Characterization of a High-Spin Non-Heme {FeNO}⁸ Complex: Implications for the Reactivity of Iron Nitroxyl Species in Biology**

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Nitric oxide (NO[•]) has long been implicated in numerous signaling and immune defense pathways in mammalian systems. More recently, its one-electron-reduced and potentially protonated form nitroxyl (NO-/HNO) has been shown to elicit a variety of biological responses, although the endogenous production of nitroxyl has not yet been established.^[1] HNO has been suggested to bind to both ferric and ferrous hemes to yield {FeNO}⁷-type species (in the Enemark-Feltham notation^[2]) in the former case and {Fe-(H)NO⁸-type species in the latter.^[3] Non-heme iron centers are an alternative target for HNO, but non-heme iron nitroxyl adducts have not been well investigated. However, since nonheme {FeNO}⁷ complexes have more positive reduction potentials than their heme counterparts, it is feasible that {FeNO}⁸ species could be formed from non-heme {FeNO}⁷ complexes under physiological conditions. Furthermore, nonheme {Fe-(H)NO}⁸-type species have also been proposed as key intermediates in the catalytic cycles of bacterial respiratory NO reductases (NorBC) and flavodiiron NO reductases (FNORs).^[4,5] Although multiple *low-spin* (diamagnetic) {FeNO}⁸ model complexes have been reported previously,^[6-8] there are no corresponding examples of model complexes for biological non-heme iron centers.

Herein, we report the isolation and characterization of the model complex $[Fe(TMG_3tren)(NO)](OTf)_2$ (1, Scheme 1), which is the first five-coordinate high-spin $\{FeNO\}^7$ complex with neutral N-donor ligands, and its one-electron-reduced form 2, which is the first example of a stable high-spin $\{FeNO\}^8$ model complex. The spectroscopic and electronic properties of 2 are then contrasted with those of low-spin $\{FeNO\}^8$ species. Finally, the biological implications of the electronic structure of high-spin $\{FeNO\}^8$ species are discussed.

The addition of excess NO gas to a colorless solution of $[Fe(TMG_3tren)(CH_3CN)](OTf)_2$ in CH₃CN causes an immediate color change to black, which is indicative of the formation of **1**. NO binding leads to the appearance of an intense absorption band at 368 nm and two lower-intensity

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Scheme 1. Structures of the ligand TMG_3 tren, the model complex [Fe(TMG_3 tren)(NO)](OTf)₂ (1), and the one-electron-reduced form 2. OTf=trifluoromethanesulfonate (triflate).

features at 569 and 800 nm in the UV/Vis spectrum (Figure 1). Complex 1 has the expected high-spin ground state, as indicated by its EPR spectrum, which exhibits a rhombic



Figure 1. UV/Vis absorption spectra of [Fe(TMG₃tren)(CH₃CN)]OTf₂ and **1** in CH₃CN, and of the bulk electrolysis of **1** (ca. 2 mM) to **2** in 0.1 m NBu₄ClO₄ in CH₃CN at -1.0 V versus Ag wire (transition from black to gray via thin lines). Inset: Spectra at a concentration of approximately 100 μ M.

S = 3/2 signal (Figure 2). The FTIR spectrum of solid **1** shows an intense v(N–O) stretching band at 1730 cm⁻¹ (see Figure S1 in the Supporting Information). The identity of **1** was further confirmed by X-ray crystallography (Figure 3).^[9] As has been observed for other iron–TMG₃tren complexes, **1** has a completely trigonal bipyramidal geometry ($\tau = 1.03$).^[10,11] The Fe–N(O) and N–O bond lengths are typical for {FeNO}⁷ species.^[4] The Fe-N-O unit has a relatively large angle of 168°.

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Figure 2. X-band EPR spectrum (shown with fit) of a frozen solution of 1 in CH_2CI_2 . Fit parameters: $g_x = g_y = 2.01$, $g_z = 2.00$, E/D = 0.070.



Figure 3. Crystal structure of 1 with ellipsoids shown at 50% probability.^[9] Hydrogen atoms, counterions, and solvent molecules have been omitted for clarity. Fe orange, C gray, N blue, O red. Selected bond distances [Å] and angles [°]: Fe–N (O) 1.748, N–O 1.154, avg. Fe–N_{guan} 2.037, Fe–N_{amine} 2.251; Fe-N-O 168.0.

which is a result of the sterically encumbering $TMG_3 tren \ ligand.^{[12]}$

The electrochemical reduction of high-spin non-heme {FeNO}⁷ species has been reported for only a few complexes.^[12-15] Importantly, the cyclic voltammograms of these complexes show quasi-reversible or irreversible {FeNO}^{7/8} redox couples, which suggests that the reduced species is highly unstable and hence cannot be isolated for further studies. In contrast, the cyclic voltammogram of 1 exhibits a nearly reversible redox event assigned to the {FeNO}^{7/8} couple at -1.34 V versus Fc/Fc⁺ (Figure 4). The nature of this reduction was first confirmed by IR spectroelectrochemistry (Figure 5; see also Figures S2 and S3). The reduction of 1 to 2 in a thin-layer electrochemical cell is accompanied by the disappearance of the $\nu(N-O)$ stretch at 1748 cm⁻¹ and a splitting of the ligand guanidinium stretches at 1548 and 1538 cm⁻¹.^[16] Concomitantly, a new broad shoulder that overlaps with ligand bands appears at 1618 cm⁻¹. This feature could be unambiguously assigned as the ν (N–O) stretch of the {FeNO}⁸ complex by comparison with the ¹⁵NO isotopomer.



Figure 4. Cyclic voltammogram of 1 in a 0.1 M solution of NBu₄ClO₄ in CH₃CN. Fc = ferrocene.



Figure 5. IR spectroelectrochemical analysis of the reduction of 1-NO to 2-NO (top) and the reduction of $1-^{15}$ NO to $2-^{15}$ NO (middle). Bottom: A difference spectrum generated by subtracting the spectrum for $2-^{15}$ NO from the spectrum for 2-NO. * denotes contributions from residual 1-NO.

Interestingly, the 130 cm⁻¹ downshift in ν (N–O) upon oneelectron reduction is significantly smaller than that observed in most low-spin systems (see below).

Bulk electrolysis of **1** produces a species for which the $\nu(N-O)$ band observed by solution IR spectroscopy is identical to that observed in the electrochemical cell (see Figure S4). The reduction leads to a color change from brown to yellow corresponding to a decrease in the intensity of the absorption feature at 365 nm and the appearance of two new bands at 500 and 700 nm in the UV/Vis spectrum (Figure 1). This reduction is a single, clean transformation, as indicated by the isosbestic points at 660 and 750 nm.

To facilitate the isolation and further study of 2, we also used chemical reductants. The treatment of 1 with equimolar $[CoCp*_2]$ produces 2 cleanly, as indicated by solution IR, EPR, and ¹H NMR spectroscopic measurements (see Figur-



es S6, S7, and S9; Cp* = pentamethylcyclopentadienyl). Because the $[CoCp*_2]OTf$ by-product produced in the reaction is diamagnetic, the spin of **2** may be readily determined by ¹H NMR spectroscopy using the Evans method. Consistent with its S = 3/2 spin state, the ¹H NMR spectrum of **1** exhibits multiple broad, paramagnetically shifted residues between 0 and 200 ppm (see Figure S8) and a magnetic moment of 3.9 μ_B , as measured by the Evans method. In contrast to low-spin {FeNO}⁸ species, which are diamagnetic, **2** also shows paramagnetically shifted residues between -15 and 110 ppm (see Figure S9) and a magnetic moment of 3.1 μ_B , which is relatively close to the spin-only value for an S = 1 system ($\mu_{eff} = 2.87 \mu_B$). Taken together, these results demonstrate the formation of the first high-spin {FeNO}⁸ model complex reported to date.

We used DFT calculations to further gain an understanding of the changes in bonding that occur upon reduction. The electronic structure of **1** is consistent with the previously proposed bonding description for high-spin {FeNO}⁷ complexes, according to which high-spin Fe^{III} is antiferromagnetically coupled to a triplet NO⁻ (S = 1).^[4,13,17] Correspondingly, in the α -spin manifold, the empty NO π^* orbitals form a weak π back bond with the occupied iron d_{xz} and d_{yz} orbitals (in a coordinate system in which the *z* axis corresponds to the Fe–N(O)bond). In the β -spin manifold, the occupied NO π^* orbitals donate strongly into the unoccupied iron d_{xz} and d_{yz} orbitals (Figure 6, left). The important question is how the electronic structure changes upon reduction, and where the additional electron is localized in **2**. The calculations show that the extra electron occupies the d_{xy} orbital of iron and thus indicate that the high-spin complex **1** undergoes a *metal-centered* reduction (Figure 6, right). The increased electron density of the iron center then causes a moderate decrease in π donation from NO⁻ to Fe^{II} in the {FeNO}⁸ complex, and this induces the observed moderate decrease in ν (N–O). This decrease in π donation in **2** is directly reflected by the increased NO π^* character of the β -spin orbitals of **2** relative to **1** (see Figure 6). In summary, these results are indicative of decreased covalency in the Fe–NO unit upon one-electron reduction, and therefore a weaker Fe–NO bond.

In contrast to these findings, the reduction of low-spin $\{FeNO\}^7$ complexes $(S = 1/2; S_{Fe} = 0, S_{NO} = 1/2)$ has been observed to be *NO-centered* and to lead to a diamagnetic species in which the coordinated NO⁻ ligand is in the singlet state. For example, in [Fe(cyclam-ac)(NO)] (cyclam-ac = 1,4,8,11-tetraazacyclotetradecane-1-acetate), reduction is almost entirely NO-centered, as reflected by its large ν (N–O) downshift of 336 cm⁻¹.^[8] In heme systems, DFT calculations have revealed an electronic structure of the {FeNO}⁸ species that is intermediate between low-spin Fe^{II}–NO⁻ and Fe^I–NO^{.[6c,d,18]} Thus, heme {FeNO}⁸ complexes show somewhat reduced downshifts of about 200 cm⁻¹ in ν (N–O); however, these shifts are still much larger than that observed for **2** (see Table S1).



Figure 6. Schematic molecular-orbital diagrams of 1 (left) and 2 (right). $S_{\alpha\beta}$ indicates the degree of spatial overlap between the α - and β -spin orbitals.

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The proposed metal-centered reduction and decreased Fe–NO covalency in high-spin non-heme {FeNO}⁸ species as compared to {FeNO}⁷ complexes has implications for the basicity and reactivity of non-heme iron nitroxyl complexes. Owing to the high degree of covalency of the Fe–NO bond, {FeNO}⁷ units are generally stable and unreactive, and cannot be protonated. Correspondingly, whereas **1** shows no reactivity upon the addition of a weak acid (see Figure S10), **2** reacts immediately with the acid to form a purple species **3**, which decays within approximately 1 min at room temperature (Figure 7). Low-temperature studies of this potential



Figure 7. UV/Vis spectra showing the rapid decomposition of **2** upon the addition of [HNEt₃][PF₆] (1.2 equiv).

{Fe-HNO}⁸ complex are currently underway. Interestingly, this result indicates that in the high-spin {FeNO}⁸ complex 2, the basicity of the NO⁻ ligand is greatly increased despite the fact that the reduction is metal-centered.^[19] In line with this increased basicity, heat annealing of the cryoreduced NO adduct of ferrous taurine:alpha-ketoglutarate dioxygenase (TauD) leads to the formation of a new species, which is proposed to be an {Fe-(H)NO}⁸ intermediate.^[20] Reduction also promotes other reactivity; for example, reduction of the stable diferrous dinitrosyl FNOR model complex [Fe2- $(BPMP)(OPr)(NO)_2](BPh_4)_2$ (BPMP = 2,6-bis][bis(2-pyridy]methyl)amino]methyl]-4-methylphenolate) to the corresponding $[{FeNO}^{8}]_{2}$ dimer leads to the rapid, quantitative production of N₂O.^[21] Overall, these results suggest that reduction serves as a potent method for the activation of stable non-heme {FeNO}⁷ units for further reactivity.

In conclusion, we have reported herein the first example of a high-spin non-heme {FeNO}⁸ model complex. In contrast to low-spin {FeNO}⁸ systems, which undergo NO-centered reduction and are diamagnetic, **1** undergoes iron-centered reduction (into d_{xy}) to an S = 1 species in which a triplet NO⁻ ligand is bound to a high-spin ferrous center. Interestingly, a similar electronic structure was proposed for the {FeNO}⁸ adduct of TauD;^[20] thus, metal-centered reduction may be a general feature of high-spin {FeNO}^{7/8} systems. Because the ligand TMG₃tren is an extremely strong donor, the reduction of **1** occurs at a relatively negative potential. However, biological donors are much weaker ligands; therefore, biological non-heme iron nitrosyl complexes are expected to undergo reduction at more positive, biologically feasible potentials. Accordingly, the non-heme {FeNO}⁷ model complexes $[Fe(BMPA-Pr)(NO)]X (X = ClO_4, OTf; BMPA-Pr =$ *N*-propanoate-*N*,*N*-bis(2-pyridylmethyl)amine) undergo reduction at approximately -300 mV versus the normal hydrogen electrode (NHE).^[13] This result suggests that nonheme iron centers could potentially function as HNO synthases in vivo, if HNO loss from these complexes was facile. Importantly, the decreased covalency in the Fe-NO unit that results from the reduction makes high-spin {FeNO}⁸ complexes reactive towards weak acids and amenable to other chemical transformations. In particular, {FeNO}⁸ species could be central intermediates responsible for the key N-N bond forming step in NO reductases.^[4]

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