

Conversion of Fe-NH₂ to Fe-N₂ with release of NH₃

John S. Anderson, Marc-Etienne Moret, and Jonas C. Peters*

Division of Chemistry and Chemical Engineering, California Institute of Technology, Pasadena, California 91125, United States

Supporting Information

ABSTRACT: Tris(phosphine)borane ligated Fe(I) centers featuring N2H4, NH3, NH2, and OH ligands are described. Conversion of Fe-NH₂ to Fe-NH₃⁺ by the addition of acid, and subsequent reductive release of NH3 to generate Fe-N2, is demonstrated. This sequence models the final steps of proposed Fe-mediated nitrogen fixation pathways. The five-coordinate trigonal bipyramidal complexes described are unusual in that they adopt S = 3/2 ground states and are prepared from a four-coordinate, S = 3/2 trigonal pyramidal precursor.

ue to the structural and mechanistic complexity of biological nitrogen fixation¹ a variety of mechanisms have been proposed that invoke either Mo or Fe as the likely active site for N₂ binding and reduction. Fe-NH₂ is an intermediate common to both limiting mechanisms (i.e., distal vs alternating) being considered for Fe-mediated N2 fixation scenarios at the FeMo-cofactor.^{2,3} Such a species could form via reductive protonation of the nitride intermediate of a distal scheme (i.e., $Fe(N) \rightarrow Fe(NH) \rightarrow Fe(NH_2) \rightarrow Fe(NH_3)$), or by reductive protonation of a hydrazine intermediate of an alternating scheme (i.e., $Fe(NH_2-NH_2) \rightarrow Fe(NH_2) + NH_3$). In this context, detection of an EPR active Fe-NH2 or possibly Fe-NH₃ common intermediate has been proposed under reducing conditions at the FeMo-cofactor from substrates including N₂, N₂H₄, and MeN=NH.^{3a}

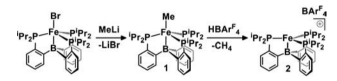
One key to realizing a catalytic cycle in either limiting scenario concerns the regeneration of Fe-N2 from Fe-NH2 with concurrent release of NH₃.⁴ While there have been recent synthetic reports demonstrating NH₃ generation from Fe(N) nitride model complexes, these studies have not provided information about the plausible downstream $Fe(NH_r)$ (X = 1, 2, 3) intermediates en route to NH₃ release, nor have these systems illustrated the feasibility of regeneration of Fe-N₂. Herein we describe a terminal, S = 3/2 Fe-NH₂ complex for which the stepwise conversion to Fe-NH₃, and then to Fe-N₂ along with concomitant release of NH3, is demonstrated (eqs 1 and 2).

$$Fe - NH_2 + H^+ \rightarrow Fe - NH_3^+ \tag{1}$$

$$Fe - NH_3^+ + e^- + N_2 \rightarrow Fe - N_2 + NH_3$$
 (2)

The addition of methyllithium to (TPB)FeBr⁶ affords the corresponding methyl complex (TPB)FeMe (1) in high yield (Scheme 1). Protonation of 1 by $HBAr_4^F \cdot 2Et_2O$ ($BAr_4^F B(3,5-C_6H_3(CF_3)_2)_4^-)$ in a cold ethereal solution releases methane to yield [(TPB)Fe][BArF4] (2) which serves as a useful synthon with a vacant coordination site.

Scheme 1



XRD data were obtained for 1 and 2 (Figure 1). The geometry of 1 is pseudotrigonal bipyramidal about Fe with an Fe-C bond length of 2.083(10) Å and an Fe-B bond length of 2.522 (2) Å. In the solid state 2 possesses a four-coordinate distorted trigonal pyramidal geometry with no close contacts in the apical site trans to boron, making this complex coordinatively unsaturated. Additionally, there is one wide P-Fe-P angle of 136°. The origin of this large angle is not clear, but a possible explanation is increased back-bonding from a relatively electron rich Fe center into the phosphine ligands that would arise from this distortion (see SI).

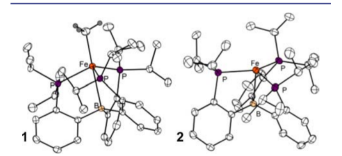


Figure 1. X-ray diffraction (XRD) structures of complexes 1 (A) and 2 (B) with hydrogen atoms and counterion (for B) omitted for clarity. See Table 1 for selected bond lengths and angles.

The Fe-B distance in 2 (2.217(2) Å) is markedly shorter than that in (TPB)FeBr (2.459(5) Å), which is noteworthy because one might expect the loss of an anionic σ -donor ligand to reduce the Lewis basicity of the metal and thus weaken the Fe-B bond. For example, the Au-B distance in (TPB)AuCl (2.318 Å) lengthens upon chloride abstraction to 2.448 Å in [(TPB)Au]^{+,7} To explain this difference we note that the boron center in four-coordinate 2 is less pyramidalized ($\Sigma(C-B-C)$ = 347.3°) than that in five-coordinate (TPB)FeBr (Σ (C-B-C) = 341.2°), pointing to a weak interaction despite the short distance. This observation suggests that the geometry of 2 might be best understood as derived from a planar three-

Received: August 3, 2012 Published: December 21, 2012 coordinate Fe(I) center distorted toward a T-shaped geometry,⁸ the unusually short Fe-B distance being due largely to the constraints imposed by the ligand cage structure. This interpretation is consistent with a computational model study: the DFT (B3LYP/6-31G(d)) optimized geometry of the hypothetical complex $[(Me_2PhP)_3Fe]^+$ (see SI for details) exhibits a planar geometry with P-Fe-P angles of 134.8°, 113.1°, and 111.7°, very close to those measured for $[(TPB)Fe]^+$ (137.5°, 113.2°, 109.1°).

When considering the bonding of the $(Fe-B)^7$ subunit of 2 to estimate the best oxidation state and valence assignment, two limiting scenarios present themselves: Fe(III)/B(I) and Fe(I)/ B(III). The structural data and computations for 2 are suggestive of a weak Fe-B interaction and indicate that this species is better regarded as Fe(I)/B(III) rather than Fe(III)/ B(I). Calculations indicate that a small amount of spin density resides on the B-atom of 2 (SI) and suggest some contribution from an Fe(II)/B(II) resonance form may also be relevant. The rest of the complexes 3-6 presented herein possess significantly longer, and presumably weaker, Fe-B interactions (vide infra) and are hence also better classified as Fe(I) species. Additional spectroscopic studies (e.g., XAS and Mössbauer) will help to better map the Fe-B bonding interaction across the variable Fe-B distances and also the spin states of the complexes. These studies would thereby help to determine the value and limitation of classically derived oxidation/valence assignments for boratranes of these types.

Table 1. Selected Metrics for Complexes 1-6

	Fe-X (Å)	Fe-B (Å)	avg Fe-P (Å)	Σ P–Fe–P	Σ С-В-С
1	2.083(10)	2.523(2)	2.40	339°	341°
2		2.217(2)	2.38	359°	347°
3	2.205(2)	2.392(2)	2.44	350°	339°
4	2.280(3)	2.433(3)	2.44	349°	341°
5	1.918(3)	2.449(4)	2.39	343°	339°
6	1.8916(7)	2.4438(9)	2.39	348°	337°

Solutions of 2 are orange in Et₂O and pale yellow-green in THF. The titration of THF into an ethereal solution of 2 results in a distinct change in the UV–vis spectrum consistent with weak THF binding. Addition of an excess of N₂H₄ to an ethereal solution of 2 results in a slight lightening of the orange color of the solution to afford [(TPB)Fe(N₂H₄)][BAr^F₄] (3) in 89% yield. Complex 3 shows a paramagnetically shifted ¹H NMR spectrum indicative of an S=3/2 Fe center that is corroborated by a room temperature solution magnetic moment, $\mu_{\rm eff}$ of 3.5 $\mu_{\rm B}$. Crystals of 3 were obtained, and XRD analysis (Figure 2A) indicates a distorted trigonal bipyramidal geometry. The Fe–N distance of 2.205(2) Å is unusually long (2.14 Å is the average quaternary N–Fe distance in the Cambridge Structural Database) reflecting its unusual quartet spin state.

Complex 3 is stable to vacuum, but solutions decompose cleanly at room temperature over hours to form the cationic ammonia complex [(TPB)Fe(NH₃)][BAr^F₄], 4, which was assigned by comparison of its ¹H NMR spectrum with an independently prepared sample formed by the addition of NH₃ to the cation 2. Analysis of additional degradation products shows only NH₃ and trace H₂ (SI). The assignment of 4 as an ammonia adduct was confirmed by XRD analysis (Figure 2B). Like 3, complex 4 shows a long Fe—N distance of 2.280(3) Å in the solid state. The complexes 2, 3, and 4 are unusual by virtue

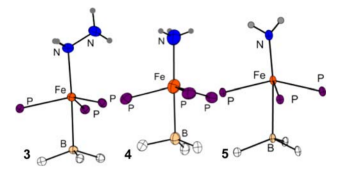


Figure 2. XRD structures of the cores of complexes 3 (A), 4 (B), and 5 (C). See Table 1 for selected distances and angles.

of their S = 3/2 spin states and underscore the utility of local 3-fold symmetry with respect to stabilizing high spin states at iron, even in the presence of strong-field phosphine ligands.

The addition of NaNH₂ to the cation **2** affords the terminal amide, (TPB)Fe–NH₂ (**5**) in ca. 85% nonisolated yield by 1 H NMR integration. The XRD structure of **5** (Figure 2C) shows an overall geometry similar to that observed in **1**, **3**, and **4**. Of interest is the short Fe–N distance of 1.918(3) Å by comparison to **4** (2.280(3) Å). The amide hydrogens were located in the difference map and indicate a nearly planar geometry about N (with the sum of the angles around N being 355°).

While the XRD data set of 5 is of high quality, we were concerned about the difficulty in distinguishing an Fe-NH₂ group from a potentially disordered Fe-OH moiety. We therefore independently characterized the hydroxo complex, (TPB)Fe-OH (6) (Scheme 2), which possesses a geometry

Scheme 2

similar to that observed in **5** with an Fe–B distance of 2.4438(9) Å and an Fe–O distance of 1.8916(7) Å. Despite the structural similarity between **5** and **6**, different spectral signatures in both their 1 H NMR and EPR (Figure 3) spectra allow for facile distinction between them. Like **2**, **3**, and **4**, both **5** and **6** are S = 3/2.

Low-temperature EPR data (Figure 3) have been obtained on complexes 1–6. All complexes show features shifted to large g-values consistent with quartet Fe species. ¹⁰ This assignment is verified by the solution magnetic moments obtained for these complexes. Variable temperature solid-state SQUID magnetic data for complexes 2–5 (SI) also establish quartet spin-state

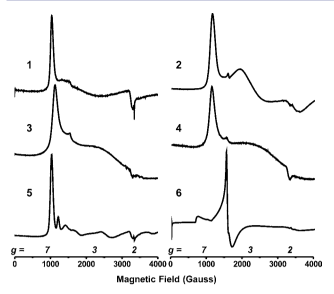


Figure 3. X-Band EPR spectra for complexes 1–6. Conditions: (1) toluene, 8 K; (2) 2:1 toluene/Et₂O, 10 K; (3) 2-MeTHF, 10 K; (4) 2-MeTHF, 10 K; (5) 2-MeTHF, 10 K; (6) toluene, 10 K.

assignments and display no evidence of spin-crossover phenomena. These data show a drop in magnetic moment in the range 50–70 K for all compounds studied. We propose that this effect is due to a large zero-field splitting in these species, which is consistent with Fe centers in related geometries. Simulations with zero-field splitting of 10–20 cm⁻¹ provide reasonable fits to the data.

Parent amide complexes of first row transition metals are rare. ¹² Noteworthy precedent for related terminal M-NH₂ species includes two square planar nickel complexes ^{12a,d} and one octahedral and diamagnetic iron complex, (dmpe)₂Fe(H)-NH₂. ^{12e} In addition to their different coordination numbers, geometries, and spin-states, (dmpe)₂Fe(H)(NH₂) and 5 show a distinct difference at the Fe–NH₂ subunit. Six-coordinate (dmpe)₂Fe(H)(NH₂) is an 18-electron species without π-donation from the amide ligand, which is pyramidalized as a result. By contrast, five-coordinate 5 accommodates π-bonding from the amide. This is borne out in its much shorter Fe–N distance (1.918(3) Å for 5 vs 2.068 Å for (dmpe)₂Fe(H)-(NH₂)), and also its comparative planarity (the sum of the angles around N is 355° for 5 vs 325° for (dmpe)₂Fe(H)-(NH₂)).

With the terminal amide 5 in hand we explored its suitability as a precursor to the previously reported N2 complex (TPB)Fe(N₂) via release of NH₃, and hence explored reduction/protonation vs protonation/reduction sequences as a means of effecting overall H-atom transfer to the Fe-NH₂ unit. Attempts to carry out the one-electron reduction of 5 were not informative. For example, electrochemical studies of 5 in THF failed to show any reversible reduction waves, but the addition of harsh reductants (e.g., tBuLi) to 5 did show small amounts of (TPB)Fe(N2) in the product profile. A more tractable conversion sequence utilized protonation followed by chemical reduction. Thus, the addition of HBAr^F₄·2Et₂O to 5 at low temperature (-35 °C) rapidly generates the cationic ammonia adduct 4. The conversion is quantitative as determined by ¹H NMR spectroscopy, and 4 can be isolated in ca. 90% yield from the solution. Subsequent exposure of 4 to 1 equiv of KC₈ under an atmosphere of N₂ releases NH₃ and generates the (TPB)FeN2 complex in similarly high yield.

In summary, an unusual series of S=3/2 iron complexes featuring terminally bonded N_2H_4 , NH_3 , NH_2 , and OH functionalities has been thoroughly characterized. These complexes are supported by a tris(phosphine)borane ligand and are best described as Fe(I) species that feature weak Fe-B bonding, though other resonance contributions to the bonding scheme warrant additional consideration. The $Fe-NH_2$ species faithfully models the reductive replacement of the terminal NH_2 group by N_2 with concomitant release of NH_3 , lending credence to such a pathway as mechanistically feasible in Fe- mediated N_2 reduction schemes. Because spectroscopic detection of a common $Fe-NH_2$ or $Fe-NH_3$ intermediate under reductive turnover of the FeMo-cofactor has been recently proposed, 3 EPR active model complexes of the types described here should prove useful for comparative purposes.

ASSOCIATED CONTENT

S Supporting Information

Detailed experimental and spectroscopic data. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

Corresponding Author

jpeters@caltech.edu

Notes

The authors declare no competing financial interest.

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