

Low-Temperature N₂ Binding to Two-Coordinate L₂Fe⁰ Enables Reductive Trapping of L₂FeN₂[−] and NH₃ Generation**

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Abstract: The two-coordinate [(CAAC)₂Fe] complex [CAAC = cyclic (alkyl)(amino)carbene] binds dinitrogen at low temperature ($T < -80^{\circ}\text{C}$). The resulting putative three-coordinate N₂ complex, [(CAAC)₂Fe(N₂)], was trapped by one-electron reduction to its corresponding anion [(CAAC)₂FeN₂][−] at low temperature. This complex was structurally characterized and features an activated dinitrogen unit which can be silylated at the β-nitrogen atom. The redox-linked complexes [(CAAC)₂Fe^I][BAR^F₄], [(CAAC)₂Fe⁰], and [(CAAC)₂Fe^{−I}N₂][−] were all found to be active for the reduction of dinitrogen to ammonia upon treatment with KC₈ and HBAR^F₄·2 Et₂O at -95°C [up to (3.4 ± 1.0) equivalents of ammonia per Fe center]. The N₂ reduction activity is highly temperature dependent, with significant N₂ reduction to NH₃ only occurring below -78°C . This reactivity profile tracks with the low temperatures needed for N₂ binding and an otherwise unavailable electron-transfer step to generate reactive [(CAAC)₂FeN₂][−].

While hundreds of transition metal/N₂ complexes have been prepared and studied,^[1] comparatively few systems afford access to productive N₂ functionalization.^[2] This is particularly true for the case of N₂ functionalization by protons and electrons to produce NH₃ (or N₂H₄). Building on extensive early molybdenum and tungsten model work,^[3] Yandulov and Schrock^[4] and Nishibayashi and co-workers^[5] have reported molybdenum-containing coordination complexes (**A** and **B**; Figure 1) which facilitate catalytic N₂ reduction to NH₃ in the presence of suitable acids and reductants. Because iron is 1) the only transition metal known to be essential to enzymatic nitrogenase function,^[6] and 2) the predominant transition-metal catalyst used in the Haber–Bosch ammonia synthesis,^[7] studying N₂ reduction chemistry for well-defined iron model complexes is of interest.^[8,9] Recently, our lab demonstrated that iron coordination complexes (**C**) are capable of modest catalytic N₂ reduction to NH₃.^[10]

The functional Fe–N₂ reduction systems we have studied to date are phosphine-supported, five-coordinate XL₃Fe–N₂ adducts. It is of interest to explore whether other donor ligands and geometries might expose similar reactivity

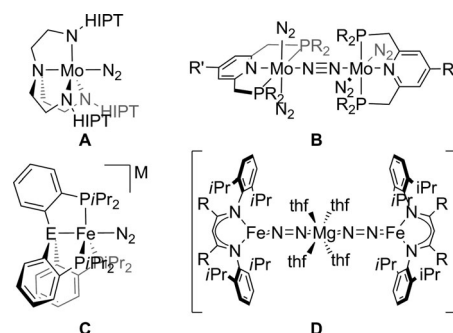


Figure 1. Representative molybdenum and iron complexes studied for catalytic N₂ reduction to NH₃. E = B, C; M = K, Na; R = *t*Bu; R' = H, OMe. HIPT = hexaisopropylterphenyl.

patterns for Fe–N₂ species.^[11,12] For example, Holland and co-workers recently reported a solution characterization of a three-coordinate iron complex with an N₂ ligand linearly bridged between iron and a solvated magnesium cation (**D**; Figure 1).^[13,14] The N₂ ligand in this complex is highly activated. However, only trace amounts of NH₃ (< 2 % per Fe) could be detected upon attempts to mediate its reductive protonation. Related low-coordinate Fe–N₂ species can undergo reductive N₂ cleavage and, when followed by a separate acidic work-up, can achieve higher ammonia yields (ca. 35 % overall).^[8c] Understanding subtle factors that lead to productive reactivity at N₂ is essential to improving molecular nitrogen fixation catalysts.

We recently reported the utilization of a π -accepting cyclic (alkyl)(amino)carbene (CAAC)^[15] to synthesize a structurally unusual two-coordinate, formally [L₂Fe⁰] complex by reduction of its corresponding two-coordinate iron(I) cation.^[16] We reasoned that the polarizability of the CAAC ligand, by comparison to typical N-heterocyclic carbenes (NHCs),^[17] might better facilitate N₂ binding and productive functionalization. Note that low-valent iron/carbene complexes^[18] have, to date, been surprisingly resistant to N₂ binding, with the tripodal NHC-containing systems reported by the groups of Meyer^[19] and Smith^[20] underscoring this point. To our knowledge, a bis(imidazol-2-ylidene)pyridine scaffold is the only iron/carbene derivative where N₂ binding has been established.^[21] Here we show that the two-coordinate [(CAAC)₂Fe] complex binds N₂, but only at low temperature, and that this binding event exposes an otherwise unavailable one-electron reduction step to form three-coordinate [(CAAC)₂Fe(N₂)][−] and desirable reactivity at the coordinated N₂.

Under an atmosphere of N₂, a solution of [(CAAC)₂Fe] in pentane exhibited drastic changes in the absorption spectrum

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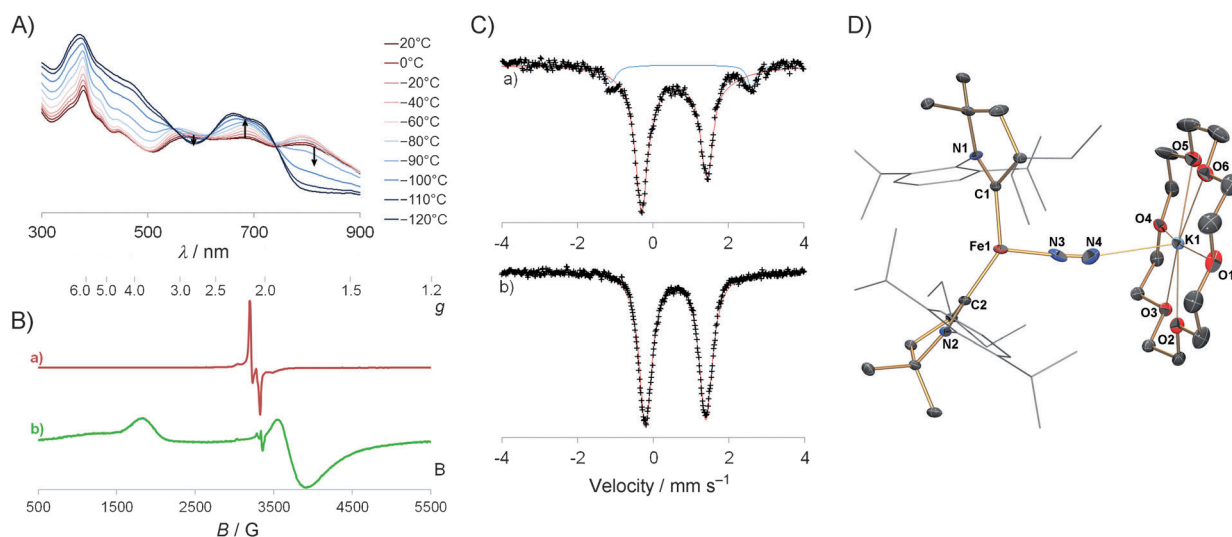
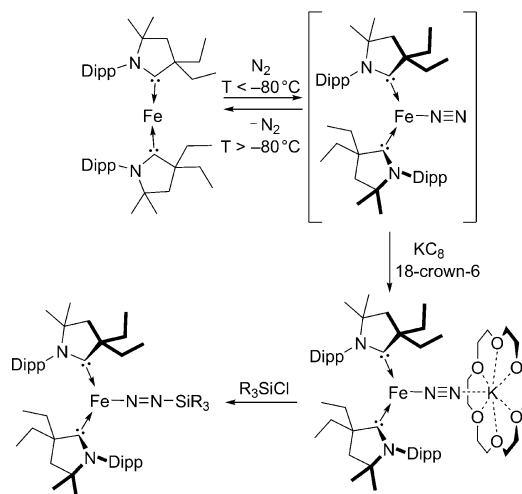


Figure 2. A) Variable-temperature UV/Vis spectra of $[(\text{CAAC})_2\text{Fe}]$ in a pentane solution under N_2 . B) X-Band EPR spectra at 10 K: a) $[(\text{CAAC})_2\text{Fe}(\text{N}_2)][\text{K}(18\text{-crown-6})]$ in a frozen Et_2O glass; b) $[(\text{CAAC})_2\text{Fe}(\text{N}_2\text{SiEt}_3)]$ in a frozen methylcyclohexane glass. C) ^{57}Fe Mössbauer spectra for microcrystalline samples obtained at 80 K in boron nitride pellets. a) $[(\text{CAAC})_2\text{Fe}(\text{N}_2)][\text{K}(18\text{-crown-6})]$; b) $(\text{CAAC})_2\text{Fe}(\text{N}_2\text{SiEt}_3)$. The solid red line corresponds to the simulated data. The solid blue line in (a) corresponds to a minor species (ca. 9% of Fe); parameters of $\delta = 0.74 \text{ mm s}^{-1}$ and $\Delta E_Q = 3.79 \text{ mm s}^{-1}$. D) Structure of $[(\text{CAAC})_2\text{Fe}(\text{N}_2)][\text{K}(18\text{-crown-6})]$ in the solid state. Ellipsoids are drawn at 50% probability. Hydrogen atoms are omitted for clarity.^[23] Selected bond lengths [Å] and angles [°]: C1–Fe1 1.924(2), C2–Fe1 1.919(2), C1–N1 1.386(3), C2–N2 1.411(3), Fe1–N3 1.778(3), N3–N4 1.035(4), N4–K1 2.726(4); C1–Fe1–C2 140.81(10), C1–Fe1–N3 101.91(11), C2–Fe1–N3 117.08(11), Fe1–N3–N4 175.5(3).

upon cooling to temperatures below -80°C (Figure 2 A). This temperature-dependent change in the optical profile of $[(\text{CAAC})_2\text{Fe}]$ under N_2 was reversible over several cooling-warming cycles. To confirm that the change in the optical spectrum was indeed related to the presence of N_2 , the solution was thoroughly degassed by five freeze-pump-thaw cycles and variable-temperature UV/Vis spectra were re-collected (see Figure S7 in the Supporting Information). In this case, the standard spectrum of $[(\text{CAAC})_2\text{Fe}]$ was recovered. Re-exposing the solution to N_2 resulted in the previously observed temperature-dependent profile. Thermodynamic parameters were extracted from a van't Hoff plot of the experimental data (see the Supporting Information). As expected, binding of dinitrogen is exothermic ($\Delta H = -22.1 \text{ kcal mol}^{-1}$) and is associated with negative entropy ($\Delta S = -8.1 \text{ cal K}^{-1} \text{ mol}^{-1}$). These parameters are similar to those recently reported for N_2 coordination at cobalt.^[22] The N_2 binding constant at room temperature is very low ($K_{\text{eq}} = 0.2$) and consistent with our initial report of a two-coordinate $[(\text{CAAC})_2\text{Fe}]$ complex.^[16]

To structurally confirm the binding of dinitrogen at low temperature, we sought to trap the putative complex $[(\text{CAAC})_2\text{Fe}(\text{N}_2)]$ (Scheme 1). The presence of an additional π -accepting ligand (N_2) facilitates the further reduction of $[(\text{CAAC})_2\text{Fe}^0]$ to a formal Fe^{-1} . Hence, reduction of $[(\text{CAAC})_2\text{Fe}]$ with KC_8 at -95°C in diethyl ether in the presence of 18-crown-6 yielded, after work-up, dark brown crystals of $[(\text{CAAC})_2\text{Fe}(\text{N}_2)][\text{K}(18\text{-crown-6})]$ in moderate yield (42%). Temperature control was crucial: reduction at room temperature resulted instead in decomposition of the starting material to intractable products, and at -78°C , only traces of $[(\text{CAAC})_2\text{Fe}(\text{N}_2)]^-$ could be detected by IR spectroscopy. An XRD study of crystals of $[(\text{CAAC})_2\text{Fe}(\text{N}_2)]$



Scheme 1. Reduction of $[(\text{CAAC})_2\text{Fe}]$ and subsequent trapping with silylating reagents. R = Me, Et. Dipp = 2,6-diisopropylphenyl.

$[\text{K}(18\text{-crown-6})]$ confirmed the binding of N_2 to a three-coordinate iron center in an end-on fashion with an additional ion-pair of the β -nitrogen atom to the $\text{K}(18\text{-crown-6})$ cation (Figure 2 D).

The structure of $[(\text{CAAC})_2\text{Fe}(\text{N}_2)][\text{K}(18\text{-crown-6})]$ reveals a distorted trigonal-planar iron center ($\Sigma_{\text{angle}} = 359.8^\circ$), with a very wide C–Fe–C angle [$140.81(10)^\circ$]. The C–N distances of the CAAC ligand are substantially elongated: they are 1.386(3) and 1.411(3) Å, compared with 1.315(3) Å for the free ligand.^[15c] This C–N lengthening is indicative of a substantial amount of spin leakage from the highly reduced iron center to the supporting CAAC ligand.^[16] Complex $[(\text{CAAC})_2\text{Fe}(\text{N}_2)][\text{K}(18\text{-crown-6})]$ represents a rare

example of a structurally characterized pseudoterminaly bonded, three-coordinate Fe–N₂ complex.^[13,14]

A sharp band was observed in the IR spectrum of [(CAAC)₂Fe(N₂)] [K(18-crown-6)] at 1850 cm^{−1} (ATR-IR; thin film), and attributed to the ν_{NN} stretch. This value is higher in energy than the transiently observed [β-diketimidatoFe(N₂)]₂Mg(thf)₄ species reported by Holland and co-workers (1818 cm^{−1}),^[13] but lower than other anionic terminal Fe–N₂ complexes where the iron center resides in higher coordination numbers (1905–1927 cm^{−1}).^[9,10a] As expected for a formal Fe^{−I} species, a room temperature solution magnetic moment for [(CAAC)₂Fe(N₂)] [K(18-crown-6)] of 1.9 μ_B in C₆D₆ was measured, and is consistent with its frozen glass (Et₂O) EPR spectrum (Figure 2B,a) and indicative of an *S* = 1/2 ground state.

Exposure of [(CAAC)₂Fe(N₂)] [K(18-crown-6)] to trimethylsilylchloride in diethyl ether resulted in a rapid color change from dark brown-red to dark green. A diagnostic broad band at 1675 cm^{−1} was observed in the IR spectrum of the crude reaction mixture, and is reminiscent of previously observed ν_{NN} stretches for iron/silyldiazenido complexes.^[9,10a] The presumed diazenido product [(CAAC)₂Fe(N₂SiMe₃)] rapidly decomposed to the [(CAAC)₂Fe] complex upon attempted work-up. A bulkier silylating reagent led to a more stable product. Hence, a similar change of color from dark brown-red to dark green was observed when triethylsilylchloride was added to the [(CAAC)₂Fe(N₂)] [K(18-crown-6)] complex (Scheme 1). The diazenido product, [(CAAC)₂Fe(N₂SiEt₃)] was isolated as an analytically pure, dark-green solid in moderate yield (52 %). Its IR spectrum exhibits a broad band at 1690 cm^{−1}. The complex [(CAAC)₂Fe(N₂SiEt₃)] is high-spin with *S* = 3/2 (room temperature solution magnetic moment of 3.9 μ_B), and consistent with its frozen glass EPR spectrum at 10 K showing broad signals at low field (Figure 2B,b). Its electronic structure is therefore distinct from previously characterized five-coordinate [XL₃Fe(N₂SiR₃)] species which are low spin. For instance, (TP^{Pr}B)Fe(N₂SiMe₃) is an *S* = 1/2 species.^[9b]

The ⁵⁷Fe Mössbauer spectra of [(CAAC)₂Fe(N₂)] [K(18-crown-6)] and [(CAAC)₂Fe(N₂SiEt₃)] were recorded at 80 K in the solid state (Figure 2C) and fits to the data provided isomer shifts at δ = 0.56 and δ = 0.59 mm s^{−1}, respectively, with corresponding quadrupole splittings of ΔE_Q = 1.67 and ΔE_Q = 1.60 mm s^{−1}. The isomer-shift value is in line with previously reported three-coordinate bis(carbene) iron complexes.^[16,24] For example, [(CAAC)₂FeCl] has an isomer shift of 0.52 mm s^{−1}. The spectrum of [(CAAC)₂Fe(N₂)] [K(18-crown-6)] displays an asymmetrical doublet presumably caused by anisotropy in the polycrystalline sample.^[25] A minor component (ca. 9 % of the total iron content), whose parameters fit the oxidation product [(CAAC)₂Fe]⁺,^[16] was also detected (Figure 2C,a).

In previous studies of five-coordinate Fe–N₂ species, we have empirically determined that the ability to successfully silylate the β-nitrogen atom of an Fe–N₂ derivative can translate to efficacy of the system towards reductive protonation.^[9c,10] Attempts to fix N₂ to NH₃ at room temperature using [(CAAC)₂Fe] in the presence of excess KC₈ and HBAR^F₄·2Et₂O in diethyl ether, regardless of the order of

addition, proved largely ineffective (Table 1; see Figure S12). This observation is consistent with a very limited degree of N₂ binding to [(CAAC)₂Fe] at room temperature, and the requirement that a [(CAAC)₂Fe(N₂)] species be present to facilitate N₂ fixation. Indeed, when [(CAAC)₂Fe] was treated at −95 °C with an excess of KC₈ (50 equiv), followed by an excess of HBAR^F₄·2Et₂O (50 equiv), the formation of (3.3 ± 1.1) equivalents of NH₃ per iron center was observed. Similar yields [(3.4 ± 1.0) equiv] were obtained by employing the two-coordinate iron(I) cation [(CAAC)₂Fe][BAR^F₄] under the

Table 1: Catalytic reduction of N₂ to NH₃.^[a]

$\text{N}_2 + \text{KC}_8 + \text{HBAr}^{\text{F}}_{24} \cdot 2 \text{Et}_2\text{O} \xrightarrow[\text{Et}_2\text{O}]{\text{cat.}} \text{NH}_3$			
Run	Catalyst	<i>T</i> [°C]	Equiv NH ₃ per Fe
1	[(CAAC) ₂ Fe]	−113	3.0 ± 0.7
2	[(CAAC) ₂ Fe]	−95	3.3 ± 1.1
3	[(CAAC) ₂ Fe]	−78	0.9 ± 0.3 ^[b]
4	[(CAAC) ₂ Fe]	−50	0.3 ± 0.2 ^[b]
5	[(CAAC) ₂ Fe]	23	0.4 ± 0.2 ^[b]
6	[(CAAC) ₂ Fe][BAR ^F ₄]	−95	3.4 ± 1.0
7	[(CAAC) ₂ FeN ₂][K(18-c-6)]	−95	2.6 ± 0.6
8	CAAC ^[c]	−95	< 0.1
9	none	−95	< 0.1

[a] Catalytic conditions: catalyst (0.002 mmol), KC₈ (0.1 mmol), HBAR^F₄·2Et₂O (0.1 mmol), Et₂O, 45 min. Yields are an average of 8 independent runs and were determined by the indophenol method (See the Supporting Information). [b] Average of 4 runs. [c] 0.004 mmol of CAAC used.

same conditions at −95 °C. The enhancement of the N₂-to-NH₃ reduction yield at very low temperature correlates with the variable-temperature optical data for [(CAAC)₂Fe], which shows that N₂ binding to [(CAAC)₂Fe] becomes favorable only at low temperature. Indeed, the yield of NH₃ is negligible at −50 °C (0.3 equiv) and is still quite low at −78 °C (0.9 equiv). Note that significantly lower temperature (−113 °C) was examined but did not improve the NH₃ yield beyond that obtained at −95 °C.

Although these NH₃ yields are low compared with our two previously reported iron catalysts,^[10] Schrock's original molybdenum system,^[4] and especially Nishibayashi's recently improved molybdenum system,^[5a] [(CAAC)₂Fe] is nevertheless a very modest catalyst for nitrogen fixation to ammonia and the yields of NH₃ per iron equivalent reported here are far greater than those which have been observed for previously studied three-coordinate Fe–N₂ complexes (< 0.2 equiv per Fe), including [β-diketimidatoFe(N₂)]₂Mg(thf)₄ species (Figure 1).^[13] We speculate that the ability of [(CAAC)₂Fe]/[(CAAC)₂Fe(N₂)] to perform nitrogen fixation may arise from the relative flexibility of the system, which is capable of switching between two- and three-coordinate geometries, and allows the formation of highly covalent Fe–N_x multiple-bond interactions. The recent isolation of a three-coordinate iron/bis(imide) complex supported by CAAC by Deng and co-workers is noted in the latter context.^[26]

For comparison, we also explored [(CAAC)₂Fe] in the context of catalytic N₂ silylation. Catalytic silylation reactions

of N_2 under strongly reducing conditions have been known for many years using metals such as chromium,^[27] titanium,^[28] and molybdenum,^[29] and recently, such silylations were extended to systems using iron precatalysts including $[Fe(CO)_5]$ and $[Cp_2Fe]$.^[30] For example, $[Fe(CO)_5]$ was shown to produce up to 25 equivalents of $N(SiMe_3)_3$ per iron in the presence of a vast excess sodium metal and $TMSCl$ at room temperature.^[30] These silylation reactions remain mechanistically ill-defined. Only trace yield of $N(SiMe_3)_3$ could be detected using Nishibayashi's conditions (Na as reductant, in THF)^[30] with $[(CAAC)_2Fe]$ as precatalyst. However, when a catalytic amount of $[(CAAC)_2Fe]$ was instead treated with a large excess of KC_8 (600 equiv) and trimethylsilylchloride (600 equiv), the formation of (24.4 ± 2.7) equivalents of $N(SiMe_3)_3$ was observed (see Table S1). Slightly lower TONs (19.4 ± 3.0) were obtained using $[(CAAC)_2Fe][BAR^F_4]$ as a precatalyst instead. The catalytic activity was significantly reduced (7.0 ± 1.0) when the reaction was performed at $-78^\circ C$. The attenuation in $N(SiMe_3)_3$ product is presumably related to slower generation of trimethylsilyl radical at low temperature.

Although it is presently unclear what the mechanistic requirements are for the N_2 to $N(SiMe_3)_3$ catalysis mediated by various iron precursors, including $[(CAAC)_2Fe]$, the N_2 to NH_3 chemistry described herein appears to require a well-defined molecular $Fe-N_2$ species. Indeed, significant N_2 to NH_3 conversion is observed only at the low temperatures where N_2 binds $[(CAAC)_2Fe]$ sufficiently favorably. The low-temperature N_2 binding event facilitates subsequent electron transfer as evidenced by the very low temperature required to synthetically reduce $[(CAAC)_2Fe(N_2)]$ and isolate highly reactive $[(CAAC)_2Fe(N_2)]^-$.

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- [1] a) B. A. MacKay, M. D. Fryzuk, *Chem. Rev.* **2004**, *104*, 385; b) M. D. Fryzuk, S. A. Johnson, *Coord. Chem. Rev.* **2000**, *200*, 379.
- [2] a) J. L. Crossland, D. R. Tyler, *Coord. Chem. Rev.* **2010**, *254*, 1883; b) M. Hidai, Y. Mizobe, *Chem. Rev.* **1995**, *95*, 1115; c) J. Chatt, J. R. Dilworth, R. L. Richards, *Chem. Rev.* **1978**, *78*, 589.
- [3] a) M. Hidai, Y. Mizobe, *Can. J. Chem.* **2005**, *83*, 358; b) A. E. Shilov, *Russ. Chem. Bull.* **2003**, *52*, 2555; c) T. A. Bazhenova, A. E. Shilov, *Coord. Chem. Rev.* **1995**, *144*, 69; d) C. J. Pickett, J. Talarmin, *Nature* **1985**, *317*, 652; e) J. Chatt, A. J. Pearman, R. L. Richards, *Nature* **1975**, *253*, 39; f) J. Chatt, G. A. Heath, R. L. Richards, *J. Chem. Soc. Dalton Trans.* **1974**, 2074.
- [4] D. V. Yandulov, R. R. Schrock, *Science* **2003**, *301*, 76.
- [5] a) S. Kuriyama, K. Arashiba, K. Nakajima, H. Tanaka, N. Kamaru, K. Yoshizawa, Y. Nishibayashi, *J. Am. Chem. Soc.* **2014**, *136*, 9719; b) K. Arashiba, Y. Miyake, Y. Nishibayashi, *Nat. Chem.* **2011**, *3*, 120.
- [6] J. Yang, X. Xie, X. Wang, R. Dixon, Y.-P. Wang, *Proc. Natl. Acad. Sci. USA* **2014**, *111*, E3718.
- [7] H. Liu, *Ammonia Synthesis Catalysts: Innovation and Practice*, World Scientific Publishing Co. Pte. Ltd, Singapore, Chemical Industry Press, Beijing, **2013**.
- [8] a) G. J. Leigh, M. Jimenez-Tenorio, *J. Am. Chem. Soc.* **1991**, *113*, 5862; b) J. D. Gilbertson, N. K. Szymczak, D. R. Tyler, *J. Am. Chem. Soc.* **2005**, *127*, 10184; c) M. M. Rodriguez, E. Bill, W. W. Brennessel, P. L. Holland, *Science* **2011**, *334*, 780.
- [9] a) J. Rittle, J. C. Peters, *Proc. Natl. Acad. Sci. USA* **2013**, *110*, 15898; b) M.-E. Moret, J. C. Peters, *J. Am. Chem. Soc.* **2011**, *133*, 18118; c) M.-E. Moret, J. C. Peters, *Angew. Chem. Int. Ed.* **2011**, *50*, 2063; *Angew. Chem.* **2011**, *123*, 2111; d) Y. Lee, N. P. Mankad, J. C. Peters, *Nat. Chem.* **2010**, *2*, 558.
- [10] a) S. E. Creutz, J. C. Peters, *J. Am. Chem. Soc.* **2014**, *136*, 1105; b) J. S. Anderson, J. Rittle, J. C. Peters, *Nature* **2013**, *501*, 84.
- [11] a) T. A. Betley, J. C. Peters, *J. Am. Chem. Soc.* **2004**, *126*, 6252; b) T. A. Betley, J. C. Peters, *J. Am. Chem. Soc.* **2003**, *125*, 10782.
- [12] K. C. MacLeod, P. L. Holland, *Nat. Chem.* **2013**, *5*, 559.
- [13] T. R. Dugan, K. C. MacLeod, W. W. Brennessel, P. L. Holland, *Eur. J. Inorg. Chem.* **2013**, 3891.
- [14] a) J. M. Smith, A. R. Sadique, T. R. Cundari, K. R. Rodgers, G. Lukat-Rodgers, R. J. Lachicotte, C. J. Flaschenriem, J. Vela, P. L. Holland, *J. Am. Chem. Soc.* **2006**, *128*, 756; b) J. M. Smith, R. J. Lachicotte, K. A. Pittard, T. R. Cundari, G. Lukat-Rodgers, K. R. Rodgers, P. L. Holland, *J. Am. Chem. Soc.* **2001**, *123*, 9222.
- [15] V. Lavallo, Y. Canac, C. Prasang, B. Donnadieu, G. Bertrand, *Angew. Chem. Int. Ed.* **2005**, *44*, 5705; *Angew. Chem.* **2005**, *117*, 5851.
- [16] G. Ung, J. Rittle, M. Soleilhavoup, G. Bertrand, J. C. Peters, *Angew. Chem. Int. Ed.* **2014**, *53*, 8427; *Angew. Chem.* **2014**, *126*, 8567.
- [17] O. Back, M. Henry-Ellinger, C. D. Martin, D. Martin, G. Bertrand, *Angew. Chem. Int. Ed.* **2013**, *52*, 2939; *Angew. Chem.* **2013**, *125*, 3011.
- [18] For reviews on low-valent iron/carbene complexes, see: a) K. Riene, S. Haslinger, A. Raba, M. P. Högerl, M. Cokoja, W. A. Herrmann, F. E. Kühn, *Chem. Rev.* **2014**, *114*, 5215; b) D. Bézier, J.-B. Sortais, C. Darcel, *Adv. Synth. Catal.* **2013**, *355*, 19; c) M. J. Ingleson, R. A. Layfield, *Chem. Commun.* **2012**, 48, 3579.
- [19] C. S. Vogel, F. W. Heinemann, M. M. Khusniyarov, K. Meyer, *Inorg. Chim. Acta* **2010**, *364*, 226.
- [20] I. Nieto, F. Cervantes-Lee, J. M. Smith, *Chem. Commun.* **2005**, 3811.
- [21] A. A. Danopoulos, J. A. Wright, W. B. Motherwell, *Chem. Commun.* **2005**, 784.
- [22] D. L. M. Suess, C. Tsay, J. C. Peters, *J. Am. Chem. Soc.* **2012**, *134*, 14158.
- [23] CCDC 1025565 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge from The Cambridge Crystallographic Data Centre via www.ccdc.cam.ac.uk/data_request/cif.
- [24] Z. Mo, Z. Ouyang, L. Wang, K. L. Fillman, M. L. Neidig, L. Deng, *Org. Chem. Front.* **2014**, *1*, 1040.
- [25] a) P. Gütllich, R. Link, A. Trautwein, *Mössbauer Spectroscopy and Transition Metal Chemistry*, Springer, Berlin, **1978**; b) V. I. Goldanskii, R. Herber, *Chemical Applications of Mössbauer Spectroscopy*, Academic Press, New York, **1964**.
- [26] H. Zhang, Z. Ouyang, Y. Liu, Q. Zhang, L. Wang, L. Deng, *Angew. Chem. Int. Ed.* **2014**, *53*, 8432; *Angew. Chem.* **2014**, *126*, 8572.
- [27] K. Shiina, *J. Am. Chem. Soc.* **1972**, *94*, 9266.
- [28] M. Mori, *J. Organomet. Chem.* **2004**, *689*, 4210.
- [29] a) Y. Nishibayashi, *Dalton Trans.* **2012**, *41*, 7447; b) H. Tanaka, A. Sasada, T. Kouno, M. Yuki, Y. Miyake, H. Nakanishi, Y. Nishibayashi, K. Yoshizawa, *J. Am. Chem. Soc.* **2011**, *133*, 3498; c) K. Komori, H. Oshita, Y. Mizobe, M. Hidai, *J. Am. Chem. Soc.* **1989**, *111*, 1939.
- [30] M. Yuki, H. Tanaka, K. Sasaki, Y. Miyake, K. Yoshizawa, Y. Nishibayashi, *Nat. Commun.* **2012**, *3*, 1254.