

Mechanistic Insight into N=N Cleavage by a Low-Coordinate Iron(II) Hydride Complex

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Abstract: The reaction pathways of high-spin iron hydride complexes are relevant to the mechanism of N_2 reduction by nitrogenase, which has been postulated to involve paramagnetic iron-hydride species. However, almost all known iron hydrides are low-spin, diamagnetic Fe(II) compounds. We have demonstrated that the first high-spin iron hydride complex, L^{tBu}FeH (L^{tBu} = bulky β -diketiminate), reacts with PhN=NPh to completely cleave the N-N double bond, giving L^{tBu}FeNHPh. Here, we disclose a series of experiments that elucidate the mechanism of this reaction. Crossover and kinetic experiments rule out common nonradical mechanisms, and support a radical chain mechanism mediated by iron(I) species including a rare η^2 azobenzene complex. Therefore, this high-spin iron(II) hydride can break N-N bonds through both nonradical and radical insertion mechanisms, a special feature that enables novel reactivity.

Introduction

Transition metal hydride complexes play a role in numerous bond transformations.¹ One of their characteristic reactions is transfer of "H-" to substrates like alkenes and ketones, which results in an overall two-electron reduction of that substrate. Because nitrogenase substrates (e.g., N₂, alkynes, CO₂, CN⁻, N₂O) are all reduced by multiples of two electrons, metal hydrides have been proposed as intermediates in the catalytic reduction of N₂ to NH₃ by nitrogenase enzymes.^{2,3} Iron is the only transition metal that is common to all nitrogenases,⁴ suggesting that an Fe-H intermediate could be an active species during nitrogenase catalysis. Consistent with the presence of a hydride intermediate, nitrogenase produces H2 from water at low substrate concentrations.5 Recent ENDOR and EPR evidence supports the presence of a paramagnetic iron-hydride species upon reduction of the α -70^{Ile} mutant of A. vinelandii nitrogenase.⁶ Although this mutant is incapable of reducing N₂, its close similarity to the wild-type nitrogenase indicates that Fe-H species should be considered as prospective intermediates during the nitrogenase catalytic cycle.

Our interest in synthetic analogues for nitrogenase⁷ motivated us to synthesize iron hydride complexes with coordination

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Figure 1. Active site "FeMoco" of nitrogenase. The "belt" region refers to the central Fe₆S₃X core, where X is an unidentified anion (probably C⁴⁻, N³⁻, or O²⁻).9

environments similar to those potentially present in nitrogenase. An ideal "model" complex would act as a functional model, reducing substrates like those reduced by nitrogenase (N₂, alkynes, NO, CO₂, HCN, diazenes, hydrazine) in a manner that is more amenable to detailed mechanistic inquiry than the enzyme.8 A hydride complex with perfect structural analogy to the "belt iron" sites implicated as active sites of the FeMoco (Figure 1)^{3,9,10} would have three weak-field sulfur donors, a high-spin electronic configuration, and a low coordination number (3, 4, 5) at iron. No complex with all of these properties is known. However, by using the bulky anionic β -diketiminate ligand L^{tBu} ($L^{R} = 1,3-R_{2}-1,3$ -bis(2,6-diisopropylphenylimido)propyl, Figure 2), we were able to synthesize the first hydride complex of iron with a coordination number less than five, $[L^{tBu}Fe(\mu-H)]_2$.^{11,12} In solution, it is in equilibrium with a three-

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Figure 2. Bulky β -diketiminate ligands used in this work.

coordinate monomer, L^{tBu} FeH, and it binds pyridine to give L^{tBu} Fe(H)(pyridine), which has a trigonal pyramidal geometry. The monomeric hydride complexes have the desired high-spin (S = 2) electronic configuration at the metal, by virtue of the low coordination number and weak ligand field at iron. Although they lack the characteristic three-sulfur coordination environment of the iron atoms in the FeMoco, diketiminate-supported iron hydrides show promise for giving hints into the relevant reactivity patterns of hydride ligands in a weak ligand field.¹³

The main function of nitrogenase is the cleavage of N–N bonds, but *in vitro* it reduces numerous substrates. For example, *A. vinelandii* iron–molybdenum nitrogenase has been reported to reduce methyldiazene and hydrazine.¹⁴ N–N single bond cleavage is promoted by nitrogenase and model complexes,^{14–16} though rarely with iron.¹⁷ In a preliminary communication, we reported that $[L^{tBu}FeH]_2$ reacts with azobenzene (PhN=NPh) to afford the amido complex $L^{tBu}FeNHPh$ (boxed reaction in Scheme 1).¹¹ This reaction was the first example of complete N=N cleavage by an iron-hydride complex, and one of the few by any iron complex.^{18–21} An unusual facet of the transformation in Scheme 1 is that the complete N=N bond cleavage of azobenzene requires no overall change in oxidation state at the iron. This contrasts with literature N=N cleavage reactions that result in four-electron oxidation of a metal (or multiple metals).²²

Given the importance of breaking nitrogen-nitrogen bonds in the formation of ammonia from N_2 by nitrogenase, the ability

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Scheme 1



of a low-coordinate iron hydride to completely break an N=N bond is remarkable. This manuscript reports investigations aimed to determine the mechanism of this unusual transformation.

Results and Discussion

Synthesis of High-Spin Iron(II) Hydride Complexes. We have reported the synthesis of dimeric $[L^{tBu}FeH]_2$, and showed that it is in equilibrium with its monomeric, trigonal-planar form $L^{tBu}FeH$.¹¹ In order to provide an alternative ligand with comparable electronic properties and more steric bulk, we have prepared the new ligand abbreviated $L^{tBu'}$ (Figure 2), which differs from L^{tBu} by the presence of an additional isopropyl group at the *para* position of each aryl ring. Creation of $L^{tBu'}$ necessitated the synthesis of 2,4,6-triisopropylaniline, which came from reduction of 2,4,6-triisopropylnitrobenzene. Incorporation of this aniline into the β -diketiminate ligand followed procedures analogous to those used in the synthesis of $L^{tBu'}$ (see the Experimental Section for details).

Hydride complexes were accessed by addition of KHBEt₃ to the chloride complex L^{tBu}FeCl or L^{tBu}′FeCl at room temperature in toluene. An immediate color change from red to brown was

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Figure 3. Solid-state structure of $[L^{1Bu'}FeH]_2$, with 50% probability thermal ellipsoids. Hydrogen atoms on the diketiminate ligands are omitted for clarity. Fe1–N11 1.988(1) Å, Fe1–N21 1.978(1) Å, Fe2–N14 2.043(1) Å, Fe2–N24 2.010(1) Å, Fe1–Fe2 2.5292(3) Å, N11–Fe1–N21 96.41-(5)°, N14–Fe2–N24 97.60(5)°.

observed, with dissolution of the starting material ($L^{tBu'}$ complexes tend to be somewhat more soluble than their L^{tBu} analogues). In each case, it is important to remove the byproduct BEt₃ immediately, because the borane reacts with the hydride complex. The details of the reactions between iron hydride complexes and boranes will be reported separately.²³

The new hydride complex [LtBu'FeH]2 was characterized by solution methods and by X-ray crystallography. The solid-state structure of [L^{tBu}/FeH]₂ (Figure 3) shows a dimer similar to [L^{tBu}-FeH]₂.¹¹ Each iron atom has a distorted tetrahedral geometry around the metal center, from binding of the diketiminate ligand in a η^2 binding mode, and coordination of two bridging hydrides. These hydrides were located in a Fourier map, and their positional parameters were refined, but their positions must be regarded with uncertainty, given the small electron density at the hydrogen atom. The β -diketiminate ligands are twisted into a distorted boat conformation, with a substantial folding along the N····N axis (21.79(9)° and 27.39(7)° in [L^{tBu'}FeH]₂; 24.3- $(3)^{\circ}$ in [L^{tBu}FeH]₂). The Fe-Fe distance of [L^{tBu}FeH]₂ is slightly shorter (2.5292(3) Å) than that of $[L^{tBu}FeH]_2$ (2.624(2) Å) despite the increased steric hindrance in the former. The nature of the iron-iron interaction is the subject of ongoing studies.

The ¹H NMR spectrum of each hydride dimer is unusually complicated, with at least 17 relatively sharp but partially overlapped peaks present over the range 80 to -130 ppm. The complicated room-temperature ¹H NMR spectrum is attributed to hindered C–C and C–N bond rotations in the dimer, because the X-ray crystal structures show that the complexes are exceptionally crowded. Despite the complex ¹H NMR spectra, these materials are analytically pure, react with 3-hexyne to give high yields of vinyl products,¹¹ and convert to monomeric forms with simple ¹H NMR spectra at elevated temperatures. The ratio

of monomer to dimer in a 17 mM solution in C_6D_6 at room temperature is slightly larger for the $L^{tBu'}$ compound (ca. 0.5) than the L^{tBu} compound (ca. 0.3). Therefore, a majority of the hydride complex is present as dimer at room temperature. However, our previous studies have shown that monomer is formed rapidly (the reported activation parameters for addition of alkyne to monomer extrapolate to a half-life of several minutes at room temperature and a few seconds at 80 °C),¹¹ so in the following discussion the dimeric hydride complexes are treated as being functionally equivalent to the monomers.

Reaction with Azobenzene. Reaction of [LtBuFeH]2 or [LtBu'-FeH]₂ with 2 equiv of PhN=NPh in diethyl ether gave the isolable hydrazido complexes LtBuFeNPhNHPh and LtBu'FeN-PhNHPh, respectively. The reactions are complete in less than 30 min at room temperature. Although the swiftness of this reaction has prevented kinetic studies, it is possible to propose a mechanism based on analogous reactions we have reported previously. We envision the first step of the reaction as dissociation of the hydride dimer into monomeric LtBuFeH or L^{tBu}FeH, which is rapid at room temperature.¹¹ This is followed by 1,2-insertion of the Fe-H bond across the N=N bond.24 We have observed $[L^{R}FeH]_{2}$ to add across C=C, C=C, C=N, and C=O bonds, and we have used kinetic isotope effects, kinetics, and computations to support a 1,2-insertion mechanism.²⁵ Therefore, it is reasonable to presume that reduction of the N=N bond of diazene to the N-N bond of the hydrazido ligand follows an analogous pathway. Note that this reaction requires 1 equiv of azobenzene per iron, twice the amount needed for the overall conversion of [L^{tBu}FeH]₂ and PhN=NPh to LtBuFeNHPh.

The spectroscopic characteristics of the hydrazido complexes are unremarkable, and the characterization of L^{tBu} FeNPhNHPh has been presented previously.¹¹ As an alternative to the preparation of the hydrazido complexes from azobenzene, they can also be accessed by addition of two equiv of PhNHNHPh to [L^{tBu} FeH]₂, releasing H₂. Samples of L^{tBu} FeNPhNHPh from the two preparatory methods gave similar rate constants in subsequent reactions (see below), but the rate constants were more reproducible when generated from the latter method.

Kinetic and Crossover Studies on the N–N Cleavage Reaction. Benzene and toluene solutions of purified L^{tBu}-FeNPhNHPh (36–48 mM) decompose at 80 °C to form 1 equiv of L^{tBu}FeNHPh and 0.5 equiv of PhN=NPh (Scheme 1).^{11,26} We followed the progress of this reaction using ¹H NMR spectroscopy in C₆D₆ and toluene- d_8 . The reaction was monitored at a number of temperatures, and rate constants were derived from exponential fits to the integrations of peaks at 32 and -72 ppm for L^{tBu}FeNPhNHPh and 28 ppm for L^{tBu}FeNHPh. Integrations were calibrated to an internal capillary containing Tp*₂Co in toluene- d_8 . No intermediates or other products were observed under these conditions. At most temperatures (see below), [L^{tBu}FeNPhNHPh] followed an exponential decay with a rate constant near that calculated for growth of [L^{tBu}FeNHPh].

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⁽²⁶⁾ When [L^{IBu}FeH]₂ is reacted with 1 equiv of PhN=NPh (i.e., 0.5 equiv of PhN=NPh per iron atom), then all of the PhN=NPh is consumed. In this case, the PhN=NPh formed in the decomposition of L^{IBu}FeNPhNHPh immediately reacts with the excess iron-hydride complex to form more L^{IBu}FeNPhNHPh, eventually giving only L^{IBu}FeNPhPhNHPh as a product.





Figure 4. (a) Top: Plot of $[L^{1Bu}FeNPhNHPh]$ vs time at 358 K. (b) Bottom: Plot of $ln[L^{1Bu}FeNPhNHPh]$ vs time at 358 K.

The exponential fit implies a first-order rate dependence on [L^{tBu}-FeNPhNHPh], and plots of ln[L^{tBu}FeNPhNHPh] versus time were linear (Figure 4).²⁷

Interestingly, the plots of $[L^{tBu}FeNPhNHPh]$ and $[L^{tBu}FeNHPh]$ versus time at the lowest temperature used (42 °C) show an induction period before the beginning of the exponential decay or growth (Figure 5). This behavior, evident in three different trials, suggests an initiation step in the reaction pathway. The length of the induction period varied from 45 to 90 min in different trials at this temperature. The implications of this observation will be described in more detail below.

The reaction path to the products was studied using a crossover experiment. A mixture of two different hydrazido compounds, L^{tBu} FeNPhNHPh and L^{tBu} FeNTolNHTol (Tol = *m*-tolyl), was heated to 80 °C for 2 h and passed through an activated alumina column to remove iron salts. Analysis of the



Figure 5. Kinetic data for the conversion of L^{tBu} FeNPhNHPh to L^{tBu} FeNHPh at 315 K: (\bullet) [L^{tBu} FeNPhNHPh]; (\times) [L^{tBu} FeNHPh].

organic byproducts by mass spectrometry shows the formation of PhN=NPh (m/z = 182) and TolN=NTol (m/z = 210), but no mixed diazene PhN=NTol (m/z = 196). This result implies that diazene is formed before N-N bond cleavage.

Finally, we used a double crossover experiment with two different diketiminate ligands and two different hydrazido ligands. A mixture of L^{tBu}FeNPhNHPh and L^{tBu}'FeNTolNHTol was heated to 80 °C for 2 h and the products were analyzed by mass spectrometry. L^{tBu}FeNHPh (m/z = 649), L^{tBu}FeNHTol (m/z = 663), L^{tBu}'FeNHPh (m/z = 733), and L^{tBu}'FeNHTol (m/z = 747) were observed in roughly equal amounts. Because the amido compounds had statistically scrambled between (diketiminate)Fe units, we conclude that *all Fe–N bonds are broken at some point in the mechanism*.

We also attempted to study the kinetic isotope effect for the reaction. For this purpose, deuterium labeled hydrazido compound, L^{tBu}FeNPhNDPh, was synthesized from addition of PhNDNDPh to [L^{tBu}FeH]₂. This compound was heated under similar conditions as used with L^{tBu}FeNPhNHPh. Unfortunately, the IR spectrum of the product shows the presence of L^{tBu}-FeNHPh ($\nu_{N-H} = 3319 \text{ cm}^{-1}$) as well as L^{tBu}FeNDPh ($\nu_{N-D} = 2499 \text{ cm}^{-1}$), suggesting that some of the deuterium label was lost through an unknown mechanism. While the fate of these deuterons remains a mystery, the complication of the proton exchange makes it impossible to interpret the relative rates as a genuine kinetic isotope effect for the reaction.²⁸

Exploring the Possible Mechanisms. 1. Disproportionation Mechanism. One possible mechanism for the formation of amido product is shown in Scheme 2. A pericyclic mechanism

Scheme 2



of this type was proposed for the disproportionation of N,N'diphenylhydrazine to azobenzene and aniline at high temperature.²⁹ However, later kinetic studies showed a first-order

⁽²⁷⁾ A linear Eyring plot (Figure S-1) derived from the rate constants at different temperatures gives *apparent* activation parameters of $\Delta H^{i} = 13 \pm 1$ kcal mol⁻¹ and $\Delta S^{i} = -36 \pm 4$ eu. However, the induction period suggests a chain mechanism (discussed at length below), in which the rate constants do not reflect elementary steps. Therefore, these *apparent* activation parameters should be interpreted only with the greatest caution.

dependence on [PhNHNHPh], ruling out this mechanism.³⁰ According to this mechanism, the products are formed through a bimolecular transition state as shown in Scheme 2. This requires the rate of our reaction to have a second-order dependence on [LtBuFeNPhNHPh]. The observed first-order dependence on iron concentration and the induction period are inconsistent with the disproportionation mechanism.

2. β-Hydride Elimination Mechanism. Next, we considered β -hydride elimination (Scheme 3) as a potential rate-limiting step in the mechanism. This pathway has some attractive features: it would be expected to give a first-order dependence on [Fe]; the azobenzene is generated without N-N bond cleavage; and the β -hydride elimination step is the reverse of the 1,2-insertion of the Fe-H bond across the N=N bond, which forms the hydrazido complex. If the reaction goes by this mechanism, LtBuFeH should be formed as an intermediate of the reaction. However, no intermediate species were detected by ¹H NMR spectroscopy during the course of the reaction, and the rate of the reaction was not affected by the addition of excess azobenzene. To account for these observations, the β -hydride elimination would have to be the rate-limiting step: in other words, the intermediate LtBuFeH has to react quickly with LtBuFeNPhNHPh.

Scheme 3



To query the intermediacy of the hydride species, the hydrazido complex was heated in the presence of 3-hexyne in varying amounts. The reaction of LtBuFeH with 3-hexyne to afford the vinyl compound, LtBuFeCEtCHEt, is very fast and irreversible at room temperature,^{11,31} so it is expected to be an effective trap. However, only a small amount (<20%) of L^{tBu}-FeCEtCHEt was formed, and the ratio of vinyl product to the amido product was independent of the concentration of 3-hexyne (see Supporting Information for details). This result is inconsistent with the intermediacy of LtBuFeH in the reaction.

It is instructive to further consider the possibility that the hydride and hydrazido react with each other so quickly that trapping is not possible. This possibility was experimentally tested by adding 1 equiv of [LtBuFeH]2 to LtBuFeNPhNHPh in C_6D_6 . This mixture does not react under ambient conditions, even though [LtBuFeH]2 is expected to form monomer within a few minutes at room temperature.¹¹ When this mixture was heated to 358 K, LtBuFeNHPh is formed with a rate constant of $(5.3 \pm 0.7) \times 10^{-4}$ s⁻¹. Because the rate constant for this reaction is roughly the same as that in the reaction without added hydride complex $[(7.1 \pm 1.3) \times 10^{-4} \text{ s}^{-1}]$ it is unlikely that the hydride is an intermediate in the transformation.

One final piece of evidence argues against the mechanism in Scheme 3: if L^{tBu}FeH reacted rapidly with L^{tBu}FeNPhNHPh, as required in Scheme 3, then it would not have been possible to isolate LtBuFeNPhNHPh during its synthesis from [LtBuFeH]2 and PhN=NPh (some LtBuFeH is present in the reaction mixture at room temperature). These combined observations suggest that the N–N cleavage reaction is unlikely to follow a β -hydride elimination pathway.

3. Ion-Pair and Acid-Catalyzed Mechanisms. The formation of LtBuFeNHPh, LtBuFeNHTol, LtBu'FeNHPh, and LtBu'-FeNHTol from heating LtBuFeNPhNHPh and LtBu'FeNTolNHTol (see above) suggests that Fe-N bonds are disrupted as part of the mechanism. If the reaction proceeds through heterolytic Fe-N bond cleavage in an ion-pair mechanism as shown in eq 1, the rate of the reaction should depend on the polarity of the solvent in which the reaction is carried out. However, the rate constant determined in THF- d_8 [$k = (4.6 \pm 0.2) \times 10^{-4} \text{ s}^{-1}$] at 352 K was not faster than that in toluene- d_8 [$k = (6.9 \pm 0.2) \times$ 10^{-4} s^{-1}] or benzene- $d_6 [k = (5.8 \pm 0.2) \times 10^{-4} \text{ s}^{-1}]$ under similar conditions, arguing against an heterolytic cleavage in the rate-limiting step of the mechanism.

$$LFeNPhNHPh \rightarrow LFe^{+} + {}^{-}NPhNHPh \qquad (1)$$

The possibility that trace acid catalyzes the reaction was tested by adding a non-nucleophilic base under the same reaction conditions. When the reaction was monitored at 359 K in the presence of lutidine (35 mM), the rate constant was (6.5 ± 0.2) \times 10⁻⁴ s⁻¹, again showing no effect on the reaction rate.

4. Radical Mechanisms. The results discussed so far are consistent with a pathway initiated by a homolytic cleavage to yield radicals that react through a chain mechanism. The crossover experiment shows that the mechanism cannot involve N-N bond homolysis along the way to PhN=NPh, and so Fe-N bond homolysis is most strongly implicated as an initiation reaction (eq 2).³² Light does not initiate the reaction, because all kinetic studies were done in the dark (inside an NMR probe).

$$LFeNPhNHPh \rightarrow LFe + \cdot NPhNHPh \qquad (2)$$

Homolytic Fe-N bond cleavage would give the N,N'diphenvlhvdrazinvl radical, which could undergo spontaneous disproportionation to PhN=NPh and PhNHNHPh (eq 3).³³ The two sides of the double bond in the PhN=NPh molecule derive from the same hydrazido complex, assuming a pericyclic mechanism.³³ Therefore, this initial step is consistent with the crossover experiments given above.34,35

$2 \cdot \text{NPhNHPh} \rightarrow \text{PhNHNHPh} + \text{PhN} = \text{NPh}$ (3)

⁽²⁸⁾ The apparent $k_{\rm H}/k_{\rm D}$ is 0.95 \pm 0.23, and the value near unity is consistent with loss of deuterium label.

⁽²⁹⁾ Holt, P. F.; Hughes, B. P. J. Chem. Soc. 1953, 1666-1669.

⁽³⁰⁾ Heesing, A.; Schinke, U. Chem. Ber. 1977, 110, 2867-2871.

⁽³¹⁾ The reaction of LtBuFeCEtCHEt and PhN=NPh at room temperature does not form L^{iBu}FeNPhNHPh. Many other control experiments were also performed to verify that the products do not react with each other or with byproducts (see Supporting Information).

⁽³²⁾ We cannot rule out that the initiation step is N-N bond homolysis, which would give LtBuFeNPh (a potential chain carrier, as discussed below) and PhNH, which would give a small amount of PhNH₂ (presumably undetected). After initiation, the chain would proceed as in Scheme 4.

^{(33) (}a) Shizuka, H.; Kayoiji, H.; Morita, T. Mol. Photochem. 1970, 2, 165– 176. (b) Van Beek, H. C. A.; Heertjes, P. M.; Houtepen, C.; Retzloff, D. J. Soc. Dyers Colour. 1971, 87, 87–92. (c) The Chemistry of the Hydrazo, Azo, and Azoxy Groups; Patai, S., Ed.; Wiley: New York, 1975

⁽³⁴⁾ However, second-order reactions between two transient radicals are relatively unlikely, a phenomenon known as the "persistent radical effect" 35 The implication is that the transient radical species will react primarily with the major species in solution, L^{1Bu}FeNPhNHPh. (a) Fischer, H. *Chem. Rev.* **2001**, *101*, 3581–3610. (b) Daikh, B. E.; Finke, R. G. J. Am. Chem. Soc. **1992**, *114*, 2938–43.

⁽³⁵⁾



What would happen to the iron(I) LtBuFe fragment formed in eq 2? First, it could react with PhNHNHPh (eq 4). This hypothesis was tested by reacting a genuine iron(I) complex, L^{tBu} FeClK(solvent)_n,³⁶ with N,N'-diphenylhydrazine (1 equiv) in C₆D₆. The amido compound L^{tBu}FeNHPh was formed within a few minutes at ambient temperature. Therefore, this step is a reasonable route to the observed product of the overall reaction.

$$2LFe + PhNHNHPh \rightarrow 2LFeNHPh$$
 (4)

The iron(I) fragment could also abstract an NHPh radical from LtBuFeNPhNHPh to give LtBuFeNHPh and LtBuFeNPh, an iron-(III) imido species (eq 5). We have spectroscopically characterized a similar complex, L^{Me} FeNAd (where Ad = 1-adamantyl).37,38 One characteristic reaction of LMeFeNAd is to abstract hydrogen atoms from hydrocarbons (e.g., 9,10-dihydroanthracene and indene), giving an iron(II) amido complex. While we have been unsuccessful in preparing LtBuFeNPh,39 the analogy to L^{Me}FeNAd suggests strongly that the transient formation of LtBuFeNPh is feasible.

> $LFe + LFeNPhNHPh \rightarrow LFeNHPh + LFeNPh$ (5)

 $LFeNPh + \cdot NPhNHPh \rightarrow LFeNHPh + PhN=NPh$ (6)

 $LFeNPh + LFeNPhNHPh \rightarrow LFeNHPh + LFeNPhNPh$ (7)

L^{tBu}FeNPh should be capable of abstracting a hydrogen atom from the hydrazinyl radical (which has a very weak N-H bond) to form LtBuFeNHPh and PhN=NPh (eq 6). This would represent a chain termination step.34 Alternatively, the transient



Figure 6. Crystal structure of LtBuFeNPhNPh. Thermal ellipsoids shown at 50% probability. Hydrogen atoms have been omitted for clarity. Selected metrical parameters follow: Fe-N11 1.987(1) Å, Fe-N1 1.953(1) Å, N1-N1A 1.398(2) Å, N11-Fe-N11A 97.52(5)°, N1-Fe-N1A 41.94(5)°.

imido species could abstract a hydrogen atom from LtBu-FeNPhNHPh to give LtBuFeNPhNPh (eq 7), which may be formulated either as an iron(II) complex with a radical ligand, or an iron(I) complex with coordinated PhN=NPh.

L^{tBu}FeNPhNPh was prepared independently by treating L^{tBu}-FeNNFeL^{tBu} with PhN=NPh at room temperature, and the solidstate structure was determined using X-ray crystallography. Figure 6 shows that the azobenzene ligand is coordinated through the N=N π -bond, and backbonding considerably weakens the N=N bond (N-N = 1.398(2) Å) compared to free trans-azobenzene (1.247(2) Å).⁴⁰ This feature indicates that the formal azobenzene-iron(I) formulation does not account for substantial backbonding into the azobenzene ligand. There are strong analogies to diketiminate-iron(I) complexes of alkenes and alkynes recently reported by our group, in which crystallographic, vibrational, Mössbauer, and theoretical evidence support substantial backbonding into the unsaturated ligand.⁴¹ Therefore, it is reasonable to think of LtBuFeNPhNPh as an iron-(II)-radical complex $L^{tBu}Fe^{2+}(N_2Ph_2)^{\bullet-}$.

In turn, L^{tBu}FeNPhNPh can lose azobenzene to regenerate the active iron(I) fragment (eq 8), which continues the radical chain (Scheme 4). Both sides of the azobenzene produced through eq 8 have come from a single molecule of LtBu-FeNPhNHPh, so this mechanism is consistent with the crossover experiment described above.

$$LFeNPhNPh \rightarrow LFe + PhN=NPh$$
(8)

Effect of Radical Traps on the Rate of the Reaction. Kinetic studies on the decomposition of L^{tBu}FeNPhNHPh in C₆D₆ at 358 K were repeated with large concentrations of the radical traps triphenylmethane (Ph₃CH) (0.46 M) and dihydroanthracene (DHA) (0.35 M). The rate constant for the reaction was not affected [(9.8 \pm 0.3) \times $10^{-4}~s^{-1}$ and (9.3 \pm 0.2) $\times 10^{-4}$ s⁻¹, respectively]. The bond dissociation energy (BDE) of the C-H bonds in Ph₃CH (81 kcal/mol) and DHA (78 kcal/mol) is higher than the BDE of the N-H bond in N,N'-

⁽³⁶⁾ Smith, J. M.; Sadique, A. R.; Cundari, T. R.; Rodgers, K. R.; Lukat-Rodgers, (30) Shini, J. W., Saduci, A. K., Culdari, T. K., Kodges, K. K., Lukar-Rodges, G.; Lachicotte, R. J.; Flaschenriem, C. J.; Vela, J.; Holland, P. L. *J. Am. Chem. Soc.* 2006, *128*, 756–769.
 (37) Eckert, N. A.; Vaddadi, S.; Stoian, S.; Lachicotte, R. J.; Cundari, T. R.; Holland, P. L. *Angew. Chem., Int. Ed.* 2006, *45*, 6868–6871.

⁽³⁸⁾ Other recently isolated iron(III) species with a terminal imido ligand: (a) 634-643. (e) Thomas, C. M.; Mankad, N. P.; Peters, J. C. J. Am. Chem. Soc. 2006, 128, 4956-4957.

⁽³⁹⁾ All attempts to create (diketiminate)Fe^{III}=N(aryl) species have given (diketiminate)Fe^{II}−NH(aryl) products, consistent with powerful H-atom abstraction ability of the putative imido. Eckert, N. A. Ph.D. Thesis, University of Rochester, 2005.

⁽⁴⁰⁾ Bouwstra, J. A.; Schouten, A.; Kroon, J. Acta Crystallogr., Sect. C 1983, 39, 1121-1123.

diphenylhydrazine (69 kcal/mol),⁴² and lower than the BDE of N–H in aniline (92 kcal/mol), so this observation is compatible with the presence of hydrazinyl radicals and argues against amido radicals. Because L^{tBu} FeNPhNHPh is decomposed by TEMPO and nitrosobenzene, these radical traps are not informative.

Which Species Are Chain Carriers? The decomposition of solutions of LtBuFeNPhNHPh at 42 °C in C7D8 shows an induction period, as described above (Figure 5). This reaction was repeated with different reagents added (12-14 mM), to learn whether they eliminate the induction period. We used LtBu-FeCl as a representative of iron(II) compounds, LtBuFeClK- $(solvent)_n$ as a proxy for the "L^{tBu}Fe" intermediate in the catalytic cycle, and the azobenzene complex LtBuFeNPhNPh, in separate experiments. Addition of LtBuFeCl did not eliminate the induction period, while addition of $L^{tBu}FeClK(solvent)_n$ or L^{tBu}FeNPhNPh resulted in formation of L^{tBu}FeNHPh without an induction period at 42 °C. These observations support the chain mechanism including LtBuFe and LtBuFeNPhNPh as chain carriers. In a different experiment, we added benzophenone (2 equiv) to LtBuFeNPhNHPh in order to trap transient iron(I).43 Heating this mixture afforded unidentified paramagnetic complexes and no LtBuFeNHPh was formed. Other traps for iron(I) gave ambiguous results.43

Which Is the Rate-Limiting Step after Initiation? This is a subject that must be approached with caution. As stated by Huyser, "Rate laws for free-radical chain reactions are of limited value in determining the general mechanism of the reaction."46 This is especially true when considering Scheme 4, in which many of the propagation and termination steps give the same products. However, the following arguments may be advanced. The post-initiation kinetics follow a first-order rate dependence on [Fe], which suggests that eq 8 (top arrow in the catalytic cycle of Scheme 4) may be rate-limiting. When LtBuFeNPh-NHPh was heated in the presence of added PhN=NPh (45 mM) in C_6D_6 at 358 K, there was no effect on the rate constant [(7.0 \pm 0.1) \times 10⁻⁴ s⁻¹]. This result suggests that eq 8 is effectively irreversible: every time LtBuFe is formed it is trapped by PhNHNHPh or L^{tBu}FeNPhNHPh rather than by PhN=NPh. In order to test this idea, 1 equiv of PhN=NPh was added to a C₆D₆ solution of L^{tBu}FeClK(solvent)_n with and without 1 equiv of LtBuFeNPhNHPh. Without LtBuFeNPhNHPh, only LtBu-FeNPhNPh was observed in the ¹H NMR spectrum, but in the presence of the hydrazido species no LtBuFeNPhNPh was evident (instead, there was slow conversion to LtBuFeNHPh). Therefore, our experiments suggest that loss of PhN=NPh from the formally iron(I) center in LtBuFeNPhNPh is the slow step of the chain.44

Although the details are tentative, the amassed experimental data are most supportive of a chain mechanism of the type shown in Scheme 4. It remains only to explain how addition of 3-hexyne to the hydrazido complex could yield a small, concentration-independent amount of L^{tBu}FeCEtCHEt (see above). To rationalize this observation, we note that L^{tBu}FeH could be formed by abstraction of a hydrogen atom from the weak N–H bond in the hydrazinyl radical •NPhNHPh by "L^{tBu}Fe" (eq 9; Scheme 4). Because the hydride complex is off the main reaction pathway, the amount of vinyl complex formed is independent of [3-hexyne].

$$LFe + \cdot NPhNHPh \rightarrow LFeH + PhN=NPh$$
(9)

$$LFeH + EtCCEt \rightarrow LFeCEtCHEt$$
 (10)

Conclusions

The reaction of L^{tBu}FeH with 0.5 equiv of PhN=NPh cleanly leads to L^{tBu}FeNHPh, formally cleaving the diazene into "NPh" units that insert into the Fe-H bond. Although the reaction stoichiometry is simple, the mechanism is complex. Fortunately, a combination of mechanistic experiments can be used to rule out most mechanisms, and to support a chain mechanism. The first step of this chain process consists of adding the Fe-H bond across the N=N bond to give a hydrazido complex L^{tBu}-FeNPhNHPh. This intermediate is susceptible to Fe-N bond homolysis to give iron(I) fragments that are capable of breaking N-N bonds. The picture that emerges is a radical chain mechanism that has odd-electron iron species as chain carriers.

One lesson to be learned from this mechanism is that iron complexes with weak ligand fields are capable of [1,2]-addition reactions through both nonradical (in the addition of Fe-H across the N=N bond) and radical (in the N-N cleaving chain reaction) pathways.^{45,46} Although there are clear differences between the coordination sphere of the diketiminate-iron complexes here and the important iron sites in the FeMoco of nitrogenase, one can use these conclusions to speculate about the numerous N-N cleavage and N-H bond forming steps that lie along the path from N₂ to ammonia. Previous workers have understandably favored simpler nonradical pathways as in the Chatt cycle,⁴⁷ and these are representative of the mechanisms of reactions at molybdenum and other second- and third-row metal ions. However, given the recent evidence supporting highspin iron as the reactive site on the FeMoco of nitrogenase,¹⁰ it is important to consider one-electron reactivity in the enzymatic system.

Experimental Section

General Procedures. All manipulations were performed under a nitrogen atmosphere by standard Schlenk techniques or in an M. Braun glove box maintained at or below 1 ppm of O₂ and H₂O. Glassware was dried at 150 °C overnight. NMR data were recorded on a Bruker

 ^{(41) (}a) Stoian, S. A.; Yu, Y.; Smith, J. M.; Holland, P. L.; Bominaar, E. L.; Münck, E. *Inorg. Chem.* 2005, *44*, 4915–4922. (b) Yu, Y.; Smith, J. M.; Flaschenriem, C. J.; Holland, P. L. *Inorg. Chem.* 2006, *45*, 5742–5751.

 ⁽⁴²⁾ Zhao, Y.; Bordwell, F. G.; Cheng, J.-P.; Wang, D. J. Am. Chem. Soc. 1997, 119, 9125–9129.
 (22) Division of the second second

⁽⁴³⁾ Diketiminate-iron(I) complexes react with ketones to give pinacol coupling products: see ref 36. We have obtained ambiguous results when using other traps for iron(I) in the conversion of L^{1Bu}FeNPhNHPh to L^{1Bu}FeNPhPh, as follows. PPh₃ and PEt₃ bind weakly to the L^{1Bu}Fe fragment (see refs 36 and 41b) and do not affect the rate of conversion. CO does bind to L^{1Bu}Fe (see ref 36), but does not affect the decomposition of L^{1Bu}FeNPhNHPh: this is consistent with the observation that addition of a small amount of L^{1Bu}Fe(CO)₂ to the L^{1Bu}FeNPhNHPh abolishes the induction period for its decomposition. Therefore, it is likely that CO binds reversibly to the L^{1Bu}Fe fragment, and does not substantially affect the chain reaction.

⁽⁴⁴⁾ Azobenzene loss may require an isomerization of the η^2 -bound ground state (Figure 6) to an η^1 isomer, a process that can be slow: see ref 220. We are unable to reconcile this rate-limiting step with the *apparent* negative value of ΔS^{ζ} in footnote 27.

⁽⁴⁵⁾ Recent review on radical organometallic reactions in diketiminate chemistry: Smith, K. M. Organometallics 2005, 24, 778–784.

⁽⁴⁶⁾ Despite the conceptual complexity of radical processes, they can lead to high yields and clean products. See, for example: Huyser, E. S. Free-Radical Chain Reactions; Wiley: New York, 1970.

⁽a) Chatt, J.; Pearman, A. J.; Richards, R. L. Nature 1975, 253, 39–40.
(b) Yandulov, D. V.; Schrock, R. R. Science 2003, 301, 76–78. (c) Betley, T. A.; Peters, J. C. J. Am. Chem. Soc. 2003, 125, 10782–10783. (d) Betley, T. A.; Peters, J. C. J. Am. Chem. Soc. 2004, 126, 6252–6254.

Avance 400 (400 MHz) or Bruker Avance 500 spectrometer (500 MHz). All peaks in the NMR spectra are referenced to residual C_6D_5H at 7.16 ppm. In parentheses are listed integrations and assignments. In some cases, it was not possible to determine integrations because of peak overlap. UV–vis spectra were measured on a Cary 50 spectrophoto meter, using screw-cap cuvettes. Solution magnetic susceptibilities were determined by the Evans method,⁴⁸ monitoring the shift in the residual solvent peak relative to a capillary of solvent. Elemental analyses were determined by Desert Analytics, Tucson, AZ.

Pentane, diethyl ether, and toluene were purified by passage through activated alumina and "deoxygenizer" columns from Glass Contour Co. (Laguna Beach, CA). Deuterated benzene and toluene were first dried over CaH₂, then over Na/benzophenone, and then vacuum transferred into a storage container. Before use, an aliquot of each solvent was tested with a drop of sodium benzophenone ketyl in THF solution. Celite was dried overnight at 200 °C under vacuum. The compounds nitro-2,4,6-triisopropylbenzene,49 LtBuFeCl,50 KBEt3H,51 [LtBuFeH]2,11 LtBuFeNPhNHPh,11 LtBuFeCEtCHEt,11 LtBuFeNNFeLtBu,36 $L^{tBu}Fe(Cl)K(solvent)_x^{36}$ were prepared by published procedures. 3-Hexyne, N,N'-diphenylhydrazine, and azobenzene were obtained from Aldrich, and *m*-azotoluene was obtained from TCI America. 3-Hexyne was degassed, vacuum transferred to a new container, and stored in the glovebox at -35 °C. Azobenzene and *m*-azotoluene were dissolved in pentane, dried over molecular sieves, filtered through Celite, and then dried under vacuum. 1,3,5-Triisopropylbenzene, lutidine, n-butyllithium, methyllithium, TMEDA, Pd/C, phosphorus pentachloride, pivaloyl chloride, and triphenylmethane were purchased from Aldrich Chemical Co. and used as received. 9,10-Dihydroanthracene and Ph₃CH were crystallized from ethanol, and the crystals were vacuum-dried. D₂O was purchased from Cambridge Isotope Laboratories, Inc.

Preparation of L^{tBu}'H. Step a. A 200 mL Paar bomb was charged with nitro-2,4,6-triisopropylbenzene (20.0 g, 80.2 mmol) dissolved in isopropanol (80 mL) and Pd/C (3.0 g). H₂ gas (~ 600 psi) was introduced, and the mixture was stirred at 60 °C for 18 h. The contents were filtered in air after cooling to room temperature. Isopropanol was removed by distillation, and the product was distilled at 80 °C (3 mbar) to afford 2,4,6-triisopropylaniline (16.5 g, 94%) as a pale yellow oil. ¹H NMR (400 MHz, CDCl₃) δ 1.1 (12, d, J = 8 Hz, o-iPr-CH₃), 1.3 (6, d, J = 8 Hz, p-iPr-CH₃), 2.9 (1, septet, p-iPr-CH), 3.0 (2, septet, o-iPr-CH), 3.8 (2, br, NH), 6.9 (2, s, m-H).

Step b. A round-bottom flask was charged with 2,4,6-triisopropylaniline (16.5 g, 75.2 mmol) and dichloromethane (50 mL). To the flask was added triethylamine (7.7 g, 75 mmol) and then a solution of pivaloyl chloride (9.3 mL, 75 mmol) in 50 mL of dichloromethane slowly (~5 drops/s). The reaction mixture was refluxed for 2 h at 40 °C, and the solvent was removed using a rotory evaporator. The resultant pink solid was washed with water and diethyl ether to yield *N*-(2,4,6triisopropylphenyl)-2,2-dimethylpropionamide as a white microcrystalline powder which was dried under vacuum (21.8 g, 96%). ¹H NMR (400 MHz, CDCl₃) δ 1.17 (12, d, *J* = 8 Hz, *o*-iPr-CH₃), 1.22 (6, d, *J* = 8 Hz, *p*-iPr-CH₃), 1.33 (9, s, tBu), 2.86 (1, septet, *p*-iPr-CH), 2.97 (2, septet, *o*-iPr-CH), 6.73 (1, s, NH), 6.98 (2, s, *m*-H).

Step c. Phosphorus pentachloride (16 g, 73 mmol) was added in small portions to a slurry of N-(2,4,6-triisopropylphenyl)-2,2-dimethylpropionamide (21.8 g, 71.8 mmol) in benzene (33 mL) under N₂ flow. During the course of the addition, HCl gas was liberated, and it was passed through aqueous Na₂CO₃. The resultant yellow reaction mixture was stirred for 18 h under the continuous flow of N₂. The solvent and byproducts were distilled off, and the product was distilled at 110 °C (3 mbar) to yield 1-chloro-1-(2,4,6-triisopropylphe-nylimino)-2,2-di(methylpropane) (17.8 g, 77%) as a colorless gel. This product was handled under N₂. ¹H NMR (400 MHz, CD₂Cl₂) δ 1.16 (12, d, J = 8 Hz, *o*-iPr-CH₃), 1.18 (6, d, J = 8 Hz, *p*-iPr-CH₃), 1.42 (9, s, tBu), 2.73 (2, septet, *p*-iPr-CH), 2.95 (1, septet, *o*-iPr-CH), 7.0 (2, s, *m*-H).

Step d. A round-bottom flask was charged with 1-chloro-1-(2,4,6-triisopropylphenylimino)-2,2-dimethylpropane (14.8 g, 46 mmol) and pentane (80 mL) inside a glovebox and cooled in a cold well (-60 °C). To this solution was added 1.6 M methyllithium (29 mL, 46 mmol) in diethyl ether. The turbid yellow reaction mixture was stirred at room temperature for 18 h, and then brought out of the glove box. Water (50 mL) was added to the reaction mixture slowly and very carefully. The aqueous layer was separated and extracted with diethyl ether. The organic fractions were combined and dried with anhydrous magnesium sulfate and filtered, and solvent was removed. The product was dried under vacuum to yield 2-(2,4,6-triisopropylphenylimino)-3,3-dimethylbutane (10.7 g, 77%) as a pale yellow solid. ¹H NMR (400 MHz, CD₂Cl₂) δ 1.2 (12, br, *o*-iPr-CH₃), 1.3 (24, br, *p*-iPr-CH₃,tBu), 1.7 (3, br, CH₃), 2.6 (2, br, *o*-iPr-CH), 2.8 (1, br, *p*-iPr-CH), 6.9 (2, s, *m*-H).

Step e. A round-bottom flask was charged with 2-(2,4,6-triisopropylphenylimino)-3,3-dimethylbutane (10.7 g, 35 mmol) and diethyl ether (60 mL). To this was added TMEDA (5.5 mL, 36 mmol), and the reaction mixture was cooled in a cold well (-60 °C). n-Butyllithium (14.6 mL, 2.5 M in hexanes, 36 mmol) was added slowly while stirring. The resultant dark orange reaction mixture was stirred in the cold well for 1 h and then stirred at room temperature for 18 h. The dark orange reaction mixture with yellow precipitate was cooled in the cold well, and then 1-chloro-1-(2,4,6-triisopropylphenylimino)-2,2-dimethylpropane (11.4 g, 35 mmol) in 20 mL of pentane was added dropwise. Once the addition was complete, the reaction mixture was stirred at 40 °C for 6 h. Water (50 mL) was added; the aqueous layer was separated and extracted with diethyl ether. All the organic fractions were combined and dried with magnesium sulfate and filtered, and the volatile components were removed under vacuum. The pale yellow solid was dissolved in hot pentane, filtered, and stored at -25 °C to afford L^{tBu'}H (15.3 g, 72%) as pale yellow crystals. ¹H NMR (400 MHz, CDCl₃) δ 1.1 (24, br, o-iPr-CH₃), 1.3 (18, br, tBu), 1.7 (12, br, p-iPr-CH₃), 2.7 (1, br, p-iPr-CH), 2.9 (2, br, o-iPr-CH), 3.3 (2, br, backbone CH), 6.9 (4, s, *m*-H). LCMS: m/z = 587. Anal. Calcd for C₄₁H₆₆N₂: C 83.89, H 11.33, N 4.77. Found: C 82.62, H 11.33, N 4.65.

Preparation of L'^{Bu'}Li(THF). A Schlenk flask was charged with L'^{Bu'}H (12.1 g, 20.6 mmol) and THF (100 mL) and cooled to -60 °C. A solution of *n*-butyllithium (9.0 mL, 2.5 M in hexanes, 22.5 mmol) was added slowly with stirring. The reaction mixture was warmed to room temperature and heated at 60 °C for 2 h. The volatile components were removed under vacuum, and the product was dissolved in warm pentane and stored at -35 °C to afford pale yellow crystals of L'^{Bu'}Li(THF) (9.6 g, 70%). ¹H NMR (400 MHz, C₆D₆) δ 0.9 (4, br, THF), 1.14 (12, d, iPr-CH₃), 1.20 (12, d, iPr-CH₃), 1.39 (18, s, tBu), 1.42 (12, d, iPr-CH₃), 2.3 (4, br, THF), 2.8 (2, septet, iPr-CH), 3.5 (4, septet, iPr-CH), 5.24 (1, s, backbone-CH), 6.96 (4, s, *m*-H).

Preparation of L^{tBu}'FeCl. A bomb flask was charged with FeCl₂-(THF)_{1.5} (1.3 g, 5.6 mmol), L^{tBu'}Li(THF) (3.7 g, 56 mmol), and toluene (70 mL). The red reaction mixture was heated at 100 °C for 18 h. The volatile components were removed under vacuum, and the red residue was transferred to a glass thimble, which was placed in a Soxhlet extractor. The residue was extracted continuously with boiling diethyl ether, until the extracting solvent was clear (24 h). The red extract was concentrated under vacuum, and the resultant red solid was isolated (2.04 g, 54%). The mother liquor was stored at -35 °C to afford L^{tBu'}-

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FeCl as red crystals (1.2 g, 32%). Total yield 3.2 g (86%). ¹H NMR (400 MHz, C₆D₆) δ 105 (1, backbone-CH), 42 (18, 0.3, tBu), 35 (2, 0.7, iPr-CH), 2.1 (4, *m*-H), -5 (12, 0.7, iPr-CH₃), -28 (12, 0.5, iPr-CH₃), -111 (12, 0.1, iPr-CH₃), -115 (4, iPr-CH). LCMS: *m*/*z* = 676. Anal. Calcd for C₄₁H₆₅N₂FeCl: C 71.71, H 9.67, N 4.14. Found: C 70.81, H 10.25, N 3.99.

Preparation of $[L^{tBu'}FeH]_2$. A Schlenk flask was charged with $L^{tBu'}$ -FeCl (0.884 g, 1.30 mmol) and toluene (30 mL). To this red solution was added a clear solution of KBEt₃H (0.180 g, 1.30 mmol) in toluene (10 mL). The dark red reaction mixture was stirred at room temperature for 45 min, and the volatile materials were removed under vacuum. The residue was extracted with pentane and filtered through Celite to give a dark red solution. This solution was concentrated, warmed to dissolve the product, and then cooled to -35 °C to afford very dark red crystals (501 mg, 78%). It is critical that the reaction is not stirred for too long because [LtBu'FeH]2 reacts with the BEt3 byproduct. Unlike its LtBu analogue, [LtBu'FeH]2 is soluble in pentane. 1H NMR ([LtBu'-FeH]₂, 22 °C, C₆D₆) 72, 42, 22, 21, 14, 13, 1.2, -1.3, -5, -8, -10, -19, -26, -29, -42, -51, -120. ¹H NMR (L^{tBu'}FeH, 80 °C, C₆D₆) δ 45 (1, backbone-CH), 31 (2, p-iPr-CH), 30 (18, tBu), 9 (4, m-H), -3 (12, iPr-CH₃), -17 (12, iPr-CH₃), -81 (4, o-iPr-CH), -82 (12, iPr-CH₃). Anal. Calcd for C₈₂H₁₃₂N₄Fe: C 76.61, H 10.35, N 4.36. Found: C 76.59, H 10.42, N 4.22.

Preparation of L^{tBu'}FeNPhNHPh. From [L^{tBu'}FeH]₂: An orange solution of azobenzene (14 mg, 77 µmol) in Et₂O (2 mL) was added to a dark red slurry of [LtBu'FeH]2 (50 mg, 38 µmol) in Et2O (8 mL). A red solution was formed over the course of an hour. The volatile materials were removed under vacuum. The residue was extracted with pentane, filtered through Celite, concentrated, and stored at -35 °C to give LtBu'FeNPhNHPh as red crystals. From LtBu'FeCl: A 20 mL scintillation vial was charged with LtBu'FeCl (172 mg, 253 µmol), azobenzene (46 mg, 253 µmol), and Et₂O (10 mL). A solution of KBEt₃H (35 mg, 253 μ mol) in Et₂O (2 mL) was added to give a dark red solution. After stirring at room temperature for 2 h, the volatile components were removed under reduced pressure. The resultant residue was extracted with pentane and filtered through Celite to give a red solution. The solution was concentrated and stored at -35 °C to afford LtBu'FeNPhNHPh as red crystals (126 mg, 60%). ¹H NMR (400 MHz, C₆D₆) δ 111 (1, backbone-CH), 43 (18, tBu), 23 (2, p-iPr-CH), -7.8 (12, p-iPr-CH₃), -8.6 (4, diketiminate m-H), -23 (6, o-iPr-CH₃), -29 (6, o-iPr-CH₃), -45 (1), -82 (6, o-iPr-CH₃), -170 (6, 0.09, o-iPr-CH₃), -181 (br). Due to the thermal instability of the solid, we have not been able to obtain successful elemental analysis.

Preparation of L^{tBu}FeNTolNHTol. A 20 mL scintillation vial was charged with L^{tBu}FeCl (102 mg, 172 μmol), azo-*m*-toluene (36 mg, 172 μmol), and Et₂O (10 mL). A solution of KBEt₃H (24 mg, 172 μmol) in Et₂O (2 mL) was added to yield a dark red solution. After stirring at room temperature for 2 h, the volatile components were removed under reduced pressure. The resultant residue was extracted with pentane and filtered through Celite to give a red solution. The solution was concentrated and stored at -35 °C to afford L^{tBu}-FeNTolNHTol as red crystals (82 mg, 62%). ¹H NMR (400 MHz, C₆D₆) δ 112 (1, backbone-CH), 43 (18, tBu), -7 (2, diketiminate *m*-H), -8 (2, diketiminate *m*-H), -23 (6, iPr-CH₃), -28 (6, iPr-CH₃), -44 (1, hydrazido *p*-H), -83 (6, iPr-CH₃), -91 (2, diketiminate *p*-H), -172 (8, iPr-CH₃ and iPr-CH).

Preparation of L^{tBu}'FeNTolNHTol. An orange solution of azo-*m*toluene (16 mg, 76 μ mol) in Et₂O (2 mL) was added to a dark red slurry of [L^{tBu}'FeH]₂ (49 mg, 38 μ mol) in Et₂O (8 mL). A red solution was formed over the course of an hour. The volatile materials were removed under vacuum. The residue was extracted with pentane, filtered through Celite, concentrated, and stored at -35 °C to give L^{tBu'}-FeNTolNHTol as red crystals (39 mg, 60%).

Preparation of PhNDNDPh. A Schlenk tube was charged with N,N'-diphenylhydrazine (215 mg, 1.17 mmol) and pentane (10 mL). A

solution of *n*-butyllithium (0.95 mL, 2.5 M in hexanes) was added while stirring. The thick yellow suspension was stirred for 18 h at room temperature. The volatile components were removed under reduced pressure. The contents were cooled in an ice bath, and D₂O (10 mL) was added under nitrogen flow to give a white precipitate. The slurry was stirred for 2 h, and the reaction mixture was allowed to settle. The yellow solution was decanted using a cannula, and the white solid was dried under reduced pressure. The solid was extracted with pentane/ diethyl ether solvent mixture, filtered through Celite, concentrated, and stored at -35 °C to afford PhNDNDPh as colorless flaky crystals (~98% deuteration by NMR). ¹H NMR (400 MHz, C₆D₆) δ 6.7 (4, *o*-H), 6.8 (2, *p*-H), 7.2 (4, *m*-H).

Preparation of L^{tBu}FeNPhNDPh. A solution of PhNDNDPh (8.3 mg, 44 μmol) in Et₂O (2 mL) was added to a dark red slurry of [L^{tBu}FeH]₂ (25 mg, 22 μmol) in Et₂O (8 mL). A red solution was formed over the course of an hour. The volatile materials were removed under vacuum. The residue was extracted with pentane, filtered through Celite, concentrated, and stored at -35 °C to give L^{tBu}FeNPhNDPh as red crystals (56 mg, 60%). ¹H NMR (400 MHz, C₆D₆) δ 112 (1, backbone-CH), 43 (18, tBu), -7 (2, diketiminate *m*-H), -8 (2, diketiminate *m*-H), -23 (6, iPr-CH₃), -28 (6, iPr-CH₃), -45 (1, hydrazido *p*-H), -87 (6, iPr-CH₃), -94 (2, diketiminate *p*-H), -171 (8, iPr-CH₃ and iPr-CH).

Preparation of L^{IB}^u**FeNPhNPh.** A 20 mL scintillation vial was charged with L^{IB}^uFeNNFeL^{IB}^u (98 mg, 86 μmol) and Et₂O (10 mL). A solution of azobenzene (31 mg, 170 μmol) in Et₂O (2 mL) was added to yield a brown solution, which turned dark green upon stirring for 30 min. After stirring at room temperature for 2 h, the volatile components were removed under reduced pressure. The resultant residue was extracted with toluene and filtered through Celite to give a dark green solution. The solution was concentrated and stored at -35 °C to afford L^{IB}^uFeNPhNPh as dark green crystals (51 mg, 40%). ¹H NMR (400 MHz, C₆D₆) δ 119 (4, br, Ph *o*-H), 72 (1, Ph *p*-H), 42 (1, Ph *p*-H), 24 (18, tBu), -11 (12, iPr-CH₃), -15 (4, diketiminate *m*-H), -17 (2, diketiminate *p*-H), -22 (2, Ph *m*-H), -50 (12, iPr-CH₃), -70 (2, br, iPr-CH). Anal. Calcd for C₄₇H₆₃N₄Fe: C 76.30, H 8.58, N 7.57. Found: C 76.51, H 8.42, N 8.00.

Crystallography. Crystals were placed onto the tip of a 0.1 mm diameter glass capillary tube or fiber and mounted on a Bruker SMART APEX II CCD Platform diffractometer⁵² for data collection at 100.0-(1) K using Mo Ka radiation (graphite monochromator). A randomly oriented region of reciprocal space was surveyed: four major sections of frames were collected with 0.30° steps in ω at four different ϕ settings and a detector position of -28° in 2θ . The intensity data were corrected for absorption.53 Final cell constants were calculated from the xyz centroids of >3500 strong reflections from the actual data collections.54 The structures were solved using direct methods and refined using SHELXL-97.55 The space groups (C2/c in each case) were determined on the basis of systematic absences and intensity statistics. All nonhydrogen atoms were refined with anisotropic displacement parameters. The bridging hydrogen atoms in [L^{tBu'}FeH]₂ were found from the difference Fourier map, and their positions were refined independently from the iron atoms, but with relative (based on Fe1) isotropic displacement parameters. It is understood that these positions are approximate. All other hydrogen atoms were placed in ideal positions and refined as riding atoms with relative isotropic displacement parameters. The final full-matrix least-squares refinement for [LtBu'-FeH]₂ converged to R1 = 0.0467 (F^2 , $I > 2\sigma(I)$) and wR2 = 0.1328 $(F^2, all data)$. The final full-matrix least-squares refinement for

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L^{tBu}Fe(N₂Ph₂) converged to R1 = 0.0384 (F^2 , $I > 2\sigma(I)$) and wR2 = 0.1035 (F^2 , all data).

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Supporting Information Available: Eyring plot and NMR data (pdf) and crystallographic data (cif). This material is available free of charge via the Internet at http://pubs.acs.org.

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