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Mixed Tb/Dy coordination ladders based on tetra(carboxymethyl)thiacalix[4]arene: a new avenue towards luminescent molecular nanomagnets†

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The macrocyclic ligand calix[4]arene (L1) and its sulphur-containing analogue thia[4]calixarene (L2) are promising precursors for functional molecular materials as they offer rational functionalization with various organic groups. Here, we present the first example of lanthanide-based coordination polymers built from the macrocyclic thiacalix[4]arene backbone bearing four carboxylic moieties, namely, ligand H_4L3 . The combination of H_4L3 with the Tb^{3+} and Dy^{3+} cations led to the formation of 1D ladder-type coordination polymers with the formula $[Ln^{III}HL3DMF_3]\cdot (DMF)$ (where DMF = dimethylformamide and Ln = Tb or Dy, denoted as HL3-Tb and HL3-Dy), which resulted from the coordination of the lanthanide cations with the partially deprotonated ligand HL33- that behaved as a T-shape connector. The coordination sphere around the metal was completed by the coordinated DMF solvent molecules. By combining both Tb^{3+} and Dy^{3+} cations, isostructural heterobimetallic solid solutions $HL3-Tb_{1-x}Dy_x$ were also prepared. HL3-Tb and HL3-Dy showed visible light photoluminescence originating from the f-f electronic transitions of pale green emissive Tb^{3+} and pale yellow emissive Dy^{3+} with efficient sensitization by the functionalized thia[4]calixarene ligand HL3. In the $HL3-Tb_{1-x}Dy_x$ solid solutions, the Tb/Dy ratio governed both the emission colour as well as the emission quantum yield, which reached even 28% at room temperature for HL3-Tb. Moreover, HL3-Dy exhibited a slow magnetic relaxation effect related to the magnetic anisotropy of the dodecahedral Dy3+ complexes, which were well isolated in the crystal lattice by expanded organic spacers.

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Introduction

Coordination polymers (CPs), ^{1,2} MOFs³ and molecular networks^{4,5} have molecular architectures that offer properties such as gas storage, ^{6,7} catalysis, ^{8,9} magnetism^{10,11} and luminescence. ¹²⁻¹⁴ The

generation of such molecular species is based on the formation of coordination bonds between organic coordinating ligands and metal centres or complexes.¹⁵ A large number of coordination polymers that display different connectivity patterns have been generated using different types of polytopic ligands and metals.

Concerning the luminescence properties⁷ of CPs, special attention has been devoted to lanthanide-based CPs,¹⁶⁻¹⁸ owing to their outstanding photoluminescence originating from the triplet excited states of f-f transitions. Lanthanide-based CPs may find applications in luminescence sensing, light emission together with photonics,¹⁹⁻²¹ and as ligands presenting a strong antenna effect,²² allowing energy transfer from an organic ligand to the metal. The use of macrocyclic ligands for the formation of such assemblies has been documented;²³ however, most of the reports are mainly related to porphyrin-based lanthanide CPs.²⁴

Concerning the magnetic properties of lanthanide-based CPs, examples of molecular nanomagnets have been reported earlier.^{25–27} In addition, LnCPs may combine both SMM behaviour and tunable photoluminescence, as shown recently.^{28–30}

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For building new CPs based on macrocycles, the use of calixarene derivatives presents a nice alternative. Calix[4] arenes³¹-³³ are macrocyclic compounds composed of four phenolic moieties bridged by methylene groups as in the "classic" calix[4] arene, CA, L1 (Fig. 1), sulphur atoms in the case of thiacalix[4] arene, TCA, L2 (Fig. 1), ³⁴ or hydroxyl or mercapto groups as in tetramercaptothiacalix[4] arene (TMTCA). ³⁵, ³⁶ These molecules can be easily modified by the functionalization of the upper and/or lower rims of the macrocyclic scaffold. They can adopt four different conformations (cone, partial cone, 1,2-alternate and 1,3-alternate), which make these compounds attractive precursors for designing molecular networks. The generation of coordination polymers with various dimensionalities using tetrasubstituted (tetramercapto)(thia)calix[4] arene derivatives that bear different coordinating sites is well documented. ³⁵

By combining functionalized calix[6]arene bearing amide groups, calix[8]arene, 38,39 unsubstituted calix[4]arene and thia-calix[4]arene derivatives in cone conformation $^{40-43}$ with lanthanides salts or mixtures of 3d/4f, $^{44-51}$ different types of clusters have been obtained. Among them, the use of calixarenes bearing carboxylic coordinating sites, for which the lanthanides cations present great affinity, has been reported for the formation of isolated complexes. 52,53

It is important to note that, to date, only a few lanthanide-based coordination networks involving calixarene derivatives, which are mainly formed by the tetrasulfonate derivatives of (thia)calix[4]arenes in cone conformation,^{54–57} have been reported. To the best of our knowledge, no examples involving carboxylate derivatives or (thia)calix[4]arenes in 1,3-alternate conformation have been reported so far.

Several calixarene derivatives bearing carboxylic/ate coordinating groups have been described. Among them, the compound H_4L3 has been used in the formation of coordination polymers with 3d metals, such as $Co(\pi)$ or $Mn(\pi)$, and ancillary ligands. ^{58,59} An analogous ligand without the *tert*-butyl groups has been used for the formation of coordination polymers with Ag^+ , Ni^{2+} , Co^{2+} , and Zn^{2+} together with K^+ . ^{60,61} TMTCA derivative-based coordination polymers using 3d metals have also been reported ⁶² and shown to form hydrogen-bonded networks. ⁶³

In this study, we report the structure of a series of isostructural new 1D lanthanide-based coordination networks,

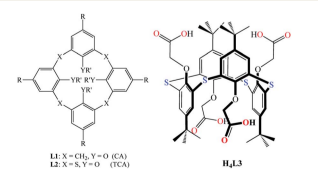


Fig. 1 Calix[4]arene (L1, CA) and thiacalix[4]arene (L2, CA) and tetramethylenecarboxylate derivative of thiacalix[4]arene (H_4L3) adopting a 1,3-alternate conformation.

derived from tetrasubstituted TCA in 1,3-alternate conformation bearing carboxylate moieties (HL3³⁻). A series of isostructural, homo and heterometallic lanthanide-based periodic architectures (Dy/Tb, because of the isostructurality of the related CPs) has been prepared. Mixing lanthanide cations in solid solution coordination networks can lead to the combination of two phosphorescent centres with large emission differences, which is a common strategy to gain self-referenced luminescence. Heteronuclear lanthanide Gd/Eu or La/Tb, Tb/ Eu or Dy/Ln polymers are usually reported for the fine-tuning (colour and brightness) of the corresponding luminescence properties.64-66 Generally, the mixed Eu/Tb compounds allow energy transfer in the mixed coordination polymers due to the optimal energy match between the lanthanide cations. 67-72 But tuneable emission can also arise when mixed Tb/Dy coordination compounds are used.73

The ligand fluorescence and Dy³⁺ and Tb³⁺ phosphorescence were thoroughly studied in the solid-state in the corresponding (Dy/Tb) solid solutions. In addition, the magnetic properties of the anisotropic Dy³⁺ compound were investigated in detail.

Results and discussion

Synthesis and structural details

The synthesis of compound $\mathbf{H_4L3}$ was achieved following a previously reported procedure that involves a nucleophilic substitution reaction between TCA and ethyl bromoacetate in the presence of $\mathrm{Cs_2CO_3}$ in acetone, followed by hydrolysis under basic conditions using LiOH in a THF/ $\mathrm{H_2O}$ mixture. The combination of compound $\mathbf{H_4L3}$ with lanthanide(III) nitrate salts (Ln = Tb or Dy) under solvothermal conditions in DMF (see Experimental part) led to the formation of isostructural 1D coordination networks. The structural investigation of both single crystals of $\mathbf{HL3-Ln}$ (Ln = Dy and Tb) was carried out using X-ray diffraction. The formula derived from the crystallographic analysis was found to be $[\mathrm{C_{48}H_{53}O_{12}S_4Ln}(\mathrm{C_3H_7NO})_3]$ · $(\mathrm{C_3H_7NO})$ ([Ln III $\mathbf{HL3DMF_3}$]·(DMF), Ln = Dy or Tb), as shown in Table 1. In the present CP, the ligand/metal ratio is 1/1.

The crystals were composed of **HL3**³⁻ in its tris-deprotonated form (see Table 2 for C–O distances), the trivalent Ln cation (Ln(III)) and four DMF solvent molecules. The description of the structure in the solid-state is reported below only for **HL3**–**Tb**. The terbium atom was octacoordinated and surrounded by oxygen atoms. Two carboxylate moieties coordinated with the lanthanide in a bidentate mode and the third carboxylate presented a monodentate coordination mode. The 3 oxygen atoms belonging to three DMF molecules completed the coordination sphere of Tb (Fig. 2c). Using the SHAPE program, ⁷⁶ the geometry around the Tb centres was found to be a deformed triangular dodecahedron, as shown in ESI.†

The crystal also contained a non-coordinated DMF molecule occupying the interstices between antiparallel 1D coordination networks without forming any specific interactions with them. The terbium atoms acted as T-shaped connectors, connecting three carboxylate groups belonging to three adjacent **HL3**³⁻ ligands, leading to a double chain ladder-like structure (Fig. 2a and b and schematically represented in d).

Table 1 Crystallographic parameters for HL3-Dy and HL3-Tb recorded at 173 K

Formula	$C_{48}H_{53}O_{12}S_4Dy(C_3H_7NO)_3,$ $C_3H_7NO,$ $C_{60}H_{81}DyN_4O_{16}S_4,$ HL3-Dy	${ m C_{48}H_{53}O_{12}S_4Tb(C_3H_7NO)_3}, \ { m C_3H_7NO}, \ { m C_{60}H_{81}TbN_4O_{16}S_4}, \ { m HL3-Tb}$
Molecular weight	1405.02	1401.44
Crystal system	Triclinic	Triclinic
Space group	$Par{1}$	$Par{1}$
$a(\mathring{A})$	15.0091(17)	15.0160(17)
$b(\mathring{A})$	15.9798(18)	15.9630(14)
$c(\mathring{A})$	16.7401(18)	16.7660(16)
α (deg)	62.217(5)	62.226(3)
β (deg)	67.626(5)	67.513(3)
γ (deg)	79.181(6)	79.132(4)
$V(\mathring{A}^3)$	3284.5(7)	3285.2(6)
Z	2	2
Colour	Colourless	Colourless
Crystal dim (mm³)	$0.090 \times 0.100 \times 0.100$	$0.090 \times 0.100 \times 0.100$
$D_{\rm calc}$ (g cm ⁻³)	1.421	1.417
F(000)	1454	1452
$\mu (\text{mm}^{-1})$	1.332	1.270
Wavelength (Å)	0.71073	0.71073
Number of data meas	83 469	36 284
Number of data with $I > 2\sigma(I)$	17 032 $[R(int) = 0.0752]$	17 822 [$R(int) = 0.0558$]
R	$R_1 = 0.0557, \text{w}R_2 = 0.1391$	$R_1 = 0.0494, \text{ w} R_2 = 0.1073$
$R_{ m w}$	$R_1 = 0.0756, \text{ w} R_2 = 0.1559$	$R_1 = 0.0708, \text{ w} R_2 = 0.1186$
GOF	1.057	1.011
Largest peak in final difference ($eÅ^{-3}$)	1.740 and −1.406	1.411 and −1.140

Consequently, the fourth carboxylic group remained protonated without interactions with the other components of the crystal.

As expected, different C–O distances were observed for the carboxylic moieties of HL3³⁻, revealing the presence of one non-coordinated carboxylic group and three coordinated carboxylate moieties (see Table 2).

Within the 1-D arrays, the shortest Tb–Tb distance was equal to 11.879(3) Å, and the packing of the 1D networks in the yOz plane (Fig. 2b, packing along the a-axis) led to a shorter Tb–Tb distance of 7.836(4) Å between two Tb atoms belonging to two different chains.

Since both compounds **HL3-Dy** and **HL3-Tb** were isostructural (which is not the case for Gd and Eu analogues, for example), solid solutions of formula $HL3-Tb_{1-x}Dy_x$ were

prepared using the same synthetic procedure as the one used for HL3-Dy and HL3-Tb (see Experimental section). Meanwhile, in our case, the related Eu compound obtained with H_4L3 was not isostructural with HL3-Tb or HL3-Dy.

Starting with different experimental Dy/Tb ratios, microcrystalline compounds of the following formula were obtained: $HL3-Tb_{0.07}Dy_{0.93}$, $HL3-Tb_{0.20}Dy_{0.80}$, $HL3-Tb_{0.42}Dy_{0.58}$, $HL3-Tb_{0.67}Dy_{0.33}$ and $HL3-Tb_{0.95}Dy_{0.05}$.

The structure of the solid solution was investigated by XRPD, as shown in Fig. 3 (for the determined cell parameters of the corresponding single crystals, see Table S1, ESI†). XRPD clearly demonstrated the structural homogeneity of the phase of the formed coordination compounds. All the $\text{HL}3\text{-Tb}_{1-x}\text{Dy}_x$ solid

Table 2 Selected bond distances (Å) for HL3-Dy and HL3-Tb

	HL3-Dy	HL3-Tb
C-O (carboxyl-ate/-ic)	1.232(5) and 1.273(5)	1.215(4) and 1.269(4)
	1.239(5) and 1.268(4)	1.253(4) and 1.258(4)
	1.254(5) and 1.261(5)	1.243(4) and 1.259(4)
	1.189(5) and 1.323(5) (carboxylic)	1.194(5) and 1.320(5) (carboxylic)
Ln-O (DMF)	2.354(3)	2.355(3)
	2.366(3)	2.369(2)
	2.380(3)	2.395(3)
Ln-O (carboxylate)	2.256(3)	2.277(2)
	2.380(3)	2.393(3)
	2.393(3)	2.411(3)
	2.404(3)	2.418(3)
	2.438(3)	2.440(2)

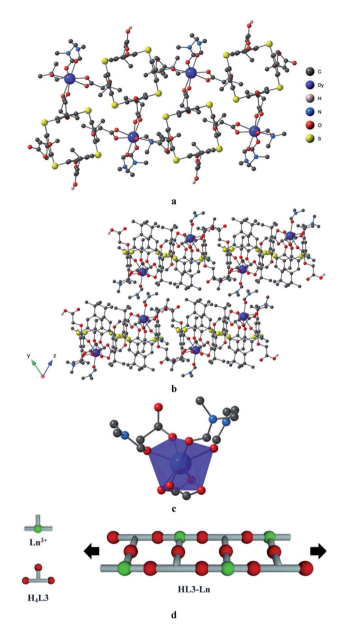


Fig. 2 (a) A portion of the crystal structure of HL3-Tb, (b) crystal packing of HL3-Tb, (c) the coordination environment of Ln^{3+} and (d) the schematic representation of connectivity in HL3-Tb.

solutions were isostructural and displayed the same structure as those of **HL3-Tb** or **HL3-Dy**. Since there was only one type of crystallographically independent Ln ion in the unit cell, owing to similar size of both lanthanide cations, one may assume a statistical distribution of the lanthanide cations in the crystal.

The composition of the $\text{HL3-Tb}_{1-x}\text{Dy}_x$ solid solutions was also checked using energy-dispersive X-ray spectroscopy (EDS), as shown in ESI.†

The experimentally determined Tb/Dy ratios measured were in good agreement with the relative quantities of the lanthanide salts used for synthesis. The slight differences were probably due to weighing inaccuracy and the overestimation of Tb³⁺ content in the EDS measurements (Table 3).

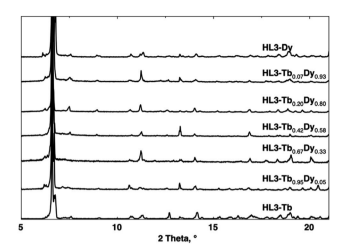


Fig. 3 Comparison of the simulated PXRD patterns for HL3–Tb (bottom) and HL3–Dy (top) and the recorded PXRD patterns for HL3– $Tb_{1-x}Dy_x$ solid solutions. Discrepancies in intensity between the observed and simulated patterns are due to the preferential orientations of the microcrystalline powders.

The luminescence and magnetic properties of this **HL3-Dy** series of compounds are presented below.

Photoluminescence properties

For all the investigated compounds, the measurements were carried out in the solid-state using polycrystalline powders.

Among the bands of the excitation spectra of **HL3-Tb** and **HL3-Dy** at RT in the 350–500 nm region with emission at $\lambda_{em}=534$ nm and 576 nm, respectively (see Fig. S1, ESI†), those attributed to the typical f-f transitions of the Tb³+ and Dy³+ cations presented lower intensities compared with the band attributed to the $\pi\to\pi^*$ transitions of **H**₄**L3** (300–350 nm). These observations clearly indicated the occurrence of an energy transfer from the excited level of ligand **HL3**³- to the emitting state of the Tb³+ (or Dy³+) cations, as seen in the case of antenna effect. The capability of calixarene carboxylate derivatives to act as antennae for Tb³+ complexes has already been documented.

The room-temperature emission spectra of the **HL3-Tb** and **HL3-Dy** coordination networks, when excited at $\lambda_{\rm ex}=350$ nm, are depicted in Fig. 4. For **HL3-Tb**, the emission spectrum is composed of the characteristic five electronic transitions, *i.e.*, ${}^5{\rm D}_4 \rightarrow {}^7{\rm F}_6$, ${}^5{\rm D}_4 \rightarrow {}^7{\rm F}_5$ (strongest), ${}^5{\rm D}_4 \rightarrow {}^7{\rm F}_4$, ${}^5{\rm D}_4 \rightarrow {}^7{\rm F}_3$ and ${}^5{\rm D}_4$

Table 3 Comparison of the observed and experimental Tb/Dy ratios in $HL3-Tb_{1-x}Dy_x$ coordination polymers

	Tb/Dy experimental ratio	Tb/Dy observed ratio (EDS)
HL3-Tb _{0.95} Dy _{0.05}	95:05	95:05
HL3-Tb _{0.67} Dy _{0.33}	60:40	67:33
HL3-Tb _{0.42} Dy _{0.58}	40:60	42:58
HL3-Tb _{0.20} Dy _{0.80}	20:80	20:80
HL3-Tb _{0.07} Dy _{0.93}	10:90	07:93

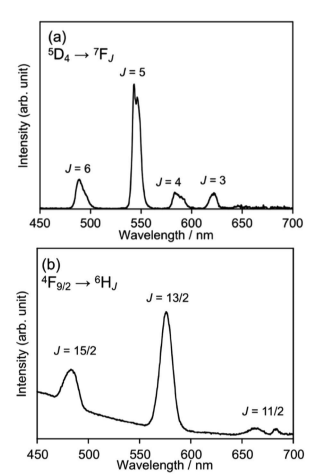


Fig. 4 Emission spectra (λ_{ex} = 350 nm) of HL3-Tb (a) and HL3-Dy (b) at RT.

 \rightarrow 7F_2 (weakest), whereas for HL3–Dy, the $^4F_{9/2}$ \rightarrow $^6H_{15/2},\,^4F_{9/2}$ \rightarrow $^6H_{13/2}$ (strongest) and $^4F_{9/2}$ \rightarrow $^6H_{11/2}$ (weakest) electronic transitions are observed, as shown in Fig. 4a and b, respectively. These were typical f–f transitions with emission in the green region for Tb³+ and yellow-white region for Dy³+. The corresponding emission spectra at 77 K are presented in Fig. S2a and b, ESI.†

The photophysical behaviour of the mixed $\text{HL3-Tb}_{1-x}\text{Dy}_x$ solid solutions was analysed at RT and 77 K. The luminescence spectra of the $\text{HL3-Tb}_{1-x}\text{Dy}_x$ solid solutions observed at RT and 77 K are shown in Fig. 5 and S3.†

At RT, the $\mathbf{HL3-Tb_{1-x}Dy_x}$ solid solutions exhibited the typical luminescence bands of the f-f transitions of $\mathbf{Tb^{3+}}$ at 545 nm and those of $\mathbf{Dy^{3+}}$ at 574 nm. The band intensity at 545 nm decreased when x was decreased, and that at 574 nm increased when x was decreased, as shown in Fig. 5a and b. The introduction of $\mathbf{Dy^{3+}}$ in the $\mathbf{HL3-Tb_{1-x}Dy_x}$ solid solutions allowed slight tuning of the emission colour ranging from pale green to pale yellow, as shown in Fig. 5a and b.

The absolute quantum yields of Tb^{3+} in the series of $HL3-Tb_{1-x}Dy_x$ solid solutions were observed (see Fig. S3a and b, ESI†), and the values at 77 K were found to be 58.9% (27.9% at RT) for HL3-Tb, 32.6% for $HL3-Tb_{0.95}Dy_{0.05}$, and 15.1% for

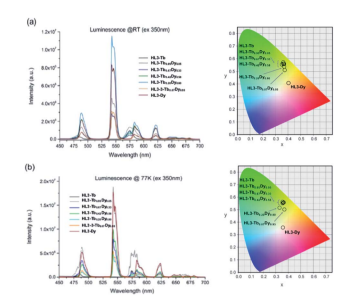


Fig. 5 Emission spectra ($\lambda_{ex}=350$ nm) of HL3–Dy (black) and HL3–Tb (brown), and the solid solutions HL3–Tb $_{0.07}$ Dy $_{0.93}$ (grey), HL3–Tb $_{0.20}$ Dy $_{0.80}$ (blue), HL3–Tb $_{0.42}$ Dy $_{0.58}$ (green), HL3–Tb $_{0.67}$ Dy $_{0.33}$ (light blue) and HL3–Tb $_{0.95}$ Dy $_{0.05}$ (orange) at (a) room temperature and (b) 77 K and the corresponding emission colours presented on the CIE 1931 chromaticity diagrams.

HL3-Tb_{0.20}**Dy**_{0.80}. Therefore, the existence of Dy³⁺ ion in the solid solution played a role in decreasing the luminescence quantum yields of the **HL3-Tb**_{1-x}**Dy**_x solid solutions.

The luminescence decay profile and fitted results are summarized in Tables S2 and S3 and Fig. S3-S5 in ESI.† As described above, the luminescence quantum yields of Dy3+ in the neat and solid solutions with Tb³⁺ were quite low, and it was not possible to obtain the decay profiles at both RT and 77 K for HL3-Dy. The lifetimes of HL3-Tb at RT and 77 K were estimated as a single component, and the τ values were 0.866 and 0.916 ms, respectively. After the addition of Dy³⁺ in the HL3-Tb_{1-x}Dy_x solid solutions, the τ values decreased, and the number of luminescence components of Tb³⁺ increased to two or three (see Tables S2 and S3 in ESI†). The luminescence decay curve of HL3-Tb_{0.2}Dy_{0.8} could be divided into two components, namely 0.378 and 0.932 ms ($\lambda = 543$ nm), at RT. The latter value corresponded to the one observed for HL3-Tb. The second and third emission lifetimes were related to the increasing number of Dy³⁺ centers surrounding the Tb³⁺ centers.

The $\tau_{\rm obs}$ values (Tables S2 and S3 in ESI†) were in accordance with those reported for other calixarene-based coordination compounds in the solid-state. ^{53,78}

Magnetic measurements

As **HL3–Dy** contains the magnetically anisotropic Dy³⁺ cations, which may lead to a potential single-molecule magnet character, its magnetic properties were investigated. The magnetic behaviour of **HL3–Tb** is not presented here because of the lack of significant anisotropy for Tb. As shown in Fig. 6, at 300 K, the $\chi_{\rm M}T$ product reached 14.1 cm³ mol⁻¹ K, which is the exact value

expected for free-ion approximation. A typical decrease in the $\chi_{\rm M}T$ product was observed upon cooling, which could be related to thermal depopulation within the $m_{\rm J}$ levels of the ground $^6{\rm H}_{15/2}$ multiplet of Dy $^{3+}$. Besides a slow decrease in the signal, the $\chi_{\rm M}T$ versus T plot did not display abrupt changes, which suggested the lack of magnetic interactions down to 1.8 K as a result of the efficient magnetic isolation of lanthanide centres in the crystal lattice. Magnetization measured at $T=1.8~{\rm K}$ increased quickly with increasing magnetic field to 10 kOe and much slower at higher fields, reaching 5.4 $\mu_{\rm B}$ at 60 kOe without saturation, as expected for anisotropic Dy $^{3+}$ (inset in Fig. 6).

To investigate the slow magnetic relaxation effects in HL3-Dy, the alternate-current (ac) magnetic characteristics were gathered at $H_{dc} = 0$ (Fig. 7a). The maxima of the frequency dependence of out-of-phase magnetic susceptibility, χ_{M} ", appeared on the edge of the measurement range, even at 1.8 K, quickly moving towards higher frequencies while the temperature increased. This indicated the presence of slow magnetic relaxation, a characteristic effect of the single-molecule magnet (SMM) behaviour. To slow down the relaxation by reducing the presumably disturbing quantum tunnelling of the magnetization (QTM) process, an external dc magnetic field was applied (see Fig. S7, ESI†). The field-variable ac magnetic characteristics were analysed using the generalized Debye model for a single relaxation.80 The extracted relaxation times were plotted against the magnetic field, and the resulting dependence was analysed by taking into account the QTM effect (Quantum Tunnelling of the Magnetisation), which was dominant at low fields, and a field-induced direct process responsible for the acceleration of the relaxation process at higher fields (see ESI, Fig. S7†).81 An equilibrium between these processes appeared at H = 1000 Oe, which was selected as the optimal dc field to investigate magnetic relaxation in HL3-Dy. The dc magnetic characteristics were gathered in the 1.8-5 K temperature range (Fig. 7b and see Fig. S8, ESI†). They were analysed using the generalized Debye model for a single relaxation, and the resulting relaxation times are presented as the function of temperature (Fig. 7c).

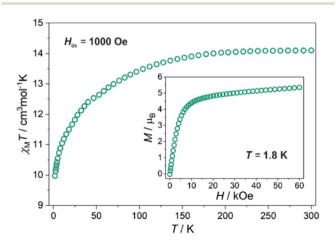


Fig. 6 The temperature dependence of the $\chi_{\rm M}T$ product of HL3–Dy at $H_{\rm dc}=1000$ Oe, and the field dependence of molar magnetization at T=1.8 K (inset).

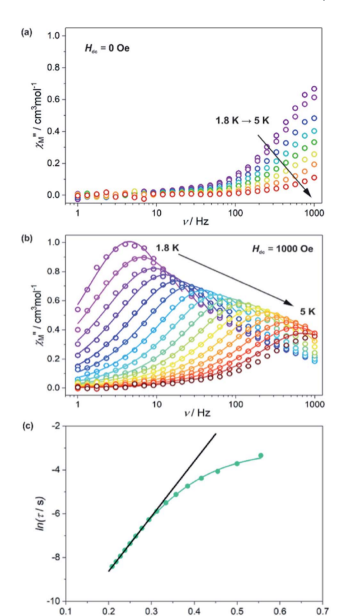


Fig. 7 For HL3-Dy: (a) Zero dc field $\chi_{\text{M}''}(\nu)$ curves in the range of 1.8-5 K, (b) dc field $\chi_{\text{M}''}(\nu)$ curves in the range 1.8-5 K under a dc field of 1 kOe, and (c) temperature dependence of the relaxation time, τ , at 1 kOe, fitted using the Arrhenius law in the limited 3.4-5 K range (extrapolated black line) and within the full 1.8-5 K range using the contributions from the Orbach, Raman, direct and QTM relaxation processes (green line; see text for details).

T-1 / K-1

In the higher temperature range of 3.4–4.8 K, the temperature dependence of the relaxation times was close to linear, and thus, it could be analysed by a simple Arrhenius law, which is presented in eqn (1):

$$\ln(\tau) = \frac{U_{\text{eff}}}{k_{\text{B}}T} + \ln(\tau_0) \tag{1}$$

where $U_{\rm eff}$ represents the effective thermal energy barrier, while τ_0 is the relaxation attempt time for spin reversal at infinite temperature. The obtained parameters $U_{\rm eff}/k_{\rm B}=25.08(9)$ K, $\tau_0=$

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 $1.2(2) \times 10^{-6}$ s were within the characteristic limits of single-molecule magnets with moderate magnetic anisotropy. More precisely, the full temperature dependence curve of the relaxation times was fitted using contributions from the Orbach and Raman relaxation processes together with direct and QTM processes taken from the previous analysis of field-dependent relaxation. Therefore, eqn (2) was used:

$$\tau^{-1} = \tau_1^{-1} \exp(-U_{\text{eff}}/k_B T) + B_{\text{Raman}} T^n + \frac{a(1+c^2 H^2)}{(1+bH^2)} + ATH^4$$

where the first term represents the Orbach process related to the Arrhenius law, the second introduces a two-phonon Raman relaxation pathway, the third denotes the QTM effect and the last one shows a direct process.82,83 To avoid overparameterization, the two last terms describing the QTM effect and direct process taken from the field-dependence analysis and the related parameters were not optimized during the fitting procedure (Fig. S8 \dagger). The obtained values U_{eff} / $k_{\rm B}=31(3)~{\rm K}$ and $\tau_0=1.2(7)\times 10^{-6}~{\rm s}$ were in good agreement with those extracted from the Arrhenius law. The Raman process had to be included with $B_{\text{Raman}} = 0.27(1) \text{ s}^{-1} \text{ K}^{-n}$ and n= 6.0(3). Thus, the last value expected for lanthanide-based SMMs would be located in the 2-9 range.³⁹ As several examples of SMMs lacking the Orbach process have been reported,84 we tried to fit the temperature dependence of the relaxation times excluding Orbach process; however, it was not possible to obtain a satisfying fit.

Based on these results, we could clearly postulate that the Dy³⁺ complexes in **HL3-Dy** exhibit single-molecule magnet characteristics with a moderated thermal energy barrier originating from the Orbach relaxation process and a strong QTM effect, partially reduced by using the external dc magnetic field.

Conclusions

In this work, we have demonstrated the possibility of growing a series of lanthanide-based isostructural homometallic coordination polymers using thiacalix[4]arene bearing carboxylic moieties. Owing to their isostructural nature, a series of unprecedented heterometallic solid solutions based on the macrocyclic thiacalix[4]arene derivatives were generated. The single crystals of the two isostructural Tb3+- and Dy3+-based coordination polymers (HL3-Tb and HL3-Dy, formula [Ln^{III}HL3DMF₃]·(DMF)) were structurally characterized using Xray diffraction, and their photophysical properties were investigated. The formation of the 1D ladder-type coordination compounds resulted in the specific binding of the Ln³⁺ cation to the partially deprotonated organic ligands that behaved as a "T-Shape" connector. In the solid solutions that combined Tb³⁺ and Dy³⁺ forming HL3-Tb_{1-x}Dy_x, the Ln³⁺ cations are randomly distributed within the crystals.

The photoluminescence properties of the coordination polymers were investigated. Ligand $HL3^{3-}$ sensitized the luminescence of the Tb^{3+} or Dy^{3+} cations in HL3-Tb and HL3-Dy at RT, and they exhibited the characteristic f-f transition due to the presence of the Ln^{3+} cations. The introduction of Dy^{3+} in

HL3-Tb allowed the tuning of the emission colour of the solid solutions, which ranged between pale yellow and pale green.

In addition, the magnetic measurements of **HL3-Dy** revealed significant magnetic anisotropy in the Dy³⁺ complexes attached to the thiacalix[4]arene moieties, leading to SMM behaviour. Therefore, this series of new coordination polymers can be considered as a novel source of bifunctional magneto-luminescent molecule-based materials.⁸⁵ We present a novel and elegant pathway for the construction of luminescent molecular nanomagnets by the incorporation of lanthanide ions in coordination chains based on functionalized thiacalix[4] arene ligands.

Experimental methods

X-ray crystallography

Data were collected at 173(2) K on a Bruker Apex-II-CCD diffractometer equipped with an Oxford Cryosystem liquid N_2 device using graphite-monochromated Mo-K α ($\lambda=0.71073$ Å) radiation. For all the structures, diffraction data were corrected for absorption. The structures were solved using SHELXS-97 and refined by full matrix least-squares on F^2 using SHELXL-97. Hydrogen atoms were introduced at calculated positions and refined using a riding model. The structures can be obtained free-of-charge from the Cambridge Crystallographic Data Centre *via* www.ccdc.cam.ac.uk/datarequest/cif. CCDC: 1898825 (HL3-Tb) and 1898826 (HL3-Dy).

Powder diffraction studies (PXRD)

The diffraction diagrams were collected on a Bruker D8 diffractometer using monochromatic Cu-K α radiation in a scanning range of 4 and 40° using a scan step size of 8° min⁻¹.

Elemental analysis

Elemental analysis was performed by the Analysis Service of the Faculty of Chemistry of the University of Strasbourg.

EDS experiments

Elemental analyses using EDS were performed on a Bruker-QUANTAX System, where the EDS Detector was equipped with a GEMINI FE-SEM ULTRA55 (Carl Zeiss). The samples were mixed with colloidal graphite in isopropanol, dried and pressed into pellets. The EDS analysis was conducted at an accelerating voltage of 15 kV.

Photophysical properties

Luminescence and excitation spectra were recorded on a Fluorolog 3-22 (Horiba Jobin Yvon).

The absolute luminescence quantum yields and luminescence lifetimes were determined using an absolute luminescence quantum yield spectrometer C9920-02 (Hamamatsu Photonics K. K.) and a Quantaurus-Tau C11367-12 (Hamamatsu Photonics K. K.), respectively, with pulsed excitation light sources.

Magnetic properties

The magnetic properties were investigated using a Quantum Design MPMS-3 Evercool magnetometer with the powder sample of **HL3-Dy** dispersed in paraffin oil to avoid the rotation of the crystals under the magnetic field. The magnetic data were corrected for the diamagnetic contributions from the sample, oil and the sample holder.

Synthesis

All reagents were purchased from commercial sources and used without further purification. H_4L3 was prepared using a previously reported procedure with some modifications.^{63,74}

Crystallization conditions

HL3–Tb. A solution containing H_4L3 (5 mg, 0.005 mmol) dissolved in DMF (1 mL) was mixed with a 0.01 M solution of $Tb(NO_3)_3 \cdot 6H_2O$ in DMF (1 mL, 0.01 mmol). Then, 2 drops of 1 M HCl were added to the mixture. In a vial closed with a cap, the solution was heated at 80 °C for 3 days, and after cooling to RT, colourless single crystals were obtained (5.3 mg, 73%). They were filtrated and washed with DMF (2 mL) and diethyl ether (2 mL). Formula: $C_{60}H_{81}TbN_4O_{16}S_4$, anal. calcd: C, 51.42%; H, 5.83%; N, 4.00%; found: C, 51.45%; H, 5.87%; N, 4.06%.

HL3–Tb_{0.95}**Dy**_{0.05}. The same procedure was applied starting from a 0.01 M solution of Tb(NO₃)₃·6H₂O in DMF (0.95 mL, 0.0095 mmol) and a 0.01 M solution of Dy(NO₃)₃·5H₂O (0.05 mL, 0.005 mmol). Yield, 4.6 mg (64%). Formula: $C_{60}H_{81}Tb_{0.05}$ -Dy_{0.95}N₄O₁₆S₄, anal. calcd: C, 51.30%; H, 5.81%; N, 3.99%; found: C, 51.38%; H, 5.85%; N, 4.05%.

HL3–**Tb**_{0.67}**Dy**_{0.33}. The same procedure was applied starting from a 0.01 M solution of Tb(NO₃)₃·6H₂O in DMF (0.75 mL, 0.0075 mmol) and a 0.01 M solution of Dy(NO₃)₃·5H₂O (0.25 mL, 0.0025 mmol). Yield, 4.5 mg (62%). Formula: $C_{60}H_{81}$ -Tb_{0.67}Dy_{0.33}N₄O₁₆S₄, anal. calcd: C, 51.38%; H, 5.82%; N, 4.00%; found: C, 51.42%; H, 5.96%; N, 4.08%.

HL3-Tb_{0.42}Dy_{0.58}. The same procedure was applied starting from a 0.01 M solution of Tb(NO₃)₃·6H₂O in DMF (0.5 mL, 0.005 mmol) and a 0.01 M solution of Dy(NO₃)₃·5H₂O (0.5 mL, 0.005 mmol). Yield, 5.2 mg (65%). Formula: $C_{60}H_{81}Tb_{0.42}Dy_{0.58}N_{4}$ -O₁₆S₄, anal. calcd: C, 51.35%; H, 5.82%; N, 3.99%; found: C, 51.52%; H, 5.85%; N, 3.98%.

HL3-Tb_{0.2}**Dy**_{0.8}. The same procedure was applied starting from a 0.01 M solution of Tb(NO₃)₃·6H₂O in DMF (0.25 mL, 0.0025 mmol) and a 0.01 M solution of Dy(NO₃)₃·5H₂O (0.75 mL, 0.0075 mmol). Yield, 5.8 mg (75%). Formula: $C_{60}H_{81}Tb_{0.2}-Dy_{0.8}N_4O_{16}S_4$, anal. calcd: C, 51.32%; H, 5.81%; N, 3.99%; found: C, 51.39%; H, 5.83%; N, 4.02%.

HL3-Tb_{0.07}Dy_{0.93}. The same procedure was applied starting from a 0.01 M solution of Tb(NO₃)₃·6H₂O in DMF (0.1 mL, 0.001 mmol) and a 0.01 M solution of Dy(NO₃)₃·5H₂O (0.9 mL, 0.009 mmol). Yield, 5.3 mg (73%). Formula: $C_{60}H_{81}Tb_{0.07}Dy_{0.93}N_{4}-O_{16}S_{4}$, anal. calcd: C, 51.30%; H, 5.81%; N, 3.99%; found: C, 51.45%; H, 5.79%; N, 4.08%.

HL3-Dy. The same procedure was applied starting from a 0.01 M solution of $Dy(NO_3)_3 \cdot 5H_2O$ in DMF (1 mL, 0.01 mmol).

Yield, 4.8 mg (68%). Formula: $C_{60}H_{81}O_{16}N_4S_4Dy$, anal. calcd: C, 51.29%; H, 5.81%; N, 3.99%; found: C, 51.35%; H, 5.86%; N, 4.02%.

Conflicts of interest

There are no conflicts to declare.

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