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Terminal dysprosium and holmium organoimides†

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Terminal rare-earth-metal imide complexes $Tp^{tBu,Me}Ln(NC_6H_3iPr_2-2,6)(dmap)$ of the mid-late rare-earth elements dysprosium and holmium were synthesized via double methane elimination of Lewis acid stabilized dialkyl precursors $Tp^{tBu,Me}LnMe(GaMe_4)$ with primary aniline derivative $H_2NC_6H_3iPr_2-2,6$ (H_2NAr^{iPr}) . Exploiting the weaker $Ln-CH_3\cdots[GaMe_3]$ interaction compared to the aluminium congener, addition of the aniline derivative leads to the mixed methyl/anilido species $Tp^{tBu,Me}LnMe(HNAr^{iPr})$ which readily eliminate methane after being exposed to the Lewis base DMAP (=N,N-dimethyl-4-aminopyridine). Under the same conditions, $[AlMe_3]$ -stabilized dimethyl rare-earth-metal complexes transform immediately to Lewis acid bridged imides $Tp^{tBu,Me}Ln(\mu_2-NC_6H_3Me_2-2,6)(\mu_2-Me)AlMe_2$ (Ln=Dy, Ho). DMAP/THF donor exchange is accomplished by treatment of $Tp^{tBu,Me}Ln(NC_6H_3iPr_2-2,6)(dmap)$ with 9-BBN in THF while the terminal imides readily insert carbon dioxide to afford carbamate complexes.

Introduction

Recent years have witnessed major progress in the field of rareearth-metal (Ln) complexes with multiply bonded (dianionic) main-group ligands, most notably (organo)imide chemistry. 1-3 Upon closer inspection this is not all that surprising, since [Ln=NR] moieties display a favourable Pearson hard/hard match and enhanced steric/electronic variability through the imido substituent R,4 compared to other fragments such as [Ln=O],⁵ [Ln=PR],⁶ or [Ln=CR₂].⁷ Of particular interest have been the synthesis of terminal rare-earth-metal imides⁸⁻¹⁵ and a fundamental understanding of the Ln-imido bonding as well as reactivity,16 and in particular small-molecule-activation scenarios.17 Closely based on Chen's synthesis protocol of the first terminal scandium imide complex L1Sc(NAriPr)(dmap) = $C_6H_3iPr_2-2.6;$ L^1 = $[Ar^{iPr}NC(Me)CHC(Me)$ N(CH₂)₂NMe₂]),⁸ a donor-promoted intramolecular methane/ tetramethylsilane elimination from mixed methyl(neosilyl)/ primary amido precursors emerged as the most efficient approach for accessing terminal Ln(III) imides (Scheme 1/A).8-13 Meanwhile, another three strategies have been successfully pursued for the early larger rare-earth metals. While anionic terminal ceric imides could be obtained by the deprotonation of a neutral primary amide complex with alkali-metal silylamides (Scheme 1/B), ¹⁴ a [Ln=CH₂(GaMe₃)₂] \rightarrow [Ln=NR(thf)₂] transformation proved feasible for generating the open-shell

terminal imides $Tp^{rBu,Me}Ln(NAr^{iPr})(thf)_2$ (Ln = Ce, Nd, Sm) (Scheme 1/C). To More recently, an anionic cerium(IV) terminal imide was accessed by a two-electron oxidation of a "Ce(II)" complex supported by a tripodal tris(amido)arene ligand using azide $N_3Ar^{CF_3}$ ($Ar^{CF_3} = C_6H_3(CF_3)_2$ -3,5) (Scheme 1/D). Less surprising, terminal imides of the extremely large divalent rare-earth-metal centres have remained elusive.

Common features of all terminal rare-earth-metal imides, reported so far, are that the imido ligand is derived from a substituted aniline, H_2NAr^R ($R=iPr,\ Me,\ CF_3$), and an

A) donor-assisted intramolecular primary amido deprotonation

B) intermolecular primary amido deprotonation

C) donor-assisted methylidene-imido transformation

D) two-electron oxidation with organic azide

$$\begin{bmatrix} \text{Ce}^{\text{II}} \end{bmatrix}^{\bigcirc} [\text{K(crypt)}]^{\textcircled{\tiny{0}}} \xrightarrow{N_3 \text{Ar}^{\text{CF}_3}} \begin{bmatrix} \text{Ce}^{\text{IV}} = \text{N} - \text{Ar}^{\text{CF}_3} \end{bmatrix}^{\bigcirc} [\text{K(crypt)}]^{\textcircled{\tiny{0}}}$$

Scheme 1 Synthesis strategies for terminal rare-earth-metal imides.

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indispensable kinetic stabilization by use of sterically demanding ancillary co-ligands. Aliphatic amines, benzylic amines, and silylamines engage in imido ligand formation as well but have been detected only as metal-bridging and Lewis acid stabilized versions.³ Successfully applied ancillaries include β-diketiminato (nacnac), 8,9,11 phosphazene, 10 TriNox, 14 and multidentate pyrazolato ligands. 12,13,15 We found that especially the bulky monoanionic scorpionato ligand hydrotris(3-tert-butyl-5-methylpyrazolyl)borato ($Tp^{tBu,Me}$) provides a useful scaffold for stabilizing terminal imides as well as phosphinidenes. 6c,12,15 However, like for all terminal Ln(III) imides, stabilization of the highly polarized Ln=N bond, which predominantly consists of non-directional ionic interactions, is still challenging as it readily reacts with solvent molecules or the ancillary ligand of the aspired complexes.^{2,3} DFT calculations performed on the yttrium compound Tp^{tBu,Me}Y(NC₆H₃-Me₂-2,6)(dmap) confirmed a large ionic character of the Y-

Given the feasibility of terminal imides of the early openshell cations Ce(III) (f^1), Nd(III) (f^3), and Sm(III) (f^5), f^5 we herein envisaged the synthesis of those of the mid-late open-shell cations Dy(III) (f^9) and Ho(III) (f^{10}). These metal centres exhibit ionic radii similar to Y(III) but would (if at all) contribute to a distinct covalent bonding. On the other hand, their much higher molar mass might promote the crystallization behaviour and hence, the single-crystal X-ray structure diffraction (SCXRD) analysis of the targeted complexes. We also examined the reactivity of the first terminal, trivalent dysprosium and holmium imide complexes.

N(imido) bond but also a significant covalent bonding pattern

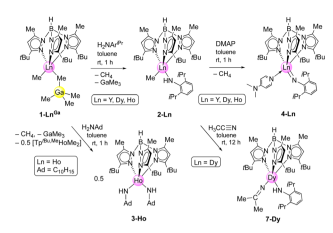
with one σ -type and two π -type interactions. ^{6c}

Results and discussion

Selection of precursors

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Our original approach toward the terminal yttrium imide Tp^{tBu,Me}Y(NC₆H₃Me₂-2,6)(dmap) involved the mixed methyl/ tetramethylgallato complex Tp^{tBu,Me}YMe(GaMe₄) as a suitable precursor. 12 There, the ease of GaMe₃ displacement proved to be crucial for the successful synthesis. Consequently, we chose the trimethylgallium-stabilized dialkyl complexes $Tp^{tBu,Me}LnMe(GaMe_4)$ (1-Ln^{Ga}; Ln = Y, 12 Dy, 6c Ho) as precursors for the present study. Like their aluminium congeners, 20 complexes 1-Ln^{Ga} are available in moderate yield via protonolysis of the homoleptic gallates Ln(GaMe₄)₃ (ref. 21) with H $[Tp^{tBu,Me}]$ (ref. 22) and precipitation from toluene or *n*-hexane solution (Scheme 2; for detailed metrics of Ln(GaMe₄)₃ (Dy, Ho) and 1-Ho^{Ga}; see the ESI, Fig. S1-S3†). The solid-state structure of the bimetallic compounds $Tp^{tBu,Me}LnMe(\mu_2-MeEMe_3)$ (1-Ln^E, E = Al, Ga), depicting one terminal methyl group and an almost linear Ln-Me-E linkage, is not reflected in the solution NMR spectra, which reveal highly fluxional methyl groups at ambient temperature, an even higher mobility in case of the gallium derivatives. The isostructural 1-LnGa show Ln-C(Me) (Dy: 2.389(3) Å, 6c Y: 2.385(3) Å, 12 Ho: 2.356(5) Å) and Ln-C(Me_{Ga}) distances (Dy: 2.736(2) Å,6c Y: 2.688(2) Å,12 Ho: 2.652(4) Å) in accordance with the distinct Ln(III) radii.23 Unexpectedly, the solid-state structures of the formally five-coordinate 1-Ln^{Ga} of



Scheme 2 Formation of $Tp^{fBu,Me}Ln(NAr^{iPr})(dmap)$ (4-Ln: Ln=Y, Dy, Ho) via reaction of $Tp^{fBu,Me}LnMe(GaMe_4)]$ (1-Ln^{Ga}) with primary aniline H_2NAr^{iPr} to afford mixed methyl/amido complexes $Tp^{fBu,Me}LnMe(HNAr^{iPr})$ (2-Ln), and subsequent addition of DMAP. Use of 1-adamantylamine led to bis(amido) complex $[Tp^{fBu,Me}Ho(HNAd)_2]$ (3-Ho). Nucleophilic attack of acetonitrile by Dy-CH₃ affords 7-Dy.

the similar sized yttrium and holmium differ in the hapticity of the Tp^{tBu,Me} ligand and in the Ln1–C26–Ga1 angle, which is more linear for the holmium derivative (174.9(2) vs. 163.3(1)°). Packing effects of co-crystallizing toluene in **1-Ho**^{Ga} have presumably a major impact on the coordination of the GaMe₄ moiety and might also cause the bending of one pyrazolyl moiety toward the rare-earth metal centre. The pyrazolyl nitrogen atoms exhibit Ho–N interatomic distances ranging from 2.334(3) to 2.376(3) Å with an additional close contact to the tilted pyrazolato ligand (Ho···N5, 2.859(3) Å, see Fig. S3†). The proposed mechanism for the formation of **1-Ln**^{Ga} includes the preformation of [Tp^{tBu,Me}Ln(GaMe₄)₂] under release of methane and trimethylgallium, and the elimination of a second molecule GaMe₃ sterically induced by the bulky Tp^{tBu,Me} ligand.

The primary aniline H_2NAr^{iPr} ($Ar^{iPr} = C_6H_3iPr_2-2.6$) has proven a privileged imido ligand precursor in rare-earth-metal chemistry. The molecule not only features the right balance of adequately acidic protons, but also a sufficient steric protection with bulky substituents in the positions 2 and 6 at the aryl group. The reaction of bis(alkyl) **1-Ln**^{Ga} with H_2NAr^{iPr} to form the mixed methyl/primary amido complexes $Tp^{fBu,Me}LnMe(HNAr^{iPr})$ **2-Ln** (Ln = Y, Dy, Ho) is clearly visible by the elimination of methane and displacement of trimethylgallium (Scheme 2).

Complexes **2-Dy** and **2-Ho** are isostructural but crystallize in different space groups (**2-Dy**: $P2_1/c$; **2-Ho**: $P\overline{1}$; for the crystal structures and detailed metrics, see the ESI, Fig. S4/S5†). Both compounds are insoluble in n-hexane, but dissolve in aromatic solvents such as toluene. The ancillary ligand coordinates in a κ^3 fashion (N, N', N'') with considerably varying Ln–N_{pz} distances (**2-Dy**: 2.3752(17)–2.5163(18) Å; **2-Ho**: 2.369(2)–2.536(2) Å). The Ln–N_{amido} (**2-Dy**: 2.212(2) Å; **2-Ho**: 2.222(2) Å) and the Ln–CH₃ (**2-Dy**: 2.436(2) Å; **2-Ho**: 2.427(3) Å) interatomic distances lie the range of the simlar mixed methyl/amido complexes Tp^{tBu,Me}LuMe(HNAr^R) (Ar^R = Ar^{Me_2} = C₆H₃Me₂-2,6: Lu–N_{amido} 2.189(2) Å, Lu–CH₃ 2.369(2) Å; Ar^R = Ar^{CF_3} = C₆H₃(CF₃)₂-3,5: Lu–N_{amido} 2.215(1) Å, Lu–CH₃ 2.360(1) Å). ¹² The

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Ln–N_{amido}–C_{ipso} angle spans a wide range from 142.5(1)° (Ln = Lu, Ar^R = Ar^{CF3}), ¹² 153.8(1)° (Ln = Lu, Ar^R = Ar^{Me2}) ¹² over 155.84(16)° (Ln = Dy, Ar^R = Ar^{iPr}) to 160.28(19)° (Ln = Ho, Ar^R = Ar^{iPr}), dependent on the Ln(III) centre and the substitution pattern of the amido ligand. As the paramagnetic nature of dysprosium and holmium impedes conclusive interpretations of their NMR spectra, the yttrium congener 2-Y was accessed from 1-Y^{Ga} and H₂NAr^{iPr}. The ¹H NMR spectrum of 2-Y shows a sharp singlet at 0.46 ppm for the methyl group and a broader singlet at 4.87 ppm for the proton of the amido ligand.

Noteworthy, the reaction of 1-Ho^{Ga} with 1-adamantylamine in a 0.9:1 ratio gave the bis(amido) holmium complex $Tp^{tBu,Me}Ho(HNAd)_2$ (3-Ho, Ad = adamantyl), even though the primary amine was added in deficit. Apparently, the single deprotonation is favored over the second deprotonation, as the less Brønsted acidic second proton is less prone to be abstracted than the first proton of a second primary aniline. According to the present synthesis protocol, so far only sufficiently acidic aniline derivatives lead to the successful isolation of terminal rare-earth-metal imides, as other, less bulky and less electronically advantageous amines only form bis(amido) complexes, 3,66c or in case 1-Ln^{Al} are employed result in trimethylaluminiumstabilized imide species.200 Like its precursors, complex 3-Ho is insoluble in aliphatic solvents, but readily dissolves in toluene and THF. The crystal structure of 3-Ho (triclinic $P\bar{1}$ space group) shows the expected κ^3 fashion (N, N', N'') of the ancillary ligand with Ho-N_{DZ} distances in the range of 2.393(3)-2.610(3) Å (Fig. 1).

The Ln–N_{amido} distances (\emptyset 2.171 Å) are longer compared to the Ln–N_{imido} interactions in Tp^{rBu,Me}Ho(μ_2 -NAd)AlMe₃ (2.087(2) Å),^{20 α} but slightly shorter compared to the mixed methyl/amido complexes **2-Ln** (Ln = Dy, Ho) described beforehand. This reflects a better electron donation of two nitrogen atoms to the metal centre, despite increased steric bulk. However, the angles Ho1–N7–C35 (144.0(3)°) and Ho1–N8–C25

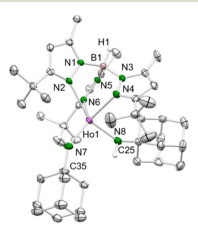


Fig. 1 Crystal structure of **3-Ho**. All atoms are represented by atomic displacement ellipsoids set at 50% probability. Solvent molecules and hydrogen atoms except for those of B–H and N–H are omitted for clarity. Selected interatomic distances (Å) and angles (°): Ho1–N7 2.172(3), Ho1–N8 2.170(3); Ho1–N7–C35 144.0(3), Ho1–N8–C25 148.5(3) (for further metrics, see ESI†).

(148.5(3)°) are significantly more bent compared to the methyl/anilido complex **2-Ho** (160.28(19)°).

Ln(III) imide synthesis

Treatment of the mixed methyl/anilido complexes 2-Ln with the Lewis base DMAP led to the isolation of the targeted terminal lanthanide imide complexes Tp^{tBu,Me}Ln(NAr^{iPr})(dmap) (4-Ln: Ln = Y, Dy, Ho). DMAP was used previously for the synthesis of terminal rare-earth-metal imide complexes,8,10-12 exploiting its strong donor capacity (versus e.g., THF) to induce the elimination of methane via inner-sphere deprotonation of the amido species. We assume, that the preceding coordination of DMAP to the metal centre affects the geometry of its coordination sphere in a way that the methyl group and the amido proton come into close proximity and finally evolve methane, being a very potent leaving group. The ¹H NMR spectrum of 4-Y evidences the deprotonation of the amido functionality via the methyl ligand since both the N-H and methyl signal disappeared (see Fig. S21†). Complexes 4-Ln are insoluble in nonpolar solvents (e.g. n-hexane), but easily dissolve in aromatic or polar solvents like toluene and THF.

While the yttrium complex 4-Y could not be obtained in very pure form and gave only poor crystal quality (connectivity structure only, see ESI Fig. S7†), the dysprosium and holmium congeners displayed good crystallization and diffraction behaviours (isotypic, monoclinic space group C2/c, Fig. 2 and S8/9†). The metal centres are pentacoordinate with the ancillary Tp^{tBu,Me} ligand coordinating in the familiar κ^3 fashion (N, N', N''). The Ln-N_{pz} distances typical of scorpinate ligands show two shorter bonds (4-Dy: 2.450(3)/2.452(3) Å, 4-Ho: 2.426(4)/ 2.436(3) Å) and one longer (4-Dy: 2.517(3) Å, 4-Ho: 2.492(4) Å). The Ln-N_{imido} distances are in line with those of other terminal imides, considering the changes in the ionic radii (Table 1). Except for the anionic ceric complex [(TriNOx)Ce(NAr^{CF₃}) [Cs(2.2.2-cryptand)] (157.3(4)°), 14a the Ln-N $_{\rm imido}$ -C $_{\rm ipso}$ angles of all trivalent terminal imides are larger than 165° and almost linear (4-Dy: 166.0(2)°; 4-Ho: 166.7(3)°; Table 1). Exchange of the donor ligand dmap for thf seems to entail a shortening of the Ln-N_{imido} bond but doesn't appear to affect the Ln-N_{imido}-C_{ipso}

In contrast, and as pointed out previously, the reaction of the primary aniline $H_2NAr^{Me_2}$ ($Ar^{Me_2} = C_6H_3Me_2$ -2,6) with the aluminium congeners $Tp^{tBu,Me}LnMe(AlMe_4)$ (1- Ln^{Al}) implies the formation of trimethylaluminium-stabilized imide complexes. ¹² These complexes cannot be converted into unsupported terminal rare-earth-metal imide complexes by applying Lewis bases such as 1,4-dioxane, pyridine, DMAP, or TMEDA (N,N,N',N'-tetramethylethylenediamine). Complex 1- Dy^{Al} is accessible from $Dy(AlMe_4)_3$ (ref. 24) and $H[Tp^{tBu,Me}]_{,22}$ in analogy to the yttrium and holmium complexes reported previously. ²⁰ In order to probe the effect of the substituents on the aniline, compounds 1- Ln^{Al} (Ln = Dy, Ho) were reacted with H_2NAr^{iPr} , H_2NAr^{Me2} , and H_2NAr^{Me3} ($Ar^{Me3} = C_6H_2Me_3$ -2,4,6) (Scheme 3).

Contrary to the reaction with the weaker coordinated trimethylgallium in complexes 1-Ln^{Ga}, the 1-Ln^{Al}/H₂NAr^{iPr}

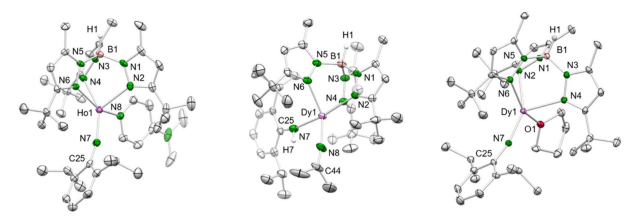


Fig. 2 Left: Crystal structure of 4-Ho. All atoms are represented by atomic displacement ellipsoids set at 50% probability. Solvent molecules, and hydrogen atoms except for that of B-H are omitted for clarity. For selected interatomic distances and angles, see Table 1 and ESI.† Middle: Crystal structure of 7-Dy. All atoms are represented by atomic displacement ellipsoids set at 50% probability. Solvent molecules and hydrogen atoms except for those of B-H and N-H are omitted for clarity. Selected interatomic distances (Å) and angles (°): Dy1-N7 2.219(3), Dy1-N8 2.148(3); Dy1-N7-C25 156.3(3), Dy1-N8-C44 165.3(3) (for further metrics, see ESI†). Right: Crystal structure of 8-Dy. All atoms are represented by atomic displacement ellipsoids set at 50% probability. Only one molecule of the asymmetric unit is shown. Solvent molecules, and hydrogen atoms except for that of B-H are omitted for clarity. For selected interatomic distances and angles, see Table 1 and ESI.†

reaction was inconclusive. Like in the case of yttrium,12 the demanding anilines sterically less gave trimethylaluminium-stabilized imides $Tp^{\ell Bu,Me}Ln(\mu_2-NAr^{Me2})$ AlMe₃ (5-Ln: Ln = Dy, Ho) and $Tp^{tBu,Me}Ho(\mu_2-NAr^{Me3})AlMe_3$ (6-Ln). The coordinated trimethylaluminium is not removable, neither under vacuum, nor with Lewis bases (e.g., DMAP, THF). Presumably, after the first methane elimination and the coordination of the primary amido functionality to the metal centre, one of the methyl groups at the $[\mu_2$ -MeAlMe₃] unit abstracts the second amido proton via release of another molecule of methane, resulting in 5-Ln. Compounds 5-Ln are insoluble in aliphatic solvents, but dissolve in aromatic and polar solvents. The isostructural complexes crystallize in different space groups (5-Dy: monoclinic, $P2_1/n$; 5-Ho: triclinic, $P\overline{1}$; 6-Ho: monoclinic, Cc; Fig. 3, S10 and S11†).

The bulky $Tp^{tBu,Me}$ ligand coordinates again in the κ^3 fashion (N, N', N'') with interatomic Ln-N distances in the range of 2.423(2)-2.492(2) Å (5-Dy), 2.407(4)-2.437(4) Å (5-Ho) and 2.394(3)-2.454(3) Å (6-Ho). The central metal ion is pentacoordinate and the Ln-N_{imido}-C_{ipso} angle is strongly bent (5-Dy: 146.68(17)°; 5-Ho: 146.7(4)°; 6-Ho: 151.0(2)°), which is due to the interaction with the Lewis acid trimethylaluminium. Hence, the electronic situation differs considerably compared to the terminal rare-earth-metal imide complexes 4-Ln (Ln = Dy, Ho) as electron density of the imido nitrogen is shifted to the empty p orbitals of the aluminium ion. This is also reflected in the Ln $m N_{imido}$ bonds of **5-Ln** and **6-Ho** which are elongated by *ca.* 0.1 Å (5-Dy: 2.129(2) Å; 5-Ho: 2.116(4) Å; 6-Ho: 2.116(4) Å), matching those in $Tp^{tBu,Me}Y(\mu_2-NAr^{Me})AlHMe_2$ (Y-N_{imido}: 2.133(2) Å; Y-N_{imido}-C_{ipso}: 145.1(2)°).25 Further structural comparison with

Table 1 Selected metrical parameters of terminal rare-earth-metal imides

Compound	$Ln{=}N_{\rm imido}/\mathring{A}$	Ln-do/Å	$Ln-N_{imido}-C_{ipso}/deg$	$IR^d/Å$ (ref. 23)	CN	Ref.
${ m Tp}^{t{ m Bu},{ m Me}}{ m Lu}({ m NAr}^{{ m CF}_3})({ m dmap})$	1.993(5)	2.377(5)	175.8(5)	0.861	5	12
$Tp^{tBu,Me}Y(NAr^{Me_2})(dmap)$	2.024(4)	2.426(4)	173.6(4)	0.900	5	12
$\operatorname{Tp}^{t\operatorname{Bu},\operatorname{Me}}\operatorname{Ho}(\operatorname{NAr}^{\operatorname{iPr}})(\operatorname{dmap})$ (4-Ho)	2.012(4)	2.429(4)	166.7(3)	0.901	5	This work
$Tp^{tBu,Me}Dy(NAr^{iPr})(dmap)$ (4-Dy)	2.017(3)	2.450(3)	166.0(2)	0.912	5	This work
$Tp^{tBu,Me}Dy(NAr^{iPr})(thf) (8-Dy)^a$	2.008(3)/2.004(4)	2.397(3)/2.392(3)	166.9(3)/165.3(3)	0.912	5	This work
$Tp^{tBu,Me}Sm(NAr^{iPr})(thf)_2$	2.067(5)	2.527(5)/2.560(4)	169.3(5)	0.958	6	15
$Tp^{tBu,Me}Nd(NAr^{iPr})(thf)_2$	2.076(4)	2.557(4)/2.594(3)	169.2(4)	0.983	6	15
$\operatorname{Tp}^{t\operatorname{Bu},\operatorname{Me}}\operatorname{Nd}(\operatorname{NAr}^{\operatorname{iPr}})(\operatorname{thf})^a$	2.036(7)/2.047(7)	2.517(6)/2.511(6)	165.8(4)/163.2(6)	0.983	5	15
$\operatorname{Tp}^{t\operatorname{Bu},\operatorname{Me}}\operatorname{Ce}(\operatorname{NAr}^{\operatorname{iPr}})(\operatorname{thf})_2$	2.101(5)	2.599(3)/2.628(3)	171.3(3)	1.01	6	15
(nacnac ^{R1})Sc(NAr ^{iPr})(dmap) ^b	1.881(8)	2.271(5)	169.6(5)	0.745	5	8
(nacnac ^{R1})Sc(NAr ^{iPr})(thf) ^b	1.852(4)	2.251(3)	168.6(3)	0.745	5	11
(nacnac ^{R2})Sc(NAr ^{iPr})(dmap) ^c	1.8591(18)	2.369(2)	167.90(17)	0.745	5	9
$[(PhN = Ph_2P)_2N]Sc(NAr^{iPr})(dmap)_2$	1.853(3)	2.379(3)/2.326(3)	168.8(3)	0.745	6	10
(BPz ₂ Py ₃)Sc(NAr ^{iPr})	1.877(3)	_ ``	173.1(3)	0.745	6	13
$[(TriNOx)Ce(NAr^{CF_3})][Cs(2.2.2-cryptand)]$	2.077(3)	_	157.3(4)	0.87	8	14 <i>a</i>
[(AdTPBN ₃)Ce(NAr ^{CF₃})][K(2.2.2-cryptand)]	2.0742(4)	_	176.3(4)	<0.87	4	18

^a Two molecules in the asymmetric unit. ^b nacnac^{R1} = $[Ar^{iPr}NC(Me)CHC(Me)N(CH_2)_2NMe_2]$. ^c nacnac^{R2} = $[Ar^{iPr}NC(Me)CHC(Me)]$ $N(CH_2)_2N(CH_2)_2NMe_2$]. ^d Effective ionic radii.

Scheme 3 Synthesis of Lewis acid stabilized, bimetallic imides $Tp^{tBu,Me}Ln(NAr^{Me2})(\mu_2-MeAlMe_3)$ (5-Ln; Ln = Dy, Ho) and $Tp^{tBu,Me}Ho(NAr^{Me3})(\mu_2-MeAlMe_3)$.

the similar holmium imides $Tp^{tBu,Me}Ho(\mu_2\text{-NR})AlMe_3$ (R = tBu, adamantyl) clearly indicate a more pronounced $Ln-N_{imido}$ interaction for the latter (Ho- N_{imido} : 2.083(2), 2.087(2) Å; Ho- N_{imido} - C_{ipso} : 140.4(2), 140.2(1)°). ^{20a} Other Lewis acid supported monomeric rare-earth-metal imide complexes include Mindiola's (PNP)Sc(μ_2 -NAr^{iPr})(μ_2 -Me)AlMe₂ [PNP=N(2-P-(CHMe₂)₂-4-methylphenyl)₂] (ref. 26) or $Tp^{tBu,Me}Ln(NAr^{iPr})(MMe_3)$ (Ln = Ce, Nd, Sm; M = Al, Ga) from our group. ^{6c}

Probing donors other than DMAP

N,N-Dimethyl-4-aminopyridine (DMAP) emerged as a most valuable donor for forcing the elimination of methane *via* innersphere deprotonation of the primary amido species in **2-Ln**-type complexes. In order to assess the impact of the donor molecule on the Ln–N_{imido} bonding, the implementation of other donor ligands was examined. In general, donor ligands might be introduced according to Scheme 1 promoting methane elimination (route A) or *via* post-imide-synthesis exchange (Scheme 4). Since imido ligand formation according to route A could not be achieved with ethereal donors such as OEt₂ or THF, complex **2-Dy** was treated with N-donors TMEDA (tetramethylethylene-diamine), TMPDA (tetramethylpropane-1,3-diamine) and

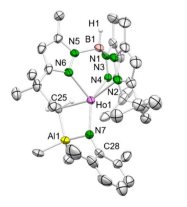
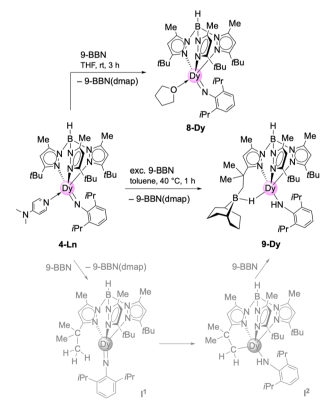


Fig. 3 Crystal structure of **5-Ho**. All atoms are represented by atomic displacement ellipsoids set at 50% probability. Solvent molecules, and hydrogen atoms except for that of B–H are omitted for clarity. Selected interatomic distances (Å) and angles (°): Ho1–N7 2.116(4), Ho1–C25 2.549(5), Al1–C25 2.105(6); Ho1–N7–C28 146.7(4) (for isostructural **5-Dy** and **6-Ho** and further metrics, see ESI†).

acetonitrile. While the potentially bidentate diamines did not undergo any reaction yielding only in the isolation of the starting compounds, acetonitrile formed the insertion complex $Tp^{tBu,Me}Dy[NC(Me)_2](HNAr^{iPr})$ (7-Dy, Scheme 2). It was previously shown that acetonitrile does react with rare-earth-metal alkyls either via C-H-bond activation/deprotonation or insertion. 27-29 While the product of the deprotonation reaction (C₅Me₅)₂La[CH(SiMe₃)₂]/CH₃CN was structurally characterized as $[(C_5Me_5)_2La(\mu\text{-CH}_2CN)]_2$, 27 the insertion product of the reaction (C5Me5)2ScCH3/CH3CN was only spectroscopically analyzed as (C₅Me₅)₂Sc[NC(Me)₂].²⁸ The SCXRD study of five-coordinate 7-Dy features distinct bonding behaviour of the primary amido and the dimethyliminato ligand (Dy-N: 2.219(3) versus 2.148(3) Å; Dy-N-C: 156.3(3) versus 165.3(3)°) (Fig. 2). Thus, the dimethyliminato coordination compares to that of imidazolin-2iminato complexes like $LY(CH_2SiMe_3)_2(thf)_2$ (L = 1,3-bis(2,6diisopropylphenyl)imidazoline-2-iminato; Y-N, 2.1255(13) Å, Y-N-C, 176.85(12)° or LYCl₂(thf)₃ Y-N, 2.1278(18) Å, Y-N-C, 174.35(16)°) supposedly featuring very short Ln-N_{iminato} bonds.30,31

The post-synthesis exchange approach (route 2, Scheme 4) was probed with **4-Dy** and the strong Lewis acid 9-borabicyclo [3.3.1]nonane (9-BBN) in THF.¹¹ Accordingly, the equimolar reaction led to the displacement of DMAP and coordination of one THF molecule to the dysprosium centre in Tp^{tBu,Me}Dy(-NAr^{iPr})(thf) (8-Dy, Scheme 4). Like the dmap adduct, **8-Dy** is soluble in toluene and THF, but insoluble in aliphatic solvents.



Scheme 4 Reactivity of $Tp^{tBu,Me}Dy(NAr^{iPr})(dmap)$ (4-Dy) (Ln = Y, Dy) towards Lewis acid 9-BBN in THF and toluene.

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It crystallized in the orthorhombic space group $\mathit{Pna2}_1$ and shows the same κ^3 coordination of the $\mathrm{Tp}^{\ell Bu,Me}$ ligand as complex 4-Dy before (Dy-N_{pz}: 2.465(4)–2.510(4)/2.450(3)–2.539(4) Å) (Fig. 2). As the coordination number did not change, the angle of the imido functionality stayed nearly the same (Dy-N_{imido}-C_{ipso}: 165.2(3)/166.9(3)°) as detected for 4-Dy. However, the Dy-N_{imido} distance of 2.004(4)/2.008(3) Å) in 8-Dy appears to be slightly shorter compared to 4-Dy (Table 1) which can be attributed to the weaker donor properties of the thf ligand.

The 9-BBN-promoted donor exchange was previously introduced by Chen, revealing that prior activation of $L^1Sc(NAr^{iPr})(dmap)$ ($L^1 = [Ar^{iPr}NC(Me)CHC(Me)N(CH_2)_2NMe_2]$) with 9-BBN led to abstraction of the donor molecule DMAP. The emerging donor-free imide intermediate $[L^1Sc(NAr^{iPr})]$ could be trapped with THF to afford $L^1Sc(NAr^{iPr})(thf)$ featuring also a shorter Sc- N_{imido} bond than the DMAP adduct (Table 1: 1.852(4) *versus* 1.881(8) Å). It was also mentioned that the direct synthesis of the THF adduct $L^1Sc(NAr^{iPr})(thf)$ is not possible by thermolysis of the mixed methyl/amido scandium complex in THF, which is the same case for 8-Dy as well. In contrast, terminal imides $Tp^{tBu,Me}Ln(NAr^{iPr})(thf)_2$ of the larger rare-earth metals can be obtained directly according to the donor(THF)-assisted methylidene \rightarrow imido transformation (Scheme 1 and Table 1).

Treatment of **4-Dy** with excess of 9-BBN in toluene gave the mixed primary amido/hydroborato complex **9-Dy** (Scheme 4 and Fig. 4). Again, this is in line with the observation made by Chen with the system $L^2Sc(NAr^{iPr})/9$ -BBN ($L^2 = [Ar^{iPr}NC(Me)CHC(Me)N(CH_2)_2N(CH_2)_2NMe_2]).^{32}$ Correspondingly, it can be hypothesized that initially the strong Lewis acid 9-BBN displaces all of the coordinated DMAP, rendering a highly reactive donor-free terminal imide $[Tp^{tBu,Me}Dy(NAr^{iPr})]$ (Scheme 4, intermediate I^1 , lower trace). Subsequent 1,2-addition of a tBu methyl group (C-H-bond activation) across the highly reactive Dy- N_{imido} bond of transient species I^1 reforms the primary amido ligand along with a 5-membered metallacycle in I^2 . Then, the highly nucleophilic

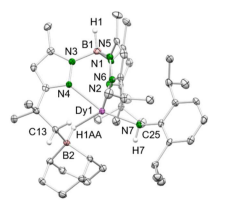


Fig. 4 Crystal structure of 9-Dy. All atoms are represented by atomic displacement ellipsoids set at 50% probability. Solvent molecules and hydrogen atoms except for those of B-H and N-H are omitted for clarity. Selected bond distances (Å) and angels (°) for 9-Dy: Dy1-N7 2.237(3), Dy 1-N2 2.452(3), Dy 1-N4 2.405(3), Dy 1-N6 2.461(3), Dy 1-B2 2.703(4); Dy1-H1AA 2.17(3), B2-H1AA 1.23(3); Dy1-N7-C35 143.4(2), N7-Dy1-B2 117.06(11), N7-Dy1-N4 161.07(10), N7-Dy1-N2 93.18(10), N7-Dy1-N6 92.04(10).

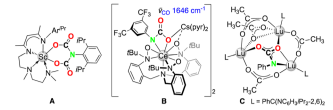


Fig. 5 Structurally authenticated CO_2 -insertion products A_1^{17a} B_2^{38} and C (ref. 17d) emerged from rare-earth-metal imides.

alkyl attached to the dysprosium attacks a second molecule of 9-BBN to afford the alkylhydroborato moiety. The resulting Dy-N_{amido}-C_{ipso} angle (143.4(2)°) and the Dy-N_{amido} distance (2.237(3) Å) of **9-Dy** are comparable to the dysprosium amide complexes discussed beforehand. The Dy-B distance of 2.703(4) Å is in the range of the Y-B distances in [(Me₃Si)₂NC(NiPr)₂]-Y[μ -H(μ -Et)₂BEt]₂(thf)₂ (2.658(4) and 2.671(4) Å)³³ and (C₅Me₅)₂YH(9-BBN) (2.767(6) Å),³⁴ but longer than those observed in (tBu₄Carb)Dy(BH₄)₂(thf) (2.473(2) and 2.487(2) Å).³⁵

In Chen's system $[L^2Sc(NAr^{iPr})]/9$ -BBN the respective $C(sp^3)$ -H bond borylation took place at the $[-CH_2NMe_2]$ side arm of the ancillary nacnac ligand $(Sc-H_{hydrido}, 2.01(3) \text{ Å}).^{32}$ Chen also reported on the reaction of $L^1Sc(NAr^{iPr})$ (dmap) with three equivalents of 9-BBN which led to mixed boroamido/hydroborato complex $L^1Sc[NAr^{iPr}(9$ -BBN-H)](9-BBN + H) $(Sc-H_{hydrido}, 1.89 \text{ Å} \text{ and } 1.20 \text{ Å}).^{11}$

Preliminary studies on the reactivity of terminal imide complex 4-Dy towards carbon dioxide (1 bar) in toluene at ambient temperature clearly indicate CO2 insertion into the Dy-Nimido bond and, hence, carbamate formation in product 10-Dy. The DRIFT spectrum of 10-Dy revealed typical carbonyl vibrations at 1701 cm⁻¹ and 1641 cm⁻¹ assigned to asymmetric and symmetric C=O stretching vibrations (Fig. S17†). Moreover, the B-H vibration of **4-Dy** at 2557 cm⁻¹, typical for the terminal B-H stretch of the Tp^{tBu,Me} ligand when coordinated in a tridentate fashion,³⁶ changed to 2431 cm⁻¹ in 10-Dy. Such a dramatic change of the position of the B-H band can be ascribed to a $\kappa^3 \rightarrow \kappa^2$ coordination switch of the Tp^{tBu,Me} ligand, likely involving a Dy···H-B interaction.37 Unfortunately, crystals of 10-Dy suitable for SCXRD analysis could not be obtained, but 10-Dy can be tentatively assigned as the dicarboxylate species [Tp^{tBu,Me}Dy{(O₂C)₂NAr^{iPr}}] on the basis of the double CO2-insertion chemistry of the terminal scandium imides L²Sc(NAr^{iPr}) (A, Fig. 5)^{17a} and (BPz₂Py₃)Sc(NAr^{iPr}) $(\tilde{\nu} = 1685 \text{ and } 1632 \text{ cm}^{-1})^{13}$ For further comparison, mono insertion of carbon dioxide was observed for the Lu₃(µ₃-NPh) moiety of the trinuclear cluster $[L^3Lu_3(\mu_2-Me)_3(\mu_3-Me)(\mu_3-NPh)](L^3$ = $PhC(NC_6H_3iPr_2-2,6; C, Fig. 5)^{17d}$ and the alkali-metal stabilized Ce(IV)-N_{imido} bond of [(TriNOx)Ce(NAr^{CF3})][Cs(2.2.2-cryptand)] (B, Fig. 5).38

Conclusion

The successful application of the donor-assisted intramolecular primary amido deprotonation protocol towards terminal imides of the rare-earth metals yttrium, dysprosium, and holmium depends on two crucial factors: first, the kinetic stabilization of the targeted imide via appropriate steric shielding of the ancillary ligand, herein supplied by the bulky ${
m Tp}^{t{
m BuMe}}$ scorpionate ligand; second, sufficient Brønsted acidity and steric demand of the employed primary amine, herein provided by H_2NAr^{iPr} ($Ar^{iPr} = C_6H_3iPr_2-2.6$). With these key factors in mind, it was possible to generate a series of terminal rare-earth-metal imides Tp^{tBu,Me}Ln(NAr^{iPr})(dmap) for mid-sized to small rareearth metals. Their reactivity parallels that of terminal scandium imides, as revealed by treatment with the strong Lewis acid 9-borabicyclo[3.3.1]nonane (9-BBN) and the heteroallene CO2. These reactions revealed effective DMAP/THF donor exchange and CO2 insertion into the Ln-Nimido bond (carbamate formation). The reactivity studies also include the isolation and structural characterization of the bis(amido) species $Tp^{tBu,Me}Ho(NAd)_2$ (Ad = adamantyl) and dimethyliminate complex $Tp^{tBu,Me}Dv[NC(Me)_2](HNAr^{iPr})$.

Data availability

Experimental, spectroscopic and structural data supporting this article have been uploaded as part of the ESI. Crystallographic data for all compounds have been deposited at the CCDC under 2312212–2312226, and can be obtained from https://www.ccdc.cam.ac.uk/structures/.

Author contributions

TR, synthesis and characterization of compounds, writing original draft; DS, synthesis and characterization of compound **1-Ho**^{Ga}, editing original draft; CM-M, crystallography, editing original draft; RA, conceptualization, supervision, writing and project administration, funding acquisition.

Conflicts of interest

There are no conflicts to declare.

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