Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2024, 15, 15160

dll publication charges for this article have been paid for by the Royal Society of Chemistry

Received 7th May 2024 Accepted 28th July 2024

DOI: 10.1039/d4sc03005b

rsc.li/chemical-science

δ -Bonding modulates the electronic structure of formally divalent nd¹ rare earth arene complexes†

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Landmark advances in rare earth (RE) chemistry have shown that divalent complexes can be isolated with non-Aufbau $4f^n\{5d/6s\}^1$ electron configurations, facilitating remarkable bonding motifs and magnetic properties. We report a series of divalent bis-tethered arene complexes, [RE(NHAr^{iPr₆})₂] (2RE; RE = Sc, Y, La, Sm, Eu, Tm, Yb; NHAr^{iPr₆} = {N(H)C₆H₃-2,6-(C₆H₂-2,4,6-iPr₃)₂}). Fluid solution EPR spectroscopy gives $g_{iso} < 2.002$ for 2Sc, 2Y, and 2La, consistent with formal nd¹ configurations, calculations reveal metalarene δ -bonding via mixing of $nd_{(x^2-y^2)}$ valence electrons into arene π^* orbitals. Experimental and calculated EPR and UV-Vis-NIR spectroscopic properties for 2Y show that minor structural changes markedly alter the metal $d_{(x^2-y^2)}$ contribution to the SOMO. This contrasts $4f^n\{5d/6s\}^1$ complexes where the valence d-based electron resides in a non-bonding orbital. Complexes 2Sm, 2Eu, 2Tm, and 2Yb contain highly-localised $4f^{n+1}$ ions with no appreciable metal-arene bonding by density functional calculations. These results show that the physicochemical properties of divalent rare earth arene complexes with both formal nd^1 and $4f^{n+1}$ configurations are nuanced, may be controlled through ligand modification, and require a multi-pronged experimental and theoretical approach to fully rationalise.

Introduction

A hallmark of trivalent lanthanide ions is the relative insensitivity of valence 4f-orbitals to the coordination environment. While this gives rise to useful physical and optical properties, such as their narrow optical emission profiles and large magnetic moments from their unquenched orbital angular momentum, the weakness of this interaction also inhibits the extent to which their properties may be tailored through molecular design. Advancements in divalent molecular rare earth and lanthanide (Sc, Y, La–Lu; collectively Ln henceforth) chemistry have shown that in coordination environments such as $[Ln(Cp^R)_3]^-$ or $[Ln(Cp^R)_2]$ $(Cp^R)_2$ substituted

cyclopentadienide ligands), Ln(II) complexes can be isolated with $4f^n5d^1$ or $4f^n{5d/6s}^1$ valence electron configurations, at least for La-Lu, while Sc and Y necessarily give 3d1 or 4d1 ions.1-23 Further work shows these non-Aufbau ions can be exploited as potential qubit candidates, 15-17 exhibit recordsetting magnetic properties, 18-20 and have produced the first examples of molecular Ln-Ln bonding outside of endohedral fullerenes.20-22 Similar advancements have also been made with actinide elements.24 However, with few exceptions, design strategies employed to isolate examples of these ions use geometries (e.g. C_{3h} or D_{5h}) that minimise or forbid, by symmetry, the mixing of 5d (or 5d/6s hybridised) valence electrons with ligand orbitals. This is useful in some applications,15-17 but in analogy to the poor tunability of 4forbitals in Ln(III) ions, it limits the extent to which the electronic structures of 4f"5d1 or 4f"{5d/6s}1 Ln(II) complexes may be controlled through ligand design.

Using ligand field principles commonplace in the d-block that also apply to $4f''5d^1$ Ln(II) ions, we may instead target molecular designs where the electronic structure and physicochemical properties are more sensitive to changes in the coordination environment. Arene ligands provide a promising route towards these goals due to their ability to act as symmetry-allowed donors into vacant d-orbitals, and further stabilise lower oxidation states through back donation. Indeed, the only examples of formally zero-valent rare earth complexes are within $[M(\eta^6-C_6R_6)_2]$ frameworks.

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[†] Electronic supplementary information (ESI) available: Details of the starting materials used, descriptions of sample preparation for spectroscopic measurements, and methodologies and coordinates for the DFT and CASSCF calculations. CCDC 2266235, 2266236, 2266243, 2266244, 2266255–2266258, 2266261–2266264, 2282041, 2295306 and 2295307. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d4sc03005b.

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Herein, we present a series of structurally analogous bistethered arene divalent rare earth complexes of the form $[M(NHAr^{iPr_6})_2]$ (2M, M = Sc, Y, La, Sm, Eu, Tm, Yb; NHAr^{iPr_6} = $\{N(H)C_6H_3-2,6-(C_6H_2-2,4,6-iPr_3)_2\}$ - the synthesis and some properties of 2Y have been reported previously.³⁸ In all cases, characterisation using SQUID magnetometry, solid and solution phase EPR spectroscopy, UV-Vis-NIR spectroscopy, density functional theory (DFT), and complete active space selfconsistent field (CASSCF) calculations support the description of these as formal Ln(II) complexes, demonstrating that this framework is robust across a range of Ln(II) ion sizes and (formal) reduction potentials. While all seven divalent complexes display close metal-arene contacts, only 2Sc, 2Y, and **2La** show mixing of a metal valence $nd_{(x^2-y^2)}$ orbital with the arene π^* to give δ -bonding interactions. Complexes 2Sm, 2Eu, 2Tm, and 2Yb instead adopt metal-localised $4f^{n+1}$ configurations in accordance with their large Ln(II) $4f^{n+1} \rightarrow 4f^n 5d^1$ promotional energies.39,40 Solid and solution phase UV-Vis-NIR and EPR for 2Y, combined with DFT calculations, reveals that the balance of metal vs. arene-centred spin-density is sensitive to small structural changes between these phases, suggesting the properties of divalent rare earth complexes with nd¹ may be tuned through ligand design.

Results and discussion

Synthesis

A reductive route was pursued for the synthesis of **2M** (M = Sc, Y, La, and Tm; Scheme 1A), which began with the synthesis of trivalent [M(NHAr^{iPr₆})₂(I)] (**1M**, M = Sc, Y, La, Tm) precursors. Complexes **1M** were synthesised by salt elimination between KNHAr^{iPr₆} (ref. 41 and 42) and the relevant rare earth tri-iodide salt [MI₃(THF)_n] (n = 4, M = La; n = 3.5, M = Y, Tm; n = 3, M = Sc) in Et₂O.⁴³ Workup and crystallisation from toluene gave fair to good crystalline yields (47–62%). ¹H NMR spectroscopy of diamagnetic **1Sc**, **1Y**, and **1La** in d₆-benzene show the M···C_{6-arene} interaction is dynamic in solution and that all four Tripp groups are in exchange (see ESI† for detailed assignments).³⁸

Reduction (KC₈, 1.2 equiv.) of **1M** (M = Sc, Y, La, Tm) in Et₂O gave, after workup and crystallization from Et₂O, dark crystals of **2M** in fair crystalline yields (shown as % yield ahead): **2Sc** (red, 53%), **2Y** (red/green, 55%), **2La** (red/brown, 45%), and **2Tm** (red/brown, 65%). Salt elimination between KNHAr^{iPr₆} and MI₂(THF)₂ (M = Sm, Eu, Yb)⁴⁴ in Et₂O gave poor to excellent isolated crystalline yields of **2Sm** (69%), **2Eu** (20%), and **2Yb** (82%) (Scheme

1B). Attempts to synthesise **1Sm** and **1Yb** using $[MI_3(THF)_n]$ precursors were unsuccessful, leading to intractable mixtures (see ESI†), and we did not attempt the synthesis of **1Eu**.

While the ¹H NMR spectra of **1Sc**, **1Y**, and **1La** showed the metal-bound and "terminal" {C₆H₂-2,4,6-*i*Pr₃} (Tripp) groups were in exchange at room temperature, the spectrum of diamagnetic 2Yb in d₆-benzene shows six doublets for the CH₃-iPr groups, along with a single N(H) resonance. Thus, 2Yb is C_2 symmetric in solution and the Tripp groups do not appreciably exchange at room temperature. This does not appear to be due to steric hindrance as the Sc(III) in 1Sc is smaller than Yb(II) (6-coordinate radii: Yb(II), 0.868 Å; Sc(III), 0.745 Å). A variable temperature ¹H NMR study in d₈-toluene shows the CH₃-iPr peaks begin to coalesce at 308 K, but full equilibrium is not reached until ca. 358 K (Fig. S40 and S41†). The 171Yb-1H HMBC NMR spectrum gave two cross-peaks at $\delta_{171}_{Yb} = -83 \text{ ppm } (\delta_{1H} = 3.53 \text{ and } 7.02 \text{ ppm}), \text{ showing the } 171 \text{Yb}$ couples to both the anilido proton and the Tripp 3,5-CH groups. The ¹H NMR spectra of paramagnetic 2M complexes were uninformative, see the ESI† for all NMR spectra.

Molecular structures

Single crystal X-ray diffraction studies on $\mathbf{1M}$ (M = Sc, Y, La, Tm) show all crystallize in the triclinic space group $P\bar{1}$ (Z'=1) and are pseudo three-coordinate – see Fig. 1. In $\mathbf{1La}$, the metal is sandwiched almost equally between two Tripp groups ($\text{La}\cdots C_{6\text{-centroid}}=2.8714(12)$ and 2.8783(12) Å; $\text{La}\cdots N_2I$ plane deviation = 0.0042(15) Å). Conversely, $\mathbf{1Sc}$ and $\mathbf{1Y}$ have a single Tripp group close to the metal ($\mathbf{M}\cdots C_{6\text{-centroid}}$: $\mathbf{1Sc}$, 2.3391(8) Å; and $\mathbf{1Y}$, 2.4949(9) Å) and hence are trigonal pyramidal. The structure of $\mathbf{1Y}$ is comparable to recently reported $[\mathbf{Y}(\mathrm{NHAr}^{\mathrm{iPr_6}})_2(\mathrm{Cl})]^{.38}$ See the ESI† for the structure of $\mathbf{1Tm}$, and a comparison of $\mathbf{1M}$ complexes. In the following sections, we shall continue to use the term " $C_{6\text{-centroid}}$ ", noting it is not strictly appropriate to define a centroid for non-planar groups.

The molecular structures of **2Sc**, **2Y**, and **2La** are also shown in Fig. 1 (see ESI† for **2Sm**, **2Eu**, **2Tm**, and **2Yb**). With the exception of **2Sc**, all **2M** complexes (M = Y, La, Sm, Eu, Tm, Yb) show two Tripp groups closely approaching the metal. When viewed along the $C_{6\text{-centroid}} \cdots C_{6\text{-centroid}}$ axis these groups are either fully eclipsed due to crystallographic C_2 symmetry (**2Y**, **2Eu**, and **2Yb**), or are pseudo-eclipsed (**2Sc**, **2La**, **2Sm**, and **2Tm**). In **2Sc**, only one Tripp group is close to the metal. The nature of this metal–arene interaction provides insight into the electronic structure vis-avis metal- or ligand-centred

$$\begin{array}{c} \textbf{A} \\ \textbf{MI}_3(\text{THF})_n + 2 \text{ KNHAr}^{\text{IP} \text{f}} \\ \textbf{M} = \text{La}, \, n = 4 \\ \textbf{M} = \text{Y}, \, \text{Tm}, \, n = 3.5 \\ \textbf{M} = \text{Sc}, \, n = 3 \\ \textbf{B} \\ \\ \textbf{B} \\ \textbf{MI}_2(\text{THF})_n + 2 \text{ KNHAr}^{\text{IP} \text{f}} \\ \textbf{M} = \text{Sm}, \, \text{Eu}, \, \text{Yb} \\ \end{array} \qquad \begin{array}{c} \textbf{Et}_2 \textbf{O} \\ \textbf{+ 1.2 KC}_8 \\ \textbf{+ 1.2 KC}_8 \\ \textbf{- KI} \\ \textbf{$$

Scheme 1 Synthesis of (Route A) [M(NHAr^{iPr₆})₂(I)] (**1M**, M = Sc, Y, La, Tm), and (Route B) [M(NHAr^{iPr₆})₂] (M = Sc, Y, La, Sm, Eu, Tm, Yb). Tripp = $\{C_6H_2-2,4,6-iPr_3\}$.

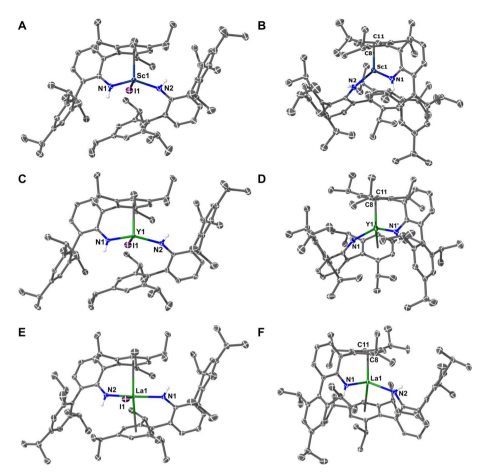


Fig. 1 Molecular structures: (A) 1Sc; (B) 2Sc; (C) 1Y; (D) 2Y; (E) 1La; (F) 2La. Thermal ellipsoids have been set at 40% probability. H-atoms except those on N–H groups, solvents of crystallization, and disordered components have been removed for clarity.

reduction. ^{45,46} Complexes **2Sc** and **2La** crystallize with a whole molecule in the asymmetric unit, and in both one metal-bound Tripp group is non-planar, showing an "open book" deformation for which a "hinge angle" (\angle arene) can be calculated – 11.43(11)° for **2Sc**, and 12.9(3)° for **2La**. In **2Sc**, the next shortest $M \cdots C_{6\text{-centroid}}$ distance (3.8304(7) Å) is too long to constitute a strong interaction; but, in **2La** the equivalent group is only *ca*. 0.4 Å further ($M \cdots C_{6\text{-centroid}} = 2.8348(12)$ Å) than the deformed arene ring, but is planar. Complex **2Sc** is similar to the Ti(v) analogue **2Ti**, ⁴⁶ but the latter exhibits \angle arene of 24.19(18)° and is diamagnetic by SQUID magnetometry, indicating the presence of a dianionic Tripp ring.

Complex 2Y is different to 2Sc and 2La as only half the molecule is present in the asymmetric unit (Z'=0.5), as previously reported;³⁸ and also to that of 2U.⁴⁷ The $\angle_{\rm arene}$ angle for both ligands in 2Y is $7.27(12)^{\circ}$, which compares well to $9.5(1)^{\circ}$ in 2U.⁴⁷ There is no clear trend in $\angle_{\rm arene}$ values (2Y < 2U < 2La < 2Sc) except that C_2 symmetric complexes (2Y and 2U) have smaller $\angle_{\rm arene}$ angles,^{38,47} and that larger values (with a single deformed arene) correlate with a greater degree of metal electron localisation (see below). The Gd(0) complex, [Gd(C₆H₃-1,3,5-tBu)₂], also displays two symmetry-equivalent distorted arene rings ($\angle_{\rm arene}=3.1(3)^{\circ}$).³³ The remaining divalent complexes, 2Sm, 2Eu, 2Tm, and 2Yb, are analogous to 2La,

except only the latter shows arene deformation. Table 1 summarizes all **2M** complexes now reported (M = Sc, Y, La, Sm, Eu, Tm, Yb, U).^{38,47}

UV-Vis-NIR spectroscopy

UV-Vis-NIR spectra were collected for **1M** (M = Sc, Y, La, Tm) and **2M** (M = Sc, Y, La, Sm, Eu, Tm, Yb) at ambient temperature as 1 mM solutions in Et₂O (Fig. 2, except for **1Tm**). The spectra of **1Sc**, **1Y**, and **1La** are uninformative, showing only a broad ligand-to-metal charge transfer (LMCT) process from ca. 20 000 cm⁻¹ (500 nm) to well into the UV region, accounting for the intense yellow colour of all three in solution. The colours and spectra of **1Tm** and **2Sm**, **2Eu**, **2Tm**, and **2Yb** are typical for these elements in their respective oxidation states (see ESI†). ^{13,14,48-51}

The spectrum of **2Sc** shows three well-resolved absorptions at 13 400 cm⁻¹ (756 nm, 805 M⁻¹ cm⁻¹), 17 400 cm⁻¹ (575 nm, 1727 M⁻¹ cm⁻¹), and 21 300 cm⁻¹ (469 nm, 2471 M⁻¹ cm⁻¹). The modest intensity of these peaks coupled with their energies suggests 3d \rightarrow 3d transitions and/or metal-to-ligand charge transfer (MLCT) bands. For **2Y**, two peaks are resolved at 13 700 cm⁻¹ (729 nm, 594 M⁻¹ cm⁻¹) and 21 600 cm⁻¹ (463 nm, 1787 M⁻¹ cm⁻¹), while a third tails in from above 26 000 cm⁻¹ (385 nm), in agreement with the previous report.³⁸ Finally, in **2La**, a single clear peak is resolved at 12 300 cm⁻¹ (816 nm,

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2.723(3)-2.870(3) 2.330(2)2.731(3) 2.723(3) 2.840(6)-3.047(6) $2.294(6)^{b}$ 2.8015(19)-2.971(2) 2.819(2)-3.118(2) 2.3169(18)2.3060(17)2.972(4)-3.176(5) $2.414(5)^{b}$ 2.955(3)-3.160(3) 2.953(3)-3.201(3) Bond lengths (Å) and angles (°) for [M(NHAr^{iPc}₅₎₂] (**2M**, M = Sc, Y, La, Sm, Eu, Tm, Yb, U)^{38,47} 2.412(2)2.425(2)2.778(16)-2.971(9) 3.047(3)-3.240(3) 2.843(13)2.903(12)2.434(3)2.395(3)2.7276(14)-2.9273(15) 2.7684(14)2.7859(15)2.2600(12)2.3913(12)-2.6304(14) 2.0678(10) 2.3913(12) 2.5418(13)2.0884(11)(1.43(11))Ring(1) Ring(2) C(8) C(11) Arene fold Fable 1 (Å or °) angle M-N

The solid-state structure is C_2 -symmetric so there is a single metal and ligand per asymmetric unit. ^b The solid-state structures show two half-molecules in the asymmetric unit, so M(1) and M(2)

each have only one unique ligand.

 $\leftarrow \lambda / nm$ 1,250 833 625 500 417 1Sc 3,000 1Y 1La 2,250 1,500 750 0 2Sc 3,000 2Y E / M-1 cm-1 2La 2,250 1,500 750 3,000 2Sm 2Eu 2,250 2Tm 2Yb 1,500 750 8,000 12,000 16,000 20,000 24,000 $E / cm^{-1} \rightarrow$ Fig. 2 Solution UV-Vis-NIR spectra of $[M(NHAr^{iPr_6})(I)]$ (1M, M = Sc, Y,

La; top) and $[M(NHAr^{iPr_6})]$ (2M, middle, M = Sc, Y, La; bottom, M = Sm, Eu, Tm, Yb) - as 1 mM solutions in Et₂O between 7000-26 000 cm⁻ (1429-385 nm) at ambient temperature

650 M^{-1} cm⁻¹), which we suggest is a 5d \rightarrow 5d transition. A broad feature at ca. 16 000 cm⁻¹ and a peak with a maximum at ca. 24 000 cm⁻¹ (417 nm) can also be seen, but background absorption precludes accurately describing these.

SQUID magnetometry

Direct current (DC) magnetic susceptibility data were collected for **1Tm**, and **2M** (M = Sc, Y, La, Sm, Eu, Tm) from 1.8 to 300 K under an applied field of 1 kOe. At 300 K, the $\chi_{\rm M}T$ ($\chi_{\rm M}$ is the molar magnetic susceptibility) values for 1Tm (6.61 cm³ mol⁻¹ K), **2Eu** $(7.27 \text{ cm}^3 \text{ mol}^{-1} \text{ K})$, and **2Tm** $(2.47 \text{ cm}^3 \text{ mol}^{-1} \text{ K})$ closely match theoretical values for $4f^{n+1}$ configurations for Tm(III), 4f¹² (³H₆, 7.15 cm³ mol⁻¹ K), Eu(II), 4f⁷ (⁸S_{7/2}, 7.88 cm³ mol⁻¹ K), and Tm(II), $4f^{13}(^{2}F_{7/2}, 2.57 \text{ cm}^{3} \text{ mol}^{-1} \text{ K})$. For $2Sm(4f^{6})$ the value at 300 K is 1.43 cm³ mol⁻¹ K, which is in the range observed for Eu(III) $(4f^6)$ complexes $(1.3 \text{ to } 1.5 \text{ cm}^3 \text{ mol}^{-1} \text{ K})$, ⁵² and is non-zero due to population of excited ⁷F_I states. ⁴⁸⁻⁵⁰ Upon cooling to 1.8 K, $\chi_{\rm M}T$ for **2Sm** lowers to 0.02 cm³ mol⁻¹ K which is consistent with a ⁷F₀ ground state; thus, a 4f⁶ configuration.

Complexes 2Sc, 2Y and 2La exhibit $\chi_M T$ values at 300 K (0.31, 0.36, and 0.29 cm³ mol⁻¹ K respectively), which are in reasonable agreement with the spin-only value for an S = 1/2 system $(0.375 \text{ cm}^3 \text{ mol}^{-1} \text{ K for } g = 2.00)$, and hence with a formal d¹ configuration. In each case, the magnetic moment is essentially invariant with temperature down to 8-10 K, where a sudden

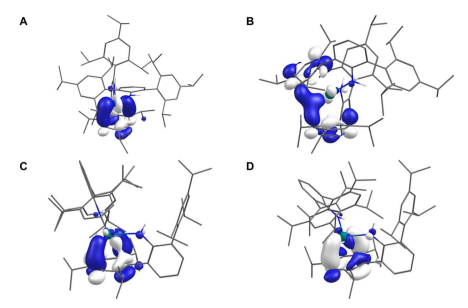


Fig. 3 SOMOs of: (A) 2Sc; (B) 2Y; (C) 2La; (D) 2Y-Et₂O (isovalues = 0.05) using geometries derived from single crystal X-ray diffraction with Hatoms optimised (2Sc, 2Y), select disordered C-atoms and all H-atoms optimised (2La), or all atom positions optimised (2Y-Et₂O). H-atoms except those of the N(H) group omitted for clarity.

drop can be seen, though this varies across independently synthesised samples (see ESI† for more details).

Electronic structure calculations

Unrestricted Kohn-Sham density-functional theory (DFT) calculations were performed on 2Sc, 2Y, 2La, 2Tm (S = 1/2), 2Sm (S=3), **2Eu** (S=7/2), and **2Yb** (S=0) using partially geometryoptimised structures (see ESI† for full details). Löwdin population and spin analyses of 2Sm, 2Eu, 2Yb, and 2Tm are consistent with experimental data and describe all four as $4f^{n+1}$ Ln(II) ions with metal-localised valence electrons. Fig. 3A and C shows the SOMOs of 2Sc and 2La, which depict M-arene δ-bonds comprised of 36% 3d (with 4% 4s) and 14% 5d (with 10% 4f and 1% 6s) metal character, respectively, with the bound Tripp ring making up 41% (2Sc) and 56% (2La) - the remainder is diffused over the rest of the molecule. Defining the M···Tripp direction as z, the d-orbital contribution is described as $d_{(xy/x^2-y^2)}$ (the superficial resemblance to d_{x^2} or d_{y^2} is due to a small degree of d_{z^2} mixing in this axis system) – this is comparable to the $e_{2\mathrm{g}}$ MO of bis-benzene transition metal complexes in D_{6h} symmetry.⁵³

In the case of 2Y (Fig. 3B), the SOMO is delocalised across the symmetry equivalent metal-bound Tripp groups such that the SOMO composition is 14% Y (12% 4d, 0.5% 5s), while the bound Tripp rings sum to 64.6% - the SOMO resembles a delocalised δ -bonding interaction. These results agree with previous work on 2Y,38 but it is an outlier compared to 2Sc and **2La.** Full geometry optimisation of **2Y** using the lower-symmetry structure of 2La as the starting point was performed in the gas phase and using an Et₂O solvent model (2Y-Et₂O henceforth). Both calculations produced geometries that have only one metal-bound Tripp group deformed (i.e. like 2Sc and 2La) and are true local minima on the potential energy surface. Fig. 3D shows the SOMO of 2Y-Et₂O, and Löwdin population analysis

shows it to be more metal-localised (22% 4d, 2% 5s, and 1% 4f total 25%) than in the C_2 -symmetric 2Y (14%), which is accompanied by a corresponding decrease in Tripp

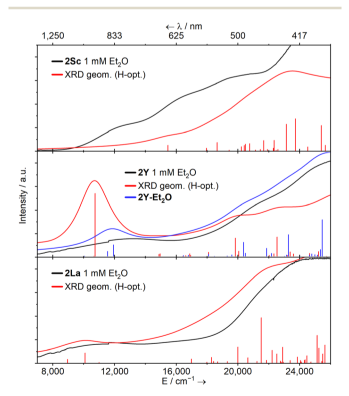


Fig. 4 Experimental (1 mM, Et₂O, black), calculated sTD-DFT transitions (red and blue vertical lines), and simulated spectra (red and blue solid lines, with Gaussian broadening and a linewidth factor of $25 \times \sqrt{E}$) UV-Vis-NIR spectra for complexes 2Sc, 2Y (and 2Y-Et₂O) and 2La calculated transitions were performed using a solvent model accounting for the dielectric constant (ε) and refractive index (n_D) of Et₂O.

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contributions to the SOMO - 65% in 2Y (over two Tripp groups) and 56% in 2Y-Et₂O (over a single Tripp group). The Löwdin spin populations at the metal in 2Y-Et₂O (0.245) and 2Y (0.142) reflect these differences.

Time-dependent DFT (TD-DFT) and simplified TD-DFT (sTD-DFT) calculations were employed to model the UV-Vis-NIR spectra of 2Sc, 2Y, 2La, and 2Y-Et₂O; here we focus on sTD-DFT with TPSSh for consistency with prior art (Fig. 4),11,12,16,23,54-56 see ESI (Fig. S102-S111†) for more details. Experimental features of 2Sc and 2La are well represented, and the Natural Transition Orbitals (NTOs) suggests the broad features in the spectrum of **2Sc** are $3d \rightarrow 3d$ transitions, the lowest energy of which resembles a $3d_{(xy/x^2-y^2)} \rightarrow 3d_{(x^2-y^2/xy)}$ transition maintaining the δ bonding interaction. For **2La**, the lowest energy feature is a $5d_{(xy)}$ x^2-y^2 \rightarrow 5d (x^2-y^2/xy) transition, and the next two lowest energy features are a combination of MLCT and 5d \rightarrow 5d transitions. There is poor agreement with all methods for 2Y (Fig. 4 middle panel, red line), however, calculations for 2Y-Et₂O are substantially better (Fig. 4 middle panel, blue line); this suggests that the structure of 2Y in Et₂O solution is similar to the solid-state structures of 2Sc and 2La. The lowest energy feature in 2Y-Et2O is comprised of two components, a $4d_{(xy/x^2-y^2)} \rightarrow 4d_{(x^2-y^2/xy)}$ transition and an MLCT process.

Complete active space self-consistent field (CASSCF) calculations (see ESI†) on 2M confirm the DFT and experimental results: 2Sc, 2Y and 2La exhibit nd1 ground states with significant orbital mixing with the arene ligand(s), and 2Sm, 2Eu, 2Tm and 2Yb have 4fⁿ⁺¹ ground states. Calculations for the lowestlying excitations in 2Sc, 2Y and 2La including multiconfigurational pair-density functional theory (MC-PDFT) corrections for dynamic correlation show nd/arene → nd/ arene excitations in the UV-Vis-NIR range, in good agreement with the experimental spectra. The character of these excitations is broadly in line with that found using (s)TD-DFT, where the lowest-lying excitations for 2Sc are mostly localised to one side of the molecule and resemble $d \rightarrow d$ transitions, while some for 2La and 2Y-Et₂O are combined MLCT and 5d → 5d transitions to the opposite Tripp ring.

EPR spectroscopy

Continuous wave (c. w.) EPR spectroscopy was used to study the orbitally non-degenerate species 2Sc, 2Y and 2La. All three are EPR active as polycrystalline solids and in solution (1 mM Et₂O or nPr₂O); spectra of 2Y in Et₂O have been reported previously.³⁸ We find better resolved frozen solution spectra in nPr_2O than in Et₂O (although the spectra are consistent; see ESI†).

X-band spectra of powders at room temperature show features around g = 2, consistent with the formal M(II) oxidation states. There is partial resolution of the metal hyperfine for 2Sc $(^{45}\text{Sc}, I = 7/2, 100\% \text{ abundant})$ and **2Y** $(^{89}\text{Y}, I = 1/2, 100\%)$ but is unresolved for 2La (139 La, I = 7/2, 100%). For 2Sc, a hyperfine octet is observed, with g = 2.000 and A = 145 MHz, for 2Y we observe a hyperfine doublet (estimated A = 14 MHz) with $g_{\perp} =$ 2.005 and $g_{\parallel}=1.995$, and for **2La**, we observe $g_{\perp}=$ 2.019 and $g_{\parallel}=1.958$. There are small changes in these parameters upon cooling to 5 K, without any improvement in resolution; the limited resolution of the powder spectra is indicative of intermolecular magnetic interactions.

Fluid solution spectra of 2Y and 2La (Fig. 5 and Table 2) give a hyperfine doublet ($A_{iso} = 46$ MHz, $g_{iso} = 1.9995$) and octet $(A_{\rm iso} = 112 \text{ MHz}, g_{\rm iso} = 1.998)$, respectively. For **2Sc** (Fig. 5 and Table 2) we also obtain an octet ($A_{iso} = 205$ MHz, $g_{iso} = 1.989$), but there is a second minor octet spectrum which differs subtly in the magnitude of the hyperfine ($A_{iso} = 186$ MHz), suggesting

Table 2 Experimental (frozen solution) and calculated EPR parameters for 2Sc, 2Y and 2La. Calculated hyperfine coupling constants in MHz; experimental A_{iso} calculated as $(A_{\parallel} + 2A_{\perp})/3$. The calculated g_1/A_1 axes are along the M-bound arene direction

		g_1	g_2	g_3	A_1	A_2	A_3	A_{iso}
2Sc	Exp.	1.990	2.002		195	210		205
	Calc.	1.990	2.009	2.015	159	171	184	171
2Y	Exp.	1.986	2.004		-36	-39		-38
2Y-Et ₂ O	Calc.	1.983	2.004	2.006	-47	-49	-50	-48
2La	Exp.	1.952	2.005	100	110		107	
	Calc.	1.954	1.971	1.993	101	106	108	105

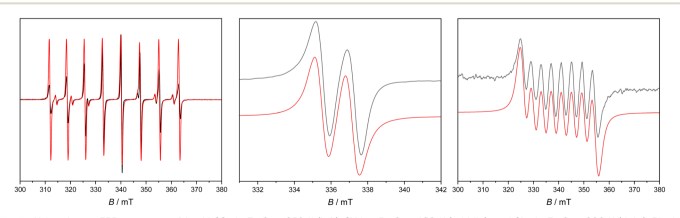


Fig. 5 X-band c. w. EPR spectrum of 1 mM 2Sc in Et₂O at 250 K (left), 2Y in nPr₂O at 180 K (middle), and 2La in Et₂O at 200 K (right). Black, experimental; red, simulations with parameters in the text.

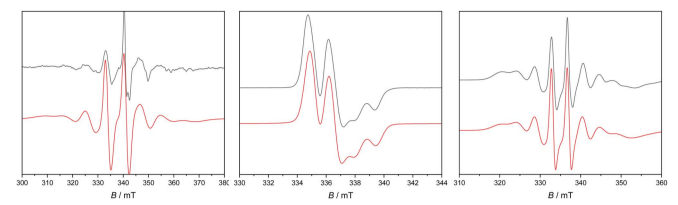


Fig. 6 X-band c. w. EPR spectrum of 1 mM 2Sc (left) in Et_2O at 130 K, 2Y (middle) and 2La (right) in nPr_2O at 60 K. Black experimental, red simulations.

two Sc(II) species in solution with a relative abundance of ca. 12:1 (similar features have been observed recently in a different Sc(II) system²³). In each case $g_{\rm iso} < g_{\rm e}$, consistent with the formal d¹ configuration. The isotropic part of the hyperfine interaction derives from s-orbital spin density, and from theoretical values of the hyperfine interaction for unit population of the valence s-orbitals⁵⁷ we estimate 7.3% (2Sc), 3.7% (2Y) and 2.0% (2La) s-orbital character of the SOMO; these are in good agreement with DFT calculations (3.6%, 1.8%, and 1.0% s-orbital character, or 3.8%, 2.0%, 1.1% Löwdin s-orbital spin populations). In summary, 2Sc has the largest metal valence s-orbital spin density, then 2Y > 2La.

Frozen solutions gave well-resolved spectra in each case (Fig. 6 and Table 2). For **2Sc** there is a dominant perpendicular hyperfine coupling $A_{\perp}\approx 210$ MHz ($g_{\perp}=2.002$), from which we can determine $A_{\parallel}\approx 195$ MHz (using $A_{\rm iso}=205$ MHz from the fluid spectra), and by simulation we find $g_{\parallel}=1.99$. However, these parameters are not well defined as there is evidence of a second species. For **2Y** and **2La** the spectra appear axially symmetric giving $g_{\perp}=2.004$, $g_{\parallel}=1.986$, with a near isotropic metal hyperfine of $A_{\perp}=-39$, $A_{\parallel}=-36$ MHz for **2Y**, while for **2La** we find $g_{\perp}=2.005$, $g_{\parallel}=1.952$ with $A_{\perp}=110$, $A_{\parallel}=100$ MHz.

The anisotropy of the g-values for 2Sc, 2Y and 2La is due to the significant d-orbital character of the SOMO. The $g_{\perp} > g_{\parallel}$ pattern indicates a dominant $d_{x^2-y^2}$ contribution (where z is the M-bound arene direction), while the greater deviation of g_{\parallel} from g_e in the series 2La > 2Y > 2Sc is in keeping with greater spinorbit coupling for the heavier elements,58 and also the trend in the lowest energy excited states (see above). A simple analysis of the anisotropic metal hyperfine interaction to give the metal $d_{x^2-y^2}$ contribution to the SOMO $[A_{\parallel}-A_{\perp}=(-6/7)a^2P_d]$, where a is the $d_{x^2-y^2}$ coefficient of the SOMO and values of P_d are tabulated in ref. 57] gives 7.3% for 2Sc, 5.6% for 2Y and 4.9% for **2La.** DFT calculations give much larger d-orbital contributions of 36%, 20%, and 14%, respectively (see above), despite also showing reasonable agreement with the measured hyperfine coupling constants (Table 2), implying an inadequacy in the simple analysis above; we have previously noted similar discrepancies in related [M^{II}L₃]⁻ systems (M = Sc, Y, La, Lu).¹⁷

Indeed, the EPR parameters and electronic structures of 2Sc, 2Y and 2La contrast to those related d¹ [M^{II}L₃]⁻ species where the trigonal crystal field instead stabilises the d_{z^2} (defined by the C_3 axis) orbital, or a d/s hybrid giving rise to characteristic $g_{\perp} < g_{\parallel} \ (\approx g_{\rm e})$ patterns.^{2,16,17} The electronic structures of the present compounds have more in common with $[Sc(Cp^{ttt})_2]^{23}$, where DFT calculations give a $d_{x^2-v^2}$ -dominated SOMO, and a similar $g_{\perp} > g_{\parallel}$ pattern can be observed from the frozen solution EPR data. The hyperfine coupling in 2Y and 2La is much weaker than in the trigonal M(II) cyclopentadienyl species: for example, [Y(Cp^R)₃] with various substituents [e.g. $(Cp^R)_3 = Cp'_3$, Cp^t_3 , $\{Cp''_2(C_5H_5)\}$] have $|A_{iso}|$ = 98-130 MHz; 17,59,60 and $[La(Cp^R)_3]^ (Cp^R = Cp', Cp'', Cp^{tt})$ have $|A_{\rm iso}|=390$ –640 MHz.^{2,17,61} Hence, there is greater metal character in the SOMOs of [M(Cp^R)₃] than in 2M. Comparing the present compounds with more symmetric sandwich compounds, [Y(Cp^{iPr₅})₂] and [La(Cp^{iPr₅})₂] have larger magnitude $|A_{\rm iso}| = 505$ and 2000 MHz, respectively, 4 while curiously $[Sc(Cp^{ttt})_2]$ has smaller magnitude $|A_{iso}| = 83$ MHz,²³ although it has been reported that [Sc(Cp*)2] (not structurally authenticated) has $|A_{iso}| = 824 \text{ MHz.}^{23}$

Conclusions

In summary, we have reported a series of room-temperature stable crystalline divalent rare earth bis-tethered arene complexes of the form $[M(NHAr^{iPr_c})_2]$ (2M; M = Sc, Y, La, Sm, Eu, Tm, Yb). In the case of Sc and La, these represent the second examples of neutral divalent complexes with these elements and are amongst just a few in any charge state. All 2M complexes feature close metal–arene contacts, which in 2Sc, 2Y, and 2La, results in an "open book" arene deformation suggestive of charge transfer, whereas in 2Sm, 2Eu, 2Tm, and 2Yb the equivalent arene groups remain planar. SQUID magnetometry and UV-Vis-NIR spectroscopy demonstrate 2Sm, 2Eu, 2Tm, and 2Yb are examples of $4f^{n+1}$ ions, and quantum chemical calculations show that the metal-localised 4f orbitals do not interact with the arene π -orbitals.

Solution-phase c. w. EPR spectroscopy of **2Sc**, **2Y**, and **2La** are consistent with formal nd^1 ions, where the SOMO has $nd_{(x^2-y^2)}$

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character with delocalisation of the spin onto the Tripp groups. Quantum chemical and ab initio calculations further support this description and reveal mixing between metal $nd_{(x^2-y^2)}$ and arene- π orbitals to give δ -bonds, which explain the arene deformation in their structures.

The electronic structures of nd¹ 2Sc, 2Y, and 2La closely resemble those of bis-benzene transition metal complexes; and, going forward, we posit that these rare $nd_{(r^2-v^2)}$ configurations afford as-yet unexplored opportunities to tune the physicochemical properties of divalent rare earth ions with formal {5d/ 6s}1 valence electron configurations. This, along with work to probe the limits of the bis-{NHAr^{iPr₆}} framework to stabilise other divalent f-block ions, remains an active area of research in our laboratory.

Data availability

The following CCDC references contain the ESI† crystal data for this article: 1Sc (2266263), 1Y (2266261), 1La (2266235), 1Tm (2266255), 1Yb (2266257), 1Y^{β} (2295306), 1La^{β} (2295307), 2Sc (2266264), 2Y (2266262), 2La (2266236), 2Sm (2266243), 2Eu (2266244), 2Tm (2266256), 2Yb (2266258), and 3 (2282041). Raw experimental data (NMR, ATR-IR, UV-Vis-NIR, SQUID Magnetometry) and computational inputs/outputs can be found freely at DOI: https://doi.org/10.48420/25245760.

Author contributions

C. A. P. G. provided the original concept. R. E. M. synthesised and characterised the complexes. R. E. M. and C. A. P. G. collected, solved, and refined the single crystal XRD data. G. F. S. W. performed final refinement and validation of XRD data. J. A. S. performed SQUID magnetometry measurements and interpreted the data. T. H. performed EPR spectroscopy measurements, E. J. L. M. and D. C. supervised the EPR measurements and data interpretation. C. A. P. G. performed DFT calculations and interpretation, N. F. C. performed CASSCF and MC-PDFT calculations and interpreted the data. C. A. P. G. wrote the manuscript with contributions from all other authors.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We thank the Royal Society for a University Research Fellowship (URF\211271 to C. A. P. G., and URF\191320 to N. F. C.); and the UoM Department of Chemistry for the funding of a PhD studentship under EPSRC DTP (EP/W524347/1 to R. E. M., and EP/T517823/1 to T. H.). We acknowledge funding from the EPSRC (EP/K039547/1, EP/V007580/1, and EP/P001386/1 for NMR spectroscopy and X-ray diffraction). We also thank the EPSRC UK EPR National Research Facility including for access to the SQUID magnetometer (EP/W014521/1, EP/V035231/1, EPS033181/1). C. A. P. G. and N. F. C. also thank the

Computational Shared Facility at the University of Manchester for support. We thank Prof. Stephen Liddle for support (J. A. S.). Elemental analyses were performed at the UoM by Mr Martin Jennings and Ms Anne Davies. We are also grateful to Prof. Aaron Odom and Prof. Selvan Demir for scientific discussions and encouragement.

References

- 1 K. R. Meihaus, M. E. Fieser, J. F. Corbey, W. J. Evans and J. R. Long, J. Am. Chem. Soc., 2015, 137, 9855-9860.
- 2 J. Liu, L. E. Nodaraki, D. O. T. A. Martins, M. J. Giansiracusa, P. J. Cobb, J. Emerson-King, F. Ortu, G. F. S. Whitehead, G. K. Gransbury, E. J. L. McInnes, F. Tuna and D. P. Mills, Eur. J. Inorg. Chem., 2023, 26, e202300552.
- 3 M. R. MacDonald, J. W. Ziller and W. J. Evans, J. Am. Chem. Soc., 2011, 133, 15914-15917.
- 4 M. R. MacDonald, J. E. Bates, M. E. Fieser, J. W. Ziller, F. Furche and W. J. Evans, J. Am. Chem. Soc., 2012, 134, 8420-8423.
- 5 M. R. MacDonald, J. E. Bates, J. W. Ziller, F. Furche and W. J. Evans, J. Am. Chem. Soc., 2013, 135, 9857-9868.
- 6 M. E. Fieser, M. G. Ferrier, J. Su, E. Batista, S. K. Cary, J. W. Engle, W. J. Evans, J. S. Lezama Pacheco, S. A. Kozimor, A. C. Olson, A. J. Ryan, B. W. Stein, G. L. Wagner, D. H. Woen, T. Vitova and P. Yang, Chem. Sci., 2017, 8, 6076-6091.
- 7 M. T. Trinh, J. C. Wedal and W. J. Evans, Dalton Trans., 2021, 50, 14384-14389.
- 8 T. F. Jenkins, D. H. Woen, L. N. Mohanam, J. W. Ziller, F. Furche and W. J. Evans, Organometallics, 2018, 37, 3863-3873.
- 9 A. J. Ryan, L. E. Darago, S. G. Balasubramani, G. P. Chen, J. W. Ziller, F. Furche, J. R. Long and W. J. Evans, Chem.-Eur. J., 2018, 24, 7702-7709.
- 10 S. A. Moehring, M. Miehlich, C. J. Hoerger, K. Meyer, J. W. Ziller and W. J. Evans, Inorg. Chem., 2020, 59, 3207-
- 11 D. N. Huh, S. R. Ciccone, S. Bekoe, S. Roy, J. W. Ziller, F. Furche and W. J. Evans, Angew. Chem., Int. Ed., 2020, 59, 16141-16146.
- 12 L. M. Anderson-Sanchez, J. M. Yu, J. W. Ziller, F. Furche and W. J. Evans, Inorg. Chem., 2023, 62, 706-714.
- 13 M. E. Fieser, M. R. MacDonald, B. T. Krull, J. E. Bates, J. W. Ziller, F. Furche and W. J. Evans, J. Am. Chem. Soc., 2015, 137, 369-382.
- 14 K. R. McClain, C. A. Gould, D. A. Marchiori, H. Kwon, T. T. Nguyen, K. E. Rosenkoetter, D. Kuzmina, F. Tuna, R. D. Britt, J. R. Long and B. G. Harvey, J. Am. Chem. Soc., 2022, 144, 22193-22201.
- 15 P. W. Smith, J. Hrubý, W. J. Evans, S. Hill and S. G. Minasian, J. Am. Chem. Soc., 2024, 146, 5781-5785.
- 16 K. Kundu, J. R. K. White, S. A. Moehring, J. M. Yu, J. W. Ziller, F. Furche, W. J. Evans and S. Hill, Nat. Chem., 2022, 14, 392-
- 17 A.-M. Ariciu, D. H. Woen, D. N. Huh, L. E. Nodaraki, A. K. Kostopoulos, C. A. P. Goodwin, N. F. Chilton,

- E. J. L. McInnes, R. E. P. Winpenny, W. J. Evans and F. Tuna, *Nat. Commun.*, 2019, **10**, 3330.
- 18 C. A. Gould, K. R. McClain, J. M. Yu, T. J. Groshens, F. Furche, B. G. Harvey and J. R. Long, *J. Am. Chem. Soc.*, 2019, 141, 12967–12973.
- 19 P.-B. Jin, Q.-C. Luo, G. K. Gransbury, I. J. Vitorica-Yrezabal, T. Hajdu, I. Strashnov, E. J. L. McInnes, R. E. P. Winpenny, N. F. Chilton, D. P. Mills and Y.-Z. Zheng, *J. Am. Chem. Soc.*, 2023, 145, 27993–28009.
- 20 C. A. Gould, K. R. McClain, D. Reta, J. G. C. Kragskow, D. A. Marchiori, E. Lachman, E. S. Choi, J. G. Analytis, R. D. Britt, N. F. Chilton, B. G. Harvey and J. R. Long, *Science*, 2022, 375, 198–202.
- 21 J. C. Wedal, L. M. Anderson-Sanchez, M. T. Dumas, C. A. Gould, M. J. Beltran-Leiva, C. Celis-Barros, D. Paez-Hernandez, J. W. Ziller, J. R. Long and W. J. Evans, *J. Am. Chem. Soc.*, 2023, **145**, 10730–10742.
- 22 K. R. McClain, H. Kwon, K. Chakarawet, R. Nabi, J. G. C. Kragskow, N. F. Chilton, R. D. Britt, J. R. Long and B. G. Harvey, J. Am. Chem. Soc., 2023, 145, 8996–9002.
- 23 J. D. Queen, L. M. Anderson-Sanchez, C. R. Stennett, A. Rajabi, J. W. Ziller, F. Furche and W. J. Evans, *J. Am. Chem. Soc.*, 2024, **146**, 3279–3292.
- 24 J. T. Boronski, J. A. Seed, D. Hunger, A. W. Woodward, J. van Slageren, A. J. Wooles, L. S. Natrajan, N. Kaltsoyannis and S. T. Liddle, *Nature*, 2021, 598, 72–75.
- 25 M. D. Straub, E. T. Ouellette, M. A. Boreen, R. D. Britt, K. Chakarawet, I. Douair, C. A. Gould, L. Maron, I. Del Rosal, D. Villarreal, S. G. Minasian and J. Arnold, *J. Am. Chem. Soc.*, 2021, 143, 19748–19760.
- 26 J. D. Cryer and S. T. Liddle, in *Comprehensive Organometallic Chemistry IV*, ed. G. Parkin, K. Meyer and D. O'Hare, Elsevier, Oxford, 2022, pp. 460–501.
- 27 J. Murillo, R. Bhowmick, K. L. M. Harriman, A. Gomez-Torres, J. Wright, R. W. Meulenberg, P. Miro, A. Metta-Magana, M. Murugesu, B. Vlaisavljevich and S. Fortier, *Chem. Sci.*, 2021, 12, 13360–13372.
- 28 R. A. Keerthi Shivaraam, M. Keener, D. K. Modder, T. Rajeshkumar, I. Zivkovic, R. Scopelliti, L. Maron and M. Mazzanti, Angew. Chem., Int. Ed., 2023, 62, e202304051.
- 29 M. Keener, R. A. K. Shivaraam, T. Rajeshkumar, M. Tricoire, R. Scopelliti, I. Zivkovic, A. S. Chauvin, L. Maron and M. Mazzanti, *J. Am. Chem. Soc.*, 2023, **145**, 16271–16283.
- 30 F.-C. Hsueh, T. Rajeshkumar, L. Maron, R. Scopelliti, A. Sienkiewicz and M. Mazzanti, *Chem. Sci.*, 2023, 14, 6011–6021.
- 31 Y. Wang, J. Liang, C. Deng, R. Sun, P.-X. Fu, B.-W. Wang, S. Gao and W. Huang, *J. Am. Chem. Soc.*, 2023, **145**, 22466– 22474.
- 32 H. S. La Pierre, A. Scheurer, F. W. Heinemann, W. Hieringer and K. Meyer, *Angew. Chem., Int. Ed.*, 2014, **53**, 7158–71562.
- 33 J. G. Brennan, F. G. N. Cloke, A. A. Sameh and A. Zalkin, *J. Chem. Soc., Chem. Commun.*, 1987, 1668–1669.
- 34 D. M. Anderson, F. G. N. Cloke, P. A. Cox, N. Edelstein, J. C. Green, T. Pang, A. A. Sameh and G. Shalimoff, J. Chem. Soc., Chem. Commun., 1989, 53–55.

- 35 W. A. King, T. J. Marks, D. M. Anderson, D. J. Duncalf and F. G. N. Cloke, *J. Am. Chem. Soc.*, 1992, 114, 9221–9223.
- 36 F. G. N. Cloke, Chem. Soc. Rev., 1993, 22, 17-24.
- 37 W. A. King, S. Di Bella, G. Lanza, K. Khan, D. J. Duncalf, F. G. N. Cloke, I. L. Fragala and T. J. Marks, *J. Am. Chem. Soc.*, 1996, 118, 627–635.
- 38 R. Jena, F. Benner, F. Delano, D. Holmes, J. McCracken, S. Demir and A. L. Odom, *Chem. Sci.*, 2023, **14**, 4257–4264.
- 39 P. Dorenbos, J. Condens. Matter Phys., 2003, 15, 575-594.
- 40 P. Dorenbos, J. Lumin., 2000, 91, 91-106.
- 41 B. R. Barnett, C. C. Mokhtarzadeh, P. Lummis, S. Wang, J. D. Queen, J. Gavenonis, N. Schüwer, T. D. Tilley, J. N. Boynton, N. Weidemann, D. W. Agnew, P. W. Smith, T. B. Ditri, A. E. Carpenter, J. K. Pratt, N. D. Mendelson, J. S. Figueroa and P. P. Power, in *Inorganic Syntheses*, 2018, pp. 85–122.
- 42 B. Twamley, C.-S. Hwang, N. J. Hardman and P. P. Power, *J. Organomet. Chem.*, 2000, **609**, 152–160.
- 43 K. Izod, S. T. Liddle and W. Clegg, *Inorg. Chem.*, 2004, 43, 214–218.
- 44 P. Girard, J. L. Namy and H. B. Kagan, *J. Am. Chem. Soc.*, 1980, **102**, 2693–2698.
- 45 S. R. Chowdhury, C. A. P. Goodwin and B. Vlaisavljevich, *Chem. Sci.*, 2024, **15**, 1810–1819.
- 46 J. N. Boynton, J. D. Guo, F. Grandjean, J. C. Fettinger, S. Nagase, G. J. Long and P. P. Power, *Inorg. Chem.*, 2013, 52, 14216–14223.
- 47 B. S. Billow, B. N. Livesay, C. C. Mokhtarzadeh, J. McCracken, M. P. Shores, J. M. Boncella and A. L. Odom, J. Am. Chem. Soc., 2018, 140, 17369–17373.
- 48 N. F. Chilton, C. A. P. Goodwin, D. P. Mills and R. E. P. Winpenny, *Chem. Commun.*, 2015, **51**, 101–103.
- 49 C. A. P. Goodwin, K. C. Joslin, S. J. Lockyer, A. Formanuik, G. A. Morris, F. Ortu, I. J. Vitorica-Yrezabal and D. P. Mills, *Organometallics*, 2015, 34, 2314–2325.
- 50 C. A. P. Goodwin, N. F. Chilton, G. F. Vettese, E. Moreno Pineda, I. F. Crowe, J. W. Ziller, R. E. P. Winpenny, W. J. Evans and D. P. Mills, *Inorg. Chem.*, 2016, 55, 10057– 10067.
- 51 C. A. P. Goodwin, N. F. Chilton, L. S. Natrajan, M.-E. Boulon, J. W. Ziller, W. J. Evans and D. P. Mills, *Inorg. Chem.*, 2017, 56, 5959–5970.
- 52 Y. Takikawa, S. Ebisu and S. Nagata, *J. Phys. Chem. Solids*, 2010, 71, 1592–1598.
- 53 V. M. Rayón and G. Frenking, *Organometallics*, 2003, 22, 3304–3308.
- 54 A. Rajabi, R. Grotjahn, D. Rappoport and F. Furche, *Dalton Trans.*, 2024, 53, 410–417.
- 55 M. E. Fieser, C. T. Palumbo, H. S. La Pierre, D. P. Halter, V. K. Voora, J. W. Ziller, F. Furche, K. Meyer and W. J. Evans, *Chem. Sci.*, 2017, 8, 7424–7433.
- 56 C. T. Palumbo, D. P. Halter, V. K. Voora, G. P. Chen, A. K. Chan, M. E. Fieser, J. W. Ziller, W. Hieringer, F. Furche, K. Meyer and W. J. Evans, *Inorg. Chem.*, 2018, 57, 2823–2833.
- 57 J. R. Morton and K. F. Preston, J. Magn. Reson., 1978, 30, 577–582.

- 58 S. Koseki, N. Matsunaga, T. Asada, M. W. Schmidt and M. S. Gordon, *J. Phys. Chem. A*, 2019, **123**, 2325–2339.
- 59 J. F. Corbey, D. H. Woen, C. T. Palumbo, M. E. Fieser, J. W. Ziller, F. Furche and W. J. Evans, *Organometallics*, 2015, 34, 3909–3921.
- 60 M. A. Angadol, D. H. Woen, C. J. Windorff, J. W. Ziller and W. J. Evans, *Organometallics*, 2019, 38, 1151–1158.
- 61 S. A. Moehring and W. J. Evans, *Organometallics*, 2020, 39, 1187–1194.