Synthesis, structure and magnetic properties of a series of Ln( iii ) complexes with radical-anionic iminopyridine ligands: effect of lanthanide ions on the slow relaxation of the magnetization

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Synthesis, structure and magnetic properties of a series of Ln(III)
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We report the synthesis, structure and magnetic properties
investigation of a series of homoleptic Ln(III) complexes
coordinated by radical-anionic iminopyridine ligands of general
formula [Ln(IPy)_4]_X-solv (IPy = iminopyridine; Ln = Tb, Dy, Er, Y, Gd).
The dysprosium analogue exhibits a zero-field Single-Molecule
Magnet (SMM) behavior.

Owing to their exceptional magnetic properties, lanthanide
coordination and organometallic chemistries have recently
taken a decisive turn in materials science. In this line of
thought, Single-Molecule Magnets (SMMs) based on
lanthanide ions have totally changed the molecular magnetism
landscape and are often considered as future candidates for
high-density information storage or quantum computing. In
such complexes, the high magnetic anisotropy of the
lanthanide ion in a particular environment gives rise to
the appearance of an energy barrier, \( \Delta \), separating two opposites
magnetic states and resulting in a superparamagnetic
behaviour.

Yet, designing high performance SMMs requires to accurately
control the coordination environment of the lanthanide ions in
order to maximize the anisotropy and crystal-field splitting,
while minimizing the Quantum Tunnelling of the
Magnetization (QTM). This latter provides underbarrier
relaxation pathways that limit the SMM performances, such as
the energy barrier and magnetic coercivity. Although the
physical origin with respect to QTM is different, Raman and
direct processes provide also underbarrier relaxations that
reduce the capacity of the SMMs. The coordination
environment around the lanthanide ion depends on both
electronic and symmetry considerations generated by the
contiguous ligands. Taking advantage of this strategy, several
lanthanide complexes exhibiting very high anisotropic barriers of
few thousands of cm\(^{-1}\) have been reported within the last
few years, while a dysprosium metalloocene complexes shows a high temperature hysteresis or that could even overcome liquid nitrogen’s boiling temperature.

Although QTM is known to depend on symmetry, dipolar and
hyperfine interactions, the presence of intermolecular
exchange interactions could greatly influence the relaxation
dynamics in a complex fashion. On one hand, sometimes the
presence of a neighbouring paramagnetic centre to the
lanthanide site can produce an internal field that could suppress the tunnelling. On the other hand, enhancement of the relaxation dynamics has also been observed in dinuclear systems exhibiting crystallographically independent lanthanide
sites due to the creation of a local transverse magnetic
moment. In addition, the strength of the exchange
interaction also influences the energy splitting between the
ground and excited states in polynuclear systems which should
be as high as large as possible to obtain large energy barrier
systems. With this in mind, several exchange biased systems
showing genuine SMM behaviour have been reported.

Due to the intern character of the \( f \) orbitals, radical ligands
could be viewed as particularly appropriate to promote
significant interactions and study the influence over the slow
relaxation of the magnetization. Many lanthanide-based
systems containing miscellaneous radical ligands, such as
nitronyl nitroxides, oxidized phthalocyanine, semiquinates,
verdazyls, triazinyl, pyridyl-pyrazine, indigo, or dinitrogen have been reported. For instance, the latest example has hold the record for the highest coercivity of 14 K for many years.

Among radical-lanthanide systems with great potentialities,
ytterbium-metalloccenes with various heterocyclic
amines have been studied. Remarkably, an exchange
coupling larger than hundreds of cm\(^{-1}\) has been reported in
several ytterbium metalloccenes complexes. Despite such
advantages, the investigation has been solely dedicated to
ytterbium-based complexes and the use of highly anisotropic
Kramers ions, such as Dy or Er, particularly suitable to
design SMMs has not been investigated. For instance, iminopyridines (IPy) belong to redox active ligands, but their
coordination chemistry with such anisotropic lanthanide ions
remains unexplored. Their great tunability in terms of both,
steric and electronic properties, allows tailoring the
coordination environment of the lanthanide site. In recent
years, the reactions of various ytterbocenes with diimino
ligands bearing various substituents were reported. Among
these complexes, a paramagnetic homoleptic Yb complex
coordinated by three iminopyridyl radical-anions has been
obtained through the oxidative cleavage of the \( \eta^5 \) Yb–Cp bond
(Cp = cyclopentadienyl). This complex shows a reduced
magnetic moment at room temperature that suggests a strong
coupling between the radical/ytterbium antiferromagnetic interaction.

Following this, we report in this communication the synthesis,
structure and magnetic studies of four homoleptic lanthanide
complexes based on iminopyridine (IPy) ligands. Remarkably,
the dysprosium analogue reveals a zero-field slow relaxation
reflecting a genuine SMM behaviour.
Complexes Ln(IPy)_3 (Ln = Tb (1), Dy (2), Er (3), Y (4), Gd(5)) were prepared by treating the potassium derivative of the related iminopyridine with LnCl_3 in 3:1 molar ratio (Scheme 1).


Complexes 1–5 were obtained as moisture and air-sensitive dark green crystalline solids in 75, 72, 70, 73 and 71 % yields, respectively. X-ray single crystal diffraction analysis (Fig. 1, Table S1) shows that compounds 1–5 crystallize in the hexagonal R3 space group as the solvates 1·0.5(THF), 2·0.5(Toluene), 3·0.5(Toluene) and 4·0.5(Toluene) and 5·0.5(Toluene). For the sake of clarity, only the crystal structure of 2 will be described. The molecular structure of 2 is shown in Figure 1. The coordination sphere is composed of six nitrogen atoms giving a hexacoordinated geometry close to a distorted octahedron as indicated by the SHAPE analysis (Table S2).

Further insights into the relaxation dynamics could be obtained by analysing the temperature dependence of ac susceptibilities (Fig. S3). Fitting of the Cole-Cole plots (Fig. S5) with a generalized Debye model show large values of the α parameter (i.e. 0.42) at low temperature, indicating an important distribution of relaxation times (Table S3). The slow relaxation of the magnetization was investigated by alternate currents (ac) measurements. Under a zero dc field, only the dysprosium analogue 2 shows a significant signal of out-of-phase susceptibility (χ″) component. The frequency dependence of χ″ at different temperatures shows the presence of a broad single peak (Fig. 2). Such out-of-phase signals could be observed up to 30 K on the temperature dependence of the ac susceptibilities (Fig. S4). The coordination sphere is composed of six nitrogen atoms giving a hexacoordinated geometry close to a distorted octahedron as indicated by the SHAPE analysis (Table S2). The Dy–N_ν distance of 2.424(2) Å is slightly longer than Dy–N_amine one of 2.392(2) Å. The bonding situation within the planar diamino fragment N_ν–C–C–N_amine in 2 is consistent with the radical anionic form of the ligand (N_amine–C = 1.339(3) Å; C–C = 1.394(2) Å; N_ν–C = 1.383(2) Å). The crystal packing shows that the shortest intermolecular distance Dy–Dy is equal to 8.306(2) Å, while no H-bonds have been detected (Fig. S1).

The radical-anionic nature of the iminopyridine ligands was further confirmed with Electron Paramagnetic Resonance (EPR) spectroscopy. While complexes 1–3 do not exhibit an observable EPR spectrum, 4 incorporating the diamagnetic Y^3+ exhibits a broad line (∆E = 62 G) at room temperature with g = 2.002 without any hyperfine splitting and visible anisotropy (Fig. S2). The g-factor value is typical for ligand-centered radicals.

The radical-anionic nature of the ligands was also corroborated by the direct current (dc) magnetic properties’ investigations of 4 incorporating the diamagnetic Y^3+ ion, which exhibits intra- and intermolecular antiferromagnetic interactions between the radicals (S = ½) (Fig. S3, see ESI for details). The χT vs T curves for 1–5 show a gradual then abrupt decrease from around 30 K, most likely associated with the thermal depopulation of the ± m_1 levels from the Ln^3+ ions combined with intra and intermolecular antiferromagnetic interactions (Fig. S3). The dc study of 5 presenting isotropic Gd^3+ ion reveals that the intramolecular radical-radical interactions seem to be much greater than the Gd^3+-radical ones.

The slow relaxation of the magnetization was investigated by alternate currents (ac) measurements. Under a zero dc field, only the dysprosium analogue 2 shows a significant signal of out-of-phase susceptibility (χ″) component. The frequency dependence of χ″ at different temperatures shows the presence of a broad single peak (Fig. 2). Such out-of-phase signals could be observed up to 30 K on the temperature dependence of the ac susceptibilities (Fig. S4). Fitting of the Cole-Cole plots (Fig. S5) with a generalized Debye model show large values of the α parameter (i.e. 0.42) at low temperature, indicating an important distribution of relaxation times (Table S3). Further insights into the relaxation dynamics could be obtained by analysing the temperature dependence of the relaxation time, τ. A clear deviation from the linearity could be observed at low temperature, pointing out a divergence from a thermally activated process. Hence, the all data range could be modelled using the following equation: τ_{ac} = τ_0 \exp(-\Delta/kT) + CT^{m} τ^{-\alpha \text{QTM}} (Eq. 1). The first term accounts for a thermally activated process, while the second and third ones stand for two-phonon Raman and QTM, respectively.
Surprisingly, the Raman process is found inoperative\(^42\) and as a consequence, the magnetization relaxes through a combination of thermally activated and QTM processes (Table 1). Considering solely a Raman and QTM processes leads to poor fitting confirming that the relaxation involves readily a thermally activated relaxation. In order to shortcut this latter, the field dependence of the relaxation time was monitored at 15 K (Fig. S6). The complex behaviour could not be modelled with the usual equation \(\tau^{-1} = DH^4T + B_1/(1 + B_2H^2) \pm K\) (Eq. 2), for which the first term accounts for the direct process (for Kramers-ion), the second one stands for the QTM, while the constant accounts for the field-independent Raman and thermally activated process. However, the data clearly shows that the field showing the highest relaxation time is 1000 Oe. The frequency dependency of the ac susceptibilities under this dc field shows the presence of a broad peak for \(\chi''\) (Fig. S7-S8, Table S4), while the temperature dependence of \(\tau\) confirms the decrease of the QTM (Fig. 3). The temperature dependence of \(\tau\) could be fitted by taking into account an additional direct process: \(\tau^{-1} = \tau_0^{-1}\exp(-\Delta/kT) + CT^{-m} + AT^{-n}\) (Eq. 4). The value of \(\Delta\) (Table 1) is found comparable with respect to the zero-field data. The large distribution of the relaxation times may be explained by the presence of the crystallographic disorder of solvent molecules and/or ligands, as well as by impact of inter and intramolecular interactions. In contrast, the analogues 1 and 3 do not show a significant out-of-phase component even in the presence of a dc field (Fig. S9-S10). The observed behaviour follows the usual trend expected by electrostatic model and considering the angular dependence of the \(4f\) electronic density for the Ln\(^3+\) ions.\(^{43}\)

### Table 1: Fit parameters of the temperature dependence of the relaxation time for 2.

<table>
<thead>
<tr>
<th>Compound</th>
<th>(\Delta) (cm(^{-1}))</th>
<th>(\tau_0) (s)</th>
<th>(m)</th>
<th>(C) (s(^{-1}).K(^{-m}))</th>
<th>(\tau_{\text{QTM}}) (ms)</th>
<th>(A) (s(^{-3}).K(^{-n}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 (0 Oe)</td>
<td>28 ± 4</td>
<td>(3 ± 1) \times 10(^{-5})</td>
<td>9*</td>
<td>-</td>
<td>1.0 ± 0.1</td>
<td>-</td>
</tr>
<tr>
<td>2 (1000 Oe)</td>
<td>34 ± 3</td>
<td>(2.1 ± 0.5) \times 10(^{-4})</td>
<td>9*</td>
<td>(2.5 ± 0.8) \times 10(^{-10})</td>
<td>-</td>
<td>43 ± 5</td>
</tr>
</tbody>
</table>

While Dy\(^{3+}\) is a Kramers’ oblate ion for which the electronic density could be stabilized by an axial crystal-field, the Er\(^{3+}\) ion Kramers ion exhibits a prolate density which could hardly be stabilized the IPy\(^-\) ligands. On the other hand, Tb\(^{3+}\) is an oblate ion, however its non-Kramers nature (considering no exchange interaction) requires a particular geometry in order to observe a slow relaxation of the magnetization.\(^{44}\) Since the presence of strong exchange interaction would modify the ground state, this suggests that the relaxation is mainly dominated by the single-ion anisotropy of the dysprosium rather than a coupled state with the radicals. This is in line with the observation of strong antiferromagnetic coupling between the radicals in 4 and 5. In contrast to the strongly coupled Nd\(^{3+}\)-radical bridged Tb complex, such lanthanide ion’s dependence has been also observed in weakly exchange systems based on \(r\) bipyrimidyl or pyridyl-pyrazine ligands.\(^{27-28}\) However, the presented systems exhibit a crystallographically imposed trigonal symmetry which should provide mixed \(m\) states. Although, zero-field slow relaxation in six-coordinate lanthanide complexes is rather scarce,\(^{45-47}\) complex 2 is the first example of a zero-field undiluted SMM with such imposed symmetry. This suggests that despite being weak, the exchange interactions may affect the relaxation.

In summary, we have shown in this communication that the use of radical anion of IPy as ligand enables the formation of homoletic magnetic complexes with a series of lanthanide ions. The study of the magnetic properties indicates that the dysprosium analogue shows a zero-field SMM behaviour for which out-of-phase signals could be still observed up to 30 K. The great tunability of such ligands in terms of both, steric and electronic factors should offer a careful tweaking of the
coordination environment in order to improve the slow relaxation features.

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Notes and references