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Rhenium is different: CO tetramerization induced by a divalent lanthanide complex in rhenium carbonyls[†]

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The reduction of $M_2(CO)_{10}$ (M = Mn, Re) with different divalent lanthanide (Ln = Sm, Yb) compounds was investigated. Depending on the steric demand of the ligand, either unusual CO tetramerization or formation of a new Re carbonyl anion occurred in the case of Re. Theoretical calculations were performed for a better understanding of the nature of bonding in the newly formed species.

The coupling of carbon monoxide in the presence of hydrogen to form short hydrocarbon chains is performed on an industrial scale in the Fischer-Tropsch process. Thus, carbon monoxide is a key C₁ feedstock for the industrial production of hydrocarbons. The primary products of the Fischer-Tropsch synthesis are alkanes, alkenes, and to a minor extent alcohols, which are all valuable starting materials in the chemical industry.¹ The product selectivity in the Fischer-Tropsch process can be varied over a wide range. Key parameters are the catalyst formulation and the reaction conditions. The chain growth probability is influenced by the catalyst that is selected for the hydrogenation process.² Industrially, mostly Fe or Co catalysts have been employed in this reaction.³ For a better understanding of the Fischer-Tropsch process, numerous metal complexes, which promote the coupling of CO have been published.⁴ For example, CO dimerization has been observed using Th-H,4d Zr-H,4e Ce-H,^{4f} or Mg-H/Ca-H^{4g,h} catalysts. However, the number of complexes forming higher coupling products (C3, C4, and C6) is very limited. Examples have been published using actinides $(C_3O_3^{2-,5}C_4O_4^{4-,5b,6}OC(CH_2-tBu)C(O)=C(O)-C(CH_2-tBu)O^{2-}),^7$ lanthanides ${(O_2C-C=C=O)^{2-}, ^8 OC(tBu)C(O)=C(O)-C(tBu)O^{2-}, ^9 OC(tBu)O^{2-}, ^9$ and a [BC(O)C(O)C(O)B] skeleton¹⁰}, tantalum $(C_6O_6^{8-})$,¹¹ iron $(C_4(OR)_4)$ ¹² and very recently $[W(CO)_6]$ and Al(I) $([C_3O_3]^{4-}$ and $[C_4O_4]^{4-}).^{4c}$

The reductive behaviour of 5d transition-metal (TM) carbonyl complexes is far less explored as compared with their lighter analogues. In 2017, Deacon and co-workers trapped the elusive $[W_2(CO)_{10}]^{2-}$ anion featuring an unsupported W-W bond by the reduction of $W(CO)_6$ with a divalent samarium mesooctaethylcalix[4]pyrrolide.¹³ Recently, single-electron reduced rhenium carbonyl complexes have shown electrocatalytic reduction of CO₂ to CO.¹⁴

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Herein, we report the reduction of Re₂(CO)₁₀ with divalent lanthanide complexes, which leads to a tetramerization of CO to give a Fischer-type rhenacycle. We also studied the effect of the ligands on the reduction ability of the lanthanide complexes and compared the reactivity of $Re_2(CO)_{10}$ to that of $Mn_2(CO)_{10}$.

Reaction of $[(Cp^*)_2 Sm^{II}(thf)_2]^{15}$ (Cp* = C₅Me₅) with Re₂(CO)₁₀ in toluene at room temperature furnished $[{(Cp^*)_2Sm}_3](\mu$ - $O_4C_4(\mu-\eta^2-CO)_2(\mu-\eta^1-CO)(CO)_5Re_2$ Sm(Cp*)₂(thf)] (1) as red coloured crystals in 18% yield (Scheme 1). Since Sm^{II} is a singleelectron reduction reagent,¹⁶ the central $[(\mu - O_4C_4)(\mu - \eta^2 - CO)_2(\mu - \eta^1 - \eta^2)]$ CO (CO)₅Re₂⁴⁻ core has been formed by a four-fold reduction process and the $[\mu-O_4C_4]$ unit can formally be considered as tetraanionic where each oxygen atom is negatively charged. The solidstate IR spectrum of 1 showed characteristic ν_{CO} stretches, which are different from the CO vibrations of $Re_2(CO)_{10}$. Notably, the low-frequency stretches at 1792(s) and 1733(s) cm⁻¹ indicate bridging isocarbonyls. No suitable NMR data could be obtained, most probably due to decomposition in solution. The solid-state structure of 1 revealed the formation of a hexametallic coordination



isomers of 1 (right), double Fischer-carbene (1a) and metallacyclopentadiene (1b)

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Fig. 1 Molecular structure of 1. Hydrogen atoms are omitted for clarity. Bond lengths and angles are given in ESI.†

cluster consisting of one $[(\mu \cdot O_4 C_4)(\mu \cdot \eta^2 \cdot CO)_2(\mu \cdot \eta^1 \cdot CO)(CO)_5 Re_2]^{4-}$, one $[(Cp^*)_2 Sm^{III}(thf)]^+$, and three $[(Cp^*)_2 Sm^{III}]^+$ moieties (Fig. 1). The Sm1, Sm2, and Sm3 atoms are coordinated by two η^5 -Cp* and two O atoms from bridging isocarbonyls or the newly formed $\mu \cdot O_4 C_4$ entity.

The Re-Re bond in 1 is slightly shorter than the Re-Re single bond in $\text{Re}_2(\text{CO})_{10}$ (2.934(3) vs. 3.041(11) Å, respectively) but theoretical investigation indicates only a weak interaction.¹⁷ The Re1-C1 (2.188(6) Å) and Re1-C4 (2.164(7) Å) bond distances are in the typical range of conjugated rhenacyclic Fischer-type carbenes.¹⁸ In principle, two different forms, either a double Fischer-carbene (Scheme 1, 1a) or a metallacyclopentadiene (Scheme 1, 1b), can describe the 5-membered rhenacycle. The short C2-C3 bond distance in the central C4O4 fragment (1.406(9) Å (C2-C3) vs. 1.445(9) Å (C1-C2) and 1.460(9) Å (C3-C4)) suggests the predominance of **1a**.¹⁸ Thus, we consider compound 1 as a cyclic double Fischer-carbene type complex. Interestingly, formation of the 5-membered Fischer-type rhenacycle in 1 occurred via an unprecedented tetramerization of CO ligands, presumably by reductive C-C coupling. It should be noted that a total of 12 CO units are present in the $[(\mu-O_4C_4)(\mu-\eta^2 CO_2(\mu-\eta^1-CO)(CO)_5Re_2]^{4-}$ moiety, more than in the $Re_2(CO)_{10}$ starting material. Since no external source of CO is present, the formation of 1 implies that more than one equivalent of $Re_2(CO)_{10}$ reacts with four equivalents of $[(Cp^*)_2Sm^{II}(thf)_2]$. To the best of our knowledge, CO tetramerization has never been observed with rhenium carbonyls. Albeit two reports describe a similar tetramerization of CO ligands, both involve reactions between trimethylsilylhalide (halide = Br, or I) or $[{(Me_3Si)_2N}BBr_2]$ and [Na₂Fe(CO)₄].¹² Furthermore, the resulting products did not feature Fischer-carbene type character but metallacyclopentadiene type character (Scheme 1, 1b). The reactivity observed is in sharp contrast to all the reports on Ln-TM carbonyl complexes discussed earlier.13,19

Recently, we have shown that using sterically demanding ligands in the coordination sphere of divalent lanthanides, different activations of main-group elements and complexes can be achieved.²⁰ We therefore investigated the reduction of TM carbonyls with [(DippForm)₂Sm^{II}(thf)₂]²¹ (DippForm = *N*,*N*'-bis(2,6-diisopropylphenyl)formamidinate), another sterically demanding divalent samarium reagent. Reaction with half of an equivalent of Re₂(CO)₁₀ resulted in the formation of [{(DippForm)₂Sm^{III}(thf)₂]{(μ - η ²-CO)₂(μ - η ¹-CO)₂(CO)₄Re₂}] (2) (Scheme 2) as red coloured crystals in 62% yield. The solid-state IR spectrum



Scheme 2 Synthesis of complex 2 (simplified view)



Fig. 2 Molecular structure of 2. Hydrogen atoms are omitted for clarity. Bond lengths and angles are given in ESI.†

of 2 showed characteristic $\nu_{\rm CO}$ stretches. The low-frequency stretch at 1804 cm⁻¹ is consistent with the occurrence of bridging isocarbonyls.^{14*a*} The solid-state structure (Fig. 2) confirmed the identity of 2, which is formed through the transfer of 2e⁻ from two molecules of [(DippForm)₂Sm^{II}(thf)₂] to one molecule of Re₂(CO)₁₀ in single-electron transfer (SET) steps, accompanied by the loss of two CO ligands. In complex 2, the [Re₂(CO)₈]²⁻ anion is entrapped between two [(DippForm)₂Sm^{III}(thf)]⁺ moieties. To the best of our knowledge, the [Re₂(CO)₈]²⁻ anion has never been reported to date. Its lighter analogue, [Mn₂(CO)₈]²⁻, has recently been isolated by reduction of Mn₂(CO)₁₀ with silylene, however, the authors failed to isolate [Re₂(CO)₈]²⁻ under similar conditions.²²

The isolation of the $[(\mu$ -CO)₄(CO)₄Re₂]²⁻ fragment in compound 2 can be traced back to the high reductive nature of divalent samarium and the sterically demanding nature of the $[(Dipp-Form)_2Sm^{III}(thf)]^+$ moieties. In the $[Re_2(CO)_8]^{2-}$ anion, each rhenium atom is coordinated by the carbon donor of two terminal and two bridging CO ligands. The Re–C51 (μ - η^2 -CO) bond lengths, 2.049(3) and 2.206(3) Å, are longer than the other Re–C bonds due to the η^2 -type bridging mode with two Re atoms. The Re–Re' bond length (2.689(3) Å) is substantially shorter than the Re–Re single bond in Re₂(CO)₁₀ (3.041(11) Å),¹⁷ suggesting a double bond character between the two Re atoms at the first glance.²³ Although the $[Re_2(CO)_8]^{2-}$ anion fits with the 36 electron count only after considering Re–Re double bond, our theoretical calculations suggest a lower bonding order.

The bonding situation in compounds **1** and **2** was investigated by theoretical methods. In many cases the bonding description nicely succeeds by calculation of the shared electron numbers (SEN) according to the population analysis of Ahlrichs and Heinzmann.²⁴ These numbers give a reliable measure of the covalent bond strength, especially even in the case of multi-centre bonding contributions. This effect is referred to as "supported metal-metal bonding" which might appear with the Re-Re contacts in 1 and 2 due to the supporting Re–C–Re bridging. To obtain SEN reference values for an unsupported Re–Re single bond, an analysis of the unbridged Re₂(CO)₁₀ molecule was performed. Here, the SEN (Re–Re) is 1.297, but due to a fourcentre contribution of the SEN (C–Re–Re–C) = 0.248, about 19% of the Re–Re bond is attributed to this multi-centre bond strengthening. This procedure is justified and was tested by Ahlrichs *et al.* for the molecule diborane.²⁵ The multi-centre adjusted value would thus be 1.297 - 0.248 = 1.049. As reference for a molecule with a triple-bridged formal Re–Re triple bond, the anion Re₂Cl₉⁻ was investigated. Here, the SEN is calculated to be 1.915, the three-centre SEN (Re–Cl–Re) is 0.273. Therefore, the pure Re–Re "triple" bond disregarding three-centre bonding effects caused by the three Re–Cl–Re bridges would result in a value of $1.915 - 3 \times 0.273 = 1.096$.

Applying this method analogously to calculate the covalent bond strengths in compounds **1** and **2**, the following conclusions can be drawn: in compound **1**, the SEN (Re–Re) = 1.071 and the three-centre contribution is SEN (Re–Re–C) = 0.290, so that for the multi-centre corrected SEN (Re–Re) of 0.491 results. In **2** SEN (Re–Re) = 1.394 and SEN (Re–Re–C) = 0.348, so that SEN (Re–Re) = 0.698, corresponding to a rather weak single bond. The results are summarized in the following way: (i) Re–Re bonding regardless of its origin evaluates in the following row (SEN values in parentheses): **1** (1.071) < Re₂(CO)₁₀ (1.297) < **2** (1.394) < Re₂Cl₉⁻ (1.915). (ii) Taking supporting effects by bridging bonds into account, we conclude that there is virtually no Re–Re bond in **1** and only a weak single two centre Re–Re bond in **2**.

To confirm the findings of the population analyses in an orienting manner, we performed a different topological approach by means of the AIM (atoms in molecules) method introduced by Bader.²⁶ The results of the AIM analysis are given in Fig. S10 (ESI†). Compound **2**, unlike **1**, consists of a bond path with a bond critical point between both rhenium atoms that undoubtedly confirms the presence of a Re–Re bond. The low value of the electron density ρ_{bcp} and the high value for the ellipticity ε ($\rho_{bcp} = 0.05$ a.u., $\varepsilon = 1.31$) calculated for this bond critical point of Re–Re indicate a rather weak bond with a high "multiple bond contribution" presumably due to the interaction with the supporting π -type Re–Re–C multicentre bonds.

The bonding properties of the rhenacycle in compound **1** are also interesting. The values for the SEN of the bonds C1–C2, C2–C3, C3–C4 of 1.514, 1.593, 1.518 correspond to strong single bonds. For pure single or double bonds, SEN values of about 1.4 or 2.28,²⁵ respectively, are expected. The SEN (Re–C) is calculated to be 0.95 which is of the same order of magnitude that is also found for the terminal Re–CO bond (1.276). Overall, the results support the view of a double Fischer carbene complex as shown in **1a** (Scheme 1).

For comparison, we investigated the reactivity between $Mn_2(CO)_{10}$ and divalent samarium complexes. The reaction of $[(Cp^*)_2Sm(thf)_2]$ with half an equivalent of $Mn_2(CO)_{10}$ in toluene resulted in the formation of $[\{(Cp^*)_2Sm(thf)\}(\mu-CO)_2\{Mn(CO)_3\}]_n$ (3) as a red precipitate (Scheme 3). The crystal structure of 3 (Fig. 3) revealed, as expected, cleavage of the Mn–Mn bond upon reduction of $Mn_2(CO)_{10}$ with $[(Cp^*)_2Sm^{II}(thf)_2]$ and formation of



Scheme 3 Synthesis of complex 3.



Fig. 3 Cutout of the polymeric structure of **3** (left) and molecular structure of **4** (Ln = Sm) and **5** (Ln = Yb) (right). Hydrogen atoms are omitted for clarity. Bond lengths and angles are given in ESI. \dagger

a $[Mn(CO)_5]^-$ anion along with $[(Cp^*)_2Sm(thf)]^+$ as cation. Due to vacant coordination sites on the samarium atom, each $[Mn(CO)_5]^-$ anion bridges two samarium atoms *via* isocarbonyl bridges, resulting in a 1D coordination polymer. The average Sm-C (Cp*) bond distance (2.723 Å) is shorter than in $[(Cp^*)_2Sm^{II}(thf)_2]$ (2.86(3) Å). The $[Mn(CO)_5]^-$ anionic fragment has a trigonal bipyramidal geometry with two bridging and three terminal CO. The Mn–C bond lengths of 1.834 Å (terminal, average) and 1.774 Å (bridged, average) are similar to that in $[\{Cp^*_2Yb\}(\mu-CO)_3\{Mn(CO)_2\}]_n$ (1.820(5) Å (terminal, average) and 1.791(13) Å (bridged, average)).^{19d}

Due to the different reactivity observed with $Re_2(CO)_{10}$ depending on the nature of the ligand on the Sm^{II} centre, the reaction between Mn₂(CO)₁₀ and [(DippForm)₂Ln^{II}(thf)₂] (Ln = Sm,²¹ Yb²⁷) was also examined. Half an equivalent of Mn₂(CO)₁₀ was reacted with $[(DippForm)_2Ln^{II}(thf)_2]$ (Ln = Sm, Yb) in toluene at 60 °C to give [{(DippForm)₂Ln^{III}(thf)}{Mn(CO)₅}] (Ln = Sm (4), Yb (5)) (Scheme 4). The low-frequency stretches at 1734 cm^{-1} (vw) for 4 and 1701 (w) cm^{-1} for 5 indicate bridging isocarbonyls.^{19c,d} Due to the different steric demand of the Cp* and DippForm ligands, only one [Mn(CO)₅]⁻ anion binds to each samarium atom, resulting in discrete molecular species. Complexes 4 and 5 are isostructural and only the structure of 4 is discussed here (see ESI⁺ for 5). In the solid-state structure of 4 (Fig. 3), each Sm atom lies in a distorted octahedral geometry, coordinated by two amidinates, one thf ligand and the oxygen atom of the bridging isocarbonyl. The $[Mn(CO)_5]^-$ fragment is in a



Scheme 4 Synthesis of complexes 4 and 5.

trigonal bipyramidal geometry. Formation of **4** and **5** occurred though cleavage of the Mn–Mn bond *via* a SET step. The [(Dipp-Form)₂Sm^{III}(thf)]⁺ and [Mn(CO)₅]⁻ fragments further assemble by formation of bridging isocarbonyls (Scheme 4).

In conclusion, we have reacted $M_2(CO)_{10}$ (M = Mn, Re) with different divalent lanthanide compounds. For $Mn_2(CO)_{10}$, the product formation is as expected and the Mn–Mn bond is cleaved upon reduction. The resulting $[Mn(CO)_5]^-$ anion and the *in situ* formed $[L_2Sm^{III}]^+$ cation (L = Cp*, DippForm) combine to give compounds 3–5. Re₂(CO)₁₀ reacts differently. The different reduction potential of Re₂(CO)₁₀ and $Mn_2(CO)_{10}$ may be one reason.²⁸ By employing $[(DippForm)_2Ln^{II}(thf)_2]$ an unexpected $[Re_2(CO)_8]^{2-}$ anion is formed. In the case of $[(Cp*)_2Sm^{II}(thf)_2]$, an unusual tetramerization of CO combined with the formation of a cyclic double Fischer-carbene type complex was seen. From previous work, we know that the ligand on the divalent samarium atom strongly influences the reactivity of the samarium reagent due to steric and electronic effects.

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Conflicts of interest

There are no conflicts to declare.

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