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Dehydrogenation of iron amido-borane and resaturation of the imino-borane complex†

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We report on the first isolation and structural characterization of an iron phosphinoimino-borane complex $Cp*Fe(\eta^2-H_2B=NC_6H_4PPh_2)$ by dehydrogenation of iron amido-borane precursor $Cp*Fe(\eta^1-H_3B-NHC_6H_4PPh_2)$. Significantly, regeneration of the amido-borane complex has been realized by protonation of the iron(II) imino-borane to the amino-borane intermediate $[Cp*Fe(\eta^2-H_2B-NHC_6H_4PPh_2)]^{+}$ followed by hydride transfer. These new iron species are efficient catalysts for 1,2-selective transfer hydrogenation of guinolines with ammonia borane.

Because of relevance to H2 storage1-10 and hydrogenation catalysis, 11-15 metal amine-borane complexes 16-18 and their dehydrogenated forms, such as amino-boranes²⁰⁻²² and iminoboranes4 are arising as a significant family in organometallic chemistry. In transition metal-catalyzed dehydrocoupling of amine-boranes and related transfer hydrogenations, the interactions between the metal and the borane fragment are essendehydrogenation and the consequent transformations. 16-20 Specifically, amino-borane complexes containing a M-H₂B=NR₂ moiety are the primary dehydrogenated species and are often identified as a resting point in the catalysis (Scheme 1a).20-22 Management of reversible dehydrogenation-regeneration reactions on a M-BH₂=NR₂ platform could provide a strategy with which to design efficient catalysts capable of operating sustainable syntheses.

Wider exploration of metal amino-borane chemistry is challenging since M-H₂B=NH₂ species are very reactive toward H₂ release. In 2010, Aldridge *et al.* reported the isolation of [(IMes)₂Rh(H)₂(η^2 -H₂B=NR₂)] and [(IMes)₂Ir(H)₂(η^2 -H₂B=NR₂)] from the metal-catalyzed dehydrogenation of R₂-HN·BH₃.^{21 α} At the same time, Alcaraz and Sabo-Etienne reported the preparation of (PCy₃)₂Ru(H)₂(η^2 -H₂B=NH_nMe_{2-n}) (n = 0-2) complexes^{22 α} by the dehydrogenation of amine-boranes with the corresponding ruthenium precursors. Subsequently, a straightforward synthesis of Ru, Rh, and Ir amino-borane complexes by reaction of H₂B=NR₂ (R = iPr or Cy) with the bis(hydrogen) complexes of M(H)₂(η^2 -H₂)₂(PCy₃)₂ or

 $[CpRu(PR_3)_2]^+$ fragments was developed. Turculet et al.

have shown that the ruthenium-alkoxide complex is able to

activate H₃B·NHR₂ producing hydrido ruthenium complex.²³

Notably, Weller and Macgregor found that dehydrocoupling of

ammonia-borane by [Ph₂P(CH₂)₃PPh₂Rh(η⁶-C₆H₅F)] affords a μ-

amino-borane bimetallic Rh complex, in which the simplest

has attracted great interest,^{24–29} iron amine-borane complexes, their dehydrogenated derivatives, and especially the catalysis

relevant to organic synthesis are largely unexplored. Recently,

Kirchner et al. reported a pincer-type iron complex generated by

protonation of the borohydride iron complex (PNP)Fe(H)(η^2 -

BH₄) with ammonium salts.³⁰ Inspired by earlier research on M-

H₂B=NR₂ chemistry, we intended to establish the reversible

Although iron-catalyzed dehydrocoupling of amine-boranes

H₂B=NH₂ moiety is trapped on a rhodium dimer.^{20a}

$$L_nM + H_3B\cdot NHR_2 \xrightarrow{-H_2} L_nM \xleftarrow{H}_B = N \xrightarrow{R}_R$$
 $M = Ru, Rh, Ir, R = Cy, ^iPr or H$

b) Hypothetical dehydrogenation-regeneration of iron amido-borane

 $\begin{tabular}{lll} Schematic & The presentation of metal-based a mine-borane dehydrogenation. \end{tabular}$

conversions of amino-borane complexes and their dehydrogenated forms in a synthetic piano-stool iron system. Herein, we report dehydrogenation of iron amido-borane complex $Cp*Fe(\eta^1-H_3B-NHC_6H_4PPh_2)$ (2) $(Cp*=Me_5C_5^-)$ to the iminoborane complex $Cp*Fe(\eta^2-H_2B=NC_6H_4PPh_2)$ (3), and

a) Fundamental step involved in catalytic amine-borane dehydrogenation

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resaturation of the imino-borane by stepwise protonation and hydride transfer (Scheme 1b). This new class of iron species is capable of catalyzing 1,2-selective transfer hydrogenation of quinolines with $H_3N \cdot BH_3$.

To synthesize the iron amido-borane complex, a new monomer, the iron tetrahydridoborate precursor Cp*Fe(η^2 -BH₄)(NCMe) (1), was prepared *in situ* by the reaction of [Cp*Fe(NCMe)₃]PF₆ with Bu₄NBH₄ in acetonitrile at room temperature for 5 min. Such ferrous borohydrides have been documented only rarely,³¹ since they are prone to form polynuclear iron borate clusters.^{32,33} The ¹¹B NMR spectrum of the reaction solution shows a quintet at δ 15.4 ($J_{\rm BH}$ = 88 Hz) for the BH₄⁻ ligand of 1, and this stands in contrast to the signal at δ –32.0 observed for Bu₄NBH₄. Upon storing the reaction mixture at –30 °C overnight, single crystals suitable for X-ray diffraction were obtained. Crystallographic analysis confirmed the structure of 1 as a piano-stool iron tetrahydridoborate compound (ESI, Fig. S1†).

Addition of phosphinoamine ligand 1,2-Ph₂PC₆H₄NH₂ to a solution of 1 in acetonitrile caused an instantaneous color change from deep blue to dark brown (Scheme 2). ESI-MS studies indicated the production of the iron amido-borane compound (2) with m/z = 481.1793 (calcd m/z = 481.1770), which was isolated in 87% yield. NMR spectra showed a boron resonance at δ -17.5, and a phosphorus resonance at δ 85.9. The ¹H NMR spectrum exhibits a characteristic hydride signal at δ –13.98, which is assigned to the bridging hydride Fe–*H*–B. Owing to exchange between the hydrogen atoms at the boron,³⁴ the terminal B-H resonances in the ¹H NMR spectrum are very broad and are obscured by the distinct Cp* signals. To assign the B-H hydride signals, the deuterated compound Cp*Fe(D₃B-NHC₆H₄PPh₂) (d-2) was synthesized from Cp*Fe(BD₄)(NCMe). In addition to the Fe-D-B signal at δ -13.98, the ²H NMR spectrum of d-2 displayed discrete peaks at δ 2.23 and 0.19 for the terminal B-D hydrides (Fig. 1).

When a C_6H_6 solution of 2 was held at 50 °C for 6 h the dehydrogenated imino-borane compound (3) was produced in 92% yield. The ESI-MS spectrum of 3 has a strong peak at m/z 479.1626 (calcd m/z=479.1637) which can be compared to the peak at m/z=481.1793 for 2. The isotopic distributions match well with the calculated values (see Fig. S3†). GC analysis shows that the reaction produced H_2 nearly quantitatively (see Fig. S4†). In solution, the ^{31}P NMR spectrum of 3 displays a sharp signal at δ 71.9, in contrast to the peak at δ 85.9 for 2. The ^{11}B resonance shifts significantly, from δ -17.5 for 2 to δ 42.7 for 3 (Fig. S16†), and is particularly diagnostic of a three-coordinate boron atom. 21,35 This result indicates the B=N

Scheme 2 Synthetic route to imino-borane complex.

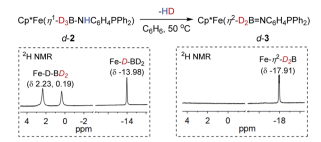


Fig. 1 2 H NMR spectra for dehydrogenation of d-2 to d-3.

double bond character in the dehydrogenated form of the amido-borane complex. In the ^1H NMR spectrum, the Fe–H–B signal was observed at δ –17.91 with the integral of 2H, and no characteristic signal for a terminal B–H hydride was found. To confirm the formation of an imino-borane compound, the hydrogen decoupling was also carried out with compound d-2 and monitored by ^2H NMR spectra. Only a deuterium signal was observed at δ –17.91 for Fe–D–B, indicating the formation of d-3 (Fig. 1). When the dehydrogenation was conducted in a J-Young tube in C₆D₆, a characteristic triplet corresponding to HD appeared at δ 4.43 ($J_{\text{HD}}=45$ Hz) in the ^1H NMR spectrum (Fig. S18†).³⁶

The structures of 2 and 3 were verified by X-ray crystallographic analysis (Fig. 2). Consistent with NMR spectroscopic analysis, the BH₃ moiety in 2 is stabilized by one of the B–H bonds binding at the Fe–NH unit to form an Fe–H–B–N four-membered metallacycle. This metal-ligand cooperative binding mode increased the B–H bond length in the bridging B–H(1) bond to 1.362 Å ν s. 1.129 Å and 1.121 Å for the two terminal B–H bonds. The B–N bond length of 1.545(3) Å in 2 is slightly shorter than that in H₃B·NH₃ ($d_{\rm B-N}=1.58(2)$ Å).³⁷ Crystallographic analysis of 3 confirmed an imino-borane complex with a Cp*Fe(η^2 -H₂B=NC₆H₄PPh₂) framework. After dehydrogenation of 2, striking structural changes were observed. The N atom

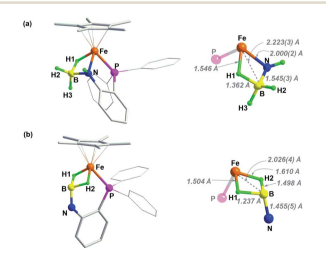


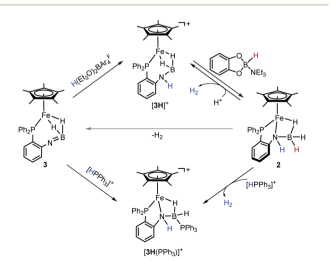
Fig. 2 Solid-sate structure (50% probability thermal ellipsoids) of (a) complex 2 and (b) 3. For clarity, hydrogen atoms of Cp* and phenyl rings are omitted.

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has been become detached from Fe, while the BH $_2$ fragment acts as a bis(σ -borane) ligand coordinated to the metal center. $^{21-23}$ The B–N bond distance of 1.455(5) Å in 3 is shorter by 0.09 Å than that in 2, and is close to that reported for the cyclic trimer borazine (1.4355(21) Å). 38 Combined with the NMR results, the B–N bond length in 3 suggests some double bond character. 21,22 As the imino-borane fragment is tethered in the coordination sphere, the boron center adopts a quasitetrahedral geometry, and the B–N bond appears to be partially sp 3 hybridized. Dehydrogenation of the amido-borane complex also caused the decrease of the Fe···B distances from 2.223(3) Å to 2.026(4) Å which is shorter than the sum of the covalent radii of Fe and B atom (2.16 Å), indicating that the borane and the metal are bonded.

Notably, the amido-borane compound 2 can be regenerated by stepwise protonation of 3 and transfer of a hydride (Scheme 3). Complex 3 reacts readily with H(Et₂O)₂BAr₄^F in C₆H₅F. The reaction solution was analyzed by ESI-MS spectroscopy, which showed an ionic peak at m/z = 480.1726 (calcd m/z = 480.1715), suggesting the formation of [3H]⁺. Alternatively, the reaction of complex 2 with H(Et₂O)₂BAr₄^F unambiguously provides [3H]⁺ and produces H₂. X-ray crystallographic analysis reveals that the resulting cationic complex [3H]+ exhibits a similar framework to its imino-borane precursor (3). The BH2 moiety retains a binding mode of the bis(σ -BH₂) fashion (Fig. 3). In contrast, the B-N distance in $[3H]^+$ (1.586(6) Å) is extended by 0.13 Å and the [3H]⁺ framework becomes much less compact than that of 3. Probably due to the fluxional structure of the seven-membered Fe-P-C-C-N-B(H) ring, the solution of [3H][BAr₄F] gives broad 1 H NMR resonances even at -60 $^{\circ}$ C. The phosphorus resonance arose at δ 72.0 as a singlet when the solution sample was cooled to -40 °C (Fig. S20 and S21†).

In [3H]⁺, the boron is coordinatively unsaturated, as manifested by its interaction with a σ -donor. For instance, treatment of 2 with [HPPh₃][BAr₄^F] (p $K_a^{\text{MeCN}} = 7.6$)³⁹ provides a Ph₃P-stabilized borane complex, [3H(PPh₃)]⁺ (m/z = 742.2620, calcd m/z = 742.2626). The ¹H NMR spectrum of [3H(PPh₃)]⁺ exhibits



Scheme 3 Conversions of iron imino-borane, amino-borane and amido-borane complexes.

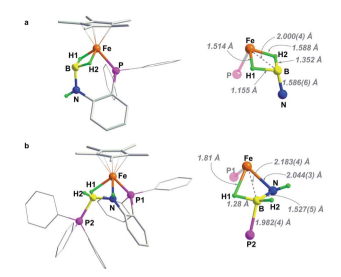


Fig. 3 Solid-state structures of (a) complex $[3H]^+$ and (b) $[3H(PPh_3)]^+$. For clarity, counterion $[BAr_4^F]^-$, hydrogen atoms of Cp* and phenyl rings have been omitted.

an N*H* resonance at δ 4.68, suggesting that protonation occurred at the N site. The distinctive upfield hydride signal for Fe–*H*–B is observed at δ –15.58. In the ³¹P NMR spectrum, two phosphorus signals at δ 78.90 and –1.26 correspond to the Fe–*P* and the B–*P* resonances, respectively. The ¹¹B signal at δ –13.72 indicates a tetracoordinated boron, which is further confirmed by crystallographic analysis of [3H(PPh₃)]⁺ (Fig. 3). In the solid-sate structure, a Ph₃P molecule is bound to the B center ($d_{\rm B-P}$ = 1.982(4) Å), leading to the formation of a new Fe–H–B–N four-membered metallacycle. As a amido-borane complex, [3H(PPh₃)]⁺ has a B–N bond length of 1.527(5) Å, somewhat shorter than 1.545(3) Å in 2.

After attaching a proton at the N atom, we subsequently explored restoration of the original borane moiety. Treatment of freshly prepared [3H][BAr₄F] in fluorobenzene with catecholborane-NEt₃ adduct ($\delta_{\rm B}=10.56, J_{\rm HB}=142.4~{\rm Hz})^{40}$ results in the regeneration of 2, as evidenced by the NMR spectra (Fig. S29 and S30†). The ¹H NMR spectrum of the reaction mixture displays a characteristic hydride signal at -13.97 ppm, indicating the recovery of the iron amido-borane complex. On the other side, concomitant formation of the borenium ion ($\delta_{\rm B}$ = 13.86) was also observed in the 11B NMR spectrum, which agrees with the hydride transfer from the organohydride reagent to $[3H]^+$. It was interesting that the ion $[3H]^+$ is stable towards 5,6-dihydrophenanthridine and Hantszch ester. These results indicate that the hydride-donating ability $(\Delta G_{\text{H}}^{-})$ of 2 is in the range of 55-59 kcal mol⁻¹.41 The reactive nature of the hydride in 2 was demonstrated by the reaction with [HPPh3] $[BAr_4^F]$, which produces $[3H(PPh_3)]^+$ and releases H_2 (Scheme 3).

The metal amine-borane complexes and their dehydrogenated derivatives are implicated throughout the catalytic cycle of amine-borane dehydrogenation. We found both the iron complexes 2 and 3 are efficient catalysts for H₃N·BH₃ dehydrogenation at room temperature. In the presence of 1 mol% catalyst, a THF solution of H₃N·BH₃ (1.0 mmol) generates about 2.2 equivalent of H₂ within 6 h based on GC quantification (Fig. S33†). More importantly, such catalytic dehydrocoupling systems allow for selective transfer hydrogenation of quinolines to dihydroquinolines, which are valuable synthons leading to many bio-active compounds. 42 For instance, addition of methyl-6-quinolineacetate (4) to the catalytic system containing one equiv. of H₃N·BH₃ and 1 mol% of 3 gave 1,2dihydro-methyl-6-quinolineacetate (5) in excellent yield within 6 h (egn (1)). The outcome of this reaction was unaffected by switching the catalyst from 3 to 2, or by use of excess reducing agent or by an increase in the reaction temperature (Table S1†).

Conclusions

By tethering the N-B unit within the coordination sphere, we have demonstrated an example of imino-borane iron(II) dehydrogenation complex isolated from its phosphinoamido-borane precursor, and have realized the regeneration of an Fe-H₃B-N(H)Ar fragment by submitting the dehydrogenated imino-borane to sequential protonation and hydride transfer reactions. Based on the dehydrogenation of ammonia-borane catalyzed by the two iron species, the catalytic reduction of quinoline to 1,2-dihydroquinoline was established. This work provides a new perspective for the studies of reversible conversions between amine-borane complexes and the dehydrogenated forms, and exploration of iron-based catalysis for important organic transformations.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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