

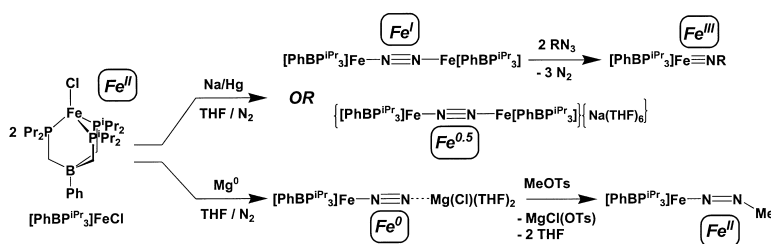
Communication

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Dinitrogen Chemistry from Trigonally Coordinated Iron and Cobalt Platforms

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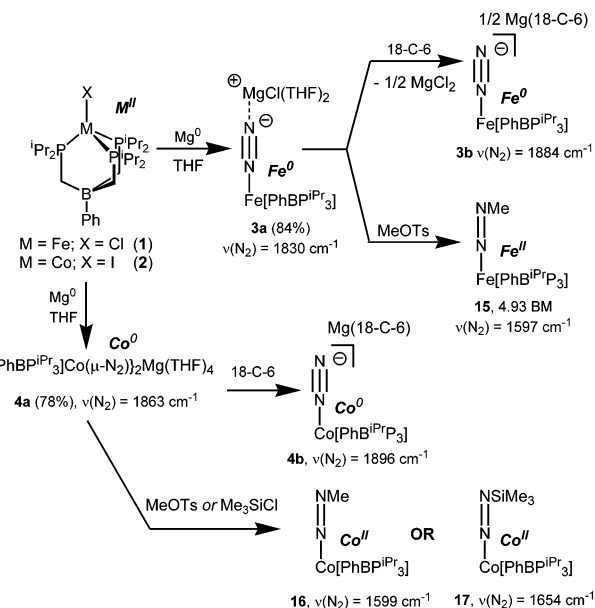
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The Chatt-type nitrogen reduction scheme is well-established for certain molecular systems that feature molybdenum.¹ For example, Schrock has recently demonstrated that a single, amido-supported molybdenum center can support various N₂-derived intermediates (e.g., M(N₂H), M(N₂H₂), M(N), M(NH)), oxidation states, and redox couples to favorably mediate the complete reduction of nitrogen to ammonia in a catalytic fashion.^{1c,d} One key feature of this and other competent N₂-reducing molybdenum systems is their ability to accommodate a π-acidic N₂ ligand, as well as π-basic functionalities derived from N₂ (e.g., N³⁻, NH²⁻), at a single site.^{1,2} To date, very few first-row ion platforms allow N₂ to be taken up and derivatized in a similar fashion (e.g., Fe(N₂) → Fe(N₂H)).^{3,4} Iron is particularly noteworthy in this regard, given its possible if not likely role in biological (and industrial) nitrogen fixation.^{5,6}

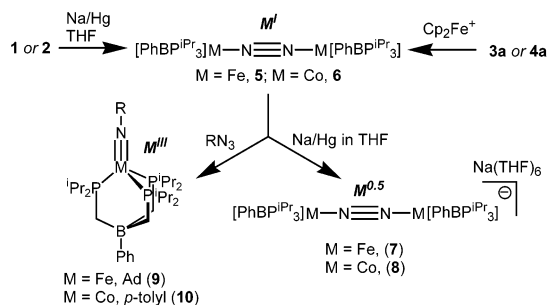
Recent work by our group has shown that trigonally coordinated iron and cobalt subunits of the form “[PhBP₃]M” ([PhBP₃] = [PhB(CH₂PPh₂)₃]⁻) will support a strongly π-donating ligand at a fourth site along their pseudo-three-fold axis (e.g., [PhBP₃]Fe≡NR).⁷ Simple electronic structure considerations suggest that these same [PhBP₃]M subunits should also accommodate strongly π-acidic ligands at the fourth site,⁷⁻⁹ a feature that might complement nitrogen reduction schemes at iron. Herein we demonstrate that second-generation “[PhBPⁱPr₃]Fe” ([PhBPⁱPr₃] = [PhB(CH₂PⁱPr₂)₃]⁻) and related cobalt systems coordinate, activate, and promote the methylation of nitrogen at the fourth binding site. Moreover, the N₂ ligand can be replaced by nitrene (NR) through an oxidative group-transfer reaction.^{7,8a} The iron–N₂ species discussed are the first thoroughly characterized, 4-coordinate complexes of their type.⁴

Entry to this reaction manifold begins with the recently reported precursors [PhBPⁱPr₃]MX (M = Fe, X = Cl (**1**); M = Co, X = I (**2**)).⁹ A THF solution of yellow **1** or green **2**, stirred under a blanket of nitrogen in the presence of Mg⁰, produced the anionic, paramagnetic dinitrogen complexes {[PhBPⁱPr₃]Fe(N₂)}⁻{MgCl(THF)₂}⁺ (**3a**, 84%) and {[PhBPⁱPr₃]Co(N₂)}⁻{Mg(THF)₄}⁺ (**4a**, 78%) as red-brown and red crystals, respectively (Scheme 1). An XRD study of single crystals of **4a** suffered from modest disorder but confirmed a Mg(THF)₄²⁺ dication sandwiched by two anionic Co(N₂)⁻ units, in accord with related structure types.^{10,11} The IR spectra of **3a** and **4a** show intense ν_{NN} bands that shift to higher energy upon addition of 18-crown-6 (18-C-6) to encapsulate the Mg²⁺ ion (ν_{NN}(¹⁵NN) in cm⁻¹ for **3a**, 1830(1769); **3b**, 1884(1822); **4a**, 1863(1802); **4b**, 1896(1842)). Gentle oxidation by ferrocenium (Cp₂Fe⁺) in THF produced the neutral, dinuclear N₂-bridged products {[PhBPⁱPr₃]Fe}₂(μ-N₂) (**5**, 88%) and {[PhBPⁱPr₃]Co}₂(μ-N₂) (**6**, 92%) (Scheme 2). Both **5** and **6** were also obtained directly from halides **1** and **2** when Na/Hg amalgam was used as the reductant in place of Mg⁰. Furthermore, extended exposure of **5** and **6** to sodium amalgam produced the mixed-valence (M⁰/M^I) complexes [(PhBPⁱPr₃]Fe)₂(μ-N₂)[Na(THF)₆] (**7**, 93%) and [(PhBPⁱPr₃]Co)₂(μ-N₂)[Na(THF)₆] (**8**, 89%). XRD data were obtained on single crystals of **7** and **8**, and the molecular structure

Scheme 1



Scheme 2



of **7** is shown in Figure 1. The structure of **8** is isomorphous and isostructural.¹¹ A crystallographically imposed inversion center reflects one-half of each dinuclear anion into the other, suggesting that these species are likely delocalized at low temperature. A modest elongation of the N₂ ligand is observed in each case (N–N' 1.171(4) for **7**, and 1.147(4) Å for **8**), and the local geometry of each metal center is perhaps best described as trigonal monopyramidal, with N₂ lying in an equatorial position.

Complexes **5** and **6** provided clean access to the “[PhBPⁱPr₃]M” fragment, as exemplified by their ability to undergo rapid oxidation upon addition of either tolyl or adamantyl azide to afford trivalent imide products. For example, burgundy [PhBPⁱPr₃]Fe≡NAd (**9**, 69%) and red [PhBPⁱPr₃]Co≡N-*p*-tolyl (**10**, 93%) were isolated and thoroughly characterized (Scheme 2), and the solid-state structure of each was determined.¹¹ The crystal structure of complex **9** (Figure 2) features a very short Fe–N bond (1.638(2) Å) and an almost linear Fe–N–C bond vector (176°), consistent with the triple-bond formulation we have proposed previously.⁷

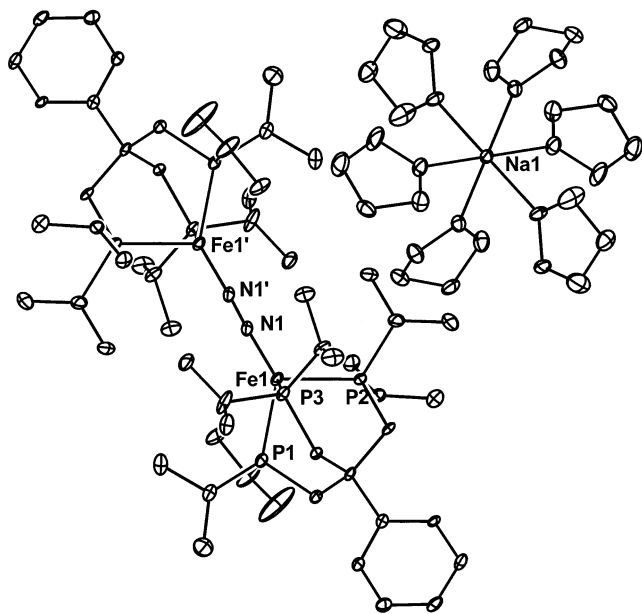


Figure 1. Displacement ellipsoid representation of $\{[(\text{PhBPiPr}_3)_2\text{Fe}]_2(\mu\text{-N}_2)\}\cdot\{\text{Na}(\text{THF})_6\}$ (**7**). Hydrogen atoms have been removed for clarity. Selected bond distances (Å) and angles (deg), for **7**: Fe–N 1.813(2), N–N' 1.171(4), Fe–P1 2.292(1), Fe–P2 2.278(1), Fe–P3 2.290(1); N'–N–Fe 178.0(3), P1–Fe–P2 96.93(4), P1–Fe–P3 97.96(4), P2–Fe–P3 97.22(4), N–Fe–P1 118.88(9), N–Fe–P2 109.66(9), N–Fe–P3 129.83(9).

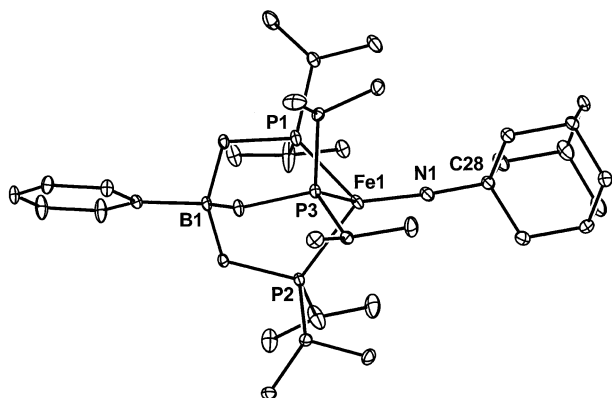


Figure 2. Displacement ellipsoid representation of $[\text{PhBPiPr}_3]\text{Fe}\equiv\text{NAD}$ (**9**). Hydrogen atoms have been removed for clarity. Selected bond distances (Å) and angles (deg), for **9**: Fe–N 1.638(2), Fe–P1 2.260(1), Fe–P2 2.297(1), Fe–P3 2.263(1); C28–N–Fe 176.0(2), P1–Fe–P2 91.05(2), P1–Fe–P3 91.56(3), P2–Fe–P3 92.39(3), N–Fe–P1 121.62(7), N–Fe–P2 130.03(7), N–Fe–P3 120.27(7).

Structural data for related **10** have been placed in the Supporting Information.

The anionic N_2 adducts **3** and **4** appear well-poised for further elaboration at the coordinated N_2 functionality. This is significant because the direct conversion of coordinated N_2 to a coordinated diazenido (N_2R^-) and/or hydrazido ($\text{N}_2\text{R}_2^{2-}$) species by simple addition of an electrophile (e.g., R^+) has, to our knowledge, not been established for a first-row transition metal ion.^{3,4} We were thus gratified to find that the simple addition of methyl tosylate (MeOTs) to a THF solution of **3** effected a modest color change from brown to gold, along with a concomitant shift in its $\nu_{\text{NN}}(^{15}\text{NN})$ vibration to 1597(1538) cm^{-1} , characteristic of a coordinated, monodentate diazenido functionality.¹ A neutral, benzene-soluble product was isolated from the reaction mixture that analyzed as the diazenido complex $[\text{PhBPiPr}_3]\text{Fe}-\text{N}=\text{NMe}$ (**11**, 56%), consistent with its magnetic susceptibility ($S = 2$, 4.93 μ_{B} , Evans). A similar protocol converted **4a** to the amber-colored complex

$[\text{PhBPiPr}_3]\text{Co}-\text{N}=\text{NMe}$ (**12**, 68%), which exhibited a very similar $\nu_{\text{NN}}(^{15}\text{NN})$ stretch 1599(1542) cm^{-1} and a solution moment indicative of low-spin cobalt(II) (1.92 μ_{B}). Complex **4a** also reacted cleanly with trimethylsilyl chloride to generate $[\text{PhBPiPr}_3]\text{Co}-\text{N}=\text{NSiMe}_3$ (**13**, 79%; $\nu_{\text{NN}}(^{15}\text{NN}) = 1654(1596) \text{cm}^{-1}$; 2.08 μ_{B}). The latter product could be obtained in good yield by a more direct procedure that involved stirring a solution of chloride **2** in the presence of Me_3SiCl and Na/Hg amalgam under nitrogen.

These data allow us to summarize several salient features of the chemistry described herein: The $[\text{PhBPiPr}_3]$ ligand has been used to isolate a single iron center in a pseudo-tetrahedral environment in which a single binding site is compatible with coordination of N_2 , diazenido (N_2Me), and imide (NR). Moreover, N_2 uptake/coordination by 4-coordinate iron has been established for the formal oxidation states M(0), M(+0.5), and M(+1). Methylation of $\text{Fe}^0(\text{N}_2)^-$ to produce $\text{Fe}^{\text{II}}(\text{N}_2\text{Me})$ constitutes a two-electron redox process at iron. The additional observation of a separate two-electron redox process, that of oxidative nitrene transfer ($\text{Fe}^{\text{I}} \rightarrow \text{Fe}^{\text{III}} \equiv \text{NR}$), establishes that the “ $[\text{PhBPiPr}_3]\text{Fe}$ ” platform will support four formal oxidation states (Fe^0 , Fe^{I} , Fe^{II} , Fe^{III}), in addition to two discrete two-electron redox couples ($\text{Fe}^{0/\text{II}}$ and Fe^{III}). An analogous set of observations has been made with the related cobalt system. These features collectively motivate further development of the present first-row systems into possible models for functional nitrogen fixation.

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Supporting Information Available: Complete experimental procedures and characterization data, and crystallographic data (PDF, CIF). This material is available free of charge via the Internet at <http://pubs.acs.org>.

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