Facile formation of a homoleptic Ce(IV) amide via aerobic oxidation[†]

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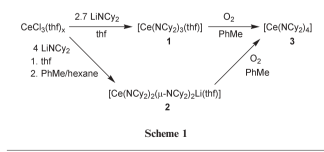
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Oxidation of $[Ce(NCy_2)_3(thf)]$ or $[Ce(NCy_2)_4Li(thf)]$ with dry air produced the first homoleptic Ce(IV) amide $[Ce(NCy_2)_4]$.

Cerium is often considered as an analogue of a group 4 metal due to its ability to form compounds in the oxidation state +4. However, homoleptic dialkyl amides, which are common and widely used for Zr and Hf,¹ are unknown for Ce. Related are Ce(IV) bis(trifluoromethanesulfonyl)amide² and porphyrinato- or phthalocyaninato-Ce complexes,³ the latter often being considered as having the Ce atom in an intermediate (between +3 and +4) oxidation state due to the high redox activity of the ligands. Each of the three heteroleptic Ce(IV) amides reported up to date, [CeI{N(SiMe₂Bu¹)CH₂CH₂}N]^{4a} and [Ce{N(SiMe₃)₂}X] (X = Cl or Br),^{4b,c} contain silyl substituents on the amido N atoms.

The initial attempts to prepare a Ce(IV) dialkylamido complex using [Ce(TMP)₃] (TMP = 2,2,6,6-tetramethylpiperidinate) failed,⁵ apparently due to the high steric demands of the TMP ligand. Thus, dicyclohexylamide (NCy₂) was chosen as a less bulky alternative to TMP for stabilisation of the Ce +4 oxidation state. Here we report the synthesis and characterisation of two new Ce(III) dicyclohexylamides and their oxidation to produce the first homoleptic Ce(IV) amide.

The starting Ce(III) amides [Ce(NCy₂)₃(thf)] (1) and [Ce(NCy₂)₄Li(thf)] (2) were prepared *via* the salt elimination route (Scheme 1) using 2.7 or 4 equivalents of Li amide, respectively.[‡] Both compounds demonstrate paramagnetically shifted ¹H NMR signals in C₆D₆ with one set of the Cy group signals for 1 and two sets (terminal and bridging NCy₂ ligands) for 2 (see ESI[†]). In a coordinating solvent (thf-d₈) complex 2 apparently dissociates into the tetrahedral anion [Ce(NCy₂)₄]⁻ (which gives only one set of slightly paramagnetically shifted Cy signals) and the [Li(thf)_x][†] cation (which gives a sharp signal in the ⁷Li NMR spectrum; no such signal was observed in C₆D₆ solvent due to paramagnetic broadening).



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The thf-solvated cerium triamide **1** was stable in solution in hydrocarbon solvents and was thermally robust up to its melting point (*ca.* 235 °C) in contrast to the bulkier analogue [Ce(TMP)₃(thf)],⁵ which can be desolvated at room temperature. The lithium tetraamidocerate(III) **2** decomposed slowly in aromatic solvents and much faster in thf, apparently due to thf C–O bond cleavage similar to that observed in the synthesis of [Ce(TMP)₃].⁶

Complexes 1 and 2 were extremely air-sensitive in solution and if the smallest amount of air was accidentally introduced during the work-up, a blue colouration appeared in the solution immediately, later yielding black microcrystals. When a measured amount of dry air was added to toluene or thf solutions of 1 or 2 the diamagnetic Ce(IV) amide [Ce(NCy₂)₄] (3)[‡] was isolated in a moderate yield (Scheme 1).

Complex **3** was only sparingly soluble in hexane, benzene, toluene or thf, producing deep blue solutions, which were stored in a vacuum-sealed tube without decomposition for several months at room temperature (in contrast to the solution instability of $[Ce{N(SiMe_{3})_2}_3X] [X = Cl, Br]^{4c}$). The solubility increased upon heating to 70 °C thus allowing recrystallisation of **3** without noticeable decomposition even at this temperature.

To confirm the oxidation state of Ce in these complexes the structures of 1, 2 and 3 were determined by single crystal X-ray diffraction.§ The molecular structures of complexes 1 and 2 are similar to those of their Sm analogues;⁷ details are given in ESI[†]. In the structure of complex 3 there are two independent molecules

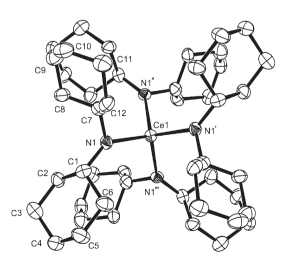


Fig. 1 Molecular structure of [Ce(NCy₂)₄] (3) (50% probability ellipsoids) showing one of the three independent molecules in the crystal. Symmetry transformations used to generate equivalent atoms: '1 - y, x, -z; "y, 1 - x, -z; "1 - y, 1 - x, z.

Compound	Ce–N	Ce–O
$[Ce(NCy_2)_3(thf)](1)$	2.299(2), 2.317(2), 2.336(2)	2.582(2)
$[Ce(NCy_2)_4Li(thf)] (2)$	2.320(2), 2.330(2)	
$\left[\operatorname{Ce}(\operatorname{TMP})_3\right]^6$	2.332(7), 2.346(7), 2.291(7)	
$[Ce(TMP)_3(thf)]^5$	2.374(2), 2.346(2), 2.363(2)	2.810(2)
$[Ce{N(SiMe_3)_2}_3]^8$	2.320(3)	
$[Ce(NCy_2)_4]$ (3) ^{<i>a</i>}	2.247(6), 2.242(6), 2.238(5), 2.240(5)	
$\left[\operatorname{CeCl}\left\{\operatorname{N}(\operatorname{SiMe}_{3})_{2}\right\}_{3}\right]^{4b}$	2.217(3)	
$\left[\operatorname{CeBr}\{\operatorname{N}(\operatorname{SiMe}_3)_2\}_3\right]^{4c}$	2.219(7)	
^a The four Ce-N bond lengths for 3 are due to the presence of 3		

ine iour Ce–N bond lengths for 3 are due to the presence of 3 different molecules in the crystal.

lying on $\overline{4}$ inversion centers and one lying on a two-fold rotation axis. Fig. 1 shows a higher-symmetry molecule of **3**, and important bond lengths for these compounds (together with the data for selected Ce(III) and Ce(IV) complexes) are given in Table 1.

Each of 1, 2 and 3 has the four-coordinate Ce atom in a distorted tetrahedral environment, which facilitates the following bond length comparisons. In complex 1 the three Ce–N distances are very similar to those in $[Ce(TMP)_3]^6$ but slightly shorter than in the overcrowded thf solvate $[Ce(TMP)_3(thf)]^5$ while the Ce–O distance is much shorter than in the latter (Table 1) paralleling the difference in the thermal stability of these complexes.

In complex **2**, two Ce–N distances to the terminal ligands are close to those in **1** and in the silylamide $[Ce{N(SiMe_3)_2}_3]$,⁸ while the two bridging Ce–N distances are longer at 2.472(2) and 2.497(2) Å. The Ce^{IV}–N distances in **3** are shorter than the terminal Ce^{III}–N distances in **1** or **2** by *ca*. 0.1 Å, but are slightly longer than the Ce^{IV}–N bonds in the less sterically hindered heteroleptic compounds $[Ce{N(SiMe_3)_2}_3X]$ (X = Cl or Br).^{4b,c} Thus the structural study (along with the NMR data) confirms the +4 cerium oxidation state in the homoleptic amide **3**.

Possible intermediates along the route to complex **3** may include Ce^{IV} superoxo-, peroxo- and oxo-complexes; the latter would disproportionate into **3** and a polymeric amido-oxo-compound of the type { $Ce(\mu-O)(NCy_2)_2$ }_x. While organolanthanides and low-valent Ln compounds with *N*-centred ligands are generally considered as decomposing completely on exposure to air, some examples of oxygen-containing products resulting from aerial oxidation have been reported, including superoxo- [$Sm{HB(3,5-Me_2pz)_3}_2(\eta^2-O_2)$],⁹ peroxo- [{ $Yb(C_5H_9C_5H_4)_2(thf)$ }₂(μ -O_2)]¹⁰ and [{ $Yb(N(SiMe_3)_2)_2(thf)$ }₂(μ -O_2)],¹¹ and complexes with oxygenated ligands.¹² The present study has demonstrated the suitability of this approach to the synthesis of oxygen-free organoamides of high-valent cerium. This oxidative route to complex **3** parallels that reported for the aerial oxidation of $Ce(\kappa^2-S_2CNEt_2)_3$ to [$Ce(\kappa^2-S_2CNEt_2)_4$].¹³

In conclusion, two new Ce^{III} amides 1 and 2 containing bulky dicyclohexylamido ligand were prepared by the salt metathesis reaction; their oxidation by air provided a route to the remarkably stable homoleptic Ce^{IV} amide [Ce(NCy₂)₄] (3).

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Notes and references

microcrystalline solvate. To this suspension a solution of LiNCy₂ (0.945 g, 5.05 mol) in thf (20 mL) was slowly added at 0 °C producing a yellow solution. The mixture was stirred for 1 h at room temperature and the solvent was removed in a vacuum. The residue was extracted with toluene (2 × 10 mL), filtered, and the filtrate was concentrated to *ca*. 2 mL and covered with hexanes. Storing at -27 °C for 2 days gave 0.811 g (57%) of 1-PhMe as large yellow crystals; mp 235–240 °C. ¹H-NMR (C₆D₆): δ 13.33, 7.00–7.13 (aromatic CH, toluene), 2.47, 2.11 (sharp s, CH₃ toluene), 1.71, 1.49 (and 1.47 sh), 1.19, 1.05, 0.27, -4.07, -7.83.

Synthesis of $[Ce(NCy_2)_4Li(thf)]$ (2): CeCl₃ (0.383 g, 1.55 mmol) was solvated with thf as described above and a solution of LiNCy₂ (1.132 g, 6.05 mol) in thf (20 mL) was slowly added at 0 °C. The mixture was stirred for 1 h at room temperature and the solvent was removed in a vacuum (if it was left as the THF solution overnight, the yield of crystalline products decreased significantly). The residue was extracted with toluene (2 × 10 mL), the combined toluene solution was concentrated to *ca.* 5 mL and covered with hexanes. Storing at 5 °C overnight gave 0.892 g (61%) of **2** as bright pink crystals; mp 146–148 °C (decomp.). ¹H-NMR (C₆D₆): δ 12.73, 11.75, 6.05, 5.50, 3.34, 1.50, 0.29, -0.35, -0.87, -1.38, -2.06, -3.44, -4.85, -6.58, -10.52, -29.74.

Synthesis of $[Ce(NCy_2)_4]$ (3): A Schlenk tube containing a degassed solution of 1·PhMe (0.303 g, 0.36 mmol) in toluene (30 mL) was connected *via* a short rubber tubing to an ampoule containing dry air (10 mL, 0.093 mmol of O₂). After stirring for 30 min at 25 °C a dark blue-violet solution was formed; the Schlenk tube was closed and stored at 5 °C overnight yielding a greenish-yellow solution with black crystals and some light brown amorphous precipitate. The crystals were washed by decantation with cold toluene and dried, yielding 0.108 g (35% based on Ce) of **3**. Complex **3** decomposed at 90–100 °C without melting. ¹H-NMR (C₆D₆): δ 4.13 (m, 1 H, NCH), 2.04 (d, 2 H), 1.88 (m, 2 H), 1.79 (m, 1 H), 1.70 (m, 2 H), 1.55 (m, 2 H), 1.30 (m, 1 H). ¹³C-NMR (C₆D₆): δ 57.44 (NCH), 39.60 (CH₂), 27.22 (CH₂), 26.08 (CH).

§ *Crystal data.* For 1·PhMe (yellow prism 0.25 × 0.20 × 0.20 mm³): [C₄₀H₇₄CeN₃O]·(C₇H₈), M = 845.28, monoclinic, space group $P2_1/c$, a = 10.3004(2), b = 23.4770(4), c = 19.3077(3) Å, $\beta = 101.189(1)$, V = 4580.29(14) Å³, Z = 4, T = 173(2) K, $\mu = 1.03$ mm⁻¹, 8044 independent reflections [$R_{int} = 0.060$], final R1 = 0.032 [for 6438 reflections with $I > 2\sigma(I)$], wR2 = 0.069 (all data). For **2** (pink prism 0.25 × 0.25 × 0.20 mm³): [C₅₂H₉₆CeLiN₄O], M = 940.39, monoclinic, space group $P2_1/n$, a = 15.1681(3), b = 19.1024(4), c = 18.1727(3) Å, $\beta = 96.822(1)$, V = 5228.21(17) Å³, Z = 4, T = 173(2) K, $\mu = 0.91$ mm⁻¹, 10277 independent reflections [$R_{int} = 0.078$], final R1 = 0.035 [for 7866 reflections with $I > 2\sigma(I)$], wR2 = 0.073 (all data). For **3** (black needle 0.10 × 0.05 × 0.05 mm³): [C₄₈H₈₈CeN₄], M = 861.34, tetragonal, space group $P\overline{4}$, a = b = 21.1876(5), c = 10.3198(3) Å, V = 4632.7(2) Å³, Z = 4, T = 173(2) K, $\mu = 1.02$ mm⁻¹, 8991 independent reflections [$R_{int} = 0.034$], final R1 = 0.048 [for 7005 reflections with $I > 2\sigma(I)$], wR2 = 0.106 (all data). CCDC numbers: 1·PhMe 609099, **2** 609100, 3 609101. For crystallographic data in CIF or other electronic format see DOI: 10.1039/b607429d

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 $[\]therefore$ Synthesis of $[Ce(NCy_2)_3(thf)]$ (1): CeCl₃ (0.453 g, 1.84 mmol) was stirred in thf (15 mL) at room temperature for 2 days to give a copious

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