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A mononuclear nonheme {FeNO}⁶ complex: synthesis and structural and spectroscopic characterization†

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While the synthesis and characterization of $\{FeNO\}^{7.8.9}$ complexes have been well documented in heme and nonheme iron models, $\{FeNO\}^6$ complexes have been less clearly understood. Herein, we report the synthesis and structural and spectroscopic characterization of mononuclear nonheme $\{FeNO\}^6$ and iron(III)—nitrito complexes bearing a tetraamido macrocyclic ligand (TAML), such as $[(TAML)Fe^{III}(NO)]^-$ and $[(TAML)Fe^{III}(NO)]^2$, respectively. First, direct addition of $NO_{(g)}$ to $[Fe^{III}(TAML)]^-$ results in the formation of $[(TAML)Fe^{III}(NO)]^-$, which is sensitive to moisture and air. The spectroscopic data of $[(TAML)Fe^{III}(NO)]^-$, such as 1H nuclear magnetic resonance and X-ray absorption spectroscopies, combined with computational study suggest the neutral nature of nitric oxide with a diamagnetic Fe center (S=0). We also provide alternative pathways for the generation of $[(TAML)Fe^{III}(NO)]^-$, such as the iron–nitrite reduction triggered by protonation in the presence of ferrocene, which acts as an electron donor, and the photochemical iron–nitrite reduction. To the best of our knowledge, the present study reports the first photochemical nitrite reduction in nonheme iron models.

Introduction

Nitrogen oxides (NO_x) , including nitric oxide (NO), nitrite anion (NO_2^-) , nitrogen dioxide (NO_2) and nitrous oxide (N_2O) , are important in environmental and atmospheric chemistries, and play critical roles in biological settings. Nitric oxide is well established in cellular signaling and it is a powerful vasodilator; ¹⁻³ NO is also intimately involved in the physiological

immune response, for example, by reacting with superoxide anion (O2. -) to generate the toxic oxidizing/nitrating species peroxynitrite (-OON=O) and/or NO2, which aid the fight against invading pathogens.4,5 Thus, the tight control of NO concentrations (i.e., homeostasis) is important and (i) it may be stored in the form of a circulating cellular nitrite pool (which can undergo reduction-protonation to give back NO),5,6 or conversely (ii) NO may be removed and transformed to innocuous nitrogen oxide forms. For the latter, hemo- or myoglobin type proteins oxidize NO to nitrate anion, in the so-called NO dioxygenase (NOD) activity.3,7,8 Bacterial nitric oxide reductases (NOR's)3,9 reductively couple two NO molecules to give N2O, and in fungi, a heme protein cytochrome P450 NOR (P450nor) effects the same reaction.3,10 As is clear from these discussions, transition metal ions are associated with NOx interactions and enzymatic reactions, for heme, and non-heme iron and copper proteins.2,11

Biomimetic synthetic models offer an attractive platform to understand physical and chemical properties of the Fe-NO moiety. For example, Ford and co-workers have reported seminal works on the characterization of Fe-NO intermediates and nitrite reduction mediated by heme models. Lehnert has demonstrated the synthesis and characterization of Fe-NO complexes in heme and nonheme models. Mascharak has synthesized and characterized non-heme complexes with a photolabile Fe-NO bond, including [(PaPy₃)Fe^{III}(NO)]²⁺. Mascharak has synthesized and characterized non-heme complexes with a photolabile Fe-NO bond, including [(PaPy₃)Fe^{III}(NO)]²⁺. Mascharak has synthesized and characterized non-heme complexes with a photolabile Fe-NO bond, including [(PaPy₃)Fe^{III}(NO)]²⁺. Mascharak has synthesized and characterized non-heme complexes with a photolabile Fe-NO bond, including [(PaPy₃)Fe^{III}(NO)]²⁺. Mascharak has synthesized and characterized non-heme complexes with a photolabile Fe-NO bond, including [(PaPy₃)Fe^{III}(NO)]²⁺. Mascharak has synthesized and characterized non-heme complexes with a photolabile Fe-NO bond, including [(PaPy₃)Fe^{III}(NO)]²⁺. Mascharak has synthesized and characterized non-heme complexes with a photolabile Fe-NO bond, including [(PaPy₃)Fe^{III}(NO)]²⁺. Mascharak has synthesized and characterized non-heme complexes with a photolabile Fe-NO bond.

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[†] Electronic supplementary information (ESI) available: Tables S1–S6 for crystallographic data, Mössbauer parameters, XAS data and computational geometric parameters and Fig. S1–S11. CCDC 1574430, 1574431. For ESI and crystallographic data in CIF or other electronic format see DOI: 10.1039/c8sc01962b

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i. protonation

NO2

ii. S-/P-triggered OAT

Fe-nitrito or
-nitro species

iii. photoirradiation

Scheme 1 Schematic illustration showing different pathways of nitrite reduction.

Recently, Pluth and Lippard reported that an Fe(III) complex with a tetraamido macrocycle TAML ligand, [Fe^{III}(TAML)]^{-,15} where the TAML scaffold is known to provide remarkable stability to higher oxidation states of iron,¹⁶ promotes NO binding and release, stated to occur without formal oxidation state changes at iron.

Instead of using NO, it has been well documented that there are two different pathways for nitrite (NO₂⁻) reduction that lead to the formation of Fe-NO complexes: addition of (i) proton resulting in the electrophilic attack of two protons on the O atom of Fe-nitrite complexes (Scheme 1, pathway i) or (ii) triphenylphosphine(PPh3)/thioether-/thio-triggered oxygen atom transfer (OAT) reactions (Scheme 1, pathway ii).12,17 However, photochemical generation of NO from Fe-bound nitrite has yet to be explored. Herein, we describe more elaborated NOx chemistry starting with the same compound [Fe^{III}(TAML)]⁻. The new aspects and advances presented here include (a) full characterization of a Fe(III)-nitrosyl complex including the X-ray crystal structure and (b) chemical and photochemical generation of the same Fe(III)-nitrosyl complex from a Fe(III)-nitrito complex (Scheme 1, pathway i and iii). Indeed, previously known Fe-NO adducts have {Fe-NO}⁷ or {Fe-NO}8 descriptions in the Enemark-Feltham notation,18 but [(TAML)Fe^{III}(NO)]⁻ (1) is a {Fe-NO}⁶ compound, like [(PaPy₃) Fe^{III}(NO)]²⁺.14

Results and discussion

Complex 1 was generated as described by Pluth and Lippard¹⁵ (see the Experimental section and Fig. S1-S4, ESI†); 1 was prepared in situ at -40 °C and excess NO(g) was removed by vacuum/argon-purge cycles. A UV-vis spectrum of 1 generated in CH₃CN at -40 °C displayed similar features with λ_{max} at 525 nm $(\varepsilon = 4500 \text{ M}^{-1} \text{ cm}^{-1}) \text{ and } 880 \text{ nm } (\varepsilon = 8000 \text{ M}^{-1} \text{ cm}^{-1}),$ respectively (Fig. S2a, ESI†). In contrast to what was found for 1 generated in CH₃OH, ¹⁴ a ¹H NMR spectrum of isolated 1 exhibited ligand peaks in the range of 0-10 ppm, and an Evans method experiment showed that there is no shift of the TMS or solvent peak. These results suggest that the complex is diamagnetic with a low-spin (S = 0) state (Fig. S3, ESI†). A cold spray ionization time-of-flight mass (CSI MS) spectrum of 1 in negative mode exhibited an ion peak at m/z of 456.1, whose mass and isotopic distribution patterns correspond to [(TAML) Fe(NO)]⁻ (calculated m/z of 456.1) (Fig. S4, ESI†). The ion peak

of 1 shifted one-mass unit upon ¹⁵NO-substitution, confirming that 1 contains one exchangeable nitrogen atom in it. Fortunately, recrystallization of 1 from CH3CN/Et2O under an Ar atmosphere afforded single crystals suitable for X-ray diffraction; the crystallographic analysis revealed average Fe1-N_{TAML}, Fe1-N5, and N5-O1 distances of 1.8843(12), 1.6131(13), and 1.1667(19) Å, respectively (Fig. 1a). The crystal structure of 1 also exhibits a linear Fe-N-O angle of 172.05(14)°, which is consistent with S = 0 ground state. It is noteworthy that while {FeNO}⁶ with a typical non-heme ligand such as TMG_3 tren exhibited S =1 ground states with a linear Fe-N-O angles of 180°, 13d {FeNO}⁶ with heme ligands displayed Fe-N-O angles ranging from 159.6(8)° to 178.3(3)°. 19 Since the TAML ligand exhibits strong π -donation that isn't seen in typical non-heme ligands, there are some geometric and electronic structural similarities between iron-nitrosyl complexes bearing TAML and heme ligands.

Zero-field Mössbauer data for 1 measured at 77 K are given in Fig. 2. Fitting of these data shows a single species (100%) with $\delta = -0.19 \ \mathrm{mm \ s^{-1}}$ and $|\Delta E_{\mathrm{Q}}| = 3.29 \ \mathrm{mm \ s^{-1}}$, in agreement with previously reported parameters for 1 ($\delta = -0.20 \text{ mm s}^{-1}$ and $|\Delta E_{\rm O}| = 3.29 \; {\rm mm \; s^{-1}}$). It should be noted that, in the previous study, the nitrosyl species only accounted for 36% of the Mössbauer spectrum. These parameters are similar to a lowspin, six-coordinate {FeNO}⁶ compound with a macrocyclic tetracarbene ligand, [(NHCL)FeNO(ONO)](OTf)₂, which has $\delta = -0.16 \text{ mm s}^{-1}$ and $|\Delta E_{\rm Q}| = 3.12 \text{ mm s}^{-1}$ (Table S4, ESI†).²⁰ They are also significantly different from the parameters obtained for a different low-spin, six-coordinate {FeNO}⁶ complex, [(PaPy₃) $[Fe^{III}(NO)]^{2+}$, which has $\delta = -0.05$ mm s⁻¹ and $|\Delta E_O| =$ 0.85 mm s⁻¹.14 Previous studies on other TAML compounds have shown that they have more negative isomer shifts for a given oxidation state, such as in the case of [(TAML)Fe^{IV}O]²⁻ $(-0.19)^{16b}$ compared to $[(N4Py)Fe^{IV}O]^{2+}(-0.04)^{2}$. This is due to the strong equatorial donation of the TAML ligand into the 4s orbital,22 and can account for the difference between the TAML and PaPy3 {FeNO}6 compounds. The tetracarbene ligand is a good equatorial donor, like TAML, which gives it similar Mössbauer parameters.

EXAFS data for **1** are shown in Fig. 3a. The best fit to the EXAFS is consistent with the crystal structure, with a short 1.62 Å Fe-NO bond length and an intense Fe-N-O multiple

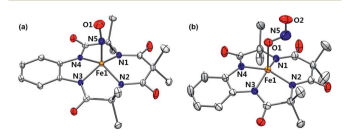


Fig. 1 X-ray crystal structures of (a) 1 and (b) 2 showing 50% probability ellipsoids. Cations such as tetraphenylphosphonium, tetrabutylammonium, hydrogen atoms and ether molecules are omitted for clarity. Colors for atoms: Fe, orange; N, blue; O, red; C, white (see Tables S1 and S2, ESI†).

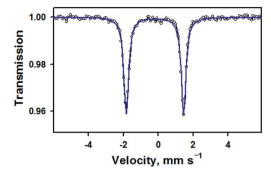


Fig. 2 Zero-field Mössbauer spectrum (black opened circles) and fit (blue line) of 1 measured at 77 K. The isomer shift of $-0.19~\rm mm~s^{-1}$ and quadrupole splitting of 3.29 mm s⁻¹ are in good agreement with parameters previously reported for this complex in ref. 14 (also see Table S3, ESI†).

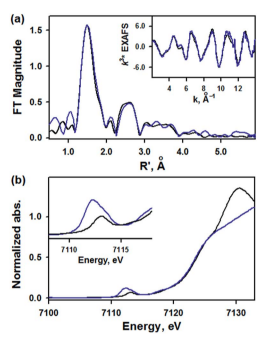


Fig. 3 (a) Fourier transform and EXAFS (inset) of 1 (blue line) with fits (black line). (b) XANES data of 1 (blue line) and $[(PaPy_3)Fe^{III}(NO)]^{2+}$ (black line) for comparison. Inset shows the expansion of the pre-edge region.

scattering path at 3.03 Å, consistent with a nearly linear Fe–N–O (Table S5, ESI†). There are also four N atoms at 1.89 Å and eight C atoms at 2.93 Å, consistent with the low-Z atoms of the TAML ring (full fit and components shown in Fig. S5, ESI†). The XANES data of 1 are shown in blue in Fig. 3b with the XANES data for $[(PaPy_3)Fe^{III}(NO)]^{2+}$ in black. The two compounds have the same rising edge energy, indicating they have the same $Z_{\rm eff}$. This is consistent with the difference in isomer shift being due to equatorial ligand donation. The pre-edge for 1 has a total intensity of 27 units, and is resolved into two peaks at 7112.1 eV (16 units) and 7113.3 eV (11 units), while the pre-edge of $[(PaPy_3)Fe^{III}(NO)]^{2+}$ has one peak at 7113.0 eV (9 units) (Table S6, ESI†). The increased pre-edge intensity of 1 is consistent with 4p

mixing into the 3d orbitals due to the loss of centrosymmetry in a 5C complex.

To understand the electronic structure of 1, DFT calculations were performed. The BP86 functional correctly reproduced the S = 0 ground state with good geometrical agreement with the crystal and EXAFS structure (Table S7, ESI†). The calculations show an electronic structure for 1 very similar to [(PaPy₃) Fe^{III}(NO)]²⁺, which was previously studied through L-edge XAS and DFT calculations.23 The L-edge XAS showed that [(PaPy3) Fe^{III}(NO)]²⁺ has backbonding equivalent to donation of an electron from the Fe to the NO for a total unoccupied metal 3d character consistent with Fe^{III}, but with unpolarized bonding to give an S = 0 Fe^{III}-NO (neutral) electronic structure, rather than antiferromagnetically coupled Fe^{III} (S = 1/2)-NO $^{\bullet}$ (S = -1/2). The unpolarized Fe^{III}-NO (neutral) description is equivalent to the "FeII-NO+ with strong backbonding" description used in previous work.^{24 α} DFT gives two (α and β) unoccupied d_{z²} and $d_{x^2-y^2}$ σ^* -bonding orbitals, and two unoccupied $\pi_y^* - d_{yz}$ and $\pi_x^* - d_{xz} \pi^*$ -bonding orbitals. While the calculated MO's are qualitatively similar to an Fe^{II}-NO⁺ electronic structure, quantitative analysis of the metal d character in the unoccupied π^* bonding orbitals (130% in 1, 112% in [(PaPy₃)Fe^{III}(NO)]²⁺) shows extensive donation from the Fe to the NO. The strong backbonding gives a Z_{eff} that is quantitatively Fe^{III}, as additionally corroborated by vibrational spectroscopy and S K-edge XAS.24 The FeIII-NO(neutral) description decouples the oxidation state from polarization to provide a more complete description. While the calculations show that 1 (338%) has a similar amount of unoccupied 3d character to [(PaPy₃) $\text{Fe}^{\text{III}}(\text{NO})$]²⁺ (342%), the calculated Mössbauer δ value agrees with the experiment in 1 being more negative than [(PaPy₃) Fe^{III}(NO)]²⁺. This results from some quantitative differences in the bonding. The TAML ligand in 1 is a much stronger σ donor into the $d_{x^2-y^2}$ orbital than the PaPy₃ ligand (55% vs. 64% 3d character), leading to greater backbonding in 1 compared to $[(PaPy_3)Fe^{III}(NO)]^{2+}$ (33% vs. 28%). The calculations also show that the d_{z^2} orbital lowers in energy going from 6C to 5C, which explains the difference in pre-edge shape (Fig. S6, ESI† for full DFT comparison). Combining the XAS, Mössbauer, and DFT analysis, the electronic structure of 1 is assigned as unpolarized Fe^{III}-NO (neutral), similar to the prior description of [(PaPy₃) Fe^{III}(NO)]²⁺ (vide supra).

Addition of a nitrite source such as sodium nitrite (NaNO₂) or tetrabutylammonium nitrite (Bu₄NNO₂) to a solution of [Fe^{III}(TAML)]⁻ in CH₃CN at -40 °C resulted in the formation of [(TAML)Fe^{III}(NO₂)]²⁻ (2), and this transformation could be easily noted by the blue-shift of the electronic absorption band from 400 nm due to 1 to 365 nm ($\varepsilon = 7500 \,\mathrm{M}^{-1} \,\mathrm{cm}^{-1}$) due to 2 (Fig. 4a). The X-band EPR spectrum showed signals identical to those found for [Fe^{III}(TAML)]⁻, at g = 3.9 and 2.0, consistent with the presence of an intermediate spin state (S = 3/2) (Fig. 4b). The CSI MS spectrum of 2 in negative mode exhibits a peak at m/z of 495.0, whose mass and distribution patterns correspond to the formulation of [(TAML)Fe(¹⁴NO₂)(Na)]⁻ (calculated m/z of 495.1) (Fig. S7, ESI†). When the reaction was carried out with isotopically ¹⁵N-labeled Na¹⁵NO₂, a mass peak corresponding to [(TAML)Fe(¹⁵NO₂)(Na)]⁻ (calculated m/z of

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495.0 496.0 Absorbance 495 500 495 500 m/z 400 600 800 1000 Wavelength, nm (b) 2.0 1000 2000 3000 4000

Fig. 4 (a) UV-vis spectra of $[Fe^{III}(TAML)]^-$ (0.10 mM, black line) and $[(TAML)Fe^{III}(NO_2)]^{2-}$ (0.10 mM, red line) under an inert atmosphere in CH₃CN at $-40\,^{\circ}$ C. Insets show the isotopic distribution patterns of the peaks at m/z of 495.0 for $2^{-14}NO_2$ (left panel) and m/z of 496.0 for $2^{-15}NO_2$ (right panel), which were generated by $Na^{14}NO_2$ and $Na^{15}NO_2$, respectively. (b) X-band EPR spectrum of 2 (1.0 mM) recorded in CH₃CN at 5 K.

Magnetic Field, G

496.1) appeared at m/z of 496.0, indicating that 2 contains one exchangeable N atom (Fig. 4a, inset). The NO₂⁻ binding to the iron(III)-TAML center is strong; variable temperature UV-vis experiments from -40 to 20 °C (Fig. S8, ESI†) indicate that the binding constant ($K_{\rm f}$) of NO₂⁻ to the Fe(III) center was 8.0(5) \times 10³ at 20 °C, whereas $K_{\rm f}$ at -40 °C was determined to be 6.0(5) \times 10⁷.

Interestingly, the X-ray crystal structure of 2 revealed an Obinding Fe(III)–nitrito complex (Fe– κ 1–ONO), in a square pyramidal geometry arising from the TAML ligand and monodentate Fe–ONO moiety (Fig. 1b). O-binding of the nitrite ligand is biologically relevant because the metal–O–nitrito binding fashion has been observed in the reaction of metmyoglobin and nitrite. ²⁵ In this nitrite complex with [Fe^{III}(TAML)]⁻, the Fe1–O1 bond distance is 2.068(4) Å, which is longer than that of Mb^{III}(ONO⁻) (1.94 Å). The angles formed from Fe1–O1–N5 of 122.4(4)° and O1–N5–O2 of 116.6(5)° are slightly larger than those for Mb^{III}(ONO⁻) (116° and 113°, respectively). ²⁵

Of importance, treatment of a solution of 2 with 2.4 equiv. of CF_3SO_3H in the presence of 1.2 equiv. of ferrocene leads to the immediate formation of 1 with over 90% yield on the basis of the extinction coefficient of isolated 1 (Fig. S9, ESI†). This result clearly showed that the nitrite reduction of 2 successfully occurred to produce NO. To rule out a possibility of the free nitrous acid (HNO₂) disproportionation that yields NO₂ and NO, we carried out control reactions by (1) adding the starting $[Fe(TAML)]^-$ complex into the Ar-saturated CH_3CN solution

containing NaNO2, ferrocene, and triflic acid and (2) a solution of NaNO2 and triflic acid was added to an Ar-saturated CH3CN solution of [Fe(TAML)]-. In these control reactions, we did not observe the formation of 1, suggesting that 1 was formed directly from the iron(III)-nitrito complex, 2, upon protonation (Fig. S10, ESI[†]). Instead of the protonation, we also endeavored to carry out (i) the thioether-/thio-triggered oxygen atom transfer (OAT) and (ii) triphenylphosphine (PPh3)-triggered OAT in the hope of finding different pathways of nitroxylation proposed by other groups; 12,13,17 unfortunately, those attempts did not lead to the formation of 1. More interestingly, preliminary results on the photoirradiation at 420 nm of a solution containing 2 (0.10 mM) in acetone under an Ar atmosphere afforded the formation of 1 within 3.5 h with over 90% yield (Fig. 5). The formation of 1 was not observed in the absence of visiblelight irradiation, indicating that the photochemical generation of 1 from 2 occurred. To investigate the photochemical transformation of free nitrite resulting in NO, we carried out a control experiment, in which we first photolyzed an Arsaturated acetone solution containing NaNO2, followed by adding the starting [Fe(TAML)] complex. However, it didn't afford the formation of 1, indicating that the photochemical transformation of 2 to 1 didn't involve the free nitrite photolysis (Fig. S11, ESI†). An X-band EPR spectrum and CSI MS of the product confirmed that 1 was successfully generated (Fig. S12, ESI†). Thus, the nitrite reduction could be triggered by adding proton or by photoirradiating 2.

Since 2 would unambiguously undergo N–O bond cleavage to yield 1 under photochemical conditions, we then pursued the fate of oxygen transfer reactions from the initial Fe(III)–nitrite complex by adding oxygen trapping substrates, such as PPh₃ and thioanisole. Otherwise, it is difficult to trap the dissociated oxygen atom in the reaction solution without adding substrates. Product analysis performed after the completion of the photochemical nitrite reduction of 2 in the presence of thioanisole or PPh₃ revealed the formation of methyl phenyl sulfoxide (~40% yield) and triphenylphosphine oxide (~45%), respectively. It has been suggested that in the case of manganese and iron porphyrin complexes, the photo-triggered oxygen atom transfer occurred from the O-bound nitrito complexes to substrates such as PPh₃, resulting in the formation of the corresponding NO complexes.²⁶ In

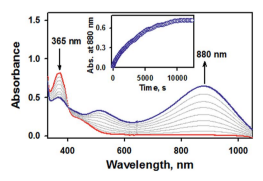


Fig. 5 UV-vis spectral changes observed in the photoirradiation reaction of 2 (0.10 mM, red line) under an inert atmosphere in acetone at $-40~^{\circ}\text{C}$.

the present case, however, the reaction rate was not affected by the nature of substrates or the concentration of substrates, consistent with the fact that the PPh_3 /thio-triggered nitrite reduction did not occur unless photoreduction of 2 does trigger oxo transfer.

Conclusions

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As demonstrated in this report (see Scheme 2), we were able to characterize novel Fe–NO₂, and Fe–NO complexes bearing TAML by using various spectroscopic methods, including UV-vis, Mössbauer, XAS, NMR, CSI MS, and IR spectroscopies, and X-ray crystallography. We also established a photochemical nitrite reduction of an Fe(III)–nitrito complex that produced a nearly quantitative yield of the corresponding Fe(III)–nitrosyl complex. Since there is a constant challenge regarding NO production and/or release related to drug delivery, the present work provides an interesting insight into producing NO from nitrite under photoirradiation. Future work will be focused on the further reactivity of Fe–NO intermediates in the formation of ammonia via hydroxylamine in the presence of protons.

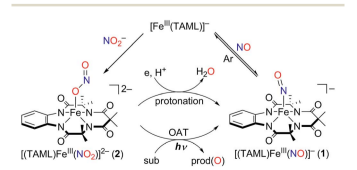
Experimental section

Materials

All chemicals, obtained from Aldrich Chemical Co. and Tokyo Chemical Industry, were the best available purity and used without further purification unless otherwise indicated. Solvents were dried according to published procedures and distilled under argon prior to use. The Normalian (>99% 15N-enriched) was purchased from ICON Services Inc. (Summit, NJ, USA) and used without further purification. Na[Fe^{III}(TAML)] and TAML-H4 (TAML = tetraamidomacrocyclic ligand, 3,3,6,6,9,9-hexamethyl-2,5,7,10-tetraoxo-3,5,6,7,9,10-hexahydro-2*H*-benzo[*e*][1,4,7,10]tetraazacyclo-tridecine-1,4,8,11-tetraide) were purchased from GreenOx Catalyst, Inc. (Pittsburgh, PA, USA), and the Na[Fe^{III}-(TAML)] complex was recrystallized from an isopropanol/H2O mixture for further use. The Parameter of the literature method. The Na Parameter of the literature method.

Instrumentation

UV-vis spectra were recorded on a Hewlett Packard 8453 diode array spectrophotometer equipped with a UNISOKU Scientific



Scheme 2 Schematic representation showing the generation of $[(TAML)Fe^{III}(NO)]^-$.

Instruments Cryostat USP-203A for low-temperature experiments. Cold spray ionization time-of-flight mass (CSI MS) spectra were collected on a JMS-T100CS (JEOL) mass spectrometer equipped with the CSI source. Typical measurement conditions are as follows: needle voltage: 2.2 kV, orifice 1 current: 50-500 nA, orifice 1 voltage: 0 to 20 V, ring lens voltage: 10 V, ion source temperature: 5 °C, spray temperature: -40 °C. The CSI MS spectra of [(TAML)Fe^{III}(NO)] and [(TAML) Fe^{III}(NO₂)]²⁻ were observed by directly infusing the reaction solution into the ion source through a pre-cooled tube under high N₂ gas pressure. X-band CW-EPR spectra were taken at 5 K using a X-band Bruker EMX-plus spectrometer equipped with a dual mode cavity (ER 4116DM). Low temperatures were achieved and controlled with an Oxford Instruments ESR900 liquid He quartz cryostat with an Oxford Instruments ITC503 temperature and gas flow controller. The experimental parameters for EPR spectra were as follows: microwave frequency = 9.647 GHz, microwave power = 1.0 mW, modulation amplitude = 10 G, gain = 1×10^4 , modulation frequency = 100 kHz, time constant = 40.96 ms and conversion time = 81.00 ms. Low temperature liquid IR spectra were recorded on a Remspec 626 FT-IR spectrometer. IR spectrum analysis was performed in the frequency range of 1000-5000 cm⁻¹ at a resolution of 4 cm⁻¹. Product analysis was performed using high performance liquid chromatography (HPLC, Waters 515). Quantitative analyses were made on the basis of comparison of HPLC peak integration between products and authentic samples. ¹H NMR spectra were measured with a Bruker model digital AVANCE III 400 FT-NMR spectrometer. All NMR spectra were recorded in CD₃CN at -40 °C.

Purification of nitric oxide

Nitric oxide gas (NO(g)) was obtained from Dong-A Specialty Gases and purified as follows (see Fig. S1†): NO(g) was first passed through two columns filled with NaOH beads and molecular sieves to remove higher nitrogen oxide and moisture impurities and collected in the frozen form in a first trap cooled at 77 K with liquid N2. Further purification was performed by the distillation of frozen NO(g) (as crystalline N2O2), which means that frozen NO_(g) was warmed up to 193 K (acetone/dry-ice mixture, 193 K) and then the sample was collected in a second trap cooled at 77 K with liquid N₂. This second trap was again warmed up to 193 K and then, the highly purified NO(g) was collected in another Schlenk flask fitted with a rubber septum (free from oxygen; after several cycles of vacuum and Ar purging). The NO(g) should be at a very high pressure in the Schlenk flask (>1 atm; the septum bulges outward due to high pressures inside the Schlenk flask). Then, NO(g) was purged into the distilled and degassed CH3CN (15 mL) in a Schlenk flask, which was already connected with an oil bubbler. NO(g) purging with vigorous stirring under 1 atm for 20 min was required to make NO-saturated CH3CN solution. The concentration of NO in CH3CN saturated solution at 293 K was approximated to be 14 mM.29

Synthesis of [(TAML)Fe(NO)] (1)

The starting complex, $[Fe^{III}(TAML)]^{-}(1)$ was bubbled with Ar for 20 min in CH₃CN at -40 °C.¹⁵ To this solution, excess NO_(g)

(~10 cm³) was injected into the head-space and stirred for

10 min as the color changed from orange to deep purple. ¹⁵Nincorporated {FeNO}⁶ complex was generated using ¹⁵NO_(g) under the identical reaction conditions. Solution IR data of both 1-NO (5.0 mM) and 1-15NO (5.0 mM) were obtained by reacting 1 with NO(g) and 15NO(g), respectively.

Synthesis of $[(TAML)Fe^{III}(NO_2)]^{2-}$ (2)

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The Fe(III)-nitrite intermediate, $[(TAML)Fe^{III}(NO_2)]^{2-}$ (2), was generated either by adding 5.0 equiv. of sodium nitrite, NaNO₂ in the presence of 10 equiv. of 15-crown ether (15C5) or by adding 5.0 equiv. of tetrabutylammonium nitrite (Bu₄NNO₂) in CH₃CN at room temperature. The solution color changed from orange to yellow. Slow diffusion of diethylether into the reaction solution allowed us to obtain single crystals. ¹⁵N-incorporated 2 was synthesized by using Na15NO2 under identical reaction conditions.

X-ray structural analysis

To improve the quality of the single crystals of 1, the cation exchange from Na⁺ to PPh₄⁺ was performed by introducing PPh₄Cl into the solution containing Na[Fe^{III}(TAML)]. Single crystals of {FeNO}⁶, (PPh₄)[(TAML)Fe(NO)], and Fe(III)-nitrite complexes, (Bu₄N)₂[(TAML)Fe^{III}(NO₂)], suitable for X-ray analysis were obtained by slow diffusion of Et₂O into the CH₃CN solution of 1 and 2, respectively. These crystals were taken from the solution using a nylon loop (Hampton Research Co.) on a handmade copper plate and mounted on a goniometer head in a N2 cryostream. The diffraction data for both 1 and 2 were collected at 100 K, on a Bruker SMART AXS diffractometer equipped with a monochromator in the Mo K α ($\lambda = 0.71073 \text{ Å}$) incident beam. The CCD data were integrated and scaled using the Bruker-SAINT software package, and the structure was solved and refined using SHELXTL V 6.12.30 Hydrogen atoms were located in the calculated positions. In the structure, the crystal lattice contains some amounts of disordered solvent molecules (most likely to be one diethyl ether lattice molecule per asymmetric unit whose occupancy factors are possibly lower than 1). The contribution of the lattice solvent molecule has been taken out using the program SQUEEZE (Spek, 2003) for the final refinement.31 All details of the SQUEEZE refinement are provided in the CIF files. CCDC 1574430 for 1 and CCDC 1574431 for 2 contain the supplementary crystallographic data for 1 and 2.†

Spectral titration for the binding constant (K_f)

The binding constant of nitrite ion to [Fe^{III}(TAML)]- was examined by varying the temperature of the CH₃CN solution containing the isolated Fe(III)-nitrite intermediate, 2. Typically, the temperature of a CH₃CN solution of 2 varied from −40 °C to 20 °C. The concentration of [Fe^{III}(TAML)]⁻ was determined from the absorption band at $\lambda = 400 \text{ nm} (\varepsilon = 8000 \text{ M}^{-1} \text{ cm}^{-1})$ and that of 2 from the absorption band at $\lambda = 365$ nm ($\varepsilon =$ $7500 \text{ M}^{-1} \text{ cm}^{-1}$).

$$\left[Fe^{III}(TAML)\right]^{-} + NO_{2}^{-} \stackrel{\textit{K}_{f}}{\rightleftharpoons} \left[(TAML)Fe^{III}(NO_{2})\right]^{2-}$$

The binding constant (K_f) is expressed by the equation below:

$$K_{\rm f} = [2]/([[{\rm Fe}^{\rm III}({\rm TAML})]^{-}] \times [{\rm NO}_{2}^{-}])$$
 (1)

Finally, the K_f values of $8.0(5) \times 10^3$ at 20 °C and $6.0(5) \times 10^7$ at -40 °C were determined from UV spectral changes with respect to the temperature (ESI, Fig. S8†).

Solution IR measurements

Solution IR signals for samples were recorded on an IR instrument (Remspec #: 626) having a very sensitive probe, attached to a sophisticated setup under an Ar atmosphere in CH₃CN at -40 °C. Solution IR data of both 1-NO (5.0 mM) and 1^{-15} NO (5.0 mM) were obtained by reacting 1 with excess of NO(g) and ¹⁵NO_(g), respectively.

Spin state measurements

The effective magnetic moment ($\mu_{\rm eff}$, BM) of the Fe^{III} complex 1 was determined using the modified ¹H NMR method of Evans at -40 °C.³²⁻³⁴ A WILMAD® coaxial insert (sealed capillary) tube containing the blank acetonitrile-d3 solvent only (with 1.0% tetramethylsilane (TMS)) was inserted into the normal NMR tube containing the complexes (1.0 mM) dissolved in acetonitrile-d₃ (with 0.10% TMS). The chemical shift of the TMS peak (and/or solvent peak) in the presence of the paramagnetic metal complexes was compared to that of the TMS peak (and/or solvent peak) in the inner coaxial insert tube. The effective magnetic moment was calculated using the following equation:

$$\mu = 0.0618(\Delta \nu T/2fM)^{1/2}$$

where f is the oscillator frequency (MHz) of the superconducting spectrometer, T is the absolute temperature, M is the molar concentration of the metal ion, and $\Delta \nu$ is the difference in frequency (Hz) between the two reference signals.32-34 The absence of shifts of the TMS peak for 1 indicated that both intermediates possess S = 0 spin state in CD₃CN solution.

X-ray absorption spectroscopy

[(TAML)Fe(NO)] XAS samples were prepared and frozen under liquid nitrogen in 2×10 mm pinhole-type Delrin cells wrapped with Kapton tape. The [(PaPy₃)Fe(NO)]²⁺ XAS sample was prepared by diluting 12.5 mg solid [(PaPy₃)Fe(NO)]²⁺ with 67.5 mg solid BN, as calculated by the program SAMPLE4 as part of the program suite EXAFSPAK,35,36 and loaded into a 2 mm thick aluminum sample holder wrapped with Kapton tape under a N2 atmosphere in a glove bag in a dark room. All XAS data were collected at beamline 7-3 (ref. 37) at the Stanford Synchrotron Radiation Lightsource (SSRL) and the samples were maintained at 10 K using an Oxford Instruments CF1208 continuous flow liquid helium cryostat. The solution samples were oriented at 45° to the incident X-ray beam and measured under ring operating conditions of 500 mA over an energy range

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of 6785–7876 eV ($k = 14 \text{ Å}^{-1}$). Fluorescence data were collected at 90° to the incident beam using a solid-state 30-element Ge detector array with Soller slits and a 6 wavelength Mn filter aligned between the Soller slits and the sample to improve the Fe Kα fluorescence signal intensity relative to that of the scattered beam.38 The solid samples were oriented at 0° to the incident X-ray beam and measured in transmission mode with N₂-filled ionization chambers under ring operating conditions of 350 mA over an energy range of 6785–8365 eV $(k = 18 \text{ Å}^{-1})$. An internal calibration was utilized with the first inflection point of an Fe foil set to 7111.2 eV.39 During measurement, the data in the Fe K-edge, K pre-edge, and extended X-ray absorption finestructure (EXAFS) regions were continuously monitored in order to ensure sample integrity by comparing each individual scan to the ones taken previously and no photodamage was observed.

Data reduction, background subtraction, and normalization were performed according to established methods³⁸⁻⁴² using the program PySpline⁴³ with the data normalized to a value of 1.0 at 7130.0 eV. The spline function through the EXAFS region was chosen to minimize any residual low frequency background but not reduce the EXAFS amplitude, as monitored by the Fourier transform intensity. For pre-edge and edge analysis, all data sets were truncated to $k = 9.5 \text{ Å}^{-1}$ and renormalized using a secondorder post-edge polynomial background and a two segment (three knot) spline for comparison between datasets and to past reference compounds.44 Normalization of the EXAFS data was accomplished using a second-order post-edge polynomial background fit over the energy range up to $k = 14 \text{ Å}^{-1}$ and a three-segment (four knot) spline.

Iron K pre-edge features were modeled with a pseudo-Voigt line shape in a 50:50 ratio of Lorentzian:Gaussian functions using the fitting program EDG_FIT45 as part of the program suite EXAFSPAK. The energy position, full width at halfmaximum (fwhm), and peak intensity were all allowed to float throughout the fitting process. A function modeling the background was empirically chosen to give the best fit, floated with all variables, and then varied with different fwhm (± 0.5 fixed from float) to establish confidence limits on pre-edge intensity. In all cases, an acceptable fit reasonably matched both the preedge data as well as those of its second derivative. A minimum of three fits with different fwhm (± 0.5 fixed from float) backgrounds were acquired over the energy ranges of 7106-7116, 7106-7117, and 7106-7118 eV to provide nine pre-edge fits. These were then averaged to give the final pre-edge energy and intensity values.

EXAFS signals were calculated using FEFF (version 7.0), and the data were fit using the program OPT as part of EXAFSPAK. In all fits the bond lengths (R) and bond variances (σ^2) were allowed to vary. The threshold energy $(k = 0, E_0)$ was also allowed to vary but was constrained as a common variable (ΔE_0) for all fit paths in a given dataset. The amplitude reduction factors (S_0^2) were fixed to a value of 1.0, and the coordination numbers (CNs) were varied systematically based on a structural model to achieve the best fit to the EXAFS data. The best choice of all available FEFF paths and the goodness of the overall fit were optimized based on a combination of weighted F-factor (F)

as well as visual fit to the EXAFS data and their Fourier transform. On the basis of studies of complexes of known structures, the uncertainties in final distances are within 0.02 Å.46 The EXAFS data for 1 were fit using the crystal structure as an initial

Mössbauer spectroscopy

Mössbauer data were collected at 77 K in a Janis model SVT-4000T cryostat using a SEE Co model W302 resonant gamma ray spectrometer. Spectral fits and simulations were performed using the Vinda software package. 47 All isomer shifts are quoted relative to Fe foil at 298 K.

DFT calculations

DFT geometry optimization calculations were performed on 1 using the Gaussian 09 (ref. 48) software package with the crystal structure as the starting structure. The BP86 (ref. 49 and 50) and B3LYP^{51,52} functionals were used with the 6-311G(d) basis set. Implicit solvation was included using the polarizable continuum model with the solvent specified as acetonitrile. Mössbauer parameters were calculated on the Gaussian optimized structures of 1 and [(PaPy₃)Fe^{III}(NO)]²⁺ using the Orca software package, 53 the same functionals, and the CP(PPP) basis set on Fe and TZVP on the remaining atoms. Isomer shifts were calculated from densities at the Fe nucleus using Neese's calibration.54 Mulliken populations were analyzed with QMForge.55

Conflicts of interest

There are no conflicts to declare.

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