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Introduction

The chemistry of lanthanides in molecular complexes was for a longtime limited to the +3 oxidation state and to one-electron transfer reactions for the few accessible redox couples ($Ce^{3+/4+}$, $Sm^{2+/3+}$, $Eu^{2+/3+}$, and $Yb^{2+/3+}$).¹ Only in 2013 all lanthanide ions (except the radioactive promethium) have been isolated in the +II oxidation state in molecular complexes expanding significantly the knowledge of this oxidation state.^{1,2} Molecular species containing rare earths in the +I oxidation state remain elusive with only an example of formal Sc(1) reported 25 years ago,³ and so far the +I oxidation state in f element chemistry has been limited to a single U(1) complex⁴ or gas phase and solidstate species.^{5,6}

Isolation and redox reactivity of cerium complexes in four redox states[†]

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The chemistry of lanthanides is limited to one electron transfer reactions due to the difficulty of accessing multiple oxidation states. Here we report that a redox-active ligand combining three siloxides with an arene ring in a tripodal ligand can stabilize cerium complexes in four different redox states and can promote multielectron redox reactivity in cerium complexes. Ce(III) and Ce(IV) complexes [(LO₃)Ce(THF)] (1) and [(LO₃)CeCl] (2) (LO₃ = 1,3,5-(2-OSi(O^tBu)₂C₆H₄)₃C₆H₃) were synthesized and fully characterized. Remarkably the one-electron reduction and the unprecedented two-electron reduction of the tripodal Ce(III) complex are easily achieved to yield reduced complexes [K(2.2.2-cryptand)][(LO₃)Ce(THF)] (3) and [K₂{(LO₃)Ce(Et₂O)₃] (5) that are formally "Ce(III)" and "Ce(II)" analogues. Structural analysis, UV and EPR spectroscopy and computational studies indicate that in 3 the cerium oxidation state is in between +II and +III with a partially reduced arene. In 5 the arene is doubly reduced, but the removal of potassium results in a redistribution of electrons on the metal. The electrons in both 3 and 5 are stored onto δ -bonds allowing the reduced complexes to be described as masked "Ce(III)" and "Ce(I)". Preliminary reactivity studies show that these complexes act as masked Ce(III) and Ce(II) in redox reactions with oxidizing substrates such as Ag⁺, CO₂, I₂ and S₈ effecting both one- and two-electron transfers that are not accessible in classical cerium chemistry.

The redox chemistry of the cerium ion in particular has received large attention,⁷ which mostly focused on Ce(π)/Ce(π) oxidation chemistry because of its application in organic synthesis, catalysis and materials science. In contrast, mononuclear Ce(π) complexes remain limited to five examples shown in Fig. 1.^{2*a*,*c*,*f*,8,9} including four cyclopentadienide derivatives (complex **A** in Fig. 1) and the aryloxide complex (complex **B** in Fig. 1). The redox reactivity of these complexes has not been so far investigated.

Three additional examples of Ce in the +2 oxidation state have been identified in multimetallic arene sandwich compounds.¹⁰ In particular, inverse sandwich complexes of cerium(II) were isolated by reduction of Ce(III) cyclopentadienyl complexes.^{10*a,b*} Remarkably, the Evans group showed that the four electrons stored in the $[K(2.2.2\text{-cryptand})]_2[(Cp'_2\text{-Ce})_2(\mu-\eta_6:\eta_6\text{-C}_6\text{H}_6)]$ complex, best described as a Ce(II)-arene^{2–}–Ce(II) species,^{10*b*} become available for the reduction of substrates suggesting that the association of cerium and redox-active ligands may provide an entry into multielectron redox chemistry usually non-accessible for lanthanides. Besides these results, the use of redox-active ligands in low valent lanthanide chemistry remains poorly developed especially in comparison to d-block metals.¹¹

Meyer and coworkers reported tripodal ligands ((($(ArO)_{3}$ -mes)³⁻) that incorporate an arene ring as part of a potentially redox-active chelating ligand and showed that these ligand were able to promote unusual reactivity and stabilize the +2 oxidation



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state in f element chemistry.¹² More recently, Evans and Meyer showed that reduced lanthanide complexes of the tripodal $(((ArO)_3mes)^{3-})$ ligand could be isolated for all lanthanide ions^{2h,13} and they act as electrocatalysts for water reduction.^{13a} Interestingly, the ligand was found to stabilize the low valent oxidation state even for highly reducing Ln²⁺ ions such as Nd²⁺,^{2h} but could also act as a redox-active ligand for some lanthanide ions. In particular the reduced Ce complex of the $((^{Ad,Me}ArO)_3mes)^{3-}$ ligand was identified as Ce(III) with a negative charge localized on the arene anchor of the tripodal ligand (complex C in Fig. 1).^{13b}

Ligand-based C–H bond activation and loss of hydrogen can also occur under reducing conditions for the aryloxide-arene tripodal ligand ($(^{Ad,Me}ArO)_3mes)^{3-}$, to form a tetraanionic ligand (($(^{Ad,Me}ArO)_3mes)^{4-}$) leading to the formation of Ln(m) hydride complexes.^{13b} Such reactivity may limit the possibility of accessing higher degrees of Ln reduction.

In our group we showed that the *tert*-butoxy siloxide ligand can stabilize the cerium cation in different oxidation states (+IV to +II), but the reduction of Ce(IV) with excess KC₈ resulted in the partial loss of the supporting ligand leading to the isolation of the tetradecker sandwich complex [K(2.2.2-crypt)]₂[{(KL₃Ce)(μ - $\eta^6:\eta^6-C_7H_8)$ }₂Ce] (complex **D** in Fig. 1). Complex **D** contains Ce(II) in a 4f² configuration and doubly reduced toluene molecules involved in delta bonds with the 4f orbitals of cerium. Inspired by the molecular and electronic structures of complexes **C** and **D** we reasoned that combining three siloxides with an arene ring in a tripodal ligand should lead to the isolation of a series of cerium in different oxidation states and enable a novel route to controlled multi-electron redox chemistry in cerium compounds. Here we report a new ligand $1,3,5-(2-\text{HOSi}(O^t\text{Bu})_2\text{C}_6\text{H}_4)_3-C_6\text{H}_3$, and its cerium complexes and show that four different redox states are accessible for this system enabling one- and two-electron transfer reactions with various substrates.

Results and discussion

Tripodal ligand

The tripodal tris(di-*tert*-butoxylsilanol) arene ligand, 1,3,5-(2-HOSi(O^tBu)₂C₆H₄)₃C₆H₃, **L(OH)**₃, was prepared from commercial products on a gram scale in a three step synthesis (Schemes S1[†] and 1) with a procedure analogous to that reported for the tri-diphenylsiloxide arene ligand.¹⁴ Colorless crystals of the tripodal ligand (**L(OH)**₃) were isolated from hexane at room temperature by slow evaporation. The solid-state molecular structure of **L(OH)**₃ as determined by an X-ray diffraction study shows that the three silanol groups are connected by hydrogen bonds that hold the three silanols in the same plane (Fig. S79[†]).

Reduction with 1 equiv. KC_8 of the ligand salt $L(OK)_3$ obtained by deprotonation of L(OH)3 with KHMDS afforded the dark bluegreen potassium salt of the ligand radical [(K(THF)₃)(L(OK)₃)]₂ which was isolated in 66% yield and crystallographically characterized (see the ESI[†]). The reduction of [(K(THF)₃)(L(OK)₃)]₂ with additional KC₈ only led to decomposition products (Fig. S17[†]). The molecular structure of [(K(THF)₃)(L(OK)₃)]₂ shows the presence of a dimeric compound where two tetra-anionic radical tripodal moieties are held together by six potassium cations binding siloxide groups from the two molecules (Fig. S80[†]). The structure is completed by two potassium cations bound to the central arene in a η⁶-fashion. The C–C bond distances are slightly longer compared to the non-reduced ligand (L(OH)₃). We found that two potassium ions can be removed by addition of 2 equiv. of 2.2.2-cryptand to [(K(THF)₃)(L(OK)₃)]₂, yielding [(K(crypt))(L(OK)₃)]₂. The solid-state structure of [(K(crypt))(L(OK)₃)]₂ was determined by X-ray diffraction studies and it shows the same dimeric structure as $[(K(THF)_3)(L(OK)_3)]_2$, but the two arene- η^6 -bound K⁺ ions have been removed from the structure and are found as outer sphere [K(2.2.2-cryptand)] cations (Fig. 2).



Scheme 1 Synthesis of the tripodal ligand $L(OH)_3$ and of the ligand radical salt $[(K(crypt))(L(OK)_3)]_2$.

Tripodal Ce(III) and Ce(IV) complexes

At first, we pursued the synthesis of the cerium(m) complex of the ligand **L(OH)**₃ by protonolysis of the $[Ce(N(SiMe_3)_2)_3]$ precursor (Scheme 2). The addition of a THF solution of $[Ce(N(SiMe_3)_2)_3]$ (1 equiv.) to a stirring THF solution of the free ligand (**L(OH)**₃) (1 equiv.) at -40 °C led to the formation of a new species and full consumption of the starting materials as indicated by ¹H NMR spectroscopy after 16 hours (Fig. S18†). Light-yellow crystals of the complex, $[(LO_3)Ce(THF)]$ (1), were isolated from a concentrated toluene solution at -40 °C. The ¹H NMR spectrum of **1** in THF-*d*₈ displayed five paramagnetic shifted resonances (Fig. S19†).

We then investigated the possibility of stabilizing Ce in the +4 oxidation state using the tripodal siloxide framework, $L(OH)_3$. Despite the numerous applications of tetravalent cerium compounds in organic synthesis and catalysis,^{15,7j} the coordination and organometallic chemistry of cerium(iv) complexes remain still underdeveloped due to the high oxidizing power of Ce(rv).^{7a,h,16}

Complex 1 did not react with one equivalent of AgCl or AgBPh₄ at room temperature or 70 °C after 2 days, as shown by the ¹H NMR spectroscopy data (Fig. S44 and S45[†]). The high stability of 1 toward silver reagents compared to the tetrasiloxide complex $[KCe(OSi(O^tBu)_3)_4]^{10c}$ which can be easily converted into the Ce(IV) analogue is probably explained by the less electron-rich character of complex 1. Therefore, we pursued the synthesis of the Ce(IV) complex by an alternative route using a Ce(n) precursor. The protonolysis reaction of the heteroleptic $[CeCl(N(SiMe_3)_2)_3]^{17}$ complex with free ligand L(OH)₃ in toluene solution at room temperature for 3 hours (Scheme 2) resulted in an orange reaction mixture and full consumption of the starting material as indicated by ¹H NMR spectroscopy (Fig. S20[†]). Light-yellow crystals of complex $[(LO_3)CeCl]$ (2) were isolated from a concentrated toluene solution at -40 °C and characterized by X-ray crystallography. Complex 2 begins immediately to decompose back to 1 in THF at room temperature but is stable in toluene solution at room temperature for up to 2 weeks



Fig. 2 Molecular structure of the monomeric unit $L(OK)_3$ in $[(K(crypt))(L(OK)_3)]_2$, with thermal ellipsoids drawn at the 50% probability level. Hydrogen atoms, methyl groups on the $-OSi(O^tBu)_2$ ligands and [K(2.2.2-cryptand)] have been omitted for clarity.



Scheme 2 Synthesis of complexes 1 and 2.

(Fig. S22 and S23[†]). These results show that the tripodal LO_3^{3-} ligand allows the access of robust heteroleptic Ce(rv) complexes that provide attractive precursors for oxidative reactivity.

Low-valent tripodal Ce complexes

With the tripodal Ce(III) complex in hand, we set out to investigate if complex **1** could be further reduced and if lower oxidation states of cerium could be in reach in the environment provided by the tripodal siloxide ligand.

The reaction of **1** with 1 equiv. of KC_8 in THF at -40 °C resulted in a dark-blue reaction mixture, which displayed only one broad resonance at 1.59 ppm in the ¹H NMR spectrum at room temperature (Fig. S25†). Attempts to isolated X-ray suitable single crystals of this new species proved unsuccessful. However, the addition of one equivalent of 2.2.2-cryptand and one equivalent of KC₈ at -40 °C to **1** (Scheme 3) resulted in a new set of eight resonances in the ¹H NMR spectrum recorded at room temperature (Fig. S29†). Dark-blue crystals of the complex [K(2.2.2-cryptand)][(LO₃)Ce(THF)] (3) were isolated from a concentrated Et₂O/THF mixture at -40 °C and characterized by X-ray crystallography. Complex **3** is stable in THF at room temperature for at least for 1 week (Fig. S29†).

In complex 3 the apical THF ligand can be easily replaced by more coordinating ligands. Notably, the reaction of 1 equiv. of Et₃PO with a THF solution of 3 at room temperature (Scheme 3) resulted immediately in the replacement of the Ce-bound THF ligand and led to the isolation of dark-blue crystals of the complex [K(2.2.2-cryptand)][(LO₃)Ce(Et₃PO)] (4).

The ¹H NMR spectrum of isolated complex 4 recorded at room temperature displays two additional broad resonances at δ 1.55 and δ 1.06 ppm compared to 3 (Fig. S34†), which were assigned to the ethyl group in the bound Et₃PO. The ³¹P NMR spectrum of 4 exhibits a singlet resonance at δ 45.06 ppm (Fig. S35†), which is in the range typically observed for phosphine-oxide complexes.¹⁸

A one-electron reduced cerium tripodal complex was isolated previously by Evans and co-workers from the reduction of the tripodal Ce(m) complex [((^{Ad,Me}ArO)₃mes)Ce] with potassium in



Scheme 3 Synthesis of complex 3–5.

the presence of 2.2.2-cryptand.^{13b} Computational and spectroscopic studies led to the assignment of the complex as a $4f^1$ Ce(m) complex of the reduced tripodal ligand with a population of 1.3 electrons on Ce and 0.7 on the mesitylene ring. Further reduction or reactivity studies with small molecules of this complex were not reported.

In contrast further reduction of complex 3 was found to proceed in a controlled manner leading to a main reduction product.

The addition of two equivalents of KC8 to a THF solution of 1 at -40 °C led to a dark-purple reaction mixture (Scheme 3). The ¹H NMR spectrum showed that a new species was formed (Fig. S37[†]). Dark-purple crystals were isolated from a concentrated Et₂O/toluene mixture at -40 °C, and X-ray crystallography revealed a doubly reduced complex $[K_2{(LO_3)Ce(Et_2O_3)}]$ (5). The ¹H NMR spectrum of isolated complex 5 recorded at -40 °C shows 6 sharp and three broad resonances paramagnetically shifted, between -59.63 and 28.51 ppm (Fig. S39[†]). Complex 5 is stable in THF at room temperature for at least for 3 days (Fig. S40[†]). The addition of 2 equiv. of 2.2.2cryptand resulted in a change in the ¹H NMR spectrum at -40° C with only three broad resonances observed between 0.85 and 3.18 ppm (Fig. S43[†]) suggesting that the potassium cations are bound in solution and can be removed by cryptand. Attempts to isolate X-ray suitable single crystals of this species proved unsuccessful.

Complexes 3 and 5 can be formally identified as Ce(II) and Ce(I) species, but reduction could occur on the arene of the tripodal ligand as previously observed for some of the mono-reduced [((^{Ad,Me}ArO)₃mes)Ln] complexes.^{13b}

EPR and UV-vis spectroscopy

To investigate the electronic structure of complexes 1, 3, 4, and 5 X-band EPR spectra were recorded for all complexes and for the ligand radical $[(K(crypt))(L(OK)_3)]_2$ (Fig. 3 and S61[†]). Complex 1 exhibits a strong and distinct EPR signal with a major feature at $g \sim 1.78$ in the frozen-solution state at 6 K (Fig. 3) in agreement with the presence of an f¹ Ce(m) species.¹⁹

On the other hand, complex 3 exhibits a EPR signal weaker and broader than 1 in the solid-state at 6 K ($g_1 \sim 3.02$ and $g_2 \sim$ 2.31) (Fig. 3, S62 and S64[†]) which disappears at higher temperatures (50 K). Since f² Ce(II) would be EPR silent, the broad and weak nature of the signal for complex 3 could be due to an f¹d¹ configuration or to the presence of a strong interaction of f^1 Ce(m) with the ligand radical. Furthermore, the EPR spectrum of 3 also shows an isotropic signal with a g_{iso} of 2.0054 which is consistent with the presence of a ligand radical. A similar signal was also observed in the EPR spectrum recorded at 96 K reported for the tripodal complex [K(2.2.2-cryptand)] [((^{Ad,Me}ArO)₃mes)Ce]^{13b} and assigned to the presence of ligandradical character. The EPR spectrum recorded before the addition of cryptand is very similar to the EPR spectrum of 3 but more intense (Fig. S65[†]) suggesting that the electronic configuration is slightly affected by the presence of bound potassium. Moreover, the replacement of the axial THF ligand by Et₃OP in complex 4 also results in a EPR spectrum similar to the one recorded for 3 (Fig. S66[†]). Finally, complex 5 shows a strong



Fig. 3 X-band EPR spectra of complexes 1 (frozen-solution), 3 (solidstate), 5 (solid-state), and 5 + cryptand (frozen-solution) at 6 K.

signal in the X band solid-state EPR at 6 K which was best fitted as a rhombic signal with $g_y = 0.870$, $g_z = 3.360$ and $g_x < 0.5$ (g_x could not be observed because it occurs at magnetic fields outside our accessible magnetic range) (Fig. 3 and S68[†]). The observed spectrum is in agreement with the assignment of complex 5 as Ce(III)(ligand)²⁻. Similar rhombic EPR spectra were previously reported for Ce(III) complexes.²⁰ Overall the EPR spectrum of 5 seems to point toward the presence of a Ce(m)component. However, the EPR spectrum is very different after the addition of cryptands showing only a very weak and broad signal and a strong radical signal at g = 2.0054 (Scheme S2,[†] Fig. 3 and S71[†]). The spectrum suggests that, in the case of the doubly reduced complex, the removal of potassium results in a significant change in the electron configuration of the cerium cation and indicates the presence of Ce(II) and a mono-reduced arene.

In previous studies of cyclopentadienyl complexes of Ln²⁺ ions, UV-vis spectroscopy has been crucial for evaluating electronic configurations.^{2a,f} We therefore decided to record UV-vis spectra for complexes **1**, **3** and **5**.

Complex 1 only shows two intense absorption bands at 210 and 250 nm (Fig. S56[†]) assigned to the ligand (Fig. S55[†]). The mono-reduced complex 3 shows the ligand absorption at 210 and 245 nm and three strong absorbances at 375, 545 and 770 nm (Fig. S58[†]) with calculated molar extinction coefficients, ε , of 7077, 3294 and 4045 M⁻¹ cm⁻¹, respectively. These λ_{max} and ε values are similar to those reported for Ln^{2+} complexes which have a $4f^{n}5d^{1}$ configuration (ε values range from 3000-7000 M^{-1} cm⁻¹). However, the UV-vis spectrum of the tripodal ligand radical anion, [(K(crypt))(L(OK)₃)]₂ (Fig. S57[†]), exhibits similar strong absorbances at 405, 520, and 625 nm with molar extinction coefficients, ε , of 14 377, 4462 and 6727 M⁻¹ cm⁻¹. The doubly reduced complex 5 shows different features compared to complex 3 with two strong absorbances at 405 and 520 nm (Fig. S60[†]) with molar extinction coefficients, ε , of 5913 and 8681 M^{-1} cm⁻¹, which again could be assigned either to a different degree of ligand reduction or to a $4f^{n}5d^{1}$ metal configuration.

Cyclic voltammetry

Cyclic voltammetry experiments were conducted at ambient temperature in 3 mM THF solutions (0.1 M $[NBu_4][B(C_6H_5)_4]$ as the supporting electrolyte) of the ligand salt $L(OK)_3$ and on complexes 1, 3, and 5 (Fig. 4). Only one reversible redox event could be clearly identified for the three complexes. The voltammogram of complex 1 shows one chemically reversible reduction wave at $E_{1/2} = -2.90$ V, corresponding to the reduction to complexes 3. The voltammogram of complex 3 confirms this assignment and has very similar $E_{1/2}$ (-2.95 V). The measured half potential is very similar to that reported for the ligand-based one-electron reduction of the [((^{Ad,Me}ArO)₃mes)Ce] complex.13a The voltammogram of 5 shows the same redox event but at slightly shifted ($E_{1/2} = -2.81$ V) potential probably due to the presence of coordinating potassium cations in the complex, which was assigned to the one-electron reduction. Indeed, the voltammogram of 5 after the addition of cryptands shows



Fig. 4 Cyclic voltammograms of complexes 1 (red), 3 (blue), 5 (purple) and 5 + cryptand (dark-blue) in THF (0.1 M [NBu₄][BPh₄]) at a 250 mV s⁻¹ scan rate *versus* Fc/Fc⁺ (arrows indicate the scan direction).

a comparable $E_{1/2}$ value at -2.91 V (Fig. 4). The redox event corresponding to the reduction of 3 to 5 is not observed under the used conditions.

No reduction waves were observed within the window permitted by THF in the voltammogram of the ligand salt $L(OK)_3$ alone or in the presence of 3 equiv. of cryptand (Fig. S76 and S77[†]), suggesting that the redox events observed in the voltammograms of **1**, **3** and **5** may be assigned to a metal centered redox reduction (Ce(m)/Ce(n)).

Structural characterization of tripodal Ce complexes

The solid-state structures of complexes **1–5** were determined by X-ray diffraction studies. The molecular structures of complexes **1–5** are presented in Fig. 5 and the selected structural parameters are in Table 1.

Complex 1 displays a mononuclear structure where the cerium cation is bound by three siloxide oxygens of the tripodal ligand and one THF molecule and display an additional η^6 arene interaction with the arene ring. The mean value of the Ce-O_{sil} distances of 2.237(3) Å is comparable to that reported for the Ce(III) complex $[Ce(OSi(O^tBu)_3)_3(THF)_2]$ (2.246(2) Å).²¹ The average values of Ce–C distances (3.068(5) Å) and the Ce– C_{centroid} distance (2.730(1) Å) are in the range of previously reported π -arene-cerium(III) interactions (average Ce-C = 2.95(4)–3.07(7) Å and Ce–C $_{\rm centroid} = 2.607–2.734$ Å). 22 The solidstate molecular structure of 2 was determined by X-ray diffraction studies and shows a mononuclear Ce(IV) complex. The Ce center is four-coordinated by three $-OSi(O^tBu)_2$ groups of the tripodal ligand and one axial chloride ligand in a pseudotetrahedron geometry. The average value of the $Ce-O_{sil}$ distances (2.102(6) Å) is significantly shorter than those found in the Ce(m) precursor (1) and reduced species (3-5), indicating the presence of a Ce(IV) center. This value is also comparable to those found for the previously reported Ce(rv) complex $[Ce(OSi(O^tBu)_3)_4]$ (2.11(3) Å).^{10c} The Ce-C_{centroid} distance (2.821(2) Å) is longer than in complexes 1 and 3-5 (2.394(3)-2.730(1) Å) indicating the absence of any cerium-arene interaction. The average C-C bond length in the centroid arene (1.40(1) Å) is comparable to those found in the free ligand $(L(OH)_3)$ (1.39(2) Å) and in complex 1 (1.402(7) Å). The Ce-Cl

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Fig. 5 Molecular structure of 1-5, with thermal ellipsoids drawn at the 50% probability level. Hydrogen atoms and methyl groups on the – OSi(O^tBu)₂ ligands and [K(2.2.2-cryptand)] (3 and 4) have been omitted for clarity.

distance (2.5709(15) Å) is similar to that found in the Ce($_{IV}$) precursor [CeCl(N(SiMe₃)₂)₃] (2.597(2) Å).¹⁷

The solid-state structure of **3** shows the same coordination geometry and ligand environment as **1**, and the structure is completed by an outer-sphere $[K(2.2.2\text{-cryptand})]^+$ cation, revealing that one-electron reduction has occurred. The structure contains two different disordered metal locations. The predominant position (Ce1) of the Ce cation (89%) is situated below the plane of the three oxygen atoms and the minor one (Ce2) is above the plane (11%).

In the following metrical data analysis, the data on the 89% components of 3 are reported. The average Ln–ligand distances were used in particular by the Evans group to differentiate the reduction of $4f^n Ln^{3+}$ complexes to $4f^n5d^1 Ln^{2+}$ complexes, resulting only in small increases in distances (0.02–0.04 Å), leading to larger differences (0.10–0.20 Å).^{2f,13b} The average Ce– O_{sil} distance of 2.258(5) Å is only 0.02 Å longer than that found

in the precursor **1** which points either to reduction to a $4f^75d^1$ Ce²⁺ complex or to an unchanged metal oxidation state.^{2f,13b} The average values of the Ce–C distances (2.933(10) Å) and the Ce–C_{centroid} distance (2.569(2) Å) are significantly shorter than those found in **1** suggesting that the Ce–arene interaction is stronger, but they remain longer than those reported for the complex [K(2.2.2-cryptand)][((^{Ad,Me}ArO)₃Mes)Ce] containing Ce(m) and a reduced arene tripodal ligand (average Ce–C = 2.804 Å and Ce–C_{centroid} = 2.413 Å).^{13b}

It is noteworthy that the central arene is almost unchanged in 3 with minimal torsion angles $(0.2(6)-5.3(6)^\circ)$ with an average C–C bond length of 1.41(1) Å, which is comparable to that observed for the Ce(m) precursor (1.402(7) Å).

Overall, the degree of reduction of the metal cannot be unambiguously established from structural parameters.

The molecular structure of complex 4 shows a similar structure to 3, but the axial THF ligand is replaced by a Et_3OP

| Table 1 Selected structural parameters of free ligand L(OH)3, ligand radical [(K(crypt))(L(OK)3)]2 a | and complexes 1- | -5 |
|--|------------------|----|
|--|------------------|----|

| Compound | L(OH) ₃ | [(K(crypt))(L(OK) ₃)] ₂ | 1 | 2 | 3 | 4 | 5 |
|-----------------------------------|--------------------|--|---------------|---------------|-----------------|-----------------|------------------|
| Ce–O (avg.) | _ | _ | 2.237(3) | 2.102(6) | 2.258(5) | 2.277(5) | 2.363(9) |
| Ce-C _{centroid} | _ | _ | 2.730(1) | 2.821(2) | 2.569(2) | 2.632(2) | 2.394(3) |
| C–C _{arene} (avg.) | 1.39(2) | 1.411(5) | 1.402(7) | 1.402(15) | 1.415(15) | 1.417(12) | 1.43(3) |
| Torsion angle (°) (central arene) | 0.3(7)-3.6(7) | 0.2(2)-2.4(2) | 0.0(3)-0.2(3) | 0.1(7)-0.4(7) | 0.2(6) - 5.3(6) | 1.0(6) - 7.3(5) | 6.9(12)-18.0(11) |

ligand. The average value of the Ce–O_{sil} distances (2.277(5) Å), O–Ce–O bond angles (118.64(11)°) and the C–C bond distances (1.417(12) Å) are comparable to that of **3**. Conversely, the average Ce–C bond lengths (2.989(10) Å) and Ce–C_{centroid} distance (2.632(2) Å) are longer than those in **3**, suggesting that the interaction between the arene and metal center becomes weaker due to the presence of the strong electron-donating Et₃OP ligand in the axial position. The Ce–O_{phosphine} distance (2.439(3) Å) is in the range of Ce(III) complexes with neutral phosphine oxide ligands (2.319(3)–2.5165(9) Å).²³

In complex 5, the Ce center is coordinated by the three siloxide groups of the tripodal ligand ligands, the distorted arene anchor and one Et_2O molecule. One K^+ cation is bound by two siloxide ligands and a second K^+ cation is held by a siloxide and by interaction with one arene side-arm.

The central arene moiety η^6 -bound to the cerium is significantly bent, with values of the torsion angles ranging from $6.9(12)^{\circ}$ to $18.0(11)^{\circ}$ suggesting the presence of a reduced arene moiety. The highest values of the torsion angles are similar to those found in the reduced arene moiety of the recently reported Th($_{IV}$) [K[2.2.2]cryptand][Th(TDA)₃(THF)] (TDA = N-(2,6diisopropylphenyl)pivalamido) complex (torsion angles: 17.4(4)° and 24.2(4)°).24 The average values of Ce-C distances (2.78(2) Å) and the Ce–C $_{\rm centroid}$ distance (2.394(3) Å) are significantly shorter than those found in the mono-reduced complex 3 revealing a stronger interaction between the arene and metal center. The average value of the Ce-O_{sil} distances (2.363(9) Å) is longer than that found in the Ce(III) precursor (1), and it is comparable to Ce-O_{sil} bond lengths (2.367(4)-2.376(4) Å) in the reduced $[K(2.2.2\text{-cryptand})]_2[{(K(OSi(O^tBu)_3)_3Ce)(\mu-\eta^6:\eta^6 C_7H_8$, and could suggest the presence of a reduced Ce ion, despite the effect of potassium binding to siloxides.

Computational studies

To get some insights into the bonding and concomitantly into the oxidation state of the cerium atom, DFT calculations (B3PW91) including dispersion corrections (see the ESI[†] for details) were carried out on complexes 1-5. The effect of alkali ion coordination on 5 was also investigated. The optimized geometries are found to be in good agreement with the experimental ones (see Tables S5, S11, S17, S24, S30 and S38 in the ESI[†]). The calculations were first carried out on complex 2, which is found to have a closed-shell singlet ground state, in line with Ce(IV). For complex 1, a doublet spin state was computed and the geometry is in good agreement with the experimental ones with the Ce-O distances reproduced with a 0.03 Å accuracy as well as the average C-C distances in the arene (1.40 Å). The SOMO of the system is a pure f orbital on Ce (see Table S10[†]) and the unpaired spin density is localized only at the cerium. This is in line with the presence of a classical Ce(m) system. Interestingly, the LUMO of the system displays a small f-arene π^* interaction so that one can anticipate that reducing complex 1 may lead to mainly the arene reduction. The reduction of 1 was carried out experimentally and it forms either complex 3 or 5 depending on the number of electrons involved in the reduction (respectively 1 or 2). Two different spin

states were considered for complex 3, namely a singlet and a triplet. As expected, the triplet is 17.1 kcal mol⁻¹ more stable than the singlet. The optimized geometry is in good agreement with the experimental one with the Ce-O and Ce-C distances well reproduced with a maximum deviation of 0.06 Å. The average C-C bond distance as well as Ce-centroid distances are also nicely reproduced computationally (1.41 and 2.55 Å). In this complex a Ce-arene interaction is found as reflected by both the C-C distances which are elongated and by the Ce-C Wiberg bond index of 0.1 (to be compared with a WBI of 0.4 for the Ce-O bonds) (Table S20[†]). It is interesting to note that the two SOMOs (Fig. 6) a pure f (Fig. 6a) and a Ce-arene δ -bond (Fig. 6c) are similar to the frontier orbitals of complex 1 (Table S10[†]) but with a larger metal contribution in the Ce–arene π^* , which is better described as a Ce-arene δ bond. Both f and d orbitals of cerium participate in the δ bonding. The unpaired electron density plot shows that the electron density is mainly located at the cerium atom with some residues on the arene (Fig. 6b). Therefore, the cerium oxidation state is in between +II and +III with an arene that is in between neutral and singly reduced. This is somewhat similar to that observed by the Meyer group for the U(II) complex of arene(tris-aryloxide ligand)^{12c} where unpaired electrons were located in the δ -bond. CASSCF calculations indicate that the triplet ground state is 85% Ce(m) and 15% Ce(II). TDDFT calculations on complex 3 indicate that the absorption bands at 375, 545 and 770 nm involve excitation from the δ -bond to ligand orbitals (Fig. S82, S83 and Tables S43, S44†).

It is noteworthy that the replacement of a THF molecule by a phosphine oxide, complex **4**, only slightly modifies the bonding situation. Indeed, the triplet is also the ground-state with some slightly longer Ce–C distances than in **3**. This longer distance, also reflected by the Ce–centroid one (2.68 comp. *vs.* 2.63 exp), implies that the Ce–arene π^* interaction in the SOMO (Table S29†) in complex **4** is weaker so that the unpaired spin density is more distributed between the cerium



Fig. 6 (a) and (c) SOMO and (b) unpaired spin density plot of complex 3.



and the arene. Therefore, in complex 4, the oxidation state of Ce is close to +III with a singly reduced arene. The observed difference compared to complex 3 suggests that it should be possible to identify axial ligands that would favor a pure Ce(II) configuration.

In the case of the two-electrons reduction of complex 1, forming complex 5, the situation is different. Indeed, two different spin states, namely a quartet and a doublet, were considered and are found to be very close in energy. The doublet is slightly more stable than the quartet by 5.3 kcal mol^{-1} but interestingly both structures are in good agreement with the experimental one (see Table S31 in the ESI[†]). This can be easily explained by checking the frontier orbitals in both spin states (Table S36[†]). In fact, in the doublet ground state, a singly occupied f orbital at the cerium center is found and the δ bonding interaction is doubly occupied. Therefore, the arene is doubly reduced with the presence of a Ce(m) center. Interestingly, removing the potassium atoms from complex 5 results in a ground state switch, with the quartet becoming the ground state (Table S37[†]). In the quartet, 2 unpaired electrons are located on two different δ bonds and the unpaired spin density is now more localized on the cerium center (Table S42[†]) so that one would conclude that the cerium is in the oxidation state +II rather than +III. This is in line with the experimental observation. However, the storage of the electrons onto the δ -interaction allows complexes 3, 4 and 5 to be described as either masked Ce(II) complexes for the two former and as masked Ce(I) for the latter.

Redox reactivity

With the low-valent tripodal complexes in hand, we performed preliminary reactivity studies to investigate if the electrons stored in the complexes **3** and **5** could become available for the reduction of substrates (Scheme 4).

At first, we reacted complex 3 with 1 equivalent of AgBPh₄, leading immediately to a color change from dark-blue to lightyellow. The ¹H NMR spectrum of the reaction mixture revealed the clean formation of **1** (Fig. S46†). The observed reactivities showed that the electron stored in the complex 3 can be transferred to oxidizing substrates.

Then, we investigated the reduction of CO_2 as an example of a less reactive substrate.

The addition of 1 atm CO_2 to a THF solution of 3 led to an immediate color change from dark-blue to colorless, full consumption of the starting material and the formation of a new species as indicated by ¹H NMR studies (Fig. S47†). Colorless crystals of the complex $[K(2.2.2\text{-cryptand})]_2[\{((LO_3) Ce)_2(\mu-C_2O_4)\}]$ 6 were isolated in 52% yield from concentrated THF at room temperature. The conversion, as measured by ¹H NMR spectroscopy (using TMS₂O as the internal standard), is 73% (Fig. S48†). The solid-state structure of 6 (Fig. 7), determined by X-ray diffraction studies showed the presence of a dinuclear complex consisting of two equivalent Ce(m) ions, bridged by one oxalate ligand. Each Ce(m) center is pentacoordinated in a distorted square pyramidal geometry and is bound by three $-OSi(O^tBu)_2R$ ($R = C_6H_4$) ligands and one bridging oxalate.

Two [K(2.2.2-cryptand)] cations complete the structure. The average Ce–O_{sil} distance of 2.235(2) Å is comparable to that found in the Ce(\mathfrak{m}) precursor (1). The Ce–O_{oxalate} (2.465(2) Å) distances are shorter than those found in previously reported oxalate-bridged cerium(\mathfrak{m}) polymeric complexes (2.515(13)–2.62(5) Å) prepared from the reaction of oxalate with cerium salts.²⁵

The C1–C1' bond length is 1.536(4) Å which is consistent with the formation of a single C–C bond (1.54 Å in ethane). Moreover, the Ce–C_{centroid} distance (3.547(1) Å) is significantly longer than that of complex 1 with a Ce–C_{centroid} distance of 2.730(1) Å, indicating that there is no interaction between the arene and metal center. The formation of complex 6 indicates that a CO₂ coupling reaction occurs after the reduction of two molecules of CO₂ by two molecules of complex 3. Notably, complex 6 is the only isolated product and the major product in the reaction mixture. This is the first example of CO₂ reduction by a low valent cerium complex and only the third example of CO₂ reduction to oxalate by a low-valent lanthanide complex.²⁶



Fig. 7 Molecular structure of **6**, with thermal ellipsoids drawn at the 50% probability level. Hydrogen atoms and methyl groups on the – $OSi(O^tBu)_2$ ligands and [K(2.2.2-cryptand)] have been omitted for clarity.

In previous studies the reduction of CO_2 by Sm(II) Yb(II) or Eu(II) complexes led either to carbonate alone²⁷ or to a mixture of carbonate and oxalate depending on the supporting ligand.²⁸

Following isolation of complex 5, the addition of 2 equivalents of AgBPh₄ to 5 led to a clean regeneration of 1, with concomitant formation of KBPh₄ and Ag (Fig. S50†) suggesting that 5 could potentially act as a two-electron reductant. Moreover, complex 5 reacted with 1 equivalent of I₂, leading immediately to a color change from dark-purple to light-yellow and the precipitation of KI. The ¹H NMR spectrum of the reaction mixture revealed the clean formation of 1 (Fig. S51†). The conversion, as measured by ¹H NMR spectroscopy (using TMS₂O as the internal standard), is 99% (Fig. S52†).

Furthermore, the reaction of 1 equiv. of S_8 with a THF solution of complex 5 at room temperature resulted immediately in a light-yellow solution. ¹H NMR studies showed that complex 1 is formed (Fig. S53†) but the reduced sulfur products could not be isolated.

However, when the same reaction was performed in the presence of 2 equiv. of 2.2.2-cryptand at room temperature, it resulted immediately in a dark-red reaction mixture. Full consumption of the starting material and the formation of the multiple species were observed in the ¹H NMR spectrum (Fig. S54†). The recrystallization of the reaction mixture from concentrated THF at room temperature afforded few orange crystals of the S₁₀ dianion [K(2.2.2-cryptand)]₂[S₁₀]²⁹ as confirmed by X-ray diffraction. These results show that complex 5 can act as a two-electron reductant masked "Ce(1)". Complex 5 was found to react also with less oxidizing substrates such as azobenzene and phenylacetylene but the isolation of the resulting products was not successful so far.

Conclusions

In conclusion, we synthesized a new tripodal *tert*-butoxysiloxide ligand that incorporates an arene ring as part of a potentially

redox-active chelating ligand and showed that this ligand allows the promotion of multielectron redox reactivity in cerium complexes and the stabilization of cerium complexes in four different redox states.

Remarkably the one-electron reduction and the unprecedented two-electron reduction of the tripodal Ce(III) complex are easily achieved to yield complexes that are formally "Ce(II)" and "Ce(1)" analogues. Structural, spectroscopic and computational studies indicated that the neutral complexes 1 and 2 show the presence of cerium in +3 and +4 oxidation states with respectively little and no interaction of the metal center with the arene ring. The assignment of the formal state of cerium is more ambiguous for the mono-reduced complex 3 and is better described as an intermediate between +2 and +3 with partial localization of electrons on the ligand and a Ce-arene δ -interaction. In complex 5 the arene is doubly reduced with the presence of a Ce(III) center, but the removal of potassium results in a $Ce(\pi)(arene^{-})$ species. However, the storage of electrons onto the δ -interaction allows the description of complexes 3 and 5 as masked "Ce(II)" and "Ce(I)" complexes respectively. Preliminary reactivity studies show that these complexes can act as masked Ce(II) and Ce(I) effecting both one- and two-electron transfer that were not accessible in classical cerium chemistry. These results demonstrate that the high stability of the arene tripodal ligand under reducing and oxidizing conditions is well poised to promote the development of f element multielectron redox chemistry.

Data availability

Synthetic details and analytical data including depictions of all spectra and coordinate data of all computationally optimised species are documented in the ESI. Crystallographic data are made available *via* the CCDC.† The data that support the findings of this study are openly available in the Zenodo repository at https://doi.org/10.5281/zenodo.7869094.

Author contributions

F.-C. H. designed and carried out all the experiments and analyzed the data; M. M. designed and supervised the project; T. R. and L. M. carried out the computational study; R. S. measured and analyzed the X-ray data; A. S. measured the EPR data. F.-C. H., L. M, and M. M. wrote the manuscript with contributions of all authors, and all authors have given approval to the final version of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- 1 J. C. Wedal and W. J. Evans, J. Am. Chem. Soc., 2021, 143, 18354–18367.
- 2 (a) L. M. Anderson-Sanchez, J. M. Yu, J. W. Ziller, F. Furche and W. J. Evans, Inorg. Chem., 2023, 62, 706-714; (b) W. J. Evans, N. T. Allen and J. W. Ziller, Angew. Chem., Int. Ed. Engl., 2002, 41, 359-360; (c) P. B. Hitchcock, M. F. Lappert, L. Maron and A. V. Protchenko, Angew. Chem., Int. Ed. Engl., 2008, 47, 1488-1491; (d) F. Jaroschik, A. Momin, F. Nief, X. F. Le Goff, G. B. Deacon and P. C. Junk, Angew. Chem., Int. Ed. Engl., 2009, 48, 1117-1121; (e) 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; (f) 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; (g) K. R. Meihaus, M. E. Fieser, J. F. Corbey, W. J. Evans and J. R. Long, J. Am. Chem. Soc., 2015, 137, 9855-9860; (h) 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; (i) 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; (j) D. N. Huh, S. R. Ciccone, S. Bekoe, S. Roy, J. W. Ziller, F. Furche and W. J. Evans, Angew. Chem., Int. Ed. Engl., 2020, 59, 16141-16146; (k) D. N. Huh, J. P. Bruce, S. G. Balasubramani, S. R. Ciccone, F. Furche, J. C. Hemminger and W. J. Evans, J. Am. Chem. Soc., 2021, 143, 16610-16620.
- 3 P. L. Arnold, F. G. N. Cloke, P. B. Hitchcock and J. F. Nixon, *J. Am. Chem. Soc.*, 1996, **118**, 7630–7631.
- 4 L. Barluzzi, S. R. Giblin, A. Mansikkamaki and R. A. Layfield, *J. Am. Chem. Soc.*, 2022, **144**, 18229–18233.
- 5 J. D. Martin and J. D. Corbett, Angew. Chem., Int. Ed. Engl., 1995, 34, 233-235.
- 6 (a) X. Chen, T. T. Chen, W. L. Li, J. B. Lu, L. J. Zhao, T. Jian, H. S. Hu, L. S. Wang and J. Li, *Inorg. Chem.*, 2019, 58, 411–418; (b) W. L. Li, T. T. Chen, W. J. Chen, J. Li and L. S. Wang, *Nat. Commun.*, 2021, 12, 6467.
- 7 (a) E. J. Schelter, Nat. Chem., 2013, 5, 348; (b) N. A. Piro, J. R. Robinson, P. J. Walsh and E. J. Schelter, Coord. Chem. Rev., 2014, 260, 21-36; (c) J. A. Bogart, C. A. Lippincott, P. J. Carroll, C. H. Booth and E. J. Schelter, Chem.-Eur. J., 2015, 21, 17850-17859; (d) L. Mathey, M. Paul, C. Coperet, H. Tsurugi and K. Mashima, Chem.-Eur. J., 2015, 21, 13454-13461; (e) T. Montini, M. Melchionna, M. Monai and P. Fornasiero, Chem. Rev., 2016, 116, 5987-6041; (f) J. R. Robinson, Y. S. Qiao, J. Gu, P. J. Carroll, P. J. Walsh and E. J. Schelter, Chem. Sci., 2016, 7, 4537-4547; (g) H. L. Yin, P. J. Carroll, B. C. Manor, J. M. Anna and

E. J. Schelter, J. Am. Chem. Soc., 2016, 138, 5984–5993; (h)
R. Anwander, M. Dolg and F. T. Edelmann, Chem. Soc. Rev.,
2017, 46, 6697–6709; (i)
R. L. Halbach, G. Nocton,
C. H. Booth, L. Maron and R. A. Andersen, Inorg. Chem.,
2018, 57, 7290–7298; (j)
Q. M. Yang, Y. H. Wang, Y. S. Qiao,
M. Gau, P. J. Carroll, P. J. Walsh and E. J. Schelter, Science,
2021, 372, 847–852.

- 8 M. A. Angadol, D. H. Woen, C. J. Windorff, J. W. Ziller and W. J. Evans, *Organometallics*, 2019, **38**, 1151–1158.
- 9 T. F. Jenkins, D. H. Woen, L. N. Mohanam, J. W. Ziller, F. Furche and W. J. Evans, *Organometallics*, 2018, **37**, 3863– 3873.
- 10 (a) Y. K. Gun'ko, P. B. Hitchcock and M. F. Lappert, Organometallics, 2000, 19, 2832–2834; (b) C. M. Kotyk, M. E. Fieser, C. T. Palumbo, J. W. Ziller, L. E. Darago, J. R. Long, F. Furche and W. J. Evans, Chem. Sci., 2015, 6, 7267–7273; (c) R. P. Kelly, L. Maron, R. Scopelliti and M. Mazzanti, Angew. Chem., Int. Ed. Engl., 2017, 56, 15663– 15666.
- 11 (a) C. Camp, V. Guidal, B. Biswas, J. Pecaut, L. Dubois and M. Mazzanti, Chem. Sci., 2012, 3, 2433-2448; (b) I. L. Fedushkin, O. V. Maslova, A. G. Morozov, S. Dechert, S. Demeshko and F. Meyer, Angew. Chem., Int. Ed. Engl., 2012, 51, 10584-10587; (c) E. J. Coughlin, M. Zeller and S. C. Bart, Angew. Chem., Int. Ed. Engl., 2017, 56, 12142-12145; (d) I. L. Fedushkin, A. N. Lukoyanov and E. V. Baranov, Inorg. Chem., 2018, 57, 4301-4309; (e) S. S. Galley, S. A. Pattenaude, R. F. Higgins, C. J. Tatebe, D. A. Stanley, P. E. Fanwick, M. Zeller, E. J. Schelter and S. C. Bart, Dalton Trans., 2019, 48, 8021-8025; (f) N. Jori, D. Toniolo, B. C. Huynh, R. Scopelliti and M. Mazzanti, Inorg. Chem. Front., 2020, 7, 3598-3608; (g) G. Nocton, W. W. Lukens, C. H. Booth, S. S. Rozenel, S. A. Medling, L. Maron and R. A. Andersen, J. Am. Chem. Soc., 2014, 136, 8626-8641.
- 12 (a) S. C. Bart, F. W. Heinemann, C. Anthon, C. Hauser and K. Meyer, *Inorg. Chem.*, 2009, 48, 9419–9426; (b) O. P. Lam, S. C. Bart, H. Kameo, F. W. Heinemann and K. Meyer, *Chem. Commun.*, 2010, 46, 3137–3139; (c) H. S. La Pierre, A. Scheurer, F. W. Heinemann, W. Hieringer and K. Meyer, *Angew. Chem., Int. Ed. Engl.*, 2014, 53, 7158–7162.
- 13 (a) D. P. Halter, C. T. Palumbo, J. W. Ziller, M. Gembicky,
 A. L. Rheingold, W. J. Evans and K. Meyer, J. Am. Chem. Soc., 2018, 140, 2587–2594; (b) 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.
- 14 R. R. Thompson, M. E. Rotella, P. Du, X. Zhou, F. R. Fronczek, R. Kumar, O. Gutierrez and S. Lee, *Organometallics*, 2019, 38, 4054–4059.
- 15 (a) V. Nair and A. Deepthi, *Chem. Rev.*, 2007, **107**, 1862–1891;
 (b) A. H. Hu, J. J. Guo, H. Pan and Z. W. Zuo, *Science*, 2018, **361**, 668–672.
- 16 Y. M. So and W. H. Leung, *Coord. Chem. Rev.*, 2017, **340**, 172–197.
- 17 O. Eisenstein, P. B. Hitchcock, A. G. Hulkes, M. F. Lappert and L. Maron, *Chem. Commun.*, 2001, 1560–1561.

- 18 T. A. Albright, W. J. Freeman and E. E. Schweizer, *J. Org. Chem.*, 1975, **40**, 3437–3441.
- 19 N. T. Rice, J. Su, T. P. Gompa, D. R. Russo, J. Telser, L. Palatinus, J. Bacsa, P. Yang, E. R. Batista and H. S. La Pierre, *Inorg. Chem.*, 2019, **58**, 5289–5304.
- 20 J. Mayans, L. Tesi, M. Briganti, M. E. Boulon, M. Font-Bardia, A. Escuer and L. Sorace, *Inorg. Chem.*, 2021, **60**, 8692–8703.
- 21 J. Friedrich, C. Maichle-Mossmer and R. Anwander, *Chem. Commun.*, 2017, **53**, 12044–12047.
- 22 (a) G. B. Deacon, T. C. Feng, C. M. Forsyth, A. Gitlits, D. C. R. Hockless, S. Qi, B. W. Skelton and A. H. White, J. Chem. Soc., Dalton Trans., 2000, 961–966; (b) M. Gorlov, L. L. Hussami, A. Fischer and L. Kloo, Eur. J. Inorg. Chem., 2008, 5191–5195; (c) A. N. Selikhov, A. V. Cherkasov, K. A. Lyssenko and A. A. Trifonov, Organometallics, 2022, 41, 820–828.
- 23 (a) J. C. Berthet, M. Nierlich and M. Ephritikhine, *Polyhedron*, 2003, 22, 3475–3482; (b) A. Bowden,
 P. N. Horton and A. W. G. Platt, *Inorg. Chem.*, 2011, 50, 2553–2561; (c) J. R. Robinson, Z. Gordon, C. H. Booth,
 P. J. Carroll, P. J. Walsh and E. J. Schelter, *J. Am. Chem. Soc.*, 2013, 135, 19016–19024; (d) R. D. Bannister,
 W. Levason and G. Reid, *Chemistry*, 2020, 2, 947–959.
- 24 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.

- 25 L. Vella-Zarb and U. Baisch, Z. Anorg. Allg. Chem., 2017, 643, 1712–1716.
- 26 (a) W. J. Evans, C. A. Seibel and J. W. Ziller, *Inorg. Chem.*, 1998, 37, 770–776; (b) W. J. Evans, J. M. Perotti, J. C. Brady and J. W. Ziller, *J. Am. Chem. Soc.*, 2003, 125, 5204–5212.
- 27 (a) N. W. Davies, A. S. P. Frey, M. G. Gardiner and J. Wang, *Chem. Commun.*, 2006, 4853–4855; (b) A. R. Willauer, A. M. Dabrowska, R. Scopelliti and M. Mazzanti, *Chem. Commun.*, 2020, 56, 8936–8939.
- 28 (a) J. Andrez, J. Pecaut, P.-A. Bayle and M. Mazzanti, Angew. Chem., Int. Ed. Engl., 2014, 53, 10448-10452; (b)
 M. Xemard, V. Goudy, A. Braun, M. Tricoire, M. Cordier, L. Ricard, L. Castro, E. Louyriac, C. E. Kefalidis, C. Clavaguera, L. Maron and G. Nocton, Organometallics, 2017, 36, 4660-4668; (c) A. R. Willauer, D. Toniolo, F. Fadaei-Tirani, Y. Yang, M. Laurent and M. Mazzanti, Dalton Trans., 2019, 48, 6100-6110; (d) A. R. Willauer, F. Fadaei-Tirani, I. Zivkovic, A. Sienkiewicz and M. Mazzanti, Inorg. Chem., 2022, 61, 7436-7447.
- 29 M. K. Mondal, L. Zhang, Z. T. Feng, S. X. Tang, R. Feng,
 Y. Zhao, G. W. Tan, H. P. Ruan and X. P. Wang, *Angew. Chem., Int. Ed. Engl.*, 2019, 58, 15829–15833.