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NEW IONIC HALF-METALLOCENES OF EARLY LANTHANIDES

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Abstract:

We present in this study a new and one-step method allowing the preparation of an unprecedented family of stable half-lanthanidocenes. X Ray analysis shows that the isolated compounds all display the same ionic Ln-Mg bimetallic structure consisting of two anionic $(Cp^R)Ln(BH_4)_3$ species and one cationic Mg(THF)₆ $(Cp^R = C_5Me_5, Ln = Nd, 1a, Ln = La, 1b; Cp^R = C_5H_5, Ln = Nd, 2a; Cp^R = C_5H_2Ph_3, Ln = Nd, 3a)$. Such complexes display high stability with respect to disproportionation in solution. Combined with dialkylmagnesium, neodymium complexes provide powerful catalysts for stereospecific isoprene polymerisation.

Keywords: polymers, elastomers, and plastics (A) • chemical synthesis (B) • catalysis (B) • crystal structure (C) • X Ray diffraction (D)

Introduction:

Monocyclopentadienyl compounds of the rare earths represent a family of complexes that interest for catalysis has spectacularly increased since very recently [1, 2]. The field of polymerisation is particularly most concerned, with the elaboration of unique controlled structure specialty polymeric materials [3, 4]. However, the potential of such compounds – also called half-sandwiches or half-metallocenes – in catalysis is still little explored compared to metallocenes, mainly because their synthesis remains a challenge [5], due to the occurrence of comproportionation reactions [6, 7]. This has been observed in particular in the early series, even in the presence of one bulky cyclopentadienyl ligand [8, 9]. Thus, the half-lanthanidocenes chemistry remains mainly limited to the smaller elements [6, 10].

Recently however, taking advantage of the modularity of hapticity of the BH₄ group, several monocyclopentadienyl complexes could be isolated by ionic metathesis (Scheme 1) [11-13].

(here scheme 1)

The more sophisticated σ -bond metathesis requires highly reactive homoleptic LnR₃ precursors (R = alkyl [6, 14], allyl [15], phenyl [16], or amido group [17]), but this method advantageously allows the formation of the expected product in one step, with a minimal experimental work-up, as the formation of inorganic salts is avoided. However, heating may be necessary to achieve the metathesis reaction, and the possible formation of hazardous by-products requires extreme attention [18]. Furthermore, ligand scrambling may be not completely excluded using this strategy [5, 16].

In the present paper, we propose an alternative method for the preparation of halfsandwiches of lanthanides in high yield, which conjugates the advantages of σ -bond metathesis, and the use of standard starting materials, the trisborohydrides Ln(BH₄)₃(THF)₃. Isolated complexes all exhibit the general ionic formula [(Cp^R)Ln(BH₄)₃]₂[Mg(THF)₆] (Cp^R is a cyclopentadienyl ligand). Polymerisation experiments show that all complexes are efficient precatalysts for isoprene polymerisation, affording highly *trans*-regular polymer in a controlled and living manner.

Synthesis of ionic complexes

General scheme

Ln(BH₄)₃(THF)₃ are common, easy to handle, and stable lanthanide compounds [19]. We first expected that they could be valuable starting materials for σ -bond metathesis since the [Ln]-BH₄ moiety is known to display the chemical behavior of an hydride in some cases [20]. However, we observed by ¹H NMR that no reaction occurred between C_5Me_5H and Nd(BH₄)₃(THF)₃ even after a prolonged period. On the other hand, when the same reaction was carried out in the presence of one half-equivalent of BEM (nbutylethylmagnesium *n*BuEtMg) at room temperature, a quantitative reaction took place within a few minutes at room temperature. Repeating the same at the bulk scale, we could isolate a light blue crystalline solid in high yield (ca 80 %). According to 1 H NMR the new compound consisted of one C₅Me₅ ligand for 3 BH₄ groups (scheme 2). X-ray structure determination allowed to establish [(C₅Me₅)Nd(BH₄)₃]₂[Mg(THF)₆] (1a) as molecular formula for the isolated complex [21]. The similar procedure was successfully followed again with cyclopentadiene with C_5H_5 , and

triphenylcyclopentadiene 1,2,4-Ph₃C₅H₃, giving rise to the formation of $[(C_5H_5)Nd(BH_4)_3]_2[Mg(THF)_6]$ (**2a**) and $[(C_5H_2Ph_3)Nd(BH_4)_3]_2[Mg(THF)_6]$ (**3a**), respectively. $[(C_5Me_5)La(BH_4)_3]_2[Mg(THF)_6]$ (**1b**) was obtained as same from $La(BH_4)_3(THF)_3$ [22].

It must be emphasized that this unprecedented synthetic method, that we call the "borohydride route", requires very mild experimental conditions, and ordinary lanthanides precursors.

(here scheme 2)

Mechanistic aspects

The synthetic strategy employed was deduced from our ongoing studies on polymerisation with borohydrido compounds of lanthanides activated by alkylating reagents [23-26], which established unambiguously that [Ln]-R species were formed in the reaction medium. Two mechanistic pathways are possible, involving such [Ln]-R species which then react with HCp^R through σ -bond metathesis (scheme 3, i), or the reaction of MgR₂ with HCp^R first to form the corresponding magnesiocene, followed by the reaction with Nd(BH₄)₃(THF)₃ (scheme 3, ii).

(here scheme 3)

X Ray studies

1a, **1b**, **2a** and **3a** are all trinuclear ionic compounds comprising two anionic halfneodymocene trisborohydride $[(Cp^R)Nd(BH_4)_3]^-$ moieties and one cationic hexa-THF magnesium $[Mg(THF)_6]^{2+}$ adduct (Figs. 1-3) that alternate in the unit cell without particular cation – anion interaction [21, 22]. This ionic trinuclear structure is comparable to that of $[Mg(THF)_6][Nd(allyl)_4]_2(2THF)$, obtained by ionic metathesis, with discrete $[Mg(THF)_6]^{2+}$ cation and allyl neodymate anions [27].

(here Figures 1, 2, 3)

The molecular structures of complexes **1a**, **1b**, **2a**, and **3a** exhibit very similar structural features:

- the [(Cp^R)Nd(BH₄)₃]⁻ anions have a pseudo tetrahedral tri-legged piano-stool geometry.
- The Mg cation exhibits the octahedral geometry with oxygen atoms of THF molecules.
- hydrogens belonging to borohydride groups could be located for 1a, 2a, and 3a. The geometric parameters (B-H, Ln-H distances, and B-H-Ln angles) are typical of monomeric borohydrido complexes bearing a tridentate Ln–(η³-H)₃B-H terminal group [25, 28, 29]. In the case of 1b, B-La-B angles and La-B distances are close (albeit slightly higher, due to the larger La vs. Nd) to these found in the Nd complexes, and thus in accordance with the same mode of bonding.
- In the case of **1a** and **3a**, one of the BH₄ ligand is distorted with one shorter Nd- $(\mu_2$ -H) distance.

In all these complexes, the absence of coordinated THF to the lanthanide atom points out the higher affinity of the lanthanide towards ionic ligands.

Stability of early half-lanthanidocenes

As far as we know, a borohydride anion $[(Cp^R)Ln(BH_4)_3]^-$ was observed just at once, in the ionic complex $\{[Sm(BH_4)_2(THF)_n]^+[(C_5Me_4^nPr)Sm(BH_4)_3]^-\}$. This compound was obtained from the reaction of $Sm(BH_4)_3(THF)_3$ with a half equivalent of $K(C_5Me_4^nPr)$ (Equation 1) [30] and crystals were isolated with n = 5 THF per metal.

(here Equation 1)

Monocyclopentadienyl of the larger lanthanides (from La to Sm) are known to easily undergo ligands redistribution reactions. Another possible evolution of such complexes is clustering, affording generally hexameric structures (scheme 4) [12].

(here Scheme 4)

With the aim to compare the stability of the anionic $[(C_5Me_5)Nd(BH_4)_3]^{-1}$ versus the neutral $(C_5Me_5)Nd(BH_4)_2(THF)_2$, a sample of each complex in C_6D_6 was kept 20 h at 75 °C in an NMR tube. We observed that the solution of **1a** prepared in our one pot procedure remained unchanged after that period, whereas blue crystals had formed from the solution of $(C_5Me_5)Nd(BH_4)_2(THF)_2$. X Ray analysis showed that the crystals corresponded to the hexamer $[(C_5Me_5)Nd(BH_4)_2]_6[31]$.

The isolation of **2a** and **3a** was more surprising. In the unsubstituted C_5H_5 series, halflanthanidocenes of the larger lanthanides are exceptional [6, 10], whereas **3a** is the first structurally characterized lanthanide complex featuring the bulky but electronwithdrawing $C_5H_2Ph_3$ ligand [33]. It is worth to be noted that the corresponding neutral half-sandwich could not be isolated pure by ionic metathesis [24].

Thus, $[(Cp^R)LnX_3]^-$ appears as a stable molecular entity in the borohydride series, with respect to comproportionation or clustering.

Isoprene Polymerisation

We showed recently that combining a lanthanide borohydride with a dialkylmagnesium is a valuable strategy to prepare, *in situ*, efficient lanthanide-based catalysts that are also highly *trans*-stereospecific towards isoprene polymerization [4, 25]. Similarly, the addition of a stoichiometric amount of BEM (1:1) to ionic neodymium complexes **1a**, **2a**, and **3a** gave rise in each case to a very valuable initiator (Table 1, runs 1-3). All these catalysts afford a highly *trans*-regular polymer (> 95 % according to NMR, see Figure 4) [32], as well as good control over the macromolecular data (fine agreement between calculated and measured molecular weights, and narrow polydispersities), typical of a single-site process. Besides, the catalytic behaviour of **1a** / 1BEM system, the most active, was found fully comparable to that one obtained from the neutral (C₅Me₅)Nd(BH₄)₂(THF)₂ (run 1 *vs.* 4). Thus, both catalysts most probably involve the same half-sandwich catalytic species (for a discussion on this question, see ref 4). It is confirmed that the more bulky the Cp^R ligand, the higher the *trans*-stereospecificity [4], whereas the activity of the catalyst decreases with the electronic richness of the cyclopentadienyl ligand, as already observed [33].

(here Table 1)

To conclude, the "borohydride route" is an elegant alternative for the preparation of half-sandwiches of early lanthanides, enabling a one pot and high yield synthesis of such complexes. Thanks to this method, it is possible to prepare very easily unprecedented lanthanide-based catalysts. It will be shown in a forthcoming paper that this approach also allows the formation of mixed magnesium lanthanidocene complexes.

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[22] Syntheses: $[Cp*La(BH_4)_3]_2Mg(THF)_6$, **1b**: 275 mg (0.5 mmol) of BEM (20 % solution in hexanes) diluted in 10 mL of toluene were added dropwise, at room temperature, to a solution of La(BH₄)₃(THF)₃ (400.0 mg, 1 mmol) and Cp*H (136 mg, 1 mmol) in toluene (20 mL). The mixture turned from white to light yellow

immediately. After 1 h of stirring at room temperature, the solvents were evaporated to dryness. 10 mL of THF were added by vacuum transfer into the flask, leading to a limpid uncoloured solution. The THF was evaporated to give an off-white sticky compound, that was redissolved in toluene (20 mL) through the same vacuum transfer process. After a filtration to eliminate insoluble residues present in small quantities (ca 30 mg), this solution was slowly concentrated, leading to a crop of well-formed white crystals (407 mg; 0.37 mmol, 74.4 %). Anal. Calc. C₄₄H₁₀₂B₆O₆La₂Mg: C, 48.26; H, 9.32. Found: C, 47.94; H, 9.23. ¹H NMR (C₆D₆) δ : 1.1 (br, 12H, BH₄, $v_{\frac{1}{2}}$ = 100 Hz), 2.36 (s, MeCp, 15H), 3.69 (s, THF, 12H), 1.27 (s, THF, 12H). [CpNd(BH₄)₃]₂Mg(THF)₆, **2a**: 235 mg (0.43 mmol) of BEM (20 % solution in hexanes) diluted in 10 mL of toluene were added dropwise, at room temperature, to a solution of Nd(BH₄)₃(THF)₃ (405.0 mg, 1.0 mmol) and freshly cracked CpH (65 mg, 1.0 mmol) in toluene (20 mL). The mixture turned from light purple to blue immediately. Progressively, a pale blue precipitate formed. After 1 h of stirring at room temperature, the solvents were evaporated off, and THF (15 mL) was distilled by vacuum transfer into the flask, leading to a limpid blue solution. The THF was evaporated to give a sticky blue compound, that was redissolved in toluene (20 mL) through the same vacuum transfer process. The toluene solution was filtrated to eliminate small quantities of insoluble residues, then slowly concentrated within 4 hours to ca. 2 mL. A first crop (30 mg) of light blue crystals, suitable for X Ray structure analysis, could be collected. A second crop was obtained after standing 24 hours at room temperature; it was washed successively with cold toluene and cold pentane, and dried under vacuum at room temperature (214 mg). From the mother liquor, 54 mg of a blue crystalline solid could be finally obtained after concentration, decantation, and drying. Total yield: 298 mg;

71.9 % (with respect to Mg). Anal. Calc. $C_{34}H_{82}B_6O_6Nd_2Mg$: C, 42.33; H, 8.57. Found: C, 42.52; H, 8.37. ¹H NMR (C_6D_6) δ : 65 (v. br, 12H, BH₄, $v_{\frac{1}{2}} > 2000$ Hz), 0.93 (THF), 0.29 (THF), -2.65 (s, Cp, 15H). [$C_5H_2Ph_3Nd(BH_4)_3$]₂Mg(THF)₆, **3a**: same procedure as for **2a**, from 275 mg (0.5 mmol) of BEM (20 % hexane solution), 405 mg (1 mmol) of Nd(BH₄)₃(THF)₃, and 294 mg (1 mmol) of HC₅H₂Ph₃. Suitable X Ray crystals could be collected from a toluene solution slowly evaporated inside the glove box. Total yield: 360 mg; 53.5 %. Anal. Calc. $C_{70}H_{106}B_6O_6Nd_2Mg$: C, 59.13; H, 7.52. Found: C, 59.39; H, 7.75.

X Ray structures analyses: Compound **1b** $(C_{88}H_{204}B_{12}La_4Mg_2O_{12})$ crystallizes in the monoclinic space group P2₁ with a = 11.077(3), b = 30.239(8), c = 17.680(5) Å, β = 106.257(4) °, V = 5685(3) Å³, and $\rho = 1.278$ gcm⁻³ for Z = 2. Data were collected at 100 K on a Bruker Smart-CCD 1K system. The structure was solved by direct methods using SHELXS-97 package (Sheldrick, 1990), and least-square refinement of the model based on 18293 reflections (I>2.0 σ (I)) converged to a final R = 6.29% (wR₂ = 15.51%). Except H pertaining to BH₄ groups, all hydrogen atoms were placed in calculated positions. Others were located in difference Fourier maps and refined with isotropic thermal parameters. Compound 2a (C₃₄H₈₂B₆MgNd₂O₆) crystallizes in the triclinic space group P-1 (2) with a = 10.016(9), b = 10.178(9), c = 12.859(12) Å, α = 110.335(14) °, $\beta = 94.702(14)$ °, $\gamma = 96.491(14)$ °, V = 1211.0(19) Å³, and $\rho = 1.323$ gcm^{-3} for Z = 1. Data were collected at 100 K on a Bruker Smart-CCD 1K system. The structure was solved by direct methods using SHELXS-97 package (Sheldrick, 1990), and least-square refinement of the model based on 4698 reflections (I>2.0 σ (I)) converged to a final R = 2.75% (wR₂ = 6.62%). Except H pertaining to BH₄ groups, all hydrogen atoms were placed in calculated positions. Others were located in difference Fourier maps and refined with isotropic thermal parameters. Compound **3a** ($C_{35}H_{53}B_{3}NdMg_{0.5}O_{3}$) crystallizes in the triclinic space group P-1 (2) with a = 10.051(5), b = 10.839(5), c = 17.695(8) Å, α = 74.629(7) °, β = 84.793(7) °, γ = 74.315(7) °, V = 1789.1(14) Å³, and ρ = 1.491 gcm⁻³ for Z = 2. Data were collected at 100 K on a Bruker Smart-CCD 1K system. The structure was solved by direct methods using SHELXS-97 package (Sheldrick, 1990), and least-square refinement of the model based on 5600 reflections (I>2.0\sigma(I)) converged to a final R = 6.03% (wR₂ = 13.62%). Except H pertaining to BH₄ groups, all hydrogen atoms were placed in calculated positions. Others were located in difference Fourier maps and refined with isotropic thermal parameters. CCDC-618594, CCDC-618595 and CCDC-618596 contains the supplementary crystallographic data corresponding to **1b**, **2a**, and **3a**, respectively. These data can be obtained free of charge via www.ccdc.cam.ac.uk/conts/retrieving.html (or from the Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB21EZ, UK; fax: (+44) 1223-336- 033; or deposit@ccdc.cam.ac.uk.

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[32] Stereospecificity was determined by both ¹H and ¹³C NMR integrations. The average variation observed is of ± 0.2 on the value of the percentage for several measurements of the same sample; % of *trans*-stereospecificity results from % of 3,4-and *cis*-defects in polyisoprenes, calculated as follows from ¹H NMR integration: (δ ppm) 5.11 (a); 4.68 (b); 1.68 (c)

%
$$cis = \frac{c/3}{a+b/2}$$
 % $3,4 = \frac{b/2}{a+b/2}$ % $trans = 100 - %cis - %3,4$

The values found are corroborated by those calculated from ¹³C NMR: (δ ppm) 16.4 (e); 18.9 (f); 23.8 (g)

% trans =
$$\frac{e}{e+f+g}$$
 % cis = $\frac{f}{e+f+g}$ % 3,4 = $\frac{g}{e+f+g}$

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$$Ln(BH_4)_3(THF)_3 + K(Cp^R) \longrightarrow (Cp^R)Ln(BH_4)_2(THF)_n + KBH_4$$
$$Cp^R = C_5H^iPr_4, C_5Me_4^nPr, C_5Me_5;$$
$$Ln = Nd, Sm$$

Scheme 1: synthesis of the monocyclopentadienyl borohydride complexes by ionic metathesis

$$Ln(BH_4)_3(THF)_3 + HCp^R + 0.5 \ ^nBuEtMg \xrightarrow{1. \ toluene} [(Cp^R)Ln(BH_4)_3][Mg(THF)_6]$$

$$Ln = Nd, La;$$

$$Cp^R = C_5Me_5 \ (1a, 1b), C_5H_5 \ (2a), C_5H_2Ph_3(3a)$$

Scheme 2. Synthesis of the ionic monocyclopentadienyllanthanide complexes by the "borohydride / alkyl route"

$$2 (BH_4)_3Nd + MgR_2 \xrightarrow{i} 2 (BH_4)_3Nd R^- + Mg^{2+} \xrightarrow{} 2 (BH_4)_3Nd R^- + Mg^{2+}$$

$$2 RH \xrightarrow{2} HCp^R$$

$$[(Cp^R)Nd(BH_4)_3^-]_2[Mg(THF)_6^{2+}]$$

$$2 Nd(BH_4)_3$$

$$2 HCp^R + MgR_2 \xrightarrow{ii} Mg(Cp^R)_2 + 2 RH$$

Scheme 3. The possible mechanistic pathways of the « borohydride/ alkyl route » (THF omitted for clarity)

$(C_5Me_4^{n}Pr)Ln(BH_4)_2(THF)_2 \xrightarrow{THF} [(C_5Me_4^{n}Pr)Ln(BH_4)_2]_6$

Scheme 4

$$Sm(BH_4)_3(THF)_3 + \frac{1}{2} KC_5 Me_4^{n} Pr \xrightarrow{1. \text{ toluene}} \frac{1}{2. \text{ THF}} \frac{1}{2} \{ [Sm(BH_4)_2(THF)_3]^+ [(C_5 Me_4^{n} Pr)Sm(BH_4)_3]^- \}$$

Equation 1

FIGURES CAPTIONS

Figure 1. One of the two entities of the asymmetric unit showing the molecular structure of $[(C_5Me_5)La(BH_4)_3]_2[Mg(THF)_6]$ (**1a**). Hydrogen atoms are omitted for clarity. La2-Mg2 7.363(4) Å, La2-B (average B4-B6) 2.70 Å, B-La2-B (average B4-B6) 105.1 °.

Figure 2. Molecular structure of $[(C_5H_5)Nd(BH_4)_3]_2[Mg(THF)_6]$ (2a). Non-BH₄ hydrogen atoms are omitted for clarity. Nd-Mg 7.225(4) Å, Nd-B (average B1-B3) 2.61 Å, B-Nd-B (average B1-B3) 104.6 °.

Figure 3. Molecular structure of $[(C_5H_2Ph_3)Nd(BH_4)_3]_2[Mg(THF)_6]$ (**3a**). Non-BH₄ hydrogen atoms are omitted for clarity. Nd-Mg 7.323(2) Å, Nd-B (average B1-B3) 2.61 Å, B-Nd-B (average B1-B3) 104.3 °.

Figure 4. (a) ¹H NMR spectrum of *trans*-polyisoprene from precatalyst **3a** (CDCl₃). * 3,4-defects: $\delta = 4.68$ ppm, CH signal; O 1,4-cis defects: $\delta = 1.68$ ppm, CH₃ signal. (b) ¹³C{¹H} NMR spectrum of *trans*-polyisoprene from **3a** (CDCl₃). * 3,4-defects: *Me* signal, $\delta = 23.8$ ppm; O 1,4-cis defects: $\delta = 18.9$ ppm, *Me* signal



Figure 1







Figure 3







Figure 4 (b)

run	Precatalyst	Yield [%]	<i>Rate of trans-</i> PI [%] ^[b]	$\overline{\overset{M_n}{(\exp)}}^{[c]}$	PDI ^[d]	$\overline{M_n}$ (calc.) ^[e]
1	1a	84	98.2	58200	1.16	56000
2	2a ^[f]	41	95.2	31600	1.57	28000
3	3a	59	96.2	35500	1.32	37300
4	$(C_5Me_5)Nd(BH_4)_2(THF)_2$	80	97.4	52300	1.18	53700

 Table 1. Isoprene polymerisation with borohydrido half-neodymocene / BEM

 catalysts.^[a]

[a] Conditions: 10 µmol Nd, 1mL toluene, [BEM]/[Nd] = 1, [isoprene]/[Nd] = 1000, T = 50 °C, t = 2 h. [b] Determined by both ¹H and ¹³C NMR integrations (average variation observed ±0.2 on the value of the percentage for several measurements of the same sample), see ref. 32. [c] Determined by Steric Exclusion Chromatography calibrated with PS standards. [d] PDI = $\overline{M_w}/\overline{M_n}$. [e] yield * [isoprene]/[Nd]. [f] t = 6 h.