See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/24443792

Dinitrosyl Iron Complexes Relevant to Rieske Cluster Nitrosylation

ARTICLE in JOURNAL OF THE AMERICAN CHEMICAL	SOCIETY · JUNE 2009	
Impact Factor: 12.11 · DOI: 10.1021/ja9030159 · Source: PubMed		
CITATIONS	READS	
10	13	

3 AUTHORS, INCLUDING:



Zachary J Tonzetich

University of Texas at San Antonio

39 PUBLICATIONS **680** CITATIONS

SEE PROFILE



Am Chem Soc. Author manuscript; available in PMC 2010 June 17.

Published in final edited form as:

J Am Chem Soc. 2009 June 17; 131(23): 7964–7965. doi:10.1021/ja9030159.

Dinitrosyl Iron Complexes Relevant to Rieske Cluster Nitrosylation

Zachary J. Tonzetich, Loi H. Do, and Stephen J. Lippard
Department of Chemistry, Massachusetts Institute of Technology, Cambridge, MA

Abstract

Reaction of the Rieske cluster model complex $(Et_4N)_2[(N_2CHPh)Fe_2S_2(S_2\text{-}o\text{-}xyl)]$ $(N_2CHPh) = \text{dianion of } 2,2'\text{-}(\text{phenylmethylene})\text{bis}(3\text{-methylindole}); S_2\text{-}o\text{-}xyl = \text{dianion of } 1,2\text{-} \text{phenylenedimethanethiol})$ with nitric oxide results in disassembly of the iron-sulfur core and formation of $\{Fe(NO)_2\}^9$ dinitrosyliron complexes. Isolation and characterization of these DNICs, including the new compound, $(Et_4N)[(N_2CHPh)Fe(NO)_2]$, demonstrates a homology between the synthetic Riekse cluster and purely thiolate bound Fe_2S_2 clusters in reactions involving NO. To model the nitrogen rich environment of Rieske cluster-derived dinitroysliron species, a new type of neutral $\{Fe(NO)_2\}^9$ DNIC was prepared containing a β -diketiminate ligand. One-electron reduction of this compound affords the isolable $\{Fe(NO)_2\}^{10}$ DNIC. These compounds represent a rare example of structurally analogous DNIC redox partners.

Iron-sulfur clusters are among the most ubiquitous cofactors in biology. The diversity of structures encountered in iron-sulfur proteins reflects the many functions nature employs for these versatile units, from electron transfer to substrate binding and activation. 1 Of the many variations of iron-sulfur clusters, only a select few have ligands other than cysteine. Among these, the best-documented examples are Rieske-type $\mathrm{Fe_2S_2}$ clusters 2 that contain a pair of histidine residues ligating one iron atom. 3,4 Recently, the first synthetic analog of a Rieske cluster was reported (1, Scheme 1). 5 We were interested to explore the nitric oxide reactivity of such a compound as part of ongoing investigations into the reactivity of non-heme iron with NO. Moreover, there is very little information about the interaction of Rieske proteins with NO, 6 which contrasts with the large body of literature regarding nitric oxide reactivity with purely cysteine-bound iron-sulfur clusters. 7 Herein we report the results of our studies of NO reactions with 1 (Scheme 1) and introduce related N-bound dinitrosyliron complexes (DNICs) containing β -diketiminate ligands.

Reaction of Rieske model complex **1** with four equiv of nitric oxide in acetonitrile solution results in the appearance of two new nitrosyliron species after workup as judged by IR spectroscopy. The first displays v_{NO} bands at 1795, 1743 and 1705 cm⁻¹ (THF solution) consistent with formation of Roussin's black salt (RBS), [Fe₄S₃(NO)₇]⁻. ⁸ We demonstrated previously that RBS can arise from reaction of NO with an Fe₂S₂ cluster containing (S₂-o-xyl)²⁻ ligands via the intermediacy of the thiolate bound DNIC. ⁹ This DNIC can be observed by IR spectroscopy in reactions of **1** with Ph₃CSNO in benzonitrile. After addition of 1 to 2 equiv of nitrosothiol, v_{NO} bands at 1739 and 1679 cm⁻¹ appear, consistent with formation of

 $[(S_2-o-xyl)Fe(NO)_2]^-$. Upon addition of further equiv of Ph₃CSNO, these IR bands diminish, giving rise to those of RBS.

The second iron nitrosyl species obtained from the reaction shown in Scheme 1 displays v_{NO} peaks at 1761 and 1692 cm⁻¹ (IR, KBr). Recrystallization from DMF/Et₂O and subsequent spectroscopic characterization identified a new DNIC, **2** (Scheme 1). The N-bound DNIC displays a rhombic EPR signal centered at 2.03 (Supporting Information, SI) consistent with its formulation as an S = ½ paramagnetic complex and a $\{Fe(NO)_2\}^9$ species in the Enemark-Feltham notation. Compound **2** also contains a quadrupole doublet with an isomer shift of 0.16(2) mm/s in the ⁵⁷Fe Mössbauer spectrum (SI), which corresponds well with the value of 0.18(2) mm/s determined previously for $(Et_4N)[(FePhS)_2(NO)_2]$. Notably, several examples of dinitrosyliron compounds containing nitrogen donor ligands have been published previously as potential models for histidine-ligated DNICs. ¹¹ The synthesis of compound **2** represents the first verification that such a DNIC can be prepared directly from reaction of an iron-sulfur cluster with nitric oxide. Compound **2** was also independently synthesized by salt metathesis of $(Et_4N)[FeCl_2(NO)_2]$ with the lithium salt of the nitrogen ligand displayed in Scheme 1. Spectroscopic features of the DNIC prepared in this fashion are identical in all respects to those observed for that prepared from **1**.

In order to prepare a more soluble version of Rieske cluster-derived DNICs such as 2 for more extensive studies, we turned our attention to sterically demanding β-diketiminate ligands. We reasoned that β -diketiminates would chelate the $\{Fe(NO)_2\}^9$ unit giving rise to stable DNICs, thereby avoiding many of the common shortcomings of thiolate ligands such as tendency toward oxidation and kinetic lability. Furthermore, \beta-diketiminate ligands have been employed previously to stabilize low valent iron. 12 As an entry into this chemistry the iodide precursor [FeI₂(NO)₂] was selected as a source of the {Fe(NO)₂} unit. ¹³ Reaction of the lithium salt of [(2,6-diisopropylphenyl)NC(Me)]₂CH (Ar-nacnac) with (PPN)[FeI₂(NO)₂] (PPN⁺ = μ nitridobis(triphenylphosphine cation), affords the new compound, [(Ar-nacnac)Fe(NO)₂] (3), after workup. Unlike typical anionic DNICs such as 2, neutral 3 is soluble in alkane and arene solvents and may be conveniently freed from salt impurities by recrystallization from pentane. The spectroscopic features of 3 ($v_{NO} = 1761$, 1709 cm⁻¹, benzene- d_6) are similar to those of 2, consistent with an $\{Fe(NO)_2\}^9$ formulation (SI). The solid-state structure of 3 (Figure 1) displays pseudo-tetrahedral geometry at iron, with the dinitrosyliron unit residing in a steric pocket created by the β-diketiminate ligand. It is interesting to contrast the pseudo-tetrahedral structure of 3 with the square-planar geometry of a related Fe(I) dicarbonyl complex. ¹⁴ The Fe-N-O metrics in 3 are similar to those observed in previously characterized DNICs. ¹⁵

The $\{\text{Fe(NO)}_2\}^9$ unit of most DNICs can be reduced by one electron to give the diamagnetic $\{\text{Fe(NO)}_2\}^{10}$ configuration. Typically this reduction process is quasi-reversible electrochemically. Only in rare instances can the reduced DNIC be isolated. With DNICs containing thiolate ligands, reduction is followed by dissociation of RS⁻ leading to formation of the reduced Roussin red ester derivatives. The cyclic voltammogram of 3 (SI) displays a quasi-reversible $(i_{pr}/i_{pf}=0.95)$ one-electron reduction at -1.34 V (vs ferrocene/ferrocenium) in THF. Chemical reduction of 3 with either decamethylcobaltocene or sodium amalgam affords the reduced $\{\text{Fe(NO)}_2\}^{10}$ DNIC. Metathesis of the cobaltocenium cation for Bu₄N⁺ produces $(\text{Bu}_4\text{N})[(\text{Ar-nacnac})\text{Fe(NO)}_2]$ (4a) after crystallization from Et₂O. Compound 4a is a stable (mp = 198 – 200 °C) diamagnetic solid as judged by sharp features in its ¹ H NMR spectrum (SI). Exposure of 4a to air immediately results in regeneration of 3 as judged by IR and UV-vis spectroscopy. The v_{NO} bands in a benzene- d_6 solution of 4a appear at 1627 and 1567 cm⁻¹, a shift of >130 cm⁻¹ from those observed for 3. Crystals of 4a proved unsuitable for X-ray determination so the PPN⁺ salt, 4b, was prepared.

The solid-state structure of the anion of **4b** (Figure 2) resembles closely that of **3**. One considerable difference are the elongated Fe-N(nacnac) bonds in **4b**, which result in a smaller displacement of the Fe atom out of the nacnac plane compared with that in **3** (0.24 vs 0.62 Å). The Fe-N(NO) bonds for **4b** are also \sim 0.03 to 0.04 Å shorter than those of **3** consistent with a greater degree of π -backbonding in the reduced $\{Fe(NO)_2\}^{10}$ core. This phenomenon is also manifest by the longer N-O bonds of **4b** (Figs. 1 and 2).

In conclusion, reaction of the cluster analog 1 with NO leads to formation of a new nitrogen containing $\{Fe(NO)_2\}^9$ DNIC. This reaction demonstrates that naturally occurring Rieske clusters may be similarly susceptible to disassembly by NO, as found for cysteine-bound Fe_2S_2 clusters. The reaction of 1 with NO (g) to form two $\{Fe(NO)_2\}^9$ units follows directly from previous work with synthetic clusters of the type $[Fe_2S_2(SR)_4]^{2^-}$ and suggests an intrinsic reactivity of the $\{Fe_2S_2\}^{2+}$ core towards NO. 9,18 β -Diketiminates are excellent ligands for dinitrosyliron complexes, allowing for isolation of a rare set of homologous DNIC redox partners. Preliminary studies (data not shown) with Fe(TPP)Cl (TPP = mesotetraphenylporphine dianion) demonstrate that these DNICs display different NO transfer properties, suggesting that control of NO transfer in biology might be redox triggered. This hypothesis is currently under investigation in our laboratory.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

This work was supported by the NSF grant CHE-0611944. Z.J.T. acknowledges postodoctoral fellowship support from the NIH General Medical Sciences (1 F32 GM082031-02). The authors thank Dr. Daniela Bucella and Dr. Peter Müller for assistance with X-ray crystallography.

References

- 1. Beinert H, Holm RH, Münck E. Science 1997;277:653-659. [PubMed: 9235882]
- 2. Rieske JS. Biochim Biophys Acta, Rev Bioenerg 1976;456:195-247.
- 3. Iwata S, Saynovits M, Link TA, Michel H. Structure 1996;4:567–579. [PubMed: 8736555]
- 4. Gurbiel RJ, Batie CJ, Sivaraja M, True AE, Fee JA, Hoffman BM, Ballou DP. Biochemistry 1989;28:4861–4871. [PubMed: 2765515]
- 5. Ballmann J, Albers A, Demeshko S, Dechert S, Bill E, Bothe E, Ryde U, Meyer F. Angew Chem Int Ed Engl 2008;47:9537–9541. [PubMed: 18972470]
- 6. Welter R, Yu L, Yu CA. Arch Biochem Biophys 1996;331:9-14. [PubMed: 8660677]
- See for example: (a)Foster MW, Cowan JA. J Am Chem Soc 1999;121:4093–4100.4100(b)Ding H,
 Demple B. Proc Natl Acad Sci USA 2000;97:5146–5150.5150 [PubMed: 10805777]
- D'Addario S, Demartin F, Grossi L, Iapalucci MC, Laschi F, Longoni G, Zanello P. Inorg Chem 1993;32:1153–1160.
- Harrop TC, Tonzetich ZJ, Reisner E, Lippard SJ. J Am Chem Soc 2008;130:15602–15610. [PubMed: 18939795]
- 10. Enemark JH, Feltham RD. Coord Chem Rev 1974;13:339–406.
- 11. (a) Reginato N, McCrory CTC, Pervitsky D, Li L. J Am Chem Soc 1999;121:10217–10218. (b) Wang X, Sundberg EB, Li L, Kantardjieff KA, Herron SR, Lim M, Ford PC. Chem Commun 2005:477–479. (c) Tsai ML, Hsieh CH, Liaw WF. Inorg Chem 2007;46:5110–5117. [PubMed: 17444639] (d) Huang HW, Tsou CC, Kuo TS, Liaw WF. Inorg Chem 2008;47:2196–2204. [PubMed: 18271533]
- 12. Holland PL. Acc Chem Res 2008;41:905–914. [PubMed: 18646779]
- 13. Connelly NG, Gardner C. J Chem Soc, Dalton Trans 1976:1525–1527.
- 14. Sadique AR, Brennessel WW, Holland PL. Inorg Chem 2008;47:784–786. [PubMed: 18171059]

- 15. Sanina NA, Aldoshin SM. Russ Chem Bull 2004;53:2428–2448.
- 16. Atkinson FL, Blackwell HE, Brown NC, Connelly NG, Crossley JG, Orpen AG, Rieger AL, Rieger PH. J Chem Soc, Dalton Trans 1996:3491–3502.
- 17. (a) Tsou CC, Lin ZS, Lu TT, Liaw WF. J Am Chem Soc 2008;130:17154–17160. [PubMed: 19053409] (b) Lu TT, Tsou CC, Huang HW, Hsu IJ, Chen JM, Kuo TS, Wang Y, Liaw WF. Inorg Chem 2008;47:6040–6050. [PubMed: 18517190]
- 18. Tsai FT, Chiou SJ, Tsai MC, Tsai ML, Huang HW, Chiang MH, Liaw WF. Inorg Chem 2005;44:5872–5881. [PubMed: 16060642]

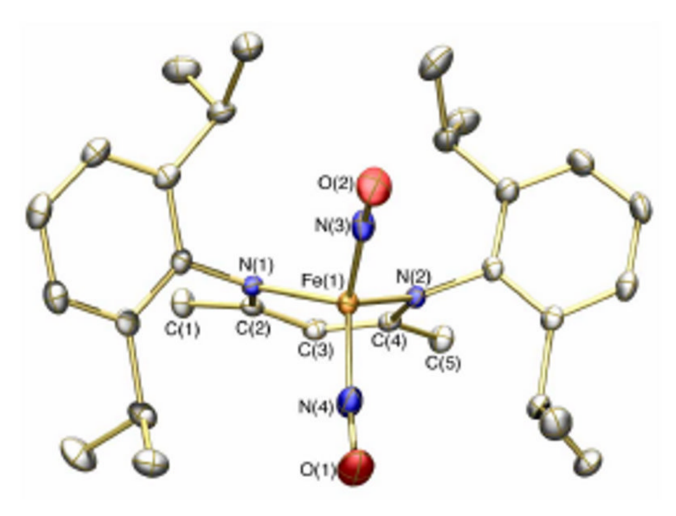


Figure 1. Structure of **3** displaying 50% thermal ellipsoids with hydrogen atoms omitted for clarity. Selected bond lengths (Å) and angles (deg): Fe(1)-N(3), 1.6964(18); Fe(1)-N(4), 1.6882(18); N(3)-O(2), 1.177(2); N(4)-O(1), 1.174(2); N(3)-Fe(1)-N(4), 114.27(9); Fe(1)-N(3)-O(2), 162.7(2); Fe(1)-N(4)-O(1), 170.1(2).

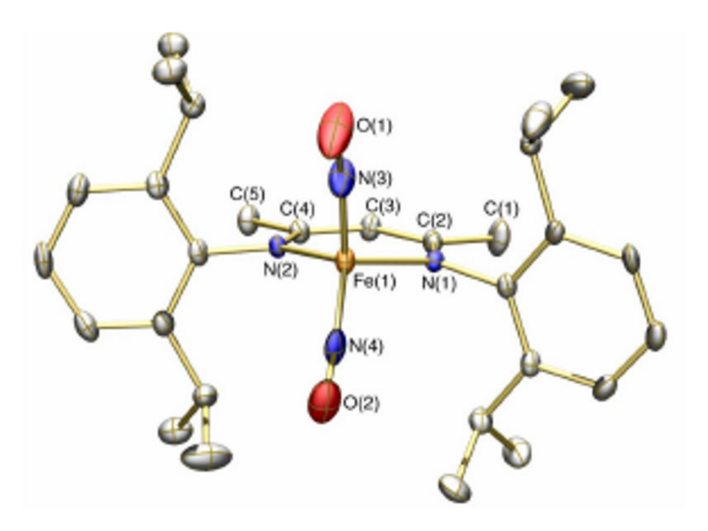


Figure 2. Structure of the anion of **4b** displaying 50% thermal ellipsoids with hydrogen atoms omitted for clarity. Selected bond lengths (Å) and angles (deg): Fe(1)-N(3), 1.668(5); Fe(1)-N(4), 1.649 (4); N(3)-O(1), 1.191(6); N(4)-O(2), 1.218(6); N(3)-Fe(1)-N(4), 109.2(2); Fe(1)-N(3)-O(1), 163.2(5); Fe(1)-N(4)-O(2), 165.1(5).

Scheme 1. Reaction of the Rieske-type cluster with NO.