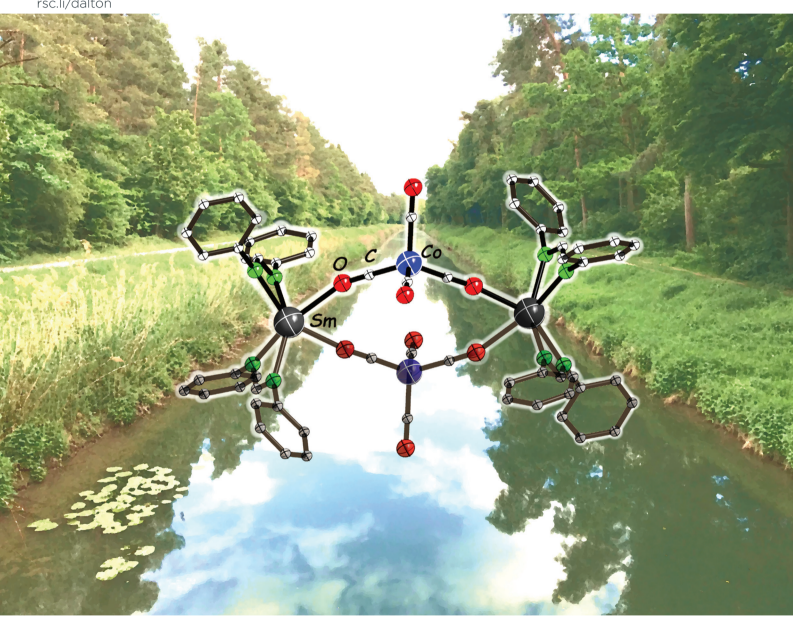
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Introduction

Heterometallic complexes have gained widespread attention due to the viable synergistic effect that may result from a judicious choice of metal centres.^{1,2} Ln-TM (TM = transition metal) heterometallic complexes have especially attracted interest due to their magnetic,³⁻⁵ and photophysical properties,^{6,7} as well as their catalytic activity.^{8,9} Among them, Ln-TM carbonyl complexes are a major class of compounds from both fundamental and application points of view.¹⁰⁻¹² For example, Dy-TM carbonyl complexes have demonstrated single-molecule magnetic behaviour.13,14 Andersen and coworkers have synthesized several Yb^{III}-TM isocarbonyl complexes, [Yb^{III}-OC-TM], by reduction of TM carbonyl complexes with [Cp*₂Yb^{II}(OEt₂)].¹⁵⁻¹⁸ Edelmann and co-workers have isolated the formally Sm^{III}-Fe⁰ complex [Cp*₂Sm(µ-OC)₂FeCp*]₂ by reducing $[Cp*Fe(CO)_2]_2$ with $[Cp*_2Sm(thf)_2]$.¹⁹ Some of us isolated the elusive $[W_2(CO)_{10}]^{2-}$ anion in a mixed-valent Sm^{II/III}

3d–4f heterometallic complexes by the reduction of transition metal carbonyls with bulky Ln^{II} amidinates†

Ravi Yadav, 🐌 ^a Md Elius Hossain, 🐌 ^b Ramees Peedika Paramban, 🐌 ^b Thomas Simler, 🐌 ^a Christoph Schoo, 跑 ^a Jun Wang, ^b Glen B. Deacon, 跑 ^c Peter C. Junk 跑 ^b and Peter W. Roesky 跑 *^a

The redox chemistry between divalent lanthanide complexes bearing bulky amidinate ligands has been studied with 3d transition metal carbonyl complexes (iron and cobalt). The reaction of $[(DippForm)_2Sm^{II}(thf)_2]$ (DippForm = N,N'-bis(2,6-diisopropylphenyl)formamidinate) with $[Co_2(CO)_8]$ resulted in the formation of a tetranuclear Sm-Co complex, $[{(DippForm)_2Sm^{III}(thf)}_2({(\mu-CO)_2Co(CO)_2}_2]$. The product of the reaction of $[(DippForm)_2Yb^{III}(thf)_2]$ and $[Co_2(CO)_8]$ gives the dinuclear Yb-Co complex $[{(DippForm)_2Yb^{III}(thf)}_{(\mu-CO)Co(CO)_3}]$ in toluene. The reaction of $[(DippForm)_2Sm^{III}(thf)_2]$ was also carried with the neighbouring group 8 carbonyl complexes $[Fe_2(CO)_9]$ and $[Fe_3(CO)_{12}]$, resulting in a pentanuclear Sm^{III}-Fe complex, $[{(DippForm)_2Sm^{III}}_2((\mu_3-CO)_2Fe_3(CO)_9]]$, featuring a triangular iron carbonyl cluster core.

calix[4]pyrrolide sandwich by the reduction of $[W(CO)_6]$ with a divalent samarium *meso*-octaethylcalix[4]pyrrolide.²⁰ Also, Ln^{II}-TM carbonyl complexes have been accessed either by redox-transmetallation between Hg salts of TM carbonyl complexes and elemental Ln⁰ or by reduction of TM carbonyl complexes with Ln/Hg amalgam.^{10,21-26} Depending on the bonding situation between the two metal centres, Ln-TM carbonyl complexes can be divided into three major categories: (i) solvent-separated ion pairs,^{22,27} (ii) Ln-TM bonded compounds,^{11,12,14,23,28,29} and (iii) compounds with isocarbonvl-linkages^{10,16,30} between both metals. Isocarbonyl bridged Ln-TM complexes are the major representatives, which can be explained by the oxophilic nature of lanthanide cations and the additional stabilisation of the negative charge on the TM centre through π back-donation from TM(d_{π}) to C–O(π^*).²⁶ Usually, the reaction of divalent lanthanide complexes with TM-carbonyls leads to isocarbonyl-bridged systems, resulting from a single-electron transfer (SET) to the TM carbonyl moiety and leading to either cleavage or formation of TM-TM bonds.

The chemistry of divalent lanthanide complexes is dominated by the use of cyclopentadienyl-based ligands.^{31–33} Investigating the effects of the nature of the supporting ligands, we have examined the reactivity of the divalent lanthanide complexes [(DippForm)₂Ln^{II}(thf)₂] (Ln = Sm,³⁵ Yb³⁶) in the reduction of main group elements and compounds. By using different electronic and steric environments around divalent lanthanide elements, a different reactivity can be

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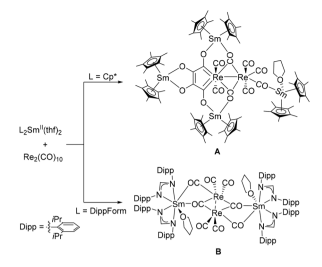
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^aInstitute of Inorganic Chemistry, Karlsruhe Institute of Technology, Engesserstr. 15, 76131 Karlsruhe, Germany. E-mail: roesky@kit.edu

^bCollege of Science and Engineering, James Cook University, Townsville, QLD, 4811, Australia

^cSchool of Chemistry, Monash University, Clayton, Vic, 3800, Australia

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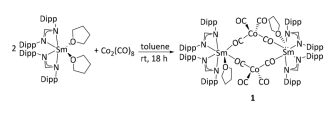
Scheme 1 Reactivity of $[{\rm Re}_2({\rm CO})_{10}]$ towards Sm(11) complexes bearing different ligands. 34

achieved with a same substrate.³⁷⁻⁴⁵ For example, Evans and co-workers have reported the formation of $[{Cp*_2Sm^{III}}_2(S_3)]$ by reaction of [Cp*₂Sm^{II}(thf)₂] with elemental sulfur,⁴⁶ whereas some of us have isolated lanthanide polysulfide coordination clusters, $[(DippForm)_3Ln_3^{III}S_{12}]$ (Ln = Sm and Yb), by using $[(DippForm)_2Ln^{II}(thf)_2]$ (Ln = Sm, Yb) as divalent lanthanide reagents.⁴² In addition, activation of white phosphorous and yellow arsenic by [(DippForm)₂Sm^{II}(thf)₂] led to the formation of $[{(DippForm)_2Sm^{III}}_2(\mu-\eta^4-Pn_4)]$ (Pn = P, As).⁴³ These products contrast with the cage-type molecules $[{Cp*_2Sm^{III}}_4(Pn)_8]$ $(Pn = P,^{37} As,^{44} and Sb^{47})$ that were obtained by reaction of [Cp*₂Sm^{II}] with white phosphorous, nanoscale arsenic, and nanoscale antimony, respectively. We were then interested to extend the study of the influence of the ligands on the reductive behaviour of divalent lanthanide complexes from main group compounds to transition metal complexes. Very recently, some of us have reported the reduction of $[Re_2(CO)_{10}]$ with $[L_2Sm^{II}(thf)_2]$ (L = Cp* or DippForm) and, depending on the nature of the ligands around samarium, were able to isolate either a Fischer-type rhenacycle (A) or a novel $[Re_2(CO)_8]^{2-}$ dianion (**B**) in the coordination sphere of $[L_2Sm^{III}]^+$ moieties (Scheme 1).³⁴ Inspired by the above results, we further explored the reactivity of group 8 and 9 metal carbonyls towards divalent lanthanide complexes supported by the bulky amidinate DippForm ligand and report herein the synthesis and characterisation of the resulting 3d-4f metal complexes obtained.

Results and discussion

Ln-Co carbonyl complexes

The reaction of $[(DippForm)_2Sm^{II}(thf)_2]$ with half an equivalent of $[Co_2(CO)_8]$ in toluene at room temperature resulted in the formation of the heterometallic complex $[{(DippForm)_2Sm^{III}(thf)}_2{(\mu-CO)_2Co(CO)_2}_2]$ (1) (Scheme 2).



Scheme 2 Synthesis of complex 1.

After a short work-up, analytically pure yellow-coloured crystals of complex 1 were grown in a 56% yield by slow evaporation of toluene. The solid-state IR spectrum of complex 1 showed characteristic $\tilde{\nu}_{CO}$ bands at 2020 (m), 1951 (br), 1935 (br), 1922 (br), 1904 (br), 1842 (s), 1819 (s) and 1782 (s) cm⁻¹. The low-frequency stretch at 1782 (s) cm⁻¹ suggests the presence of bridging isocarbonyls between the Sm and Co atoms. The terminal CO stretches from 2020 to 1819 cm⁻¹ and the lowfrequency isocarbonyl stretch at 1782 cm⁻¹ are apparent for complex 1. Similar low-frequency stretches at 1798 (m) and 1761 (s) cm⁻¹ were observed for $[{(Cp^*)_2 Yb^{III}(thf)}]{(\mu-CO)Co(CO)_3}]^{.15}$ Furthermore, the molecular structure of complex 1 was determined by single crystal X-ray crystallography. Complex 1 crystallizes in the monoclinic space group $P2_1/n$ with half a molecule in the asymmetric unit. The solid-state structure showed that two [(DippForm)₂Sm^{III}(thf)]⁺ moieties are bridged by two $[(\mu-CO)_2Co(CO)_2]^-$ units (Fig. 1). The formation of complex 1 can be rationalized by single-electron transfer steps from two [(DippForm)₂Sm^{II}(thf)₂] molecules to one molecule of $[Co_2(CO)_8]$, resulting in the homolytic cleavage of the Co-Co bond and the formation of two $[Co(CO)_4]^-$ anions along with two [(DippForm)₂Sm^{III}(thf)]⁺ cations.

Each samarium atom is heptacoordinated, surrounded by two bidentate amidinate ligands, two bridging isocarbonyls,

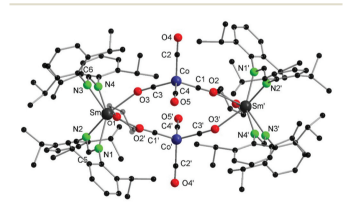
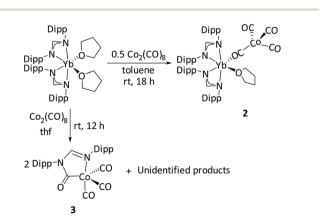


Fig. 1 Molecular structure of 1. Hydrogen atoms are omitted for clarity. Selected bond distances (Å) and angles [°]: Sm–O1 2.422(2), Sm–O2 2.488(3), Sm–O3 2.552(3), Sm–N1 2.431(3), Sm–N2 2.449(3), Sm–N3 2.409(3), Sm–N4 2.449(3), Co–C1 1.733(4), Co–C2 1.787(4), Co–C3 1.747(4), Co–C4 1.782(4), O2–C1 1.175(4), O3–C3 1.170(4), O4–C2 1.138(5), O5–C4 1.145(5), N1–C5 1.322(4), N2–C5 1.318(4), N3–C6 1.330(5), N4–C6 1.326(5); O2'–Sm–O3 67.27(8), N1–Sm–N2 55.35(9), N1–Sm–N4 143.10(10), C1–Co–C2 108.8(2), C1–Co–C3 116.1(2), C1–Co–C4 110.1(2), C3–Co–C2 109.7(2), C3–Co–C4 105.8(2), C4–Co–C2 105.8(2), N2–C5–N1 118.4(3), N4–C6–N3 120.3(3).

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and one coordinating thf. The average Sm-N bond length (2.434(3) Å) in complex **1** is significantly shorter than that in [(DippForm)₂Sm^{II}(thf)₂] (Sm–N(average) 2.573(3) Å). This shortening is consistent with a decrease in the ionic radius of the metal cation upon oxidation of Sm^{II} to Sm^{III}.⁴⁸ Besides, the shortening of the Sm-O1(thf) bond length, from 2.560(3) Å in $[(DippForm)_2Sm^{II}(thf)_2]$ to 2.422(2) Å in 1, further supports the oxidation of the Sm centre to the +3 oxidation state. The cobalt centre in the bridging tetracarbonylcobaltate anion has a distorted tetrahedral coordination geometry with C-Co-C angles ranging from $105.8(2)^{\circ}$ to $116.1(2)^{\circ}$. Owing to the coordination to the $[(DippForm)_2Sm^{III}(thf)]^+$ moiety, the O2-C1 (1.175(4) Å) and O3-C3 (1.170(4) Å) bond distances are longer than the O4-C2 (1.138(5) Å) and O5-C4 (1.145(5) Å) analogues involving terminal CO ligands. A further effect of the formation of bridging isocarbonyls can be seen by analysis of the Co-C bond lengths: the Co-C(bridging) bonds are shortened and strengthened as compared to the Co-C(terminal) bonds (Co-C1 (1.733 (4) Å) and Co-C3 (1.747(4) Å) vs. Co-C4 (1.782(4) Å) and Co-C2 (1.787(4) Å), respectively).¹⁵ To the best of our knowledge, complex 1 is a rare example of a Sm^{III}-Co carbonyl complex featuring bridging isocarbonyls between the two metal centres. Mountford and co-workers have reported the Sm^{III}-Co carbonyl complex $[Sm^{III}{(\mu-CO)Co(CO)_2(PCy_3)}Co(CO)_3(PCy_3)} (thf)_3]$ (Cy = cyclohexyl) by a reaction between SmI_2 with K[Co $(CO)_3(PCy_3)(thf_2)]^{.25}$ Although the reduction of $[CO_2(CO)_8]$ by $[Cp_{2}^{*}Sm^{II}(thf)_{2}]$ has been studied by Evans *et al.*, no solid-state structure of the product has been reported.²⁷ Besides, the reaction between $[Sm^{II}I_2(thf)_2]$ and $[Co_2(CO)_8]$ in thf led to the solvent-separated ion pair, [Sm^{III}I₂(thf)₄][Co(CO)₄], which was structurally characterized.²⁷

The ytterbium analogue of complex 1 was synthesized following a similar procedure (Scheme 3) and orange-coloured crystals were isolated in 63% yield. The IR spectrum of complex 2, [{(DippForm)₂Yb^{III}(thf)}{(μ -CO)Co(CO)₃}], shows $\tilde{\nu}_{CO}$ stretches in the carbonyl region from 2031 to 1748 cm⁻¹, in a similar range to those of [{Cp*₂Yb^{III}}(thf)(μ -CO)Co(CO)₃] (2023 to 1761 cm⁻¹).¹⁵ The lowest stretch at 1748 cm⁻¹ indicates bridging isocarbonyls between the tetracarbonylcobaltate anion and the [(DippForm)₂Yb^{III}(thf)]⁺ moiety. Similar $\tilde{\nu}_{CO}$



Scheme 3 Synthesis of complex 2 and 3 (3 is not fully characterized).

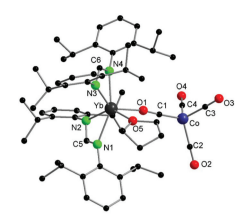


Fig. 2 Molecular structure of 2. Hydrogen atoms are omitted for clarity. Selected bond distances (Å) and angles [°]: Yb–O1 2.248(2), Yb–O5 2.304(2), Yb–N1 2.307(2), Yb–N2 2.337(2), Yb–N3 2.292(2), Yb–N4 2.324(2), Yb–C5 2.697(3), Yb–C6 2.686(3), Co–C1 1.705(3), Co–C2 1.783(5), Co–C3 1.787(4), Co–C4 1.763(3), O1–C1 1.200(3), O2–C2 1.143(5), O3–C3 1.152(5), O4–C4 1.140(4), N1–C5 1.317(3), N2–C5 1.315(3), N3–C6 1.319(3), N4–C6 1.315(3); O1–Yb–O5 81.70(8), N1–Yb–N2 58.23(7), N1–Yb N4 99.80(8), N3–Yb–N1 112.08(8), N3–Yb–N2 114.88(8), C1–Co–C2 111.8(2), C1–Co–C3 115.8(2), C1–Co–C4 109.6(2), C2–Co–C3 107.5(2), C4–Co–C2 104.6(2), C4–Co–C3 106.8(2).

stretches (2017 to 1780 cm⁻¹) have been observed in the related complex [Cp₂Lu(thf){Co(CO)₄}].⁴⁹ The solid-state structure was established by single-crystal X-ray crystallography (Fig. 2). Complex 2 crystallizes in the triclinic space group $P\bar{1}$ with one molecule in the asymmetric unit cell. In contrast to the dimeric form of complex 1, the solid-state structure of complex 2 reveals a monomeric arrangement. A possible reason for this difference could be a combined effect of the smaller ionic radius of Yb³⁺ as compared to Sm³⁺ and the sterically demanding nature of the DippForm ligands.⁴⁸ The ytterbium centre is in a distorted octahedral environment, coordinated by two chelating amidinate ligands, one isocarbonyl O donor, and one thf ligand. In complex 2, the shortening of the Yb-N bond distances involving the amidinate ligands (2.315(2) Å, average) and of Yb-O5 (2.304(2) Å), in comparison to the corresponding separations in $[(DippForm)_2Yb^{II}(thf)_2]$ (Yb-N (2.447 Å) and Yb-O (2.461 Å), average),³⁶ is consistent with the oxidation of the Yb metal centre from the +2 to the +3 oxidation state.⁴⁸ As observed in 1, the cobalt centre in the tetracarbonylcobaltate anion of 2 has a distorted tetrahedral coordination environment with C-Co-C angles varying from 104.6(2)° to 115.8(2)°. The Co–C1 bond length (1.705(3) Å) in 2 is almost the same as the Co–C(bridged) (1.699(3) Å) bond distance in [{Cp*₂Yb^{III}}(thf)(µ-CO)(Co(CO)₃].¹⁵ The average Co-C (terminal) bond lengths in 2 and [{Cp*₂Yb^{III}}(thf)(µ-CO)Co(CO)₃] are also similar (ca. 1.77 Å in both cases).

In a different approach to synthesize complex 2, $[(DippForm)_2Yb(thf)_2]$ was treated with cobalt carbonyl at room temperature in thf for 12 hours. In this case, we did not obtain the bimetallic species 2. Instead, crystals of the metallacyclic cobalt(1) complex [(DippFormCO)(CO)_3Co] (3) were isolated (Scheme 3). Despite our attempts, complex 3 could not

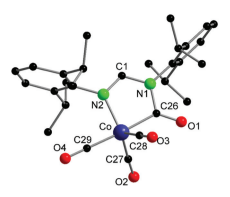
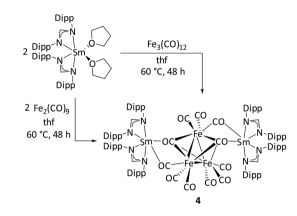


Fig. 3 Molecular structure of 3. Hydrogen atoms and solvent of crystallisation have been omitted for clarity. Selected bond lengths (Å) and angles [°]: Co1–N2 1.968(3), Co1–C26 1.930(4), Co1–C27 1.755(6), Co1– C28 1.755(6), Co1–C29 1.799(5), C1–N1 1.346(5), C1–N2 1.285(3), N1– C26 1.430(5), C26–O1 1.201(5), C27–O2 1.135(6), C28–O3 1.134(7), C29–O4 1.138(6); C26–Co1–N2 82.7(2), C27–Co1–N2 121.6(2), C28– Co1–N2 117.7(2), C29–Co1–N2 93.6(2), C26–Co1–C27 86.2(2), C26– Co1–C28 86.0(2), C26–Co1–C29 176.1(2), C27–Co1–C28 118.5(3), C28–Co1–C29 96.7(2), N1–C1–N2 118.3(4).

be obtained in analytically pure form. However, the isolation of 3 gave some insight into occurring side reactions. Complex 3 was unambiguously identified by single crystal X-ray diffraction studies. Complex 3 crystallizes in the monoclinic space group $P2_1/n$, with one molecule in the asymmetric unit (Fig. 3). This complex consists of a five-coordinate cobalt centre, connected to three carbonyl groups (C=O), one -C(=O)N donor and a nitrogen atom from the N-C-N backbone of the methanide ligand. Previously, Jones and coworkers have reported a very similar complex, [Co{C(=O)N $(Dipp)C(tBu)N(Dipp)(CO)_3]$, by reacting CO gas with an amidinate stabilized cobalt(I) complex.50 In the IR spectrum of complex 3 taken from the reaction mixture bands at 2066 (s), 2009 (s), and 1983 (m) cm⁻¹ can be attributed to C-O stretching frequencies of carbonyl groups. These values have close resemblance to the IR spectrum reported by Jones' group, where they identified 2064 (s), 2004 (s), and 1970 (br) cm^{-1} as C-O stretching frequencies. The bond lengths and angles of complex 3 and [Co{C(=O)N(Dipp)C(*t*Bu)N(Dipp}(CO)₃] are very similar.⁵⁰ The ligand in complex 3 showed a band at 1666 (s) cm⁻¹ corresponding to the ketonic carbonyl -C(=O)N stretching. Complex 3 presumably arises from [(DippForm) Yb^{III}(µ-DippForm)(µ-OC)Co(CO)₃], the latter formed by partial transmetallation from the bimetallic complex 2. The bridging DippForm is bound to the cobalt(1) centre through the imine N atom, providing an 18 electron Co species which undergoes CO insertion into the Co-N bond resulting in complex 3, where the new N,N'-(Dipp)formimidamidomethanide (C(==O)N (Dipp)C(H)N(Dipp) now C,N'-chelates the cobalt.

Sm-Fe carbonyl complex

Further reactivity studies of $[(DippForm)_2Sm^{II}(thf)_2]$ were carried out with iron-carbonyl complexes. The reaction between $[(DippForm)_2Sm^{II}(thf)_2]$ and half an equivalent of $[Fe_3(CO)_{12}]$ or one equivalent of $[Fe_2(CO)_9]$ in thf at



Scheme 4 Synthesis of complex 4 via two routes.

60 °C resulted in the formation of [{(DippForm)₂Sm^{III}}₂ $\{(\mu_3-CO)_2Fe_3(CO)_9\}$ (4) (Scheme 4). Red-coloured crystals were isolated in 33% yield after a short work-up. The solidstate IR spectrum showed characteristic $\tilde{\nu}_{CO}$ absorption bands at 2011(vs), 1979 (vs), 1967 (s), 1878 (m), 1830 (w), and 1696 (w) cm⁻¹. Similar $\tilde{\nu}_{CO}$ absorption bands were observed for $[(NEt_4)_2]$ Fe₃(CO)₁₁] at 1938, 1910, 1890, and 1670 cm⁻¹.⁵¹ The low-frequency band at 1696 cm⁻¹ is characteristic of the occurrence of bridging isocarbonyls. The solid-state structure of complex 4 was unambiguously determined by single crystal X-ray diffraction studies, revealing two [(DippForm)₂Sm^{III}]⁺ moieties associated together by a $[(\mu_3-CO)_2Fe_3(CO)_9]^{2-}$ fragment via bridging isocarbonyls (Fig. 4). In complex 4, each Sm atom is hexacoordinated, surrounded by two bidentate amidinate ligands and two bridging isocarbonyls. Notably, the [(DippForm)₂Sm^{III}]⁺ moiety is free of coordinating thf molecules, which may result from the crystallisation of the complex from hot toluene and also the steric crowding around

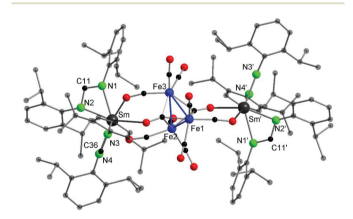
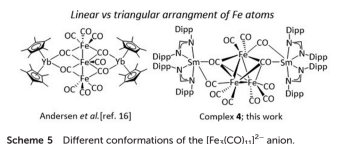


Fig. 4 Simplified view of the molecular structure of 4 in the solid state. H atoms are omitted for clarity. Selected bond distances (Å) and angles [°]: N1–Sm 2.409(2), N2–Sm 2.426(2), N3–Sm 2.381(2), N4–Sm 2.418(2), N1–C11 1.329(3), N2–C11 1.329(3), N3–C36 1.336(3), N4–C36 1.323(3); N1–C11–N2 117.87(2), N4–C36–N3 117.9(2), N1–Sm–N2 56.20(6), N1–Sm–N4 142.11(6), N3–Sm–N1 108.16(6), N3–Sm–N2 109.53(6), N3–Sm–N4 56.65(6). Discussion of the bond lengths and angles for the [Fe₃(CO)₁₁]^{2–} moiety is precluded due to the disorder.



the Sm metal centre. The average Sm-N bond length (2.408(2) Å) in complex 4 is significantly shorter than in $[(DippForm)_2Sm^{II}(thf)_2]$ (Sm-N(average) 2.573(3) Å).³⁵ The $[(\mu_3-CO)_2Fe_3(CO)_9]^{2-}$ moiety is formed by a two-electron reduction of [Fe₃(CO)₁₂] or [Fe₂(CO)₉] by two [(DippForm)₂Sm^{II}(thf)₂] molecules through SET steps. The $[(\mu_3 - CO)_2 Fe_3(CO)_9]^{2-}$ moiety features three Fe atoms arranged in a triangular shape. Upon reaction, formally one CO group in $[Fe_3(CO)_{12}]$ is replaced by a di-negative charge leading to the $[(\mu_3-CO)_2Fe_3(CO)_9]^{2-1}$ anion. In contrast, the related Yb^{III} complex [(Cp*₂Yb^{III})₂ $((\mu_2 - CO)_4 Fe_3(CO)_7)$] isolated by Andersen and co-workers exhibits a linear arrangement for the three Fe atoms (Scheme 5).¹⁶ The possible reason for this different arrangement in the solid state may be the sterically demanding nature of the DippForm ligands. Due to disorder of the $[(\mu_3 - CO)_2 Fe_3(CO)_9]^{2-}$ moiety the bond lengths and angles cannot be precisely discussed (Fig. S8, ESI[†]). Despite several attempts, the product formed upon the reaction between $[Fe_3(CO)_{12}]$ and [(DippForm)₂Yb^{II}(thf)₂] could not be crystallized and its true identification remains unknown. The structural motive of $[(\mu_3-CO)_2Fe_3(CO)_9]^{2-}$ observed in complex 4 is very rare as compared to the $[(\mu-CO)(\mu_3-CO)Fe_3(CO)_9]^{2-}$ unit observed in $[(NEt_4)_2][Fe_3(CO)_{11}]^{.51}$ Previously, the $[(\mu_3-CO)_2Fe_3(CO)_9]^{2-1}$ moiety has been observed in coordination with main group elements such as Li,⁵² Ca,²⁵ C,⁵³ and B.⁵⁴ For example, the Ca complex $[Ca{(\mu_3-CO)_2Fe_3(CO)_9}(MeCN)_4]_n$, was obtained by reduction of [Fe3(CO)12] with Ca amalgam in liquid ammonia.²⁵ In contrast, the reactions of $[Fe_3(CO)_{12}]$ with Yb metal, resulted in $[{(MeCN)_3YbFe(CO)_4}_2 \cdot MeCN]_n$ featuring a Yb-Fe bond,⁵⁵ and that of $[Fe_3(CO)_{12}]$ with $[(Cp*_2Yb(OEt_2)],$ gave [(Cp*₂Yb^{III})₂((µ₂-CO)₄Fe₃(CO)₇)] (Scheme 5).¹⁶

Conclusions

In conclusion, we have studied the reactivity of two divalent lanthanide (Sm and Yb) complexes coordinated by bulky amidinate ligands towards 3d transition metal carbonyl complexes of groups 8 and 9. The reaction of $[(DippForm)_2Sm^{II}(thf)_2]$ with $[Co_2(CO)_{10}]$ resulted in a tetranuclear Sm^{III}–Co heterometallic complex (1) representing a rare example of a Sm^{III}–Co carbonyl complex featuring a bridging isocarbonyl. An analogous reactivity was also observed when $[(DippForm)_2Yb^{II}(thf)_2]$ was reacted with $[Co_2(CO)_{10}]$ in toluene resulting in a dinuclear Yb^{III}–Co heterometallic complex (2). Interestingly, the reaction between $[(DippForm)_2Yb^{II}(thf)_2]$ and $[Co_2(CO)_{10}]$ in thf resulted in a mononuclear cobalt(1) complex (3). The possible reason for the formation of complex 3 when using thf as solvent may be due to solvent induced displacement of DippForm from the Yb centre. The reaction of $[(DippForm)_2Sm^{II}(thf)_2]$ was also carried out with two iron carbonyl complexes, $[Fe_2(CO)_9]$ and $[Fe_3(CO)_{12}]$. In both cases, a pentanuclear Sm^{III}–Fe heterometallic complex (4) was isolated. Complex 4 exhibits a $[(\mu_3-CO)_2Fe_3(CO)_9]^{2-}$ anion with a triangular Fe core. The $[(\mu_3-CO)_2Fe_3(CO)_9]^{2-}$ anion is sandwiched between two $[(DippForm)_2Sm^{III}]^+$ moieties. Interestingly, the $[(\mu_3-CO)_2Fe_3(CO)_9]^{2-}$ anion in complex 4 is in a triangular arrangement which is in sharp contrast to the linear arrangement observed in $[(Cp*_2Yb^{III})_2((\mu_2-CO)_4Fe_3(CO)_7)]^{.16}$

Experimental

General procedures

All the manipulations of air- and water-sensitive reactions were performed with rigorous exclusion of oxygen and moisture in flame-dried Schlenk-type glassware either on a dual manifold Schlenk line, interfaced to a high vacuum (10^{-3} Torr) line or in an argon-filled MBraun glove box. Tetrahydrofuran was distilled under nitrogen from potassium benzophenoneketyl before storage in vacuo over LiAlH₄. Hydrocarbon solvents were dried by using an MBraun solvent purification system (SPS 800) and degassed and stored in vacuo over LiAlH₄. Elemental analyses were carried out with an Elementar vario Micro cube. IR spectra were obtained on a Bruker Tensor 37 spectrometer equipped with a room temperature DLaTGS detector and a diamond ATR (attenuated total reflection) unit. [(DippForm)₂Sm^{II}(thf)₂]^{35,41} and [(DippForm)₂Yb^{II}(thf)₂]³⁶ were prepared according to literature procedure. $[Fe_3(CO)_{12}]$, $[Fe_2(CO)_9]$, and $[Co_2(CO_8)]$ were purchased from commercial suppliers and used as received. No reasonable NMR could be obtained due to low solubility and paramagnetic character.

Synthesis of [{(DippForm)₂Sm^{III}(thf)}₂{(µ-CO)₂Co(CO)₂}₂] (1)⁵⁶

To a mixture of [(DippForm)₂Sm^{II}(thf)₂] (205 mg, 0.20 mmol) and [Co₂(CO)₈] (34 mg, 0.10 mmol) was condensed toluene (15 mL) at -78 °C and then the reaction mixture was stirred for 18 hours at room temperature. The reaction mixture was filtered through P4 frit in a double ampule and flame sealed. Yellow-coloured crystals were grown by slow evaporation of toluene. Crystals were washed carefully with cold toluene and dried under vacuum. Yield: 127 mg, (based on crystals), 0.056 mmol, 56%. Anal. calcd for C₁₁₆H₁₅₆N₈O₁₀Co₂Sm₂ (2241.16): C, 62.17; H, 7.02; N, 5.00. Found: C, 62.34; H, 6.84; N, 4.81. IR (ATR) $\tilde{\nu}$ (cm⁻¹): 2959 (s), 2927 (m), 2866 (m), 2020 (m), 1951 (br), 1935 (br), 1922 (br), 1904 (br), 1842 (s), 1819 (s), 1782 (s), 1665 (vs), 1636 (m), 1587 (m), 1527 (m), 1518 (m), 1464 (s), 1457 (s), 1439 (s), 1383 (s), 1361 (m), 1332 (m), 1314 (m), 1289 (m), 1272 (m), 1255 (m), 1236 (s), 1185 (m), 1107 (m), 1098 (m), 1057 (m), 1043 (m), 1016 (m), 934 (m), 912 (m),

865 (br), 800 (s), 753 (vs), 673 (s), 565 (w), 553 (s), 550 (s), 531 (s) 522 (s), 506 (s) 435 (w).

Synthesis of [{(DippForm)₂Yb^{III}(thf)}{(µ-CO)Co(CO)₃}] (2)⁵⁶

Following the procedure described above for 1, the reaction of $[(DippForm)_2Yb^{II}(thf)_2]$ (209 mg, 0.20 mmol) and $[Co_2(CO)_8]$ (34 mg, 0.10 mmol) afforded orange-coloured crystals of 2. Yield: 146 mg, (based on crystals), 0.127 mmol, 63%. Anal. calcd for $C_{58}H_{78}N_4O_5CoYb$ (1143.27): C, 60.93; H, 6.88; N, 4.90. Found: C, 61.30; H, 6.89; N, 4.90. IR (ATR) $\tilde{\nu}$ (cm⁻¹): 2960 (s), 2927 (m), 2869 (m), 2031 (m), 2016 (m), 1915 (br), 1792 (m), 1748 (w), 1665 (vs), 1636 (m), 1587 (s), 1521 (m), 1465 (m), 1458 (m), 1439 (s), 1383 (s), 1361 (m), 1331 (m), 1331 (m), 1319 (m), 1290 (m), 1268 (m), 1255 (w), 1236 (m), 1186 (m), 1107 (m), 1107 (w), 1097 (w), 1058 (w), 1043 (w), 1025 (w), 1007 (w), 934 (w), 883 (w), 871 (w), 822 (w), 799 (s), 767 (w), 753 (w), 712 (w), 673 (w), 564 (w), 551 (vs), 510 (m), 434 (m), 418 (m).

Synthesis of [Co(DippFormCO)(CO)₃] (3)

THF (30 mL) was added to a Schlenk flask charged with $[Yb(DippForm)_2(thf)_2]$ (418 mg, 0.40 mmol) under purified nitrogen $Co_2(CO)_8$ (160 mg, 0.47 mmol) was added to the orange solution with stirring and the mixture was stirred for 12 hours at ambient temperature. The colour of the solution turned from orange to dark brown. The solution was filtered to remove any solid materials and the volume reduced under vacuum. The solution was then stored at 3 °C for 3 days during which time needle like crystals formed. IR (Nujol) $\tilde{\nu}$ (cm⁻¹): 2066 (s), 2009 (s), 1983 (m), 1897 (w), 1666 (s), 1587 (m), 1287 (m), 1234 (w), 1180 (m), 1098 (w), 1058 (w), 1000 (w), 935 (w), 821 (w), 799 (m), 753 (w), 767 (m), 722 (m). Complex 3 could not be obtained as analytically pure material.

Synthesis of $[{(DippForm)_2Sm^{III}}_2{(\mu_3-CO)_2Fe_3(CO)_9}] (4)^{56}$

To a mixture of $[(DippForm)_2Sm^{II}(thf)_2]$ (205 mg, 0.20 mmol) and [Fe₃(CO)₁₂] (51 mg, 0.10 mmol) or [Fe₂(CO)₉] (72 mg, 0.20 mmol) was condensed thf (15 mL) at -78 °C and the reaction mixture was stirred for 48 hours at 60 °C. All the volatiles were removed in vacuo. Toluene (15 mL) was added to the residue and refluxed for five minutes and the hot reaction mixture was filtered. Dark red-coloured crystals were obtained upon slowly cooling the filtrate to room temperature. The mother liquor was decanted off and the product was dried under vacuum. Yield: 74 mg (based on crystals), 0.033 mmol, 33% (average). Anal. calcd for C₁₁₁H₁₄₀N₈O₁₁Fe₃Sm₂ (2230.64): C, 59.77; H, 6.33; N, 5.02. Found: C, 59.77; H, 6.20; N, 4.81. IR (ATR) $\tilde{\nu}$ (cm⁻¹): 2962 (vs), 2926 (m), 2869 (m), 2011 (vs), 1979 (vs), 1967 (s), 1878 (m), 1830 (w), 1696 (w), 1667 (s), 1640 (s), 1636 (m), 1586 (s), 1512 (s), 1464 (s) 1457 (m), 1437 (m), 1384 (s), 1362 (m), 1346 (m), 1332 (m), 1314 (s), 1278 (s), 1368 (m), 1254 (s), 1236 (m), 1186 (m), 1112 (w), 1098 (m), 1054 (m), 1042 (m), 1023 (w), 1004 (w), 947 (w), 935 (m), 823 (s), 800 (s) 768 (m), 753 (s), 681 (s), 645 (m), 612 (s), 585 (s), 508 (w), 474 (w), 458 (w), 444 (w), 438 (w), 433 (w), 420 (w).

Conflicts of interest

There are no conflicts to declare.

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