Increasing the effective energy barrier promoted by the change of a counteranion in a Zn–Dy–Zn SMM: slow relaxation via the second excited state


The trinuclear complex [ZnCl₂(L)Dy(μ-L)ClZn]PF₆ exhibits a single-molecule magnetic behaviour under zero field with a relatively large effective energy barrier of 186 cm⁻¹. 

Ab initio calculations reveal that the relaxation of the magnetization is symmetry-driven (the DyIII ion possesses a C₂ symmetry) and occurs via the second excited state.

In recent years, much attention has been paid to zero-dimensional lanthanide coordination compounds exhibiting slow relaxation of the magnetization and magnetic hysteresis below a certain temperature known as the blocking temperature (Tₖ). These nanomagnets, called single-molecule magnets (SMMs), present potential future applications in fields such as molecular spintronics, and ultra-high density magnetic information storage and as qubits for quantum computing at the molecular level. The SMM behaviour arises from the existence of an anisotropic energy barrier (U) that prevents magnetization reversal below Tₖ when the magnetic polarizing field is removed (the magnetization is frozen parallel to the magnetic field). The larger the energy barrier, the higher is the Tₖ value. Therefore, to increase the energy barrier and to improve the SMM properties, systems with large magnetic anisotropy must be designed.

Lanthanide ions exhibit strong magnetic anisotropy due to the combination of strong spin–orbit coupling and crystal-field effects promoted by the surrounding ligand and therefore are excellent building blocks to be used in the construction of coordination compounds with SMM properties. In fact, numerous 3d/4f and 4f (and 5f) mononuclear and polynuclear coordination compounds, most of them containing DyIII ions, have been shown to exhibit SMM behaviour. Interestingly, mononuclear and low-nuclearity 4f metal complexes have been shown to possess energy barriers that are an order of magnitude higher than that observed in 3d and 3d/4f polynuclear clusters. The SMM behaviour observed for low-nuclearity 4f metal complexes generally is due to the individual LnIII ions rather than the whole molecule. Moreover, Ln···Ln interactions frequently favour fast quantum tunnelling of the magnetization (QTM) in the ground state, which reduces the energy barrier to an effective value, Uₑff. Nevertheless, when the complex containing paramagnetic LnIII ions is diluted in an isostructural diamagnetic lattice, the Ln···Ln interactions are reduced and the energy barrier often increases. It should be noted that although the Uₑff values of Ln-based species are very large, they rarely show an open hysteresis loop at zero field, which is due to QTM. Theoretical and experimental studies carried out by us and others have shown that the incorporation of diamagnetic metal ions such as ZnII in a DyIII complex is a good strategy to enhance Uₑff. This fact could be due to two factors: (i) the quenching of the Ln···Ln interactions promoted by the presence of diamagnetic ZnII ions (some kind of internal magnetic dilution) and (ii) the increase of electron density on the phenoxide oxygen donor atoms connecting the ZnII and DyIII ions provoked by the coordination to the ZnII ions. By exploiting these features of the ZnII–DyIII systems and playing with the distribution of donor atoms on the DyIII coordination environment following the guidelines of the simple prolate–oblate electrostatic model, we were able to rationally design a Zn–Dy–Zn SMM, [ZnCl₂(L)Dy(μ-L)ClZn][ZnCl₄(CH₃OH)₂] with a large effective energy barrier at zero field.

In this communication, we have shown how the replacement of the counteranion in the complex [ZnCl₂(L)Dy(μ-L)ClZn][ZnCl₄(CH₃OH)₂] (H₂L = N,N'-dimethyl-N,N'-bis(2-hydroxy-3-formyl-5-bromo-benzyl)ethylenediamine; Fig. 1) by PF₆⁻ affects the DyIII coordination environment leading to an important enhancement of the Uₑff.

The reaction of H₂L with ZnCl₂ and subsequently with Dy(NO₃)₃·5H₂O and KPF₆ in methanol affords well shaped
yellow prismatic crystals of [ZnCl(μ-L)Dy(μ-L)(Cl)Zn]PF$_6$ (I). The X-ray crystal structure of I (Fig. 1; selected bond distances and angles are given as the ESI$^+$) consists of [ZnCl(μ-L)Dy(μ-L)(Cl)Zn]$^+$ cations and PF$_6^-$ anions. Within the trinuclear Zn-Dy-Zn cation, two [ZnCl(μ-L)]$^+$ units are coordinated to the central Dy ion through the phenoxide and aldehyde oxygen atoms of the compartmental ligand L$^{-}$, giving rise to a DyO$_6$ coordination polyhedron with Dy-Ophenoxide distances of 2.340(3) Å and 2.275(3) Å and Dy-O_alddehyde distances of 2.386(3) Å and 2.393(3) Å. The two halves of the trinuclear cation are symmetrically related by a C$_2$ axis passing through the Dy and P atoms.

Calculations of the degree of distortion of the DyO$_6$ coordination polyhedron with respect to an ideal eight-vertex polyhedron using the continuous shape measure theory and the SHAPE software (see Table S3 in the ESI$^+$) pointed out that the DyO$_6$ coordination polyhedron can be considered as an intermediate between square antiprism, triangular dodecahedron and biaugmented triangular prism ideal polyhedra, but closer to the former geometry. In the square antiprism description, the average distance between the upper and lower planes containing the four oxygen atoms (two phenoxo and two aldehyde) is 2.565 Å, whereas the average distance of the sides of the squares defined by the oxygen atoms in each plane is 2.782 Å and therefore the distorted square-antiprism is compressed. Compared to [ZnCl(μ-L)Dy(μ-L)(Cl)Zn][ZnCl$_2$(CH$_3$OH)], compound I shows a Dy$_3$O$_9$ geometry that is closer to the square antiprism (see Table S3, ESI$^+$) and presents a different distribution of the phenoxide oxygen atoms on the Dy$_{III}$ coordination sphere (see below and Fig. S5, ESI$^+$). Zn$_{II}$ ions exhibit a distorted square-pyramid CN$_4$Cl coordination environment, in which the nitrogen and phenoxide oxygen atoms occupy the basal positions with Zn-N bonds distances of 2.106 Å and 2.134 Å and Zn-O bond distances of 2.066 Å and 2.138 Å, whereas the axial position is occupied by the Cl atom at a longer distance of 2.217 Å. The Zn····Dy distance is 3.550(3) Å, whereas the Zn····Dy····Zn angle is 109.48(1)$^\circ$. The shortest Dy····Dy distance between neighbouring [ZnDy(Zn)]$^+$ trinuclear cations is 9.199(2) Å.

Direct-current (dc) magnetic susceptibility measurements were carried out on a randomly oriented polycrystalline sample of I in the 2–300 K temperature range and under an applied magnetic field of 0.1 T (Fig. 2, top left). The room temperature

$\gamma_M T$ value for I of 14.45 cm$^3$ K mol$^{-1}$ is compatible with the calculated value of 14.17 cm$^3$ K mol$^{-1}$ for the ground state of the Dy$_{III}$ ion (4f$^7, J = 15/2, S = 5/2, L = 5, g = 4/3$ $^1H_{15/2}$) in the free-ion approximation. Upon cooling, the $\gamma_M T$ product decreases steadily to a value of 12.40 cm$^3$ K mol$^{-1}$ at 5 K. Below this temperature, the $\gamma_M T$ product decreases sharply, reaching a value of 10.15 cm$^3$ K mol$^{-1}$ at 2 K. This behaviour is associated with the magnetic anisotropy of the Dy$_{III}$ ion and possible weak intermolecular interactions, which could be responsible for the sharp decrease in $\gamma_M T$ below 5 K. The field dependence of the magnetization at 2 K shows a relatively rapid increase at low field, reaching almost saturation for magnetic fields larger than 2 T. The saturation value of 5.10 N $(\mu_B)^2$ agrees well with those observed for Dy$_{III}$ complexes with strong magnetic anisotropy.$^5$

In order to know if I exhibits SMM properties, dynamic alternating current (ac) magnetic measurements were performed as a function of both temperature and frequency on a microcrystalline powder sample (Fig. 2, top right and Fig. S1–S3 in the ESI$^+$). At zero dc field, I exhibits a strong frequency dependence of the out-of-phase susceptibility ($\chi''_M$) signals below 35 K, which clearly indicates the occurrence of slow relaxation of the magnetization and SMM behaviour in this compound. This behaviour is not surprising in view of the easy-axes anisotropy of the Dy$_{III}$ ion (see below for ab initio calculations). The fact that both $\chi_M$ and $\chi''_M$ components do not go to zero below the maxima at low temperature, points out to the existence of fast relaxation of the magnetization via a QTM mechanism. The Cole–Cole plot (Fig. S2 in the ESI$^+$) shows semicircular shapes with z values in the 0.28 (17 K)–0.04 (25 K) range, thus suggesting the existence of multiple relaxation processes.

The relaxation times extracted from the frequency-dependent susceptibility data in the 18–27 K range were fitted to the Arrhenius law, affording an effective energy barrier for the reversal of the
magnetization of 186 cm\(^{-1}\) with \(\tau_0 = 4.98 \times 10^{-10}\) s (Fig. 2, bottom left).

When the ac measurements on 1 were performed in the presence of a small external dc field of 1000 Oe (this field was chosen because under its application the relaxation process was shown to be the slowest) to partly or fully suppress the QTM, the tails at low temperatures almost disappeared and the temperature peaks remained at similar temperatures and with comparable intensities to those observed under zero dc applied field (Fig. 2, inset). The fit of the relaxation times in the 18–27 K range to the Arrhenius law leads, as expected, to an increase of the thermal energy barrier and a decrease of \(\tau_0 (U_{\text{eff}} = 222\text{ cm}^{-1}\) with \(\tau_0 = 5.66 \times 10^{-11}\) s). In the high temperature region, the \(x\) values extracted from the Cole–Cole plot (Fig. S2 in the ESIF) are in the 0.22 (19 K)–0.08 (25 K) range, which points to the existence of a distribution of the relaxation times.

The energy barrier for 1 is approximately twice that of [ZnCl[μ-L]Dy[μ-L]C2Zn][ZnCl2(CH3OH)]\(^{2-}\). With the aim of rationalizing this surprising increase of the energy barrier promoted by the change of the counteranion, we have performed \textit{ab initio} electronic calculations based on the CASSCF + RASSI/SINGLE/ANISO\(^{20}\) method using the MOLCAS 7.8 code\(^{3}\) (Table S5, ESIF). The energy spectrum of the eight Kramer’s doublets (KDs) spans up to 491.8 cm\(^{-1}\) and the energy gap between the ground and first excited state is 144 cm\(^{-1}\), which are rather normal values for Dy\(^{3+}\) complexes. The ground Kramer’s doublet (KD1) is almost pure (\(m_I = \pm 1/2\)) with effective \(g_e\) and \(g_{xy}\) values approaching 20 and zero, respectively. Therefore KD1 is an almost ideal Ising state which favours the slow relaxation of the magnetization and the SMM behaviour, in accordance with the zero-field SMM properties observed for 1. It is worth mentioning that the \textit{ab initio} calculations accurately reproduce the direct current (dc) magnetic susceptibility and magnetization data for 1 (Fig. 2 top left), which gives confidence in the calculated low-lying KDs energies. The main magnetic axis of the KD1 lies close to the two Dy–O2A vectors (Fig. 3), which shows the tendency of this axis to point to the donor atoms with greater electron density and shorter Dy–O distances (O2A phenoxide bridging atoms of the ligands with Dy–O2A = 2.275 Å). This orientation of the magnetic moment forces the oblate electron density of the Dy\(^{3+}\) to be perpendicular to the O2A atoms thus diminishing electrostatic repulsions. The O2A phenoxide atoms are roughly located at axial positions on the DyO\(_6\) coordination sphere, thus emphasizing the suitability of an axially repulsive coordination environment for achieving SMM properties in Dy\(^{3+}\) compounds, as qualitatively anticipates the oblate-prolate model.\(^{19}\) DFT calculations of the electrostatic potential maps provoked by the ligands projected on the Dy\(^{3+}\) position (Fig. 3 right) agree with the above qualitative electrostatic predictions.

Thus, the oblate beta electron density of the ground state (which is formed by mixing in the ground RASSI wave function the two first, almost degenerate, states in the CASSCF step; Fig. S4 and Table S6, ESIF) is hosted in the region with less repulsion (yellow-green in Fig. 3, right) whereas the magnetic moment is pointing towards the strongest repulsion regions (red in Fig. 3, right).

It should be noted that the extracted effective energy barrier at zero dc field is 40 cm\(^{-1}\) larger than the energy gap between the ground and first excited KDs and therefore the relaxation of the magnetization cannot take place \textit{via} the first excited state. However, the relaxation could occur through the second excited KD, which would be located 243.3 cm\(^{-1}\) above the ground KD. In such a case, the extracted \(U_{\text{eff}}\) would be lower than that predicted using electronic structure calculations, which could be justified by the existence of fast QTM relaxation. In fact, the value of the energy barrier extracted from the ac measurements under an applied dc field of 1 kOe (\(U_{\text{eff}} = 222\text{ cm}^{-1}\)), when the QTM relaxation is almost suppressed, is close to the energy of the second excited state. The relaxation \textit{via} the first excited state is suppressed because the main anisotropy axes of the ground and first excited states (the latter is an almost pure Ising state with \(m_I = \pm 13/2, g_e \sim 17\) and \(g_{xy} < 0.05\)) are not only Ising like, but also almost parallel (the angle is 2.4°). The collinearity of the principal anisotropy axes up to the first excited KD, which is due to the presence of a \(C_2\) axis around the Dy\(^{3+}\) ion, blocks Orbach relaxation.\(^{18,20}\)

In order to probe the magnetization relaxation mechanism in 1, we have computed the transition magnetic moment matrix elements between the connecting pairs of opposite magnetization (Fig. 4 and Fig. S6, ESIF). Owing to the almost pure Ising nature of the ground state, QTM is expected to be very weak, which is supported by the low magnitude of the transition magnetic moment element between the ground state KDs (\(\sim 10^{-4}\) \(\mu_B\)). This must be the reason why 1 exhibits SMM behaviour at zero field. The same argument can be used to justify the absence of thermal-assisted quantum tunnelling of magnetization (TA-QTM) through the first excited state. However, the TA-QTM mechanism \textit{via} the second excited KD (\(\pm 3\) states) becomes dominant as it exhibits the largest value of the transition magnetic moment element (1.5 \(\mu_B\)). As expected for the collinearity of the main anisotropy axes of the ground and first excited KDs, the off-diagonal term connecting these KDs (related with the Orbach process) is very small (\(\sim 10^{-7}\) \(\mu_B\)). Nevertheless, the Orbach process is operative \textit{via} the second excited state as the transversal magnetic moments are moderate (0.33 and 0.58 \(\mu_B\)). Therefore, the relaxation of the magnetization occurs through the second excited state \textit{via} Orbach/TA-QTM processes.

Although the Dy–O distances in 1 and [ZnCl[μ-L]Dy[μ-L]C2Zn][ZnCl2(CH3OH)]\(^{2-}\) are very similar, the fact that the former exhibits a \(C_2\) symmetry axis around the Dy\(^{3+}\) ion forces the
Fig. 4 Lowest three Kramer’s doublets (KDs) and the ab initio computed relaxation mechanism in 1. The thick black lines imply KDs as a function of their magnetic moment along the main anisotropy axis. The red lines correspond to the ground state QTM and TA-QM via the first and second excited KDs, blue lines show possible Orbach relaxation processes. The values indicated close to the arrows indicate the matrix elements of the transition magnetic moment.

phenoxyide oxygen atoms to adopt a different distribution in the DyIII coordination sphere with respect to that observed for the latter (in 1 the phenoxyide oxygen atoms are closer to each other and the Zn-Dy-Zn angle is much smaller, 109.48° vs. 141.7°; see Fig. 1 left and Fig. S5, ESIF), which would ultimately be responsible for the relaxation mechanism through the second excited KD.

At 2 K and using the sweep rate accessible in a conventional SQUID magnetometer, compound 1 exhibits a butterfly shaped hysteresis loop with almost negligible magnetization at zero field (Fig. 2 bottom, right), which is consistent with the QTM generally found for 4f containing complexes and with the tail that 1 exhibits at low temperature in the $M^s$ vs. $T$ plot at zero applied field.

In summary, we have shown how the replacement of the counteranion in a Zn–Dy–Zn trinuclear complex provokes a change towards a more symmetric structure where there exists a C2 axis, or axis of the DyIII ion. The presence of the C2 axis imposes collinearity of the anisotropic axes of the two lowest KDs, so that the thermal activated relaxation is suppressed via the first excited KD and takes place via the second excited state, giving rise to a dramatic increase in the effective energy barrier. This is one of the few cases where this type of symmetry-driven relaxation mechanism has been observed [3,10], and the first case where it is due to a structural change promoted by the replacement of the counteranion.

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Notes and references


