} ions ($S = 5/2$, $L = 5$, ${}^6H_{15/2}$, $g = 4/3$, $C = 14.17$ cm³·K·mol⁻¹). When reducing the temperature, the $\chi_M T$ value gradually decreases before a sharp drop below 15 K, reaching the values of 9.07, 12.01 and 11.56 cm³·K·mol⁻¹ for **1-OH**, **2-Cl** and **3-Br**, respectively. The high temperature decrease can be attributed to the depopulation of the

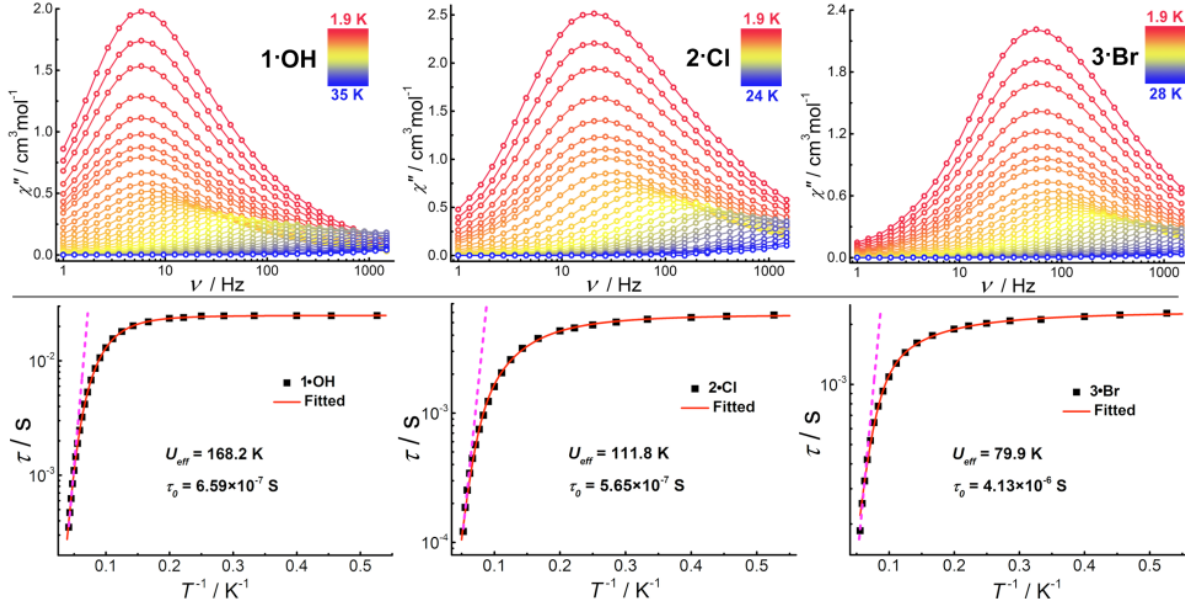


Figure 3. Frequency-dependent out of phase ac susceptibilities (top) and plots of τ vs. T^{-1} (bottom) for **1-OH** (left), **2-Cl** (middle) and **3-Br** (right) under zero dc field. The red line represents the fitting to eq. 1, pink dash line represents the Arrhenius fitted result.

excited m_J sublevels of the ground J states of the Dy^{III} ions, while the sharp drop at lower temperature region suggests non-negligible dipolar interactions between the neighboring Dy^{III} ions, especially for **1-OH** (shortest Dy...Dy distance of 7.68 Å). The isothermal molar magnetization (M) vs. H plots for **1-OH**, **2-Cl** and **3-Br** (Figures S3-S5) show sharp increase with increasing H at low fields and then linear increases at high fields, reaching the values of 5.32, 5.31 and 5.48 μ_B at 5 T, for **1-OH**, **2-Cl** and **3-Br**, respectively, which are very close to the expected value of 5 μ_B for Dy^{III} ions in a pure $m_J = |\pm 15/2\rangle$ ground state.

Variable temperature and frequency alternating current (ac) susceptibility measurements were also performed to probe the slow relaxation of the magnetization and quantum tunneling effects within these molecules, utilizing a 3 Oe oscillating field and a zero-applied direct current. Both the in-phase (χ') and out-of-phase (χ'') susceptibilities display a temperature (Figure S6) and frequency (Figures 3 and S7-S9) dependence below 35 K for **1-OH** and 25 K for **2-Cl** and **3-Br**, indicating a typical SMM behavior. Temperature-dependent out-of-phase (χ'') magnetic susceptibility signals exhibit peak at 27 K for **1-OH** and 19 K for **2-Cl** and **3-Br** (Figure S6). Upon cooling, χ'' signals

increase rapidly at low temperature region, which could be attributed to quantum tunneling effects at zero dc field. Frequency-dependent magnetic susceptibility data under zero applied dc field show frequency-independent regime at 6.4, 24.1 and 62.9 Hz for **1-OH**, **2-Cl** and **3-Br**, respectively (Figure 3), confirming classical quantum tunneling. The quantum tunneling relaxation times τ_{QTM} were extracted as 24.9, 5.78 and 2.33 ms for **1-OH**, **2-Cl** and **3-Br**, respectively. When increasing the temperature, we find that the relaxation follows a thermally activated mechanism and the maxima of the out-of-phase χ'' peaks for all three complexes shift toward higher frequency. The τ vs. $1/T$ plots at high temperature were fitted with Arrhenius law [$\tau = \tau_0 \exp(U_{eff}/T)$], yielding anisotropy barriers U_{eff} of 168.2, 111.8 and 79.9 K for **1-OH**, **2-Cl** and **3-Br**, respectively. Accordingly, we fixed τ_{QTM} and U_{eff} to the above values to avoid the over parameterization and then fitted the entire temperature range using the following equation:⁷³⁻⁷⁷

$$\frac{1}{\tau_{obs}} = \frac{1}{\tau_{QTM}} + AH^2T + CT^n + \tau_0^{-1} \exp\left(\frac{-U_{eff}}{T}\right) \quad (1)$$

where $1/\tau_{\text{QTM}}$, AH^2T , CT^n and $\tau_0 \cdot \exp(U_{\text{eff}}/T)$ represent quantum tunneling, direct, Raman and Orbach relaxation processes,^{78, 79} respectively. Herein, the direct process is cancelled since the measurements were performed in the absence of a dc field. The best fits gave pre-exponential factors τ_0 of 6.59×10^{-7} (**1-OH**), 5.65×10^{-7} (**2-Cl**) and 4.13×10^{-6} s (**3-Br**). Other parameters obtained from the fitting are listed in Table S3.

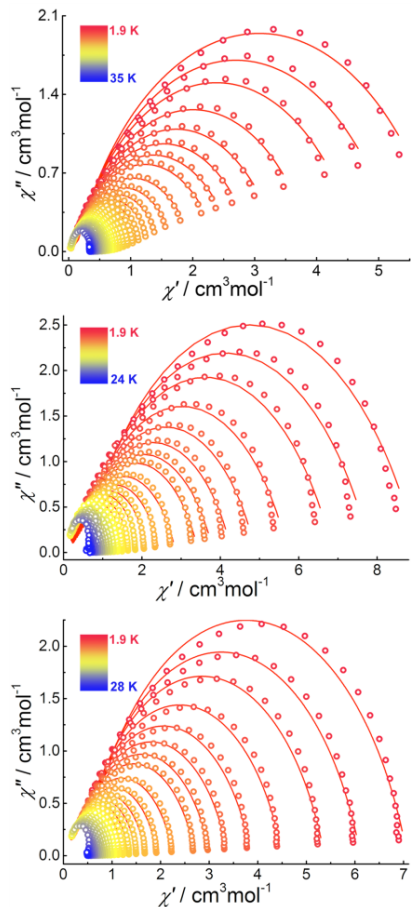


Figure 4. Cole-Cole plots for complexes **1-OH** (top), **2-Cl** (middle) and **3-Br** (bottom) at indicated temperatures under zero dc field. The solid lines indicate the best fits to the experiments with the generalized Debye model.

Cole-Cole plots of χ'' vs. χ' (Figure 4) reveal semi-circular profiles, indicating a single relaxation process is operative for all complexes. The plots were fitted to a generalized Debye model with relatively small α parameters being in the range 0.04-0.26 (**1-OH**), 0.07-0.32 (**2-Cl**) and 0.08-0.24 (**3-Br**), indicating a very narrow distribution of relaxation times for each complex.⁸⁰⁻⁸²

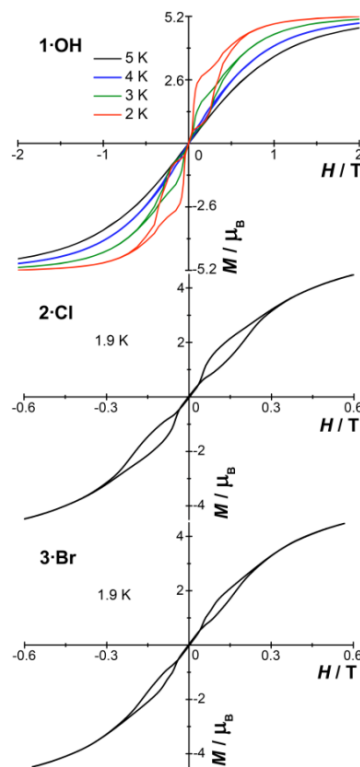


Figure 5. Magnetic hysteresis loops for **1-OH** (top), **2-Cl** (middle) and **3-Br** (bottom) with clear openings at $H \neq 0$.

We then probed the magnetic relaxation on polycrystalline samples of **1-OH**, **2-Cl** and **3-Br** via M vs. H hysteresis measurements, using a conventional SQUID magnetometer. As shown in Figure 5, all three complexes display clearly butterfly shaped magnetic hysteresis loops with openings up to 4 K for **1-OH** and 1.9 K for **2-Cl** and **3-Br** at $H \neq 0$, indicating the presence of magnetic blocking. The loss of the magnetization at zero field indicates the existence of quantum tunneling effect, which was apparent on the fast time scale (τ_{QTM}) of the dynamic ac experiment. The less opening ($H \neq 0$) of hysteresis loops at 1.9 K for **2-Cl** and **3-Br** suggest the much faster quantum tunneling than the one for **1-OH**. This is also coincident with the shorter quantum tunneling time τ_{QTM} for **2-Cl** and **3-Br** at zero field.

In order to reduce the dipole-dipole interactions between magnetic centers and slow down relaxation, magnetic dilution studies were performed on complexes **1-OH** and **2-Cl**. AC susceptibility measurements show great enhancement of the magnetic relaxation (Figures S10-S11), giving the effective energy barrier of 236.7 and 123.1 K for **1-OH** and **2-Cl**, respectively (Figure 6), which are much larger than the undiluted samples. The extracted τ_{QTM} are as long as 338.7 and 80.4 ms for **1-OH** and **2-Cl**, respectively, indicating the quantum tunneling at low temperature is also effectively suppressed. In addition, magnetic hysteresis measurements were also conducted on the diluted samples of **1-OH** and **2-Cl**. As expected, significant

improvements were observed (Figure 6). While, the hysteresis loops are still closed at zero field, suggesting the intrinsic fast quantum tunneling, which is probably induced by the electronic structure and Hyperfine-Interaction.^{49, 83}

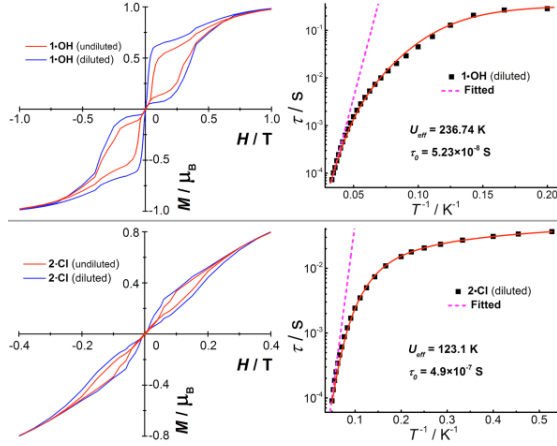


Figure 6. Magnetic hysteresis loops (left) and plots of τ vs. T^{-1} (right) for the diluted samples of **1-OH** (top) and **2-Cl** (bottom) at 1.9 K.

Table 2. Geometry parameters and energy barriers for the square antiprism dysprosium derivatives.

	O-Dy-O / °	Dy-O / Å	$\pi \cdots \pi$ / Å	ϕ / °	CShM (D_{4d})	τ_{QTM} / ms	U_{eff} / K
1-OH	149.018	2.203	3.826	58.646	1.327	24.9	168.2
2-Cl	142.855	2.247	3.596	46.946	0.643	5.78	111.8
3-Br	142.958	2.239	3.590	46.942	0.725	2.33	79.9

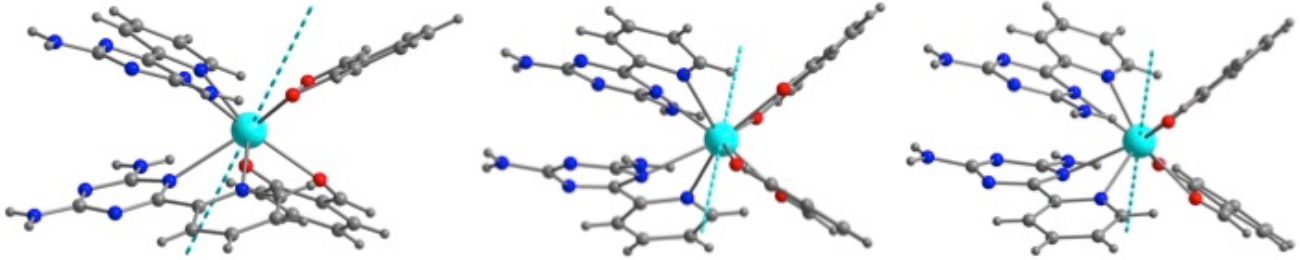


Figure 7. Orientation of the calculated main magnetic axes of the ground Kramers doublets on Dy^{III} centers in complexes **1-OH** (left), **2-Cl** (middle), and **3-Br** (right).

Ab initio Calculations. Further insights into the electronic structure was performed by *ab initio* CASSCF/RASSI-SO calculations on the X-ray structures of **1-OH**, **2-Cl** and **3-Br** (see computational details). From the calculations, the energies of the eight Kramer doublets, the anisotropic tensors as well as the directions of the ground doublets were determined (see Tables S5-S7). As expected, in the three complexes, the multiplet ground state is ${}^6H_{15/2}$, with almost pure $m_J = |\pm 15/2\rangle$ ground state. The isolated ground states and the minimum mixing of the m_J 's in the first excited states are in line with the perfect Ising-type anisotropy of Dy^{III} center under the pseu-

The coexistence of magnetic blocking and quantum tunneling in these complexes are possibly related with the structural features of each complex. As already inferred from the crystal structures, the variation of Dy-O_{sali} distances, $\angle$ O_{sali}-Dy-O_{sali} angles and $\pi \cdots \pi$ stacking effects probably have critical impact on the electronic structure of the complex, thus should be responsible for the distinct magnetic anisotropy (Table 2). In complex **1-OH**, the relatively short Dy-O_{sali} distance and large $\angle$ O_{sali}-Dy-O_{sali} angle profit for a strong axial crystal field, thus inducing a large energy barrier of magnetization reversion. While the $\pi \cdots \pi$ stacking effects between two Lz ligands might introduce large transversal components and short-cut fast relaxation process. For complexes **2-Cl** and **3-Br**, the Dy^{III} ions are in similar coordination environment, while the stronger $\pi \cdots \pi$ stacking effect would probably aggravate the quantum tunneling. Indeed, the quantum tunneling relaxation rates of **2-Cl** (2.33 ms) and **3-Br** (5.78 ms) are faster than that of **1-OH** (24.9 ms).

do- D_{4d} coordinating crystal field. Moreover, the calculated $\chi_M T$ vs. T curves well reproduce the experimental ones (Figure 2). The calculated easy magnetic axes of ground state for the three compounds are represented in Figure 6. As expected, they lie in the most negatively charged direction *i.e.* almost parallel to O_{sali}-O_{sali} direction and correspond to a direction perpendicular to the pseudo C₂ axis. The deviation angle (β) between the magnetic axes of ground state and first excited state for **1-OH** (1.73°) is slightly smaller than those in **2-Cl** (2.27°) and **3-Br** (4.68°), suggesting a more axial ligand field and highlighting the importance of short Dy-O_{sali} distances and large $\angle$ O_{sali}-

Dy-O_{sali} angles in **1**·OH. The computed magnetization blocking barriers in **1**·OH, **2**·Cl and **3**·Br are given in Figures S12–S14. They feature clearly thermally activated processes (Raman+Orbach) through the first and second excited states. The energy splitting of the ${}^6H_{15/2}$ ground multiplet is much larger in **1**·OH (229.32 cm⁻¹ for first excited state) than in **2**·Cl (186.52 cm⁻¹, idem) and **3**·Br (166.61 cm⁻¹, idem), and the transition probability (through the barrier) are clearly smaller in **1**·OH than in the two other congeners. This could probably be ascribed to the less $\pi\cdots\pi$ stacking effect enhancing the energy splitting of the ${}^6H_{15/2}$ ground multiplet, which was demonstrated in our previous work.⁵⁷ Herein, the transition probability is directly related to the fast quantum tunneling relaxation. The much larger transversal components are coincident with the faster quantum tunneling relaxation in **2**·Cl and **3**·Br than in **1**·OH (*vide supra*).

CONCLUSION

In conclusion, three closely related dysprosium SMMs with mixed salicylaldehyde/pyridin-triazine ligands and discriminative counter-anion, **1**·OH, **2**·Cl and **3**·Br, are structurally and magnetically characterized. We find that the three complexes all possess similar square-antiprismatic geometry (D_{4d}) but display different dynamic magnetic properties. Each complex displayed a unique anisotropy barrier and quantum tunneling relaxation, with U_{eff} and τ_{QTM} in the ranges of 80–168 K and 2.3–25 ms, respectively. Structural and magnetic studies reveal that the magnetic anisotropy is dominated by the axial ligand field, highlighted by short Dy-O_{sali} distances and large $\angle$ O_{sali}-Dy-O_{sali} angles, while the quantum tunneling relaxation is probably dictated by the $\pi\cdots\pi$ stacking of the Lz ligands which results in an axial constriction of the coordinating plane. *Ab initio* calculations support the different magnetic behaviors in these complexes. It appears that strong axial ligand field and less stacking of the auxiliary ligands may provide the key for unlocking even higher excited state and prevent the magnetic quantum tunneling. Notably, the counter ion facilitates the modulation of the square-antiprismatic coordination geometry in this system, therefore, providing a possibility to promote anisotropy barriers and suppress the quantum tunneling effect. This work highlights the importance of axial ligand field and ligands stacking in obtaining larger energy barriers and controlling the quantum tunneling of Dy^{III} based SMMs with square-antiprismatic geometry.

ASSOCIATED CONTENT

Supporting Information.

Physical measurements; crystallography; computational details; structural, magnetic and calculated tables and figures. This material is available free of charge via the Internet at <http://pubs.acs.org>.

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Notes

The authors declare no competing financial interest.

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Axial ligand field with short Dy-O_{sal} distance and less π - π stacking found in closely related dysprosium compounds with discriminative counter-anion where the metal in square-antiprismatic geometry suppress quantum tunneling and enhance the magnetic relaxation.

