

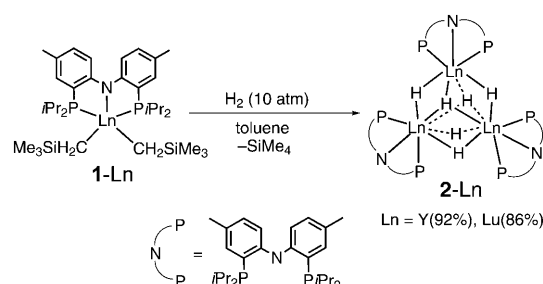
Rare-Earth Polyhydride Complexes Bearing Bis(phosphinophenyl)amido Pincer Ligands**

Jianhua Cheng, Takanori Shima, and Zhaomin Hou*

Rare-earth (Group 3 and lanthanide) metal polyhydride complexes consisting of the dihydride species “ LLnH_2 ” with one ancillary ligand per metal center are receiving intense current interest because of their fascinating structure and reactivity, which are exquisitely different from their monohydride relatives “ L_2LnH ” with two supporting ligands.^[1] The nuclearity of the rare-earth polyhydride complexes is generally dependent on the steric bulk of the ancillary ligands. An increase of ligand hindrance usually leads to a decrease of the nuclearity and increase of the reactivity of the resulting hydride clusters. To date, several hexanuclear,^[1a,2a] tetranuclear,^[2a,3] and trinuclear^[2b,4,5] polyhydride complexes have been reported. However, despite recent progress in this area, the number of well-defined rare-earth polyhydride complexes is still very limited. Most of the ancillary ligands employed to date for rare-earth polyhydride complexes are either sterically demanding cyclopentadienyl derivatives such as $\text{C}_5\text{Me}_4\text{SiMe}_3$ ^[3] or scorpionate tris(pyrazolyl)hydroborate units,^[2] although the use of tetraazacycloamido^[4] or pyridylamido^[5] ligands was also described. Well-defined cationic rare-earth polyhydrides have remained scarce,^[6] and a binuclear polyhydride complex consisting of “ LLnH_2 ” has not been reported to date. Herein, we report the synthesis and structural characterization of a new family of rare-earth polyhydride complexes supported by the rigid bis(phosphinophenyl)amido (PNP) ligands, including the first examples of cationic trinuclear and binuclear rare-earth polyhydrides.

As a possible precursor to a rare-earth polyhydride complex bearing a PNP ligand, we first chose the yttrium dialkyl complex $[(\text{PNP}^{\text{Ph}})\text{Y}(\text{CH}_2\text{SiMe}_3)_2(\text{thf})]$ ($\text{PNP}^{\text{Ph}} = \{2-(\text{Ph}_2\text{P})-\text{C}_6\text{H}_4\}_2\text{N}$), which has shown novel polymerization activity.^[7] However, the hydrogenolysis reaction of $[(\text{PNP}^{\text{Ph}})\text{Y}(\text{CH}_2\text{SiMe}_3)_2(\text{thf})]$ with H_2 seemed very complicated, and no characterizable product was obtained. In contrast, when an analogous dialkyl complex with a bis(diisopropylphosphinophenyl)amido ligand $[(\text{PNP}^{\text{iPr}})\text{Y}(\text{CH}_2\text{SiMe}_3)_2]$ ($\text{PNP}^{\text{iPr}} = \{2-(\text{iPr}_2\text{P})-\text{C}_6\text{H}_4\}_2\text{N}$) was used, clean

formation of the trinuclear polyhydride complex $[(\text{PNP}^{\text{iPr}})\text{YH}_2]_3$ was observed, as indicated by the ^1H NMR spectrum (6.04 ppm, quartet, $^1J_{\text{Y-H}} = 16.8$ Hz, in C_6D_6). Unfortunately, however, a single crystal suitable for X-ray structure determination was not obtained because of the high solubility of this hydride compound in all common organic solvents. To make a hydride compound with better crystallinity, the dialkyl complex with a methyl-substituted analogue of the PNP^{iPr} ligand $[(\text{Me-PNP}^{\text{iPr}})\text{Y}(\text{CH}_2\text{SiMe}_3)_2]$ (**1-Y**, $\text{Me-PNP}^{\text{iPr}} = \{4\text{-Me-2-(iPr}_2\text{P)-C}_6\text{H}_3\}_2\text{N}$)^[8] was then used. To our delight, the hydrogenolysis of **1-Y** in toluene under 10 atm H_2 successfully afforded the corresponding trinuclear hexahydride complex $[(\text{Me-PNP}^{\text{iPr}})\text{YH}_2]_3$ (**2-Y**) as pale yellow crystals in 92 % yield of isolated product (Scheme 1). The



Scheme 1. Synthesis of PNP-ligated trinuclear polyhydride complexes.

analogous Lu hydride cluster **2-Lu** was also obtained in a similar way.

X-ray crystallographic studies revealed that the complexes **2-Y** and **2-Lu** are isostructural. The ORTEP drawing of **2-Y** is shown in Figure 1; selected interatomic distances are listed in Table 1. Each Y atom in **2-Y** is bonded to a PNP

Table 1: Selected interatomic distances [Å] for **2-Y** and **3-Y**.

	2-Y	3-Y
Y1...Y2	3.4829(6)	3.4789(6)
Y1...Y3	3.4678(6)	3.4411(6)
Y2...Y3	3.1648(7)	3.4812(6)
Y(1,2,3)–H1	2.31(4), 2.22(4), 2.32(4)	2.37(4), 2.30(4), 2.43(4)
Y(1,2,3)–H2	2.06(4), 2.37(4), 2.34(4)	2.37(4), 2.30(4), 2.33(4)
Y(1,3)–H3	2.12(4), 2.27(4)	2.11(3), 2.07(3)
Y(1,2)–H4	2.12(4), 2.21(4)	2.08(3), 2.11(3)
Y(2,3)–H5	2.07(4), 2.11(4)	2.03(4), 2.05(4)
Y(2,3)–H6	2.07(4), 2.24(4)	
Y–N (av.)	2.367(3)	2.309(3)
Y–P (av.)	2.948(1)	2.900(1)

[*] Dr. J. Cheng, Dr. T. Shima, Prof. Dr. Z. Hou
Organometallic Chemistry Laboratory and Advanced Catalyst
Research Team, RIKEN Advanced Science Institute
2-1 Hirosawa, Wako, Saitama 351-0198 (Japan)
Fax: (+81) 48-462-4665
E-mail: houzu@riken.jp

[**] This work was supported by Grants-in-Aid for Scientific Research (S) (No. 21225004) and for Young Scientists (B) (No. 21750068) from JSPS. J.C. is grateful to RIKEN for a FPR fellowship. We thank Dr. H. Koshino for technical assistance in ^{11}B NMR spectroscopy measurements.

Supporting information for this article is available on the WWW under <http://dx.doi.org/10.1002/anie.201006812>.

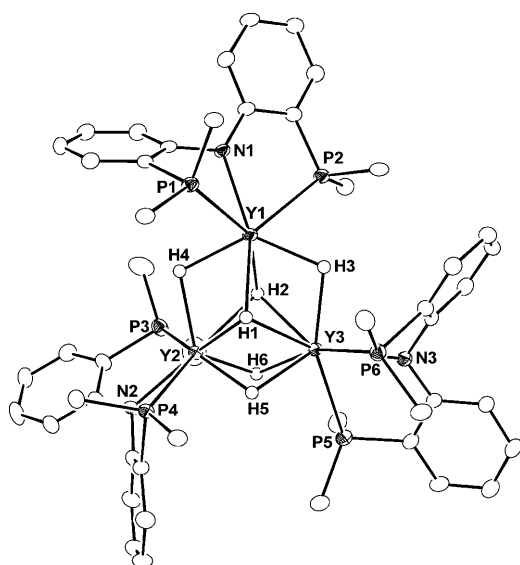


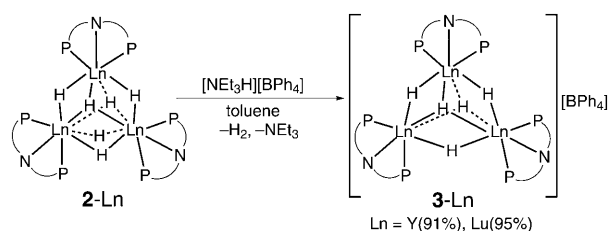
Figure 1. ORTEP plot of **2-Y** with thermal ellipsoids at the 30% probability level. The methyl groups and hydrogen atoms in the PNP ligands have been omitted for clarity.

ligand in a typical κ^3 -(*P,N,P*) mode. Two of the six hydride ligands adopt a μ_3 bonding mode, each of which caps one side of the Y_3 plane. The remaining four hydride ligands bridge the three edges of the Y_3 triangle in a μ_2 fashion, and therefore, one of the three $Y \cdots Y$ edges is bridged by two μ_2 -H ligands, while the other two $Y \cdots Y$ edges are each bridged by only one μ_2 -H ligand, in addition to the μ_3 -H capping ligands. As a consequence, the $Y2 \cdots Y3$ edge (3.1648(7) Å) bridged by two μ_2 -H ligands is significantly shorter than the other two $Y \cdots Y$ edges (3.4829(6) and 3.4678(6) Å, Table 1). This $Y2 \cdots Y3$ edge represents the shortest $Y \cdots Y$ distance ever reported, as far as we are aware. The Y_3H_6 core structure in **2-Y** is thus in contrast to those of other reported trinuclear hexahydrides, such as $[(Tp^{iPr2})YH_2]_3$ (Tp^{iPr2} = tris(3,5-diisopropyl-1-pyrazolyl)hydroborate), which possesses one μ_3 -H and five μ_2 -H ligands,^[2b] and $[(Me_3TACD)YH_2]_3$ (Me_3TACD -H = 1,4,7-trimethyl-1,4,7,10-tetraazacyclododecane),^[4] in which all six hydride ligands adopted the same μ_2 -H bonding mode. The $Y \cdots Y$ distances in these previous examples range from 3.3423(3) to 3.6841(2) Å.

The 1H NMR spectrum of **2-Y** in C_6D_6 showed a quartet ($^1J_{Y-H}$ = 17.0 Hz) at δ = 6.08 ppm corresponding to the hydride ligands, thus suggesting that the structure is fluxional and that all six hydrides are equivalent on the time scale of NMR spectroscopy. The hydrides in **2-Lu** showed a broad singlet at δ = 9.70 ppm. These results are comparable with those observed in other trinuclear polyhydride complexes.^[2b,4]

On treatment with an equivalent of $[NEt_3H][BPh_4]$ in toluene (or THF) at room temperature, both **2-Y** and **2-Lu** gave the corresponding cationic pentahydrides $[(Me-PNP^{iPr})_3Y_3H_5][BPh_4]$ (**3-Ln**; Ln = Y, Lu) in high yields, with concomitant release of H_2 and NEt_3 (Scheme 2).

The cationic complexes **3-Y** and **3-Lu** are soluble in THF and C_6H_5Cl and partly soluble in toluene and benzene but almost insoluble in hexane. Single crystals suitable for X-ray



Scheme 2. Synthesis of cationic trinuclear pentahydride complexes.

diffraction studies were obtained from saturated toluene solutions. Similar to the neutral precursors **2-Ln**, the cationic hydride clusters **3-Ln** are also isostructural. The $\{Ln_3H_5\}$ core structure of **3-Y** is shown in Figure 2 (right) together with its

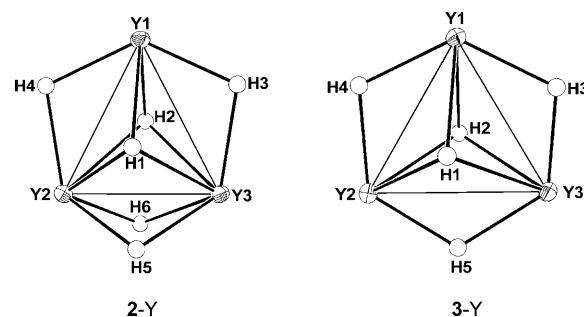


Figure 2. The metal hydride core structures of **2-Y** and **3-Y**.

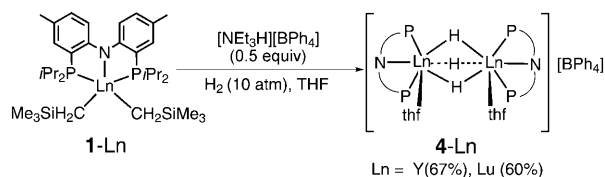
neutral counterpart **2-Y** (left) for comparison. Some selected interatomic distances are summarized in Table 1. All three Y atoms in the cationic complex **3-Y** are almost equally connected by the hydride ligands. Two of the five hydride ligands adopt a μ_3 bonding mode, and the remaining three hydride ligands are bonded in a μ_2 fashion, each bridging one $Y \cdots Y$ edge. Therefore, the three $Y \cdots Y$ edges in **3-Y** are almost equal, in contrast with its neutral precursor **2-Y**, in which one of the $Y \cdots Y$ edges is much shorter than the other two because of the additional bridging hydride ligand (Figure 2 and Table 1). The hexagon formed by the three Y atoms and the three μ_2 -H atoms in **3-Y** is almost planar, with a maximum deviation of 0.129 Å. The two μ_3 -H atoms are located 1.213 and 1.248 Å above and below the hexagonal plane. The $Y-(\mu_3-H)$ bonds in **3-Y** (av. 2.35(4) Å) are longer than the $Y-(\mu_2-H)$ bonds (av. 2.08(4) Å), similar to the situation in **2-Y**. The $Y-N$ and $Y-P$ bonds in the cationic complex **3-Y** are significantly shorter than those in the neutral precursor **2-Y** (Table 1).

It is also noteworthy that no direct bonding interaction was observed between the cation $[(Me-PNP^{iPr})_3Ln_3H_5]$ and the anion $[BPh_4]$ in **3-Y** despite steric and electronic unsaturation (compared to **2-Y**) at the metal center. Moreover, the incorporation of a thf ligand into **3-Y** was not observed even when the complex was mixed with THF. These results are in sharp contrast with what was observed in the tetranuclear cationic polyhydride complex $[(C_5Me_4SiMe_3)_4Y_4H_7][B(C_6F_5)_4]$, which adopted a contact-ion-pair structure in the THF-free form and gave the solvent-

separated ion pair $[(C_5Me_4SiMe_3)_4Y_4H_7(thf)_2][B(C_6F_5)_4]$, upon reaction with THF.^[6] Complex **3-Y** (as well as **3-Lu**) represents the first example of a structurally characterized cationic trinuclear rare-earth polyhydride complex and also the first example of a cationic rare-earth polyhydride complex bearing a non-cyclopentadienyl ligand.

The five hydride ligands of the cationic complex **3-Y** showed two sets of 1H NMR signals in C_6D_5Cl , with a broad singlet at $\delta = 5.53$ ppm for the two μ_3 -H ligands and a triplet at $\delta = 6.52$ ppm ($^1J_{Y-H} = 31.7$ Hz) for the three μ_2 -H ligands, in contrast with its neutral precursor **2-Y**, which exhibited only a quartet ($\delta = 5.97$ ppm, $^1J_{Y-H} = 16.9$ Hz) for all six hydrides under similar conditions. Similarly, the hydrides in **3-Lu** showed two broad signals at $\delta = 8.43$ and 11.85 ppm. These results suggest that the Ln_3H_5 core structure in the cationic complexes **3-Ln** is more rigid than the Ln_3H_6 unit in the neutral complexes **2-Ln**. Similar phenomena were also observed in the $C_5Me_4SiMe_3$ -ligated tetranuclear polyhydride analogues.^[6]

Surprisingly to us, when the hydrogenolysis reaction of the dialkyl complexes **1-Ln** was carried out in the presence of 0.5 equivalent $[NEt_3H][BPh_4]$ in THF, the binuclear cationic trihydride complexes $[(Me-PNP^{iPr})_2Y_2H_3(thf)_2][BPh_4]$ (**4-Ln**; $Ln = Y, Lu$) were obtained in 60–67% yields of isolated product (Scheme 3), in sharp contrast with the reaction in the



Scheme 3. One-pot synthesis of cationic binuclear trihydride complexes.

absence of $[NEt_3H][BPh_4]$, which yielded selectively the trinuclear polyhydrides **2-Ln** as described above.^[9] The three hydride ligands in **4-Y** showed a triplet at $\delta = 5.72$ ppm with $^1J_{Y-H} = 23.1$ Hz in the 1H NMR spectrum in $[D_8]THF$. Complex **4-Lu** gave a singlet at $\delta = 8.45$ ppm.

Single crystals of **4-Y** and **4-Lu** were grown from THF/hexane. These two complexes are isostructural. Figure 3 depicts the ORTEP structure of the cation part of **4-Y**. The two Y atoms are bridged by three hydride ligands, and each Y atom is also bonded to a κ^3 -PNP ligand and a thf ligand. The Y_2H_3 core adopts a slightly distorted triangular bipyramid geometry, with the three hydride ligands occupying the equatorial positions and the two Y atoms in the apical sites. The Y–H bond lengths vary from 2.08(5) to 2.21(5) Å, but fall in the range of typical Y–(μ_2 -H) distances. The Y...Y distance in **4-Y** (3.3297(5) Å) is significantly shorter than those in the trinuclear cationic complex **3-Y** (av. 3.4671(6) Å). Complexes **4-Y** and **4-Lu** represent the first examples of structurally characterized cationic binuclear rare-earth polyhydride complexes.

The present formation of **4-Ln** could formally be viewed as a combination of a neutral dihydride species such as $[(Me-$

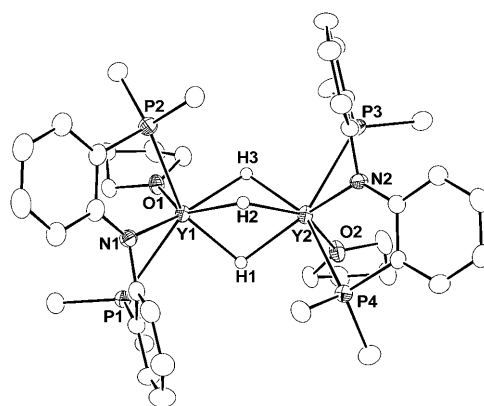


Figure 3. ORTEP plot of the cationic part of **4-Y** with thermal ellipsoids at the 30% probability level. The methyl groups and hydrogen atoms in the PNP ligands have been omitted for clarity. Selected interatomic distances [Å] and angles [°]: Y1–H1 2.08(5), Y1–H2 2.14(4), Y1–H3 2.17(4), Y1–N1 2.314(3), Y1–P1 2.9492(12), Y1–P2 2.9293(11), Y1–O1 2.372(2), Y2–H1 2.21(4), Y2–H2 2.09(4), Y2–H3 2.12(4), Y2–N2 3.321(3), Y2–P3 2.9349(12), Y2–P4 2.9266(12), Y2–O2 2.351(3), Y1...Y2 3.3297(5); Y1–H1–Y2 102.0(12), Y1–H2–Y2 104.0(12), Y1–H3–Y2 102.1(12).

$PNP^{iPr})LnH_2(thf)_x]$, formed by the hydrogenolysis of the dialkyl complex **1-Ln**, with a cationic monohydride species $[(Me-PNP^{iPr})LnH(thf)_x][BPh_4]$ resulting from the hydrogenolysis of the cationic monoalkyl species $[(Me-PNP^{iPr})Ln(CH_2SiMe_3)(thf)_x][BPh_4]$ that could be formed by reaction of **1-Ln** with $[NEt_3H][BPh_4]$ in THF.

In summary, we have demonstrated that the rigid bis(phosphinophenyl)amido moiety $[4-Me-2-(iPr_2P)-C_6H_3]_2N$ can serve as an excellent supporting ligand for rare-earth polyhydride complexes, leading to formation of a new family of structurally characterizable trinuclear and binuclear polyhydride complexes, including the first examples of cationic trinuclear and binuclear rare-earth polyhydrides. Studies on the reactivity of these hydride clusters and on the analogous rare-earth polyhydride complexes with other non-cyclopentadienyl ancillary ligands are in progress.^[10]

Received: October 29, 2010

Published online: January 14, 2011

Keywords: cationic complexes · hydrides · N,P ligands · rare earths

- [1] Recent reviews: a) Z. Hou, M. Nishiura, T. Shima, *Eur. J. Inorg. Chem.* **2007**, 2535–2545; b) M. Nishiura, Z. Hou, *Nat. Chem.* **2010**, 2, 257–268; c) J. Cheng, K. Saliu, M. J. Ferguson, R. McDonald, J. Takats, *J. Organomet. Chem.* **2010**, 695, 2696–2702.
- [2] a) J. Cheng, K. Saliu, G. Y. Kiel, M. J. Ferguson, R. McDonald, J. Takats, *Angew. Chem.* **2008**, 120, 4988–4991; *Angew. Chem. Int. Ed.* **2008**, 47, 4910–4913; b) J. Cheng, M. J. Ferguson, J. Takats, *J. Am. Chem. Soc.* **2010**, 132, 2–3.
- [3] a) O. Tardif, M. Nishiura, Z. Hou, *Organometallics* **2003**, 22, 1171–1173; b) K. C. Hultsch, P. Voth, T. P. Spaniol, J. Okuda, *Z. Anorg. Allg. Chem.* **2003**, 629, 1272–1276; c) D. Cui, O. Tardif, Z. Hou, *J. Am. Chem. Soc.* **2004**, 126, 1312–1313; d) O. Tardif, D. Hashizume, Z. M. Hou, *J. Am. Chem. Soc.* **2004**, 126,

- 8080–8081; e) D. Cui, M. Nishiura, Z. Hou, *Angew. Chem.* **2005**, *117*, 981–984; *Angew. Chem. Int. Ed.* **2005**, *44*, 959–962; f) D. Cui, M. Nishiura, Z. Hou, *Macromolecules* **2005**, *38*, 4089–4095; g) Y. Luo, J. Baldamus, O. Tardif, Z. Hou, *Organometallics* **2005**, *24*, 4362–4366; h) T. Shima, Z. Hou, *J. Am. Chem. Soc.* **2006**, *128*, 8124–8125; i) M. Yousufuddin, M. J. Gutimann, J. Baldamus, O. Tardif, Z. Hou, S. A. Mason, G. J. McIntyre, R. Bau, *J. Am. Chem. Soc.* **2008**, *130*, 3888–3891; j) Y. Takenaka, T. Shima, J. Baldamus, Z. Hou, *Angew. Chem.* **2009**, *121*, 8028–8031; *Angew. Chem. Int. Ed.* **2009**, *48*, 7888–7891; k) T. Shima, Z. Hou, *Dalton Trans.* **2010**, *39*, 6858–6863.
- [4] M. Ohashi, M. Konkol, I. Del Rosal, R. Poteau, L. Maron, J. Okuda, *J. Am. Chem. Soc.* **2008**, *130*, 6920–6921.
- [5] D. M. Lyubov, C. Doring, G. K. Fukin, A. V. Cherkasov, A. S. Shavyrin, R. Kempe, A. A. Trifonov, *Organometallics* **2008**, *27*, 2905–2907.
- [6] X. Li, J. Baldamus, M. Nishiura, O. Tardif, Z. Hou, *Angew. Chem.* **2006**, *118*, 8364–8368; *Angew. Chem. Int. Ed.* **2006**, *45*, 8184–8188.
- [7] L. Zhang, T. Suzuki, Y. Luo, M. Nishiura, Z. Hou, *Angew. Chem.* **2007**, *119*, 1941–1945; *Angew. Chem. Int. Ed.* **2007**, *46*, 1909–1913.
- [8] [(Me-PNP^{Pr})Lu(CH₂SiMe₃)₂] has been reported: J. D. Masuda, K. C. Jantunen, O. V. Ozerov, K. J. T. Noonan, D. P. Gates, B. L. Scott, J. L. Kiplinger, *J. Am. Chem. Soc.* **2008**, *130*, 2408–2409.
- [9] The hydrogenolysis of **1**-Y in THF in the absence of [NEt₃H][BPh₄] did not give a characterizable product. Neither did that of **1**-Y in toluene in the presence of [NEt₃H][BPh₄].
- [10] Structurally characterizable cationic binuclear trihydrides bearing amidinate ancillary ligands, such as [(PhC(NAr*)₂)₂Y₂H₃-(thf)₄][BPh₄] (Ar* = 2,6-diisopropylphenyl), were also obtained by the hydrogenolysis of [{PhC(NAr*)₂}Y(CH₂SiMe₃)₂(thf)] with 0.5 equivalents of [NEt₃H][BPh₄] in a THF solution.