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Magnetic anisotropy of endohedral lanthanide ions: paramagnetic NMR study of MSc₂N@C₈₀-I_h with M running through the whole 4f row†

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Paramagnetic and variable temperature ¹³C and ⁴⁵Sc nuclear magnetic resonance studies are performed for nitride clusterfullerenes $MSc_2N@C_{80}$ with icosahedral $I_n(7)$ carbon cage, where M runs through all lanthanides forming nitride clusters. The influence of the endohedral lanthanide ions on the NMR spectral pattern is carefully followed, and dramatic differences are found in peak positions and line widths. Thus, 13 C lines broaden from 0.01–0.02 ppm in diamagnetic MSc₂N@C₈₀ molecules (M = La, Y, Lu) to several ppm in TbSc₂N@C₈₀ and DySc₂N@C₈₀. Direction of the paramagnetic shift depends on the shape of the 4f electron density in corresponding lanthanide ions. In TmSc₂N@C₈₀ and ErSc₂N@C₈₀ with prolate 4f-density of lanthanide ions, ¹³C signals are shifted down-field, whereas ⁴⁵Sc peaks are shifted up-field versus diamagnetic values. In all other MSc₂N@C₈₀ molecules lanthanide ions have oblateshaped 4f electron density, and the lanthanide-induced shift is negative for ¹³C and positive for ⁴⁵Sc peaks. Analysis of the pseudocontact and contact contributions to chemical shifts revealed that the pseudocontact term dominates both in ¹³C and ⁴⁵Sc NMR spectra, although contact shifts for ¹³C signals are also considerable. Point charge computations of the ligand field splitting are performed to explain experimental results, and showed reasonable agreement with experimental pseudocontact shifts. Nitrogen atom bearing large negative charge and located close to the lanthanide ion results in large magnetic anisotropy of lanthanide ions in nitride clusterfullerenes with quasi-uniaxial ligand field.

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Introduction

The molecules of endohedral metallofullerenes (EMFs) comprise the fullerene cage encapsulating one to three metal ions or a more complex hybrid cluster. 1 Shielding magnetic ions (such as lanthanides) inside the carbon cage protects their spin states from the environment making EMFs potentially useful for spintronics applications. Numerous studies of magnetic properties of EMFs reported before 2012 did not reveal any deviation from paramagnetic behaviour, although the influence of the magnetic anisotropy of endohedral lanthanide ions on the measured magnetic moments was proposed.2 However, in 2012 it was discovered that DySc₂N@C₈₀ exhibits slow magnetization reversal at helium temperatures.3 The presence of hysteresis in the magnetization curves measured below 5 K indicated that this EMF can be classified as a single molecule magnet (SMM). In subsequent studies, the SMM behaviour was also found in Dy₂ScN@C₈₀ (ref. 4) and HoSc₂N@C₈₀,⁵ and more EMF-based SMM are likely to be discovered in the near future.

 $\pm 15/2$) and the first predicted to be 373-Leibniz Institute for Solid State and Materials Research, 01069 Dresden, Germany.

Importantly, both DySc₂N@C₈₀ and Dy₂ScN@C₈₀ exhibit unusually long attempt times, which are orders of magnitude longer than in other lanthanide-based SMMs.

One of the most important factors determining the SMM behaviour of lanthanide molecular magnets is the single ion magnetic anisotropy.6 Reliable elucidation of the ligand-field (LF) splitting of the 4f" total momentum states is crucial (although not sufficient) for the correct description of magnetic properties of lanthanide SMMs. Despite the understanding that the magnetic anisotropy is an important factor determining magnetic properties of EMF, a consistent description of the LF effects in EMF has not been provided yet. The first dedicated studies of the LF in nitride clusterfullerenes have been published only recently and were motivated by the discovery of SMM behaviour in DySc₂N@C₈₀ and Dy₂ScN@C₈₀. According to the ab initio computations at the CASSCF level,7 the LF splitting in DySc₂N@C₈₀ is so large that its magnetization behavior at low temperature is determined solely by the ground state of the $^6\mathrm{H}_{15/2}$ manifold. Namely, the gap between the ground $(m_{\mathrm{I}} =$ $\pm 15/2$) and the first excited ($m_{\rm J}=\pm 13/2$) magnetic states was predicted to be 373-415 cm⁻¹ (ref. 7a) or 485 cm⁻¹ (ref. 7b). Likewise, magnetization curves of DySc₂N@C₈₀ and HoSc₂-N@C80 measured by SQUID below 10 K are also described by only one state with $m_{\rm I}=\pm 15/2$ (Dy) or $m_{\rm I}=\pm 8$ (Ho).^{4,5} The information about excited magnetic states might be revealed

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from magnetizations studies at higher temperatures (e.g., the population of the $m_{\rm J}=\pm 13/2$ state in DySc₂N@C₈₀ near room temperature is expected to be ca. 10%). However, with the increase of the temperature the total magnetization of the sample is decreasing dramatically and can hardly be measured reliably when only limited amount of sample is available (as it is often the case with EMF SMMs). An alternative might be the use of spectroscopic techniques, such as analysis of the fine structure in the lanthanide-based luminescence spectrum. Unfortunately, the fullerene cages of EMFs absorb light in the visible range and hence block the possibility to observe the lanthanide-based luminescence for a majority of 4f elements. So far, metal-based luminescence in EMFs could be detected only for endohedral $\rm Er^{3+}$ ions emitting in the near-IR range.

Solution nuclear magnetic resonance (NMR) studies can provide complimentary information on the magnetic anisotropy of lanthanide ions for the temperatures not well accessible by direct magnetization measurements. The chemical shift of a nuclei in a paramagnetic compound can be described as a sum of diamagnetic and paramagnetic terms, $\delta^{\text{exp}} = \delta^{\text{dia}} + \delta^{\text{para}}$. In due turn, the paramagnetic shift has two major contributions, contact $\delta^{\rm con}$ and pseudocontact $\delta^{\rm pc.10}$ The contact (Fermi) shift results from the interaction between the nuclear spin (of ¹³C or ⁴⁵Sc atoms in this work) and the spin-polarized electron density of the molecule (in particular in the region close to the nuclei of interest). As such, the contact shift is proportional to the hyperfine coupling constant weighed with the expectation value of the spin operator S_z of the lanthanide. The pseudocontact shift is caused by dipolar through-space interactions of the nuclear and electronic magnetic dipoles. For an i-th atom in a lanthanide-containing molecule the pseudocontact shift can be computed as:

$$\delta_{i}^{\text{pc}} = \frac{1}{12\pi R_{i}^{3}} \left(\left(3\cos^{2}\theta_{i} - 1 \right) \left(\chi_{zz}^{\text{Ln}} - \frac{\chi_{xx}^{\text{Ln}} + \chi_{yy}^{\text{Ln}}}{2} \right) + \frac{3}{2} \left(\chi_{xx}^{\text{Ln}} - \chi_{yy}^{\text{Ln}} \right) \sin^{2}\theta_{i} \cos 2\varphi_{i} \right)$$
(1a)

where $\chi_{\alpha\alpha}^{\rm Ln}$ are components of the magnetic susceptibility tensor of the lanthanide, whereas R_i , θ_i , and φ_i are polar coordinates of the i-th atom in the coordinate system centred on the lanthanide ion. In particular, R_i is a distance between the atom of interest and lanthanide ion, and θ_i is an angle between quantization axis z and the vector connecting the lanthanide ion and the i-th atom. If $\chi_{xx}^{\rm Ln} = \chi_{yy}^{\rm Ln}$ (i.e. the ligand field is uniaxial), eqn (1a) is simplified to:

$$\delta_i^{\text{pc}} = \frac{(3\cos^2\theta_i - 1)}{12\pi R_i^3} \left(\chi_{zz}^{\text{Ln}} - \frac{\chi_{xx}^{\text{Ln}} + \chi_{yy}^{\text{Ln}}}{2} \right)$$
 (1b)

As can be seen in eqn (1), the pseudocontact shift contains information on the molecular structure and on the magnetic properties of the paramagnetic center. Paramagnetic NMR is therefore a popular structure elucidation tool for metal complexes, polymers or biomolecules. ^{10b,c,11} At the same time, it can be also used to determine or verify LF parameters in lanthanide complexes. ¹² The prerequisite for both types of

application of NMR spectroscopy in the studies of paramagnetic molecules is the possibility to separate contact and pseudocontact contributions, is since only the latter brings necessary information about molecular structure and magnetic anisotropy.

In the field of endohedral metallofullerenes, paramagnetic NMR was mainly used for the structural studies. Several Ce-EMFs, 14 anion of Pr@C₈₂, 15 three isomers of Tm@C₈₂, 16 Sm@C₈₀,¹⁷ and Sm@C₈₂ (ref. 18) were characterized by ¹³C NMR. The variable-temperature 45Sc NMR study was also reported for CeSc₂N@C₈₀. 19 The dominant contribution of the pseudocontact shift in Ce-EMFs was postulated based on the analysis of the temperature dependence in the framework of Bleaney's theory. The size of the contact term contribution in the paramagnetic NMR spectra of EMFs remains unclear. Our group recently reported paramagnetic NMR study of $HoM_2N@C_{80}$ and $Ho_2MN@C_{80}$ (M = Sc, Y, Lu). 20 13 C and 45 Sc signals of these molecules could be identified in spite of the severe broadening induced by Ho3+ ions. This study showed that paramagnetic NMR spectroscopy when combined with the analysis of the ligand field splitting, can provide information on the magnetic state of lanthanide ions in EMFs.

In this work we report on a systematic paramagnetic NMR study of MSc₂N@C₈₀ compounds with all lanthanides forming nitride clusterfullerenes (M = La, Ce, Pr, Nd, Gd, Tb, Dy, Ho, Er, Tm) and concomitant calculations of the ligand field splitting in these molecules. The main goal is to provide a consistent and uniform description of the magnetic states of lanthanide ions in these molecules based on the ligand field computations verified by experimental NMR data. The manuscript is organized in several parts. First, we describe the results of experimental NMR measurements, including the temperature dependence and relaxation times. Then, we perform a thorough analysis of the NMR data to separate contact and pseudocontact contributions to paramagnetic shifts. This section is followed by the brief analysis of the molecular structures. Then point-charge calculations of the ligand field splitting are described, which give the desired description of the single ion magnetic anisotropy in the series of MSc₂N@C₈₀ molecules across the whole 4f row. Reliability of the computed LF patterns is then verified by comparison of the computed and experimental pseudocontact shifts.

Experimental and computational details

In this work we will consider exclusively $MSc_2N@C_{80}$ compounds with the $I_h(7)$ carbon cage (Fig. 1). Hereafter the compounds will be abbreviated as 1M, where M denotes the corresponding lanthanide (e.g., $LaSc_2N@C_{80}$ is denoted hereafter as 1La). All studied compounds were synthesized using the arc-discharge method in the presence of nitrogen source such as NH_3 (ref. 21) or guanidine thiocyanate, 22 and then separated using chromatography as described in the ESI Fig. S1–S5.† The synthesis of 1Ce, 23 1Nd, 24 1Pr, 23 1Dy, 24 1Ho, 20b and $1Lu^{24}$ in our group was described earlier. 1La, 1Pr, 1Tb, 1Er and 1Tm were

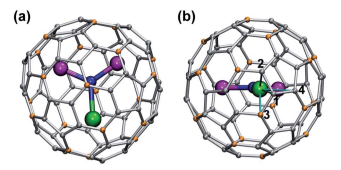


Fig. 1 DFT-optimized molecular structure of representative $MSc_2N@C_{80}$ - I_h (1Pr). Praseodymium ion (shown green) is at the distance of 2.225 Å from the central nitrogen atom (blue), whereas two Sc-N distances are 1.939 and 1.936 Å. Carbon atoms are shown in grey (pentagon-hexagon-hexagon junctions, PHHJs) and orange (triple hexagon junctions, THJs). In (b), the plane of the cluster is normal to the paper. Thin cyan lines denote shortest Pr-C distances: 2.508 Å (C1), 2.553 Å (C2), 2.591 Å (C3), and 2.658 Å (C4).

synthesized for this work using guanidine thiocyanate as a nitrogen source, whereas **1Tb** and **1Y** were synthesized using NH₃ as a reactive gas. Note that the synthesis and characterization of **1Y**,²⁵ **1La**,²⁶ **1Ce**,¹⁹ **1Tb**,²⁷ and **1Er**²⁸ was also reported by other groups using somewhat different conditions. Gd also forms mixed-metal nitride clusterfullerenes with Sc,²⁹ but neither ¹³C nor ⁴⁵Sc NMR signals could be detected for **1Gd**, so this molecule will not be discussed hereafter.

The 125 MHz 13 C NMR and 121.5 MHz 45 Sc NMR measurements were performed on a Bruker Avance 500 spectrometer equipped with a multiprobe head 1152Z. The measurements were performed for compounds dissolved in CS_2 with d^6 -acetone placed in a coaxial tube as a lock; in special cases d^4 -ortho-dichlorobenzene was also used as a solvent. Typically the spectra were measured at each 10 K in the temperature rage form 268 to 308 K.

Longitudinal relaxation rates (R_1) were measured using inversion recovery pulse sequence with different delays between π and $\pi/2$ pulses. The values were determined from the slope of the linear fit of $\ln(I_{\rm max}-I)$ versus τ (delay). Transversal relaxation rates (R_2) were measured using Carr–Pucell–Meiboom–Gill (CPMG) pulse sequence.³⁰ The values were determined from slopes of linear fit of $-\ln(I)$ versus τ (delay). In case of broad line width, R_2 cannot be determined by CPMG-pulse sequence because the relaxation is too fast. Conservative estimation of R_2 values in such cases can be done by measuring the line width at half peak maximum. However, R_2^* determined by this procedure includes the line broadening by inhomogeneity of the magnetic field $(R_2^* = \frac{y\Delta B_0}{2} + R_2)$ and therefore the T_2 time determined from the line width is shorter than the real one.

DFT optimization of molecular coordinates was performed at the B3LYP level using the Firefly³¹ code. The basis sets were def2-SVP $\{5,1,1/3,1/1\}$ for carbon atoms,³² def2-TZVP $\{6,2,1,1/4,1,1/1,1/1\}$ for nitrogen,³² and Stuttgart–Cologne effective core potentials for Sc (ECP10MDF)³³ and lanthanides (4f-in-core ECP-MWB-II)³⁴ with $\{3,1,1,1,1,1/2,2,1,1,1/4,1,1/1,1/1\}$ and $\{3,1,1,1,1,1/3,1,1,1,1/2,1,1,1,1/1,1,1/1,1/1\}$ valence parts, respectively. Spin density and hyperfine

coupling constants in GdSc₂N@C₈₀ were computed at the PBE level using the specially tailored SARC basis set of TZVP quality³⁵ combined with either DKH or ZORA scalar relativistic corrections implemented in the ORCA package.³⁶ To compute the Bader (QTAIM) charges, full electron calculations were performed at the PBE/SARC-TZVP level with DKH scalar relativistic corrections using ORCA. To avoid poor reliability of the full electron DFT treatment of the 4f elements, Y was used to model the charges of the lanthanides from Tb to Tm, whereas La was used to model Ce, Pr, and Nd. QTAIM computations of the atomic charges were then performed with the AIMAll³⁷ code. Point charge calculations of the ligand field splitting, magnetic susceptibility tensors, and 4f electron densities were performed using so1ion (Cfield) routine in McPhase code.³⁸ Molecular structures and isosurfaces were visualized with the help of VMD.³⁹

Results and discussion

NMR spectroscopy

¹³C and ⁴⁵Sc NMR spectra at 288 K. Fig. 2 and 3 show ¹³C and ⁴⁵Sc NMR spectra of **1M** compounds measured at 288 K (the temperature is chosen as the center of the temperature interval used in the variable temperature NMR studies, see below).

Fast rotation of the MSc_2N cluster inside the highly symmetric C_{80} - I_h carbon cage results in the averaging of the carbon signals, leaving only two ^{13}C NMR peaks with a 3:1 intensity ratio. The higher intensity peaks corresponds to the 60 carbon atoms at the pentagon–hexagon–hexagon junctions (PHHJ), whereas the lower intensity peak is due to the 20 atoms at the triple hexagon junctions (THJ, see Fig. 1). The chemical shift of the PHHJ carbons in diamagnetic 1M compounds is bracketed by the 1La and 1Lu values, 144.76 and 143.99 ppm, respectively. Likewise, the THJ signal varies from 138.08 ppm in 1La to 136.90 ppm in 1Lu. Chemical shifts of the THJ and PHHJ

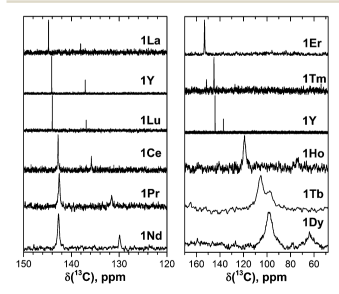


Fig. 2 13 C NMR spectra of **1M** compounds measured in CS₂ at 288 K. Note the different scale in the left and right panels; for a better comparison, the spectrum of **1Y** is shown in both panels.

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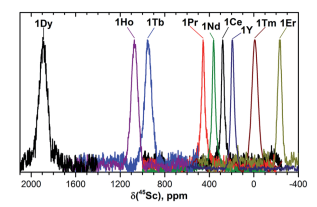


Fig. 3 45 Sc NMR spectra of **1M** compounds measured in CS₂ at 288 K. The $\delta(^{45}$ Sc) value of diamagnetic **1Y** is 191 ppm.

carbon atoms in **1Y** are intermediate between those of **1La** and **1Lu** (see Table 1). Thus, depending on the size of the endohedral cluster, the diamagnetic ¹³C values are subject to change within the range of *ca.* 1 ppm (see also ref. 40) which should be taken into account in the analysis of the paramagnetic shifts in other **1M** compounds. The δ^{dia} values listed in Table 1 and used to compute paramagnetic shifts were obtained by approximating the ¹³C chemical shifts by a quadratic function of the Shannon radii of metal ions (see ESI Fig. S6†).

The measurement of the ¹³C NMR spectra of paramagnetic EMFs is complicated by the considerable line broadening. However, thanks to the simple two-line spectrum of the C₈₀ carbon cage, the signals are detectable even when they are severely broadened as in **1Tb** or **1Dy**. In **1Ce**, **1Pr**, **1Nd**, **1Tb**, **1Dy**, and **1Ho** the ¹³C signals are shifted up-field with respect to the diamagnetic values. The magnitude of the paramagnetic shift varies from few ppm in **1Ce**, **1Pr**, and **1Nd** to tens of ppm in **1Tb**, **1Dy**, and **1Ho** (Fig. 2 and Table 1). On average, lanthanide-induced shifts of the THJ carbons are more pronounced than those of the PHHJ carbons.

In **1Er** and **1Tm** the lanthanide-induce shift is positive (down-field). The PHHJ and THJ ¹³C signals of **1Er** are coinciding near the room temperature and appear at 153.2 ppm, but can be resolved at higher or lower temperatures as discussed below. In **1Tm**, the stronger lanthanide-induced shift of the THJ signal pushes it to a lower field than that of the PHHJ (in all other **1M** compounds the PHHJ signal appears at lower field than that of the THJ carbons). Direction of the lanthanide-induced paramagnetic shift is determined by a magnetic anisotropy of the lanthanide ion (see ref. 41 for recent examples), and hence experimental NMR data gives information at least on the sign of the anisotropy. This question will be discussed in more details below.

In ⁴⁵Sc NMR spectra, all **1M** compounds exhibit one relatively broad peak (Fig. 3). The ⁴⁵Sc chemical shifts of the diamagnetic **1La** and **1Lu** are 198 and 200 ppm, respectively, whereas in paramagnetic **1M** molecules the ⁴⁵Sc signal position spans the range from –233 ppm in **1Er** to 1892 ppm in **1Dy**. The peak width varies from 30–40 ppm for the diamagnetic **1Lu** and **1La** and paramagnetic **1Ce**, **1Pr**, **1Nd** and **1Er**, to *ca.* 70–80 ppm for the **1Tb**, **1Dy**, **1Ho**, and **1Tm**.

Variable temperature NMR studies. Temperature dependence of paramagnetic NMR shifts can be used as an additional parameter to distinguish the contact and pseudocontact contributions as well as to clarify the extent of the magnetic anisotropy. Fig. 4 and 5 show variations of ^{13}C and ^{45}Sc NMR spectra in the temperature range 268–308 K. To quantify the temperature variation, we will use the $\Delta_{268}^{308}(\delta)$ parameter, which is the difference of the chemical shifts measured at 308 and 268 K.

Temperature variation of the peak positions is in line with the single-temperature δ^{para} values. Namely, the compounds with strong paramagnetic shifts exhibit enhanced variation of the peak position with the temperature and *vice versa*. Likewise, the sign of the $\Delta^{308}_{268}(\delta)$ values is opposite to the sign of the δ^{para} shifts. The only exclusion is the PHHJ signal of **1Tm**, for which both δ^{para} and $\Delta^{308}_{268}(\delta)$ parameters are positive. Since this signal

Table 1 Shannon ionic radii of lanthanides and chemical shifts of 1M molecules measured at 288 Ka

		¹³ С-РННЈ				¹³ C-THJ							
1M	$R(M^{3+})$	δ	$\Delta v_{1/2}$	$\delta^{{ m dia}b}$	$\delta^{ m para}$	Δ_{268}^{308}	δ	$\delta^{{ m dia}b}$	$\delta^{ m para}$	Δ_{268}^{308}	45 Sc δ	$\delta^{\mathrm{para}c}$	Δ_{268}^{308}
Y	0.900	144.11	1.5	144.11	_	_	137.11	137.11	_	_	191	_	_
La	1.032	144.76	2.0	144.76	_	_	138.08	138.08	_	_	198	_	_
Ce	1.010	142.69	15.1	144.63	-1.94	0.73	135.73	137.89	-2.16	0.57	279	79	_
Pr	0.990	142.55	41.8	144.51	-1.96	0.72	131.59	137.73	-6.14	0.94	455	255	-43
Nd	0.983	142.69	50.5	144.47	-1.78	0.25	129.91	137.67	-7.76	1.48	361	161	-15
Tb	0.923	105.6	579	144.20	-38.6	8.6	98.0	137.25	-39.3	7.5	949	749	-174
Dy	0.912	97.5	905	144.15	-46.7	10.2	61.6	137.18	-75.6	_	1892	1692	-290
Но	0.901	119.1	284	144.11	-25.0	_	74.2	137.12	-62.9	_	1072	872	-165
Er	0.890	153.23	${\sim}45$	144.08	9.15	-2.12	153.23	137.05	16.18	-3.12	-233	-433	109
Tm	0.880	144.83	17.2	144.04	0.79	0.34	151.97	137.00	14.97	-2.97	-10	-210	32
Lu	0.861	143.99	1.0	143.99	_		136.90	136.90	_		200	_	

^a Ionic radii are in Å, chemical shifts are in ppm, line width $\Delta \nu_{1/2}$ in Hz ¹³C chemical shifts are determined with precision to the second decimal except for **1Tb**, **1Dy**, and **1Ho**, which are determined to the first decimal. ^b Diamagnetic shifts were estimated from the polynomial fit of the δ(**1M**)-versus-R(M³⁺) data for **1Y**, **1La**, and **1Lu**. ^c The values are obtaining by subtracting the ⁴⁵Sc chemical shift of **1La** (**1Ce**, **1Pr**, **1Nd**) or **1Lu** (other compounds).

1Pr 1Tb 1Nd 268 268 288 288 140 136 δ(¹³C), ppm 140 135 130 δ(¹³C), ppm 100 δ(13C), ppm 1Tm 1Er moun 158 156 154 152 150 δ(¹³C), ppm 156

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Fig. 4 Variable-temperature 13 C NMR spectra of selected **1M** compounds measured in CS₂ in the 268–308 K range. Vertical dash lines denote chemical shifts at 288 K and are shown to guide an eye. At 288 and 308 K two carbon signals in **1Er** are not resolved, and therefore the measurements were performed in o-DCB in a broader temperature range (268–328 K). Low solubility of **1Dy** and **1Tb** at 268 K resulted in poor signal-to-noise ratio (the noise features in the spectrum of **1Dy** are marked by asterisks).

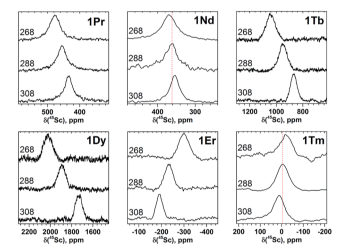


Fig. 5 Variable-temperature 45 Sc NMR spectra of selected **1M** compounds measured in CS₂ in the 268–308 K range. To guide an eye, vertical lines in the spectra or **1Nd** and **1Tm** indicate signal positions at 288 K.

has the smallest δ^{para} value (0.79 ppm), we cannot exclude that uncertainties in the estimation of the δ^{dia} value may play a role here. More likely, however, is that the pseudocontact and contact contributions have opposite signs and compensate each other in this molecule.

Line width and relaxation times. Variation of the line width, which is especially well pronounced in the $^{13}\mathrm{C}$ NMR spectra, points to the substantial change of relaxation times across the lanthanide row. A transverse relaxation time T_2 can be estimated using the relation $T_2=(\pi\Delta\nu_{1/2})^{-1}$, where $\Delta\nu_{1/2}$ is the peak width at its half maximum. The $\Delta\nu_{1/2}$ values determined for

PHHJ signals at 288 K are listed in Table 1. For 1Tm we have also measured T_2 and T_1 times directly using Carr-Purcell-Meiboom-Gill pulse sequence and inversion recovery method, which gave 50 ms and 66 ms at 288 K, respectively (see ESI Fig. S7 and S8† for other temperatures). As might be anticipated, T_1 is longer than T_2 , and the T_2 * value of 19 ms determined by the line width method underestimates the real T_2 time but gives correct order of the value. Direct measurements of shorter relaxation times is beyond the possibilities of our spectrometer. In diamagnetic 1M compounds, the nuclei spin relaxation times are much longer.⁴² For instance, T_2 and T_1 times of the PHHJ carbons in 1Y are 17 and 23 s, respectively. Note that T_2 determined from the CPMG pulse sequence for diamagnetic 1Y is significantly longer than might be estimated from the line width $(T_2^* = 0.2 \text{ s})$, which shows that the line width for such narrow signals is limited by the instrument.

Comparison of the $\Delta \nu_{1/2}$ values for 13 C peaks in **1M** series (Fig. 1 and Table 1) reveals dramatic influence of the lanthanide on the nuclei spin relaxation. For instance, 13 C nuclei spins in **1Dy** ($T_2 \approx 0.3$ ms) relax $ca. 5 \times 10^4$ times faster than in diamagnetic **1Y** ($T_2 = 17$ s). The plot of $\Delta \nu_{1/2}$ versus δ^{para} for the PHHJ carbons along the **1M** series can be well fit by a parabola (see ESI Fig. S9†). This fact points to the prevailing contribution of the pseudocontact shift in the total paramagnetic shift since the relaxation rate in paramagnetic molecules has R^{-6} dependence on the distance between the nuclei and paramagnetic center, whereas the pseudocontact shift scales as R^{-3} (eqn (1)).

The broadening of ⁴⁵Sc signals along the **1M** series is not so pronounced as for carbon signals because the ⁴⁵Sc quadrupole moment broadens the peaks even in diamagnetic molecules. The ⁴⁵Sc signals in **1Dy**, **1Tb**, **1Ho**, and **1Tm** are roughly two times broader than in the **1Y** (Fig. 3), which can be ascribed to a paramagnetic effect. At the same time, in **1Ce**, **1Pr**, **1Nd**, and **1Er** the line width is comparable to that of diamagnetic **1Y**, which means that the spin relaxation rate in these molecules is still determined by a quadrupole effect.

The line width exhibits substantial decrease at lower temperatures (Fig. 4 and 5). For instance, 45 Sc signals measured at 268 K are roughly twice broader than at 308 K (Fig. 5). In some previous works, temperature dependence of the 45 Sc line width was used to estimate rotation barrier of the endohedral clusters. 19,43 If the line width is directly proportional to the rotational correlation time, the plot $\ln(\Delta \nu_{1/2})$ -versus- T^{-1} gives a straight line with the slope proportional to the activation energy. In this work, none of the 1M molecules gave straight line in the $\ln(\Delta \nu_{1/2})$ -versus- T^{-1} coordinates, preventing thus further analysis of the data. The broadening at low temperature should result from the interplay of the slowing of the cluster dynamics and the electron spin contribution, which can hardly be separated at this moment.

Separation of contact and pseudocontact shifts

The localized "buried" nature of the 4f electrons in lanthanides implies that the contact contribution to the paramagnetic shift should be rather small. Typically, the values become negligible when the nucleus of interest is separated by more than four

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then described as:

bonds from the lanthanide ion. However, in the 1M molecules both Sc and carbon atoms are located rather close to the lanthanide, and hence the contact term cannot be ignored. With systematic NMR data along the whole lanthanide series in hand, we can address this problem. Separation of the contact and pseudocontact shifts is a long standing problem in paramagnetic NMR, and several schemes were proposed to solve it. 11c,13,44 Unfortunately, all methods allowing analytical deviation of linearizable equations are based on Bleaney's assumption⁴⁵ that LF splitting is less than the thermal energy and hence expansion of the magnetic susceptibility in reciprocal temperature series is limited to T^{-2} . This assumption is definitely not fulfilled for 1M molecules, and hence the results of these calculations may give only qualitative estimation at best (see also ref. 41a for a detailed discussion of the limitations of Bleaney's theory). Another assumption usually applied is that the system is either purely axial (LF operator described in phenomenological B_q^k parameters is limited to the B_0^2 term), or its rhombic anisotropy can be described solely by a B_2^2

$$\delta_{ij}^{\text{para}} = F_i \langle S_z \rangle_j + C_i (B_0^2 G_i + 6B_2^2 H_i)$$
 (2)

The first term is the contact shift, where F_i is proportional to a Fermi coupling constant, and $\langle S_z \rangle_j$ is the expectation value of the spin projection operator S_z for a given lanthanide. The second term is the pseudocontact shift, C_j is a numerical factor specific for each lanthanide and tabulated by Bleaney *et al.*, ^{45b} whereas G_i and H_i are structural factors (compare to eqn (1a)). If rhombic anisotropy is negligible, the second term reduces to $\delta_{ii}^{pc} = C_i B_0^2 G_i$ (compare to eqn (1b)).

parameter. The sum of contact and pseudocontact shifts can be

Reilley's method. Following the method proposed by Reilley *et al.*, 13 eqn (2) can be written in the form:

$$\frac{\delta_{ij}^{\text{para}}}{\langle S_z \rangle_j} = F_i + \frac{C_j}{\langle S_z \rangle_j} \left(B_0^2 G_i + \sqrt{6} B_2^2 H_i \right) \tag{3}$$

The plot $\delta_{ij}^{\mathrm{para}}/\langle S_z \rangle_j$ versus $C_j/\langle S_z \rangle_j$ along the lanthanide series is expected to give a straight line, if all molecules are isostructural, and the F_i value and LF parameters are not changing along the series. Contact and pseudocontact shifts can be then computed from the intercept and the slope.

Fig. 6 shows $\delta_{ij}^{\text{para}}/\langle S_z\rangle_j$ versus $C_j/\langle S_z\rangle_j$ plots for two NMR signals measured for **1M** molecules. ¹³C PHHJ and ⁴⁵Sc shifts can be roughly linearized if **1Tm** values are excluded from the set; the δ^{con} and δ^{pc} shifts computed from the linear fits are listed in Tables 2 and 3. The ¹³C THJ shifts cannot be fitted by a straight line in these coordinates (see ESI Fig. S10†).

The absolute values of $\delta^{\rm con}$ shifts for the PHHJ carbons are 2–4 times smaller than those of $\delta^{\rm pc}$. Furthermore, the $\delta^{\rm con}$ and $\delta^{\rm pc}$ shifts of **1Ce**, **1Pr**, **1Nd**, **1Er**, and **1Tm**, have opposite signs. For **1Nd** this analysis predicts mutual compensation of $\delta^{\rm con}$ and $\delta^{\rm pc}$ shifts, but uncertainties of 1–2 ppm make the analysis of the small values rather ambiguous.

The ratio between $\delta^{\rm con}$ and $\delta^{\rm pc}$ for the ⁴⁵Sc chemical shifts is smaller than for the ¹³C counterparts, and the contact term

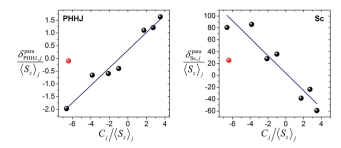


Fig. 6 Correlations between $\delta_{ij}^{\text{para}}/\langle S_z \rangle_j$ and $C_j/\langle S_z \rangle_j$ for $^{13}\text{C-PHHJ}$ (left) and ^{45}Sc (right) signals used to distinguish contact and pseudocontact contributions in **1M** molecules by Reilley's approach. **1Tm** values are shown as red dots; they were not included in the linear fit. The quality of linear fit is $R^2 = 0.97$ for $^{13}\text{C-PHHJ}$ and $R^2 = 0.90$ for ^{45}Sc .

contribution drops to less than 10% for the majority of lanthanides; the largest relative $\delta^{\rm con}$ values of ca. 20% are found in **1Nd** and **1Ho**. Thus, although uncertainty of the fit is rather high, the dominance of the pseudocontact term is beyond any doubt.

Deviations from the straight line in eqn (3) are usually considered to be an indication of the loss of the isostructurality along the series. This can hardly be the case for the **1M** series since molecular structures determined by single-crystal X-ray diffraction for some of them are very similar (subject to vary with lanthanide contraction, see below). ^{19,26–28} More likely, other assumptions used in the model are not valid. In particular, Reilley's approach implies that the LF parameters are not changing along the lanthanide row, which is most probably not correct for the **1M** molecules. ^{7b}

Parameter-free models. The requirement of the constant LF parameters is one of the obvious weak points of Reilley's approach, which can be circumvented by combining the data on two or more nuclei and excluding B_0^2 and B_2^2 terms in eqn (3). For an uniaxial system ($B_2^2 = 0$), the 2-nuclei parameter-free model is obtained by plotting $\delta_{ij}^{\text{para}}/\langle S_z \rangle_j$ versus $\delta_{kj}^{\text{para}}/\langle S_z \rangle_j$, which should give a straight line for the isostructural series (ref. 46):

$$\frac{\delta_{ij}^{\text{para}}}{\langle S_z \rangle_i} = \left(F_i - F_k \frac{G_k}{G_i} \right) + \frac{G_k}{G_i} \frac{\delta_{kj}^{\text{para}}}{\langle S_z \rangle_j}$$
(4a)

The slope of the line is a ratio of the structural factors G_i/G_k , and the intercept is a difference of F_i and F_kG_i/G_k . For the molecules with a rhombic anisotropy ($B_2^2 \neq 0$), the parameter-free model can be obtained in a similar fashion by simultaneous analysis of chemical shifts of three different nuclei. In the latter case linearization gives the equation of a plane (ref. 47):

$$\frac{\delta_{ij}^{\text{para}}}{\langle S_z \rangle_j} = (F_i - F_k D_{ikl} - F_l E_{ikl}) + D_{ikl} \frac{\delta_{kj}^{\text{para}}}{\langle S_z \rangle_j} + E_{ikl} \frac{\delta_{lj}^{\text{para}}}{\langle S_z \rangle_j}$$
(4b)

here coefficients D_{ikl} and E_{ikl} are functions of the structural parameters G_i and H_i from eqn (3), see ref. 47a for full details.

Fig. 7 visualizes analysis of all chemical shifts for **1M** molecules within parameter free models. The 3D plot corresponds to the 3-nuclei method, whereas projections on the coordinate planes allow evaluation of the 2-nuclei method for three different combination of nuclei. In the 3-nuclei model, all data

Table 2 Contact and pseudocontact ¹³C chemical shifts in 1M molecules

			РННЈ						ТНЈ			
			$\delta^{ m con}$		$\delta^{ m pc}$				$\delta^{ m con}$	$\delta^{ m pc}$		
1M	$\langle S_z \rangle_j^{\ a}$	$C_j^{\ a}$	Reilley	DFT^b	Reilley	DFT^c	PCM^d	Δ_{268}^{308e}	DFT^b	DFT^c	PCM^d	Δ_{268}^{308e}
Ce	0.98	-6.48	0.3	0.4	-2.2	-2.3	-1.2	0.2	0.2	-2.4	-3.4	0.5
Pr	2.97	-11.41	1.0	1.2	-4.0	-3.1	-2.7	0.4	0.8	-6.9	-7.7	1.2
Nd	4.49	-4.46	1.4	1.8	-1.5	-3.6	-2.1	0.5	1.1	-8.9	-6.0	1.4
Tb	-31.82	-86.84	-10.3	-12.7	-30.1	-25.9	-17.6	3.1	-8.1	-31.2	-61.2	10.9
Dy	-28.54	-100	-9.2	-11.4	-34.6	-35.2	-24.2	4.3	-7.3	-68.3	-73.0	12.9
Но	-22.63	-39.25	-7.3	-9.0	-13.6	-16.0	-12.5	3.4	-5.8	-57.2	-47.3	12.0
Er	-15.37	32.4	-5.0	-6.1	11.2	15.3	5.8	-1.3	-3.9	20.1	21.0	-4.6
Tm	-8.21	52.53	-2.6	-3.3	18.2	4.1	5.9	-1.1	-2.1	17.1	17.8	-2.8

 $_d^a \langle S_z \rangle_j$ and C_j values are adopted from ref. 44a. $_d^b$ PBE/SARC-TZVP calculations. $_d^c$ The values are obtained as $\delta^{pc}(DFT) = \delta^{para}(exp) - \delta^{con}(DFT)$. $_d^a$ "PCM" stands for point charge model. $_d^e$ Δ^{308}_{268} values are computed using PCM approach and include only δ^{pc} contribution.

Table 3 Contact and pseudocontact ⁴⁵Sc chemical shifts in **1M** molecules

	$\delta^{\rm con}$		δ^{pc}			
1M	Reilley	2-nuc ^a	Reilley	2-nuc ^b	PCM^c	Δ_{268}^{308d}
Ce	4	0	95	79	87	-14
Pr	11	-1	168	257	197	-31
Nd	17	-2	66	163	150	-36
Tb	-122	14	1278	735	1430	-255
Dy	-110	13	1472	1679	1799	-319
Но	-87	10	578	862	1060	-290
Er	-59	7	-477	-440	-458	100
Tm	-32	4	-773	-213	-394	62

^a Obtained using the fit of eqn (4a) together with ¹³C-PHHJ shifts, and then subtracting $\delta^{\rm con}(^{13}{\rm C})$ values estimated by Reilley's approach; **1Pr** value was not included in the fit; inclusion of **1Pr** shift worsened the fit from $R^2=0.96$ to $R^2=0.87$, but gave the values closer to those obtained by Reilley's approach. ^b The values are obtained as $\delta^{\rm Pc}(2\text{-nuc}) = \delta^{\rm para}(\exp) - \delta^{\rm con}(2\text{-nuc})$. ^c "PCM" stands for point charge model. ^d Δ^{308}_{268} values are computed using PCM approach and include only $\delta^{\rm pc}$ contribution.

Fig. 7 Three-nuclei 3D plot in $\delta_{ij}^{\mathrm{para}}/\langle S_z \rangle_{j}$ coordinates for $^{45}\mathrm{Sc}$, $^{13}\mathrm{C}$ -THJ and $^{13}\mathrm{C}$ -PHHJ chemical shifts in $1\mathrm{M}$ series ($T=288\,\mathrm{K}$). Red spheres are 3D data; blue, green, and black circles are projections on coordinate planes. Red grid shows a fitted plane (1Pr was not included in the fit).

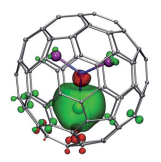
points except for **1Pr** are close to the plane obtained by the least square fitting of eqn (4b). Analysis of the 2D plots in Fig. 7 shows that a good linear correlation exists only for the PHHJ-vs.-Sc plot (**1Pr** value is deviating again). Substituting the $\delta^{\rm con}$ values obtained for the PHHJ carbons using eqn (3) into eqn (4a) for the PHHJ-vs.-Sc correlation allows estimation of the ⁴⁵Sc $\delta^{\rm con}$ shifts. The values obtained by this way are much smaller than those obtained using Reilley's approach. It should be also noted that the results of the $\delta^{\rm con}$ calculations for Sc using the 2-nuclei method are rather erratic and change significantly with small variation of the input parameters. At the same time, both methods agree in that the ⁴⁵Sc $\delta^{\rm con}$ shifts are considerably smaller than the pseudocontact contributions.

The fact that the data points for THJ carbons fit well into the 3-nuculei model but do not give linear correlations in the 2-nuclei model might indicate that the rhombic anisotropy contribution is considerable here, but the limited amounts of points does not allow more solid conclusions.

DFT calculations of contact shifts. Independent computations of contact shifts are possible using quantum chemical approaches. DFT computations of lanthanide-containing molecules for systems other than $4f^7$ cannot be reliable (unless 4f-in-core effective core potentials are used, but this approach cannot be used if spin properties related to the 4f electrons are of interest). However, the F_i value usually remains the same for the whole lanthanide row, and hence computations can be performed for Gd analogues. Recently, such calculations provided accurate estimation of the contact shifts in a series of Tb complexes. 48

Fig. 8 shows the spin density distribution in **1Gd** computed using the scalar-relativistic DKH2 approximation, PBE functional, and SARC-TZVP basis set. Negative spin polarization of the central nitrogen atoms and several carbon atoms close to the lanthanide can be well seen (red-colored lobes). At the same time, the sign of the spin density on the Sc atoms and carbon

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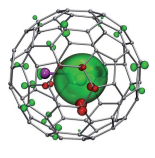


Fig. 8 Spin-density distribution in $GdSc_2N@C_{80}$ (two orientations of the molecule are shown). To visualize small spin polarization effects, the isosurfaces are plotted at relatively small spin density values of ± 0.0006 a.u. (green (+) and red (–)).

atoms further away from the Gd ion is the same as for the Gd itself.

Hyperfine coupling constants for carbon atoms in $GdSc_2N@C_{80}$ span the range from -0.041 MHz to 0.096 MHz, and the averaging over the whole carbon cage yields the values of 0.015 and 0.010 MHz for the PHHJ and THJ carbons, respectively. The pseudocontact shifts computed for ^{13}C atoms using these constants are listed in Table 2. Remarkably, the DFT-computed values for the PHHJ carbons are quite close to those obtained by Reilley's method. If we tentatively suggest that the agreement between the theory and experiment is equally well for the THJ carbons, their contact shifts should be 35% smaller than those of PHHJs. Furthermore, relative contributions of the contact term in total paramagnetic shifts of the THJ carbons are much smaller than for the PHHJ carbons.

For the ⁴⁵Sc hyperfine coupling constant, the PBE-DKH approximation with tailored SARC-TZVP basis set gives the average value of -0.27 MHz. An increasing the basis set to def2-TZVP or inclusion of the spin-orbit coupling corrections give the values of -0.26 MHz and -0.28 MHz, respectively, whereas switching to ZORA scalar relativistic method reduces the ⁴⁵Sc hyperfine coupling constant to -0.21 MHz. If hybrid PBE0 functional is used instead of PBE with SARC-TZVP basis, predicted DKH and ZORA values are -0.35 and -0.49 MHz, respectively. The use of PBE-DKH/TZVP constants for estimation of the 45Sc contact shifts in 1M molecules yields unrealistically high values (625 ppm for 1Tb, 560 ppm for 1Dy, etc.). It should be noted that computation of the 45Sc hyperfine coupling constants is less straightforward than for 13C nuclei because of the subtle balance between the valence- and coreshell spin polarizations, which often have opposite signs in transition metals.⁴⁹ Due to this reason, prediction of the ⁴⁵Sc hyperfine coupling constants meets serious difficulties even for the Sc-based EMF radicals such as Sc₃N@C₈₀⁻ (ref. 50), where coupling constants are large and dominated by SOMO contribution. Reliable prediction of the core-shell spin polarization becomes even more problematic for paramagnetic molecules, in which the spin density on the Sc atoms is largely induced by spin polarization (such as in 1M). We have to admit that DFT prediction of small ⁴⁵Sc coupling constants in **1M** can hardly be reliable at this moment and refrain from further discussion of DFT-computed $\delta^{\text{con}}(^{45}\text{Sc})$ values.

Temperature dependence of chemical shifts. If Bleaney's conditions are satisfied, the temperature dependence of chemical shifts can be also used to distinguish contact and pseudocontact terms: whereas the contact term scales as T^{-1} , the pseudocontact term has T^{-2} dependence. If one of the terms dominates, plot δ^{para} versus T^{-1} or T^{-2} should give a straight line, whose intercept is close to δ^{dia} . In due turn, a large deviation of the intercept from the reference δ^{dia} value signals that the corresponding contribution is small. This simple yet efficient approach was employed before to prove the dominance of pseudocontact shift in Ce-based EMFs. However, this approach can give reliable results only if the pseudocontact term has real T^{-2} dependence, and if one the terms dominates. If LF splitting is larger than the thermal energy, contribution of T^{-3} and higher terms cannot be ignored,51 and results of the linear fitting become inconclusive. In the 1M series, only for 1Ce is the intercept of the $\delta^{\text{para}}(T^{-2})$ plot close to δ^{dia} for all three nuclei (see ESI Table S1†). For all other lanthanides, the intercepts of $\delta^{\text{para}}(T^{-1})$ and $\delta^{\text{para}}(T^{-2})$ linear fits give rather irregular set of data, sometimes with large deviations from δ^{dia} for both T^{-1} and T^{-2} plots (ESI Table S1†). Thus, large magnetic anisotropy of lanthanide ions in 1M molecules prevents the use of temperature dependence for distinguishing contact and pseudocontact contributions to paramagnetic chemical shifts. Good results obtained for the **1Ce** in $\delta^{\text{para}}(T^{-2})$ fits are in part due to the small $\langle S_z \rangle_{Ce}$ value (Table 2), which makes the contact term negligible.

Geometry parameters of 1M molecules

For the further analysis of the magnetic state and the LF splitting of lanthanide ions in the 1M compounds it is necessary to take into account a variation of the molecular geometry parameters along the series. Namely, a significant decrease of the ionic radii of lanthanides from La to Lu cannot be ignored, especially when magnetic anisotropy is large. Therefore, to obtain a consistent set of the molecular geometry parameters along the whole 4f row for the subsequent LF calculations (see next section), we performed DFT optimization of all structures using the B3LYP functional and 4f-in-core basis set for lanthanides. The M-N and averaged Sc-N bond lengths are listed in ESI Table S2† and are plotted versus Shannon ionic radii of M³⁺ ions in Fig. 9. As expected, the decrease of the ionic radii from La to Lu results in the gradual decrease of the M-N distance (from 2.241 Å in 1La to 2.133 Å in 1Lu). At the same time, since the carbon cage remains the same, the increase of the lanthanide size also leads to the simultaneous decrease of the Sc-N bond lengths (compare 1.929 Å in **1La** to 1.979 Å in 1Lu) so that the cluster size remains more or less the same along the whole series. Single-crystal X-ray structures were reported earlier for 1La,26 1Ce,19 1Gd,27 1Tb,27 and 1Er,28 and experimental bond length values also confirm this trend. Namely, in the 1La-1Ce-1Gd-1Tb-1Er the lanthanidenitrogen bond is decreasing as 2.196(4)-2.184(2)-2.149(10)-2.126(11)-2.089(9) Å. The averaged Sc-N distance is varying as 1.932(7)-1.938(2)-1.918(9)-1.949(8)-1.968(6) Å, respectively, i.e. there is a gradual decrease from 1La/1Ce to 1Tb to 1Er (note

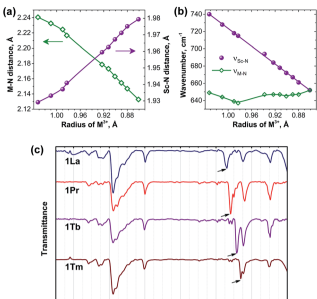


Fig. 9 (a) Correlation between DFT-computed M–N and Sc–N bond lengths in 1M molecules and ionic radii of lanthanides; (b) correlation between the metal–nitrogen stretching mode frequencies and ionic radii of lanthanides (green – M–N bonds, purple – Sc–N bonds); (c) FTIR spectra of selected 1M molecules: 1La, 1Pr, 1Tb, and 1Tm; arrows mark the ν_{Sc-N} modes.

1200

1000

800

600

400

1600

1400

that **1Gd** deviates from the trend). A comparison to the computed data shows that DFT overestimates the M–N distances by *ca.* 0.04–0.06 Å, whereas the error for the Sc–N distances is smaller than 0.01 Å. Overall, DFT-optimized bond lengths give reasonable estimation of the experimental values.

Significant variation of the bond lengths in nitride clusters can be also inferred from the analysis of the FTIR absorption spectra. In the frequency range of 500–800 cm⁻¹, all nitride clusterfullerenes have a characteristic antisymmetric metalnitrogen stretching vibration with medium to strong absorption intensity, whose frequency correlates with the metal-nitrogen bond length and therefore can be used for the analysis of the structural correlations.^{29c,52} This mode corresponds to the inplane motion of the nitrogen atom and is two-fold degenerate for the M_3N cluster with the frequency of $(3k_{M-N})^{0.5}m_N^{-0.5}$, where k_{M-N} is the force constant of the metal-nitrogen bond, and m_N is the mass of the nitrogen atom. In the mixed-metal cluster MSc_2N , the degeneracy is lifted and the frequencies of two resulting vibrations are:

$$\nu_{\text{Sc-N}} = (3k_{\text{Sc-N}})^{0.5} m_{\text{N}}^{-0.5}$$
 (5a)

$$\nu_{\rm M-N} = (2k_{\rm M-N} + k_{\rm Sc-N})^{0.5} m_{\rm N}^{-0.5}$$
 (5b)

Fig. 9c compares the IR spectra for **1La**, **1Pr**, **1Tb**, and **1Tm** (IR spectra of other **1M** compounds were reported by our group earlier^{20b,24,29c}). In all spectra both the $\nu_{\text{M-N}}$ and $\nu_{\text{Sc-N}}$ components can be clearly identified in the 600–800 cm⁻¹ range. Comparison of the $\nu_{\text{M-N}}$ values along the **1M** series is not

straightforward, because the frequency *versus* bond length correlation does not hold exactly for different metals and also because of the admixture of the $k_{\rm Sc-N}$ force constant in eqn (5b). In fact, the $\nu_{\rm M-N}$ frequency remains in the range of 635–650 cm⁻¹ in the whole **1M** series (Fig. 9b). The $\nu_{\rm Sc-N}$ frequencies are better suited for the structural correlations: Fig. 9b shows that the $\nu_{\rm Sc-N}$ frequencies vary from 652 cm⁻¹ in **1Lu** to 740 cm⁻¹ in **1La** and exhibit perfect linear correlation with the lanthanide ionic radii. Very good correlation also exists between the DFT-predicted Sc-N bond lengths and experimental $\nu_{\rm Sc-N}$ frequencies (not shown), thus confirming reliability of the DFT-computed structural parameters.

Ligand field in 1M molecules

Reliable determination and verification of the cluster bond lengths discussed in the previous section is crucial for the analysis of the LF in nitride clusterfullerenes. Recently we showed that the point charge model employing scaled Bader charges can reasonably reproduce the ab initio (CASSCF) computed ligand field splitting in DySc2N@C80,7a,20a which enables the use of this simple approach for other 1M compounds. To calculate the LF splitting in this work we used B3LYP-optimized coordinates discussed above and Bader charges for nitrogen, scandium, and four closest carbon atoms (denoted C1-C4 in Fig. 1; since the cage carbon atoms have very small contribution to the LF,20a it is sufficient to consider only few nearest atoms, their QTAIM charges are listed in the ESI Table S2†). The charges were scaled by a factor of 0.754 to match the ab initio LF splitting in 1Dy from ref. 7a (see ESI Table S3†). Fig. 10 shows an overview of the $m_{\rm I}$ levels in all studied 1M molecules (the values and assignment are listed in the ESI Table S4†).

The central nitrogen atom gives the main contribution to the LF in nitride clusterfullerenes because of its large negative charge (Bader charge -1.7) and close distance to the lanthanide

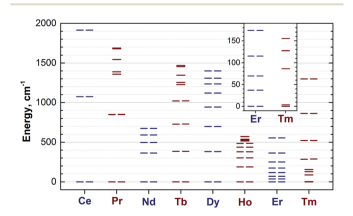


Fig. 10 Ligand-field splitting of the $m_{\rm J}$ levels in the 1M compounds computed using point-charge model. The inset shows low-energy levels in 1Er and 1Tm in the range of $0-190~{\rm cm}^{-1}$. Each degenerate $\pm m_{\rm J}$ level of Kramers ions (Ce, Nd, Dy, Er) and quasi-degenerate (within $3~{\rm cm}^{-1}$) levels of non-Kramers ions are shown as double lines. To guide an eye, Kramers and non-Kramers ions are denoted in dark blue and wine, respectively.

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ion (see ESI Table S3† for the LF point charge calculations using different parts of the MSc₂N@C₈₀ molecule). Therefore, correct description of the molecular geometry is crucial since even small variations of the metal-nitrogen distance lead to considerable changes in the ligand field.20a Two Sc ions produce the second largest contribution to the LF after the nitride ion. Their positive charges (Bader charge of ca. +1.7) reduce the effect of the nitride ion (e.g., in 1Dy the LF splitting is reduced by ca. 35% when Sc contribution is "turned on") and introduce a deviation from the uniaxial symmetry. The charges of carbon atoms are much smaller than those of N and Sc (below -0.1), and hence only few carbons close to the lanthanide ion have non-negligible contribution to the LF (in comparison to the M+ N + 2Sc system, addition of four carbons increases the splitting energies by ca. 12%, see ESI Table S3† for 1Dv and ref. 20a for a more detailed analysis of 1Ho). Thus, the LF experienced by lanthanide ions in 1M molecules has roughly a uniaxial character induced by a large negative point charge of the nitride ion, which results in rather simple LF splitting pattern despite the low molecular symmetry. A similar situation was found for a series of other Dy-based SMMs with low molecular symmetry.⁵³

Lanthanide ions with oblate 4f-density. The ground magnetic state of the ions with oblate shape of the 4f electron density (Ce, Pr, Nd, Tb, Dy, Ho; see ref. 54 and Fig. 11a for 4f density in 1Dy) has the largest J_z value (5/2 in 1Ce, 4 in 1Pr, 9/2 in 1Nd, 6 in 1Tb, 15/2 in 1Dy, and 8 in 1Ho), and the energy of the other $m_{\rm I}$ states is gradually increasing with the decrease of the $|m_{\rm I}|$ values. The gaps between the neighboring $m_{\rm I}$ states are also getting smaller with the decrease of the $|m_I|$ value. The deviation from the uniaxial symmetry induced by the two Sc ions is well seen for non-Kramers ions (Pr, Tb, Ho). For these ions, degeneracy of the $\pm m_{\rm I}$ states is not enforced, but remains rigorous in the uniaxial ligand field produced by a single nitride ion. When the contribution of the Sc ions is also taken into account, an approximate degeneracy of the $\pm m_{\rm I}$ states holds only for large projections of the total momentum, whereas for the levels with $|m_1| = 1-2$ the splitting becomes significant. A mixing of the m_1 states is also substantial for small $|m_1|$ values showing that the $m_{\rm I}$ is not a "good" quantum number anymore.

The practical importance of the states with small $|m_J|$ values in ions with oblate 4f density is rather limited because the LF splitting in **1M** is very large (Fig. 10). The smallest overall LF splitting, 571 cm⁻¹, is found in **1Ho**, whereas the largest LF splitting is reaching 1915 cm⁻¹ in **1Ce**. The gap between the ground state and the first excited state in **1Ce** (1075 cm⁻¹), **1Pr** (849 cm⁻¹), **1Nd** (364 cm⁻¹), **1Tb** (384 cm⁻¹), and **1Dy** (382 cm⁻¹) is so large that even at room temperature the ground magnetic state with the largest J_z projection is predicted to have predominant contribution to the magnetic properties. As a result, 4f density in such **1M** molecules exhibits only small variation with temperature. For instance, 4f densities in **1Dy** calculated at 2 K and 288 K can hardly be distinguished by eye (Fig. 11a).

The two-fold degenerate ground state with the large J_z projection and a significant gap to the first excited state are prerequisites for the SMM behaviour,^{54b} and hence the data in Fig. 10 show that slow magnetization reversal can be expected

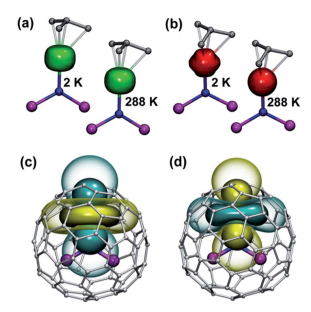


Fig. 11 (a and b) isosurfaces of 4f electron density in 1Dy (a, green) and 1Er (b, red) computed at 2 and 288 K. Only the nitride cluster and four carbon atoms nearest to the lanthanide ion are shown. Whereas the shape of the lobe remains almost the same in 1Dy, in 1Er the increase of the temperature from 2 to 288 K changes the shape of the 4f density distribution to a more spherical one. (c and d) Pseudocontact shift isosurfaces in 1Dy (c) and 1Er (d) at 288 K computed using the point-charge model (cyan – positive, yellow – negative). Solid/transparent surfaces correspond, respectively, to $\pm 6000/\pm 1500$ ppm isovalues in 1Dy and $\pm 2000/\pm 500$ ppm isovalues in 1Er.

for all **1M** compounds (except for the **1Gd**, **1Er** and **1Tm**). The SMM behaviour with exceptionally long attempt time is already discovered for **1Dy**.^{3,4} Field-induced SMM behaviour is also found in non-Kramers **1Ho**,⁵ albeit on a much shorter timescale than in **1Dy**. The rigorous protection of the two-fold degeneracy of the ground magnetic state in Kramers ions makes them more attractive for single ion magnetism, and we expect **1Ce** and **1Nd** to be the next candidates for the SMM behaviour in nitride clusterfullerenes as already detected in some organometallic complexes of Ce⁵⁵ and Nd.⁵⁶ It should be however noted that the magnetization reversal barrier in **1Dy** is smaller than expected from its LF splitting,⁴ which indicates that the mechanism for the magnetization reversal in **1M** compounds is more complex and may include, for instance, Raman-like process.⁵

Lanthanide ions with prolate 4f-density. The LF splitting pattern in **1Er** and **1Tm** is completely different because of the prolate shape of the 4f electron density of these lanthanides (Fig. 11b). The ground state for these lanthanides has the lowest J_z projection ($m_J = \pm 1/2$ in **1Er** and $m_J = 0$ in **1Tm**). The energy is increasing with the increase of the $|m_J|$ values, and the density of states is higher in the low energy part of the spectrum, whereas at higher energies the energy levels are distributed sparser. As a result of such level distribution, variation of the temperature has profound effect on the magnetic anisotropy of **1Tm** and **1Er**. As can be seen in Fig. 11b, an increase of the temperature dramatically changes 4f electron density distribution in **1Er** and makes it almost isotropic. Thus, the room-temperature magnetic properties of **1Er** and **1Tm** result from the contribution of several m_J states.

1Er is the only compound in the whole **1M** series for which spectroscopic data on the lanthanide-based luminescence are available. At helium temperatures the luminescence spectra of **1Er** exhibit the fine structure due to the Er-based transitions in the ${}^4I_{13/2} \rightarrow {}^4I_{15/2}$ manifold. 9b,57 Unfortunately, emission from several molecular/cluster sites complicates the fine structure and makes assignment less straightforward. The energy gap between the two lowest energy LF states determined in ref. 57, 28/37 cm⁻¹, compares reasonably to the 37 cm⁻¹ gap between the $m_{\rm J}=\pm 1/2$ and $m_{\rm J}=\pm 3/2$ states in our calculations, however the overall splitting determined in the optical measurements, 330 cm⁻¹, is considerably smaller than the value of 554 cm⁻¹ predicted by the point-charge model.

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Computed versus experimental pseudocontact chemical shifts

LF calculations allow direct estimation of the magnetic susceptibility tensor at different temperatures using Van Vleck formulae, and therefore enable computations of the pseudocontact chemical shifts using eqn (1a) or (1b) given the atomic coordinates are known (e.g. from DFT calculations) and they are not changing with temperature. Comparison between experimental and computed chemical shifts can be used then to evaluate reliability of the computed ligand field splitting levels. Rotation of the nitride cluster inside the carbon cage means that the values computed for individual carbon atoms should be averaged. Recently we found that the static model (simple averaging of the values) and the averaging of the values computed over molecular dynamics trajectory give comparable results for **1Ho**, ^{20a} and therefore only static model is applied in this work.

In the **1M** molecules with oblate 4f-density eqn (1a) and (1b) give identical values of the pseudocontact shift because χ_{xx} and χ_{yy} components of the magnetic susceptibility tensor are virtually equal. In the **1Er** and **1Tm** molecules, the difference between χ_{xx} and χ_{yy} values is rather large, and hence eqn (1a) is to be used. The pseudocontact shifts computed for the THJ and PHHJ carbons as well as for the Sc atoms at T=288 K are listed in Table 2 and 3 along with the computed $\Delta_{268}^{308}(\delta)$ values. Fig. 11 plots pseudocontact shift isosurfaces in **1Dy** and **1Er** as examples of lanthanides with oblate and prolate 4f density shapes, respectively.

The reliability of the point charge model in calculations of the magnetic susceptibility can be evaluated comparing experimental and computed δ^{pc} shifts (Fig. 12). As experimental values we use the results of Reilley's approach for $^{13}\text{C-PHHJ}$ signal and the values from the 2-nuclei fitting for ^{45}Sc shifts. Details for the $^{13}\text{C-THJ}$ shifts are not discussed in the text but can be found in the ESI Fig. S11.†

As can be seen in Fig. 12, reasonably good correlation exists between experimental and computed data. If **1Tm** and **1Tb** are not included in the set, the linear correlation with the intercept set to zero gives R^2 values of 0.98 and 0.99 for the PHHJ and ⁴⁵Sc shifts, respectively. Note also that the ⁴⁵Sc δ^{pc} shifts from the 2-nuclei model are very close to the total δ^{para} shifts. Thus, we can tentatively use ⁴⁵Sc chemical shifts of paramagnetic mixedmetal nitride clusterfullerenes to evaluate the size of the

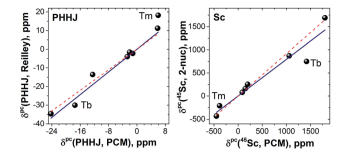


Fig. 12 Correlation between computed and experimental chemical pseudocontact shifts: 13 C-PHHJ (left) and 45 Sc (right). Experimental $\delta^{\rm pc}(^{13}$ C) shifts are obtained by Reilley's approach, whereas 45 Sc values are estimated using 2-nucleus method, see text for further details. Solid lines is a linear fit for a complete set of data ($R^2 = 0.94$ for both PHHJ and Sc), whereas red dashed lines were obtained for fitting without 1Tm and 1Tb values. The intercept was set to zero in linear fits.

magnetic anisotropy of the endohedral lanthanide ions without rather complicated and sometimes hardly possible estimation of the contact term.

It should be also noted that the slope of the linear fit in Fig. 12 is close to 0.9 for ⁴⁵Sc but is near 1.4 for ¹³C-PHHJ. The slope for an analogous fit for the 13C-THJ signals (see ESI Fig. S11†) is close to 0.9. As follows from eqn (1) and our earlier studies of the Ho-based NCFs, 20a the structural factor also plays an important role in determining the δ^{pc} values, and dynamic nature of the ¹³C shifts in **1M** molecules (*i.e.* averaging over the cluster rotation) makes calculation less straightforward than for the ⁴⁵Sc shifts. For instance, in **1Ho** the term $(3\cos^2\theta_i - 1)/(1+\cos^2\theta_i)$ $12\pi R_i^3$ (denoted hereafter as G_i) computed in this work at the B3LYP level is $-3.30 \times 10^{-4}/-1.29 \times 10^{-3} \text{ Å}^{-3}$ for the PHHJ/ THJ carbons, respectively. At the same time, our recent PBE computations using Y as a model for Ho gave the values of $-7.37 \times 10^{-4}/-5.59 \times 10^{-4} \text{ Å}^{-3}$, respectively. Along the 1M series, the G_i values for carbon atoms vary by 15–20%, from $-3.91 \times 10^{-4}/-1.13 \times 10^{-3} \text{ Å}^{-3} \text{ in 1Ce to } -3.16 \times 10^{-4}/-1.31$ \times 10⁻³ Å⁻³ in 1Er. Since the slope of the linear fit between experimental and calculated $\delta^{\rm pc}$ values for THJ carbons is closer to 1, we propose that B3LYP level gives correct values for the THI carbons, whereas G_i factors for the PHHJs are underestimated (and therefore the slope is close to 1.4). Thus, the use of paramagnetic 13C chemical shifts for analysis of the magnetic anisotropy of the endohedral lanthanide ions meets certain difficulties caused by the uncertainties in the structural factor when endohedral cluster is rotating on the NMR time scale. On the contrary, the G_i term for ⁴⁵Sc is almost independent on the level of theory employed to calculate it. Moreover, its variation along the whole 1M series is below 1% because the shortening of the M-N distance in the cluster is balanced by the elongation of the Sc-N distance. Such indifference to the structural factor makes the ⁴⁵Sc chemical shifts more suitable for the analysis of anisotropy magnetic in mixed-metal clusterfullerenes.

Computed Δ^{308}_{268} values reasonably agree with experimental data for THJ carbon and Sc signals, but are rather far from the experimental values of the PHHJ carbons. The latter is likely to

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be due to the larger relative contribution of the contact shift for the PHHJ carbons, which is not taken into account in PCM computations of Δ_{268}^{308} . This is especially well seen for the 1Tm: based on its magnetic anisotropy, negative Δ_{268}^{308} value is expected for the pseudocontact shift, but positive for the contact shift. A small positive Δ_{268}^{308} value observed experimentally for the 1Tm-PHHJ carbon is therefore due to the compensation of both terms. On the contrary, the pseudocontact contribution in the 1Tm-THJ chemical shift is much larger than the contact counterpart, and the experimental Δ_{268}^{308} value is negative in good agreement with the results of computations.

1Tm and 1Tb are important deviations from the general trend (Fig. 12). Especially noticeable is the 1Tb value for the ⁴⁵Sc shift, which is predicted to be much more positive than experimentally observed (Fig. 12). In other words, whereas the point charge model predicts that the LF splitting in 1Tb and 1Dy is quite similar (Fig. 10), from the ⁴⁵Sc NMR spectra it follows that the LF splitting in 1Tb should be considerably smaller. Interestingly, recent ab initio CASSCF calculations predict that the LF splitting in 1Tb is indeed smaller than in 1Dy,7b which would give better agreement with experimental NMR shifts. Similarly, correct prediction of the experimental values for 1Tm require reliable estimation of a considerable number of the LF splitting levels (Fig. 10), rather than only 1-2 excited states needed for other 1M molecules. Inability of the point charge model to correctly treat all states may be also the reason of the noticeable deviations well seen for 1Tm values in Fig. 12. Therefore, both 1Tb and 1Tm values provide convenient training set to be considered in future computational studies. At the current moment we can conclude that the point charge model gives qualitatively correct predictions of the LF splitting in lanthanide EMFs but fails to describe more subtle effects. More advanced approaches would be needed to address this problem,⁵⁸ but this task goes beyond the scope of this work.

Conclusions

In this work we have performed the first systematic paramagnetic NMR study of $MSc_2N@C_{80}$ - I_h molecules with M running through all lanthanides capable of forming nitride clusterfullerenes (M = La, Ce, Pr, Nd, Tb, Dy, Ho, Er, Tm, Lu). Analysis of the whole set of data enabled separation of the contact and pseudocontact contributions to the paramagnetic shifts. We showed that the contact shift of 13 C nuclei may be rather large, although pseudocontact term is still larger. For the 45 Sc shifts, the pseudocontact term dominates. Since pseudocontact term is directly dependent on the magnetic anisotropy, the 45 Sc NMR spectra are found to be especially useful for the studies of magnetic properties of lanthanide ions in clusterfullerenes.

Interpretation of the paramagnetic NMR data required modelling of the ligand filed splitting in $MSc_2N@C_{80}$ molecules, which was accomplished using the point charge model. Although quite simple, this approach provided semi-quantitative agreement with experimental data. Considerable deviations between experimental and computed chemical shifts are found only for $TmSc_2N@C_{80}$ and $TbSc_2N@C_{80}$. We showed that the

main contribution to the ligand field is from the nitride ion, which results in an almost uniaxial ligand field. For the lanthanides with oblate shape of the 4f density (i.e. all lanthanides except for Er and Tm) the ground magnetic state in MSc₂N@C₈₀ has the largest J_z projection with rather large gap to higher energy states (with smaller m_J values). It shows that MSc₂N@C₈₀ molecules with Kramers lanthanide ions are especially promising in the field of single molecule magnetism. Yet, more refined treatment of the ligand field splitting might be needed to fully account for all experimental data.

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