Synthesis, Structure, and Catalytic Activity of Mononuclear Iron and (μ-Oxo)diiron Complexes with the Ligand 2,6-Bis(N-methylbenzimidazol-2-yl)pyridine

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Iron complexes including polyimidazole and exchangeable ligands are studied with the aim of modeling the structural and functional features of the non-heme iron centers of dinuclear proteins, such as methane monoxygenase. In [Fe2O(L(MeOH))2(NO3)2][NO3]2 (1) (L = 2,6-bis(N-methylbenzimidazol-2-yl)pyridine), each Fe(III) is in a distorted octahedral environment and has a donor set of N 3O3 which includes three N atoms from L and three O atoms from a nitrate, μ-oxo, and methanol. In complex [FeLCl3]2 (L = 2,6-bis(N-methylbenzimidazol-2-yl)pyridine), Fe(III) is coordinated to three nitrogen atoms from L and three chloride ions. Complex 1 efficiently catalyzed the oxidation of cyclohexane with 51% conversion to cyclohexanol. It also catalyzed the epoxidation of styrene, cyclohexane, 2-methyl-2-butene, and cis- and trans-2-heptene with 51–84% conversions and high selectivity (71–99%) for epoxide products. Complex 2, however, has no specific reactivity toward these substrates. From the alcohol/ketone (A/K) ratio close to 1 in the oxidation of cyclohexane, the low KIE (kinetic isotope effect Kd/Kt ratio = 1.8) for cyclohexanol formation, and the nonstereospecificity of the oxidation of cis-dimethylcyclohexane, it can be concluded that long-lived alkyl radicals are involved in the oxidation catalyzed by complex 1. On the other hand, the stereospecific epoxidation of alkenes, the stereoselective oxidation of cumene, and the high degree of retention of configuration in the oxidation of cis- and trans-2-heptene suggest that a nonradical species, probably a metal-based intermediate, is involved in the oxidation of alkenes and cumene.

Introduction

Oxo- and hydroxo-bridged diiron sites occur in several proteins involved in reversible dioxygen binding or activation.1 Significant examples of this class include the following: hemerythrin (Hr), which binds dioxygen reversibly; the R2 subunit of ribonucleotide reductase (RNR) from Escherichia coli; the soluble form of methane monoxygenase from Escherichia coli; the soluble form of methane monooxygenase; and functional features of the non-heme iron centers of dinuclear proteins, such as methane monoxygenase. In

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2,6-Bis(N-methylbenzimidazol-2-yl)pyridine


8 These studies include the following systems: [Fe3O(TPA)2(H2O)]4+/t-BuOOH (H2O2); 9 [Fe3O(L)Cl3(NO3)2][NO3]2 (L = 2,6-bis(N-methylbenzimidazol-2-yl)pyridine).
Experimental Section

Reagents. All chemicals were of reagent grade unless otherwise noted. The solvents were purified by standard procedures. The ligands 2,6-bis(N-methylbenzimidazol-2-yl)pyridine (L)17 (Et4N)2[Fe-

OC18]18 and [Fe2O4Ac3(H2O)3]NO319 were synthesized according to published procedures.

Preparation of [Fe2O(OH)(NO3)3(CH3OH)][NO3]2 (1). Method A: A mixture of L (165 mg, 0.5 mmol) and Fe(NO3)3·9H2O (202 mg, 0.5 mmol) in methanol (40 mL) was refluxed for 3 h. The resulting orange-red solution was filtered to remove insoluble solids. The filtrate was allowed to stand at room temperature for several hours, after which time it yielded orange-red crystals (yield: 81%).

Analysis. Calcd for Cu3H2Fe2N2O4C: C, 47.23; H, 3.76; N, 17.53. Found: C, 47.38; H, 3.55; N, 17.08. FT-IR (KBr pellet), selected peaks: 1481 (s), 1385 (s), 1315 (s), 842 (s) and 820 (cm)2 UV/vis (in CH3OH, nm), max 293 (εmax 5700 M−1 cm−1), 355 (εmax 1200 M−1 cm−1), 542 (εmax 280 M−1 cm−1), 705 (εmax 80 M−1 cm−1) nm. Raman (in MeOH): νasym (Fe–O–Fe) 352 cm−1. Mössbauer (solid state): δ = 0.46 mm s−1, ΔE0 = 1.27 mm s−1.

Method B. A mixture of L (509 mg, 1.5 mmol) and [Fe2O(OAc)3(H2O)]3[NO3] (327 mg, 0.5 mmol) in 50 mL of methanol/water (1:1 v/v) was refluxed for 1 h. I was obtained as orange-red crystals (yield: 61% based on the ligand used) upon cooling at room temperature.

Preparation of Fe(L)Cl3 (2). (Et4N)2[Fe2O4Cl6] (115 mg, 0.19 mmol) was added to a solution of L (130 mg, 0.38 mmol) in methanol (25 mL) while stirring at room temperature. An orange-red precipitate immediately formed. After the mixture was stirred for 2 h, the precipitate was collected by filtration, washed with ethanol and dried. The precipitate was dried in a vacuum over P4O10. Anal. Calcd for Cu3H2Fe2N2O4C: C, 47.23; H, 3.76; N, 17.53. Found: C, 47.38; H, 3.55; N, 17.08. FT-IR (KBr pellet), selected peaks: 1481 (s), 1385 (s), 1315 (s), 842 (s) and 820 (cm)2 UV/vis (in CH3OH, nm), max 293 (εmax 5700 M−1 cm−1), 355 (εmax 1200 M−1 cm−1), 542 (εmax 280 M−1 cm−1), 705 (εmax 80 M−1 cm−1) nm. Raman (in MeOH): νasym (Fe–O–Fe) 352 cm−1. Mössbauer (solid state): δ = 0.46 mm s−1, ΔE0 = 1.27 mm s−1.

Oxidation of Hydrocarbon Substrates. All manipulations were performed under a dioxygen-free N2 atmosphere. In a typical reaction, complex I or 2 (0.02 mmol) in 10 mL of CH3CN and 45 mmol of substrate were mixed. After the mixture was stirred for 10 min, the reaction was started by the addition of 50 mmol of t-BuOOH (5–6 M in nonane) or H2O2 (30% in water) and stirred under N2 atmosphere at 35 °C for 2–3 h. The organic products were identified by GC, using authentic compounds as internal standards, and further verified by GC/MS and 1H NMR spectra.

Kinetic Isotope Effect Determination. In the same procedure described above, a cyclohexane/D2 cyclohexane (ratio 1:1) mixture was used. We quantitated the products by GC (acetophenone as standard). The KIE was determined by comparing the yields for cyclohexanol and (D11) cyclohexanol and corrected for the relative concentrations of cyclohexane and (D11) cyclohexane.

Collection and Reduction of X-ray Data. Single crystals of I were obtained directly from the reaction medium, while single crystals of 2-CH3OH were obtained by slow vapor diffusion of diethyl ether into the methanol solution. The crystals of I and 2-CH3OH are both air stable. All crystal data for complexes 1 and 2-CH3OH were collected at 298 K on an Enraf-Nonius CAD 4 diffractometer using graphite-monochromated Mo Kα (λ = 0.71073 Å) radiation. The unit cell dimensions were obtained from a least-squares fit of 25 reflections with 5° < 2θ < 24° for I and 8° < 2θ < 25° for 2-CH3OH. A summary of the crystal data, intensity measurement, and solution refinements is given in Table 1. Five standard reflections monitored periodically during data collection showed that the variations for I and 2-CH3OH are less than 2.2% and 1.2%, respectively. All data were corrected for Lorentz and
Table 1. Data Collection and Parameters for Complexes 1 and 2-CH$_3$OH

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<tr>
<td>R$_w$$^a$</td>
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</table>

$^a$ R = ∑(|F$_o$| − |F$_c$|)/∑|F$_o$|, R$_w$ = [∑w(|F$_o$| − |F$_c$|)$^2$/$^2$]$/^2$∑w$^2$F$_o$$^2$, w = 1/σ$^2$($^2$F$_o$$^2$).

polarization effects. An empirical absorption correction based on the ϕ plot was applied.$^{20}$

Structure Solution and Refinement. The structures for 1 and 2-CH$_3$OH were solved by direct methods and successive Fourier-difference syntheses and refined by full matrix least-squares methods. All the hydrogen atoms were generated geometrically and included isotropically in the structure factor calculations, but they were not refined. Atomic scattering factors were taken from The International Tables for X-ray Crystallography.$^{21}$ All calculations were performed with the SHELXLT program.$^{22}$

Physical Measurements. Microanalyses (C, H, N) were performed with a Carlo-Eiba 1106 elemental analyzer. The IR spectra of samples in KBr disks were recorded on a Nicolet-170 SX FT-IR spectrometer in the 4000−400 cm$^{-1}$ range. The 1H NMR spectra were recorded with a Bruker AM-80 spectrometer. The Raman spectra were recorded on a 14031 laser spectrometer with 457.9 nm excitation. Mass spectra were obtained on an AEI-MS-902 mass spectrophotometer. All UV/vis spectra of samples in CH$_3$CN were recorded on a Shimadzu model UV-240 spectrophotometer in the 190−900 nm region.

Mössbauer Spectroscopy. The Mössbauer spectrum was obtained at room temperature using a Hadler MR-351 type spectrometer. A 10 mCi (3.7 × 10$^8$ Bq) $^{57}$Co (Rh) source was used. Each spectrometer was calibrated with respect to standard stainless steel at room temperature. The spectrum was fitted by Lorentzian line shapes using least-squares methods.

Magnetic Studies. Magnetic susceptibility measurements were carried out using a Cahn-200 Faraday-type magnetometer operating at 50 KG (5T) in the 3−300 K range. A powder sample of complex 1 was placed in a Kelf bucket, which had been calibrated independently. Diamagnetic contributions were evaluated using Pascal’s constants.$^{23}$ The data were fitted to the equation for isotropic exchange (−2JS$_1$S$_2$) in a S$_1$ = S$_2$ = 5/2 spin system$^{24}$

modified to allow for a mixture of paramagnetic impurities (P%). The temperature-independent paramagnetic (TIP) susceptibility term results from the mixing of nonthermally populated wave functions of low-lying excited states into the ground-state wave functions. The value minimized was $R = ∑(μ_{exp} - μ_{calc})^2/n∑(μ_{exp})^2$, where n is the number of data used.

Results and Discussion

Synthesis. The μ-oxo-bridged diiron complex 1 with methanol and nitrate as monodentate terminal ligands was obtained readily by refluxing a reaction mixture of L with Fe(NO$_3$)$_3$·3H$_2$O, or with [Fe$_2$O$_2$(OAc)$_2$(H$_2$O)$_2$](NO$_3$)$_2$ in methanol or a methanol−water solution. The yields were 81% and 61% (based on the ligand used), respectively. Attempts to synthesize the μ-oxo-bridged diiron complex [Fe$_2$O(L)$_2$Cl$_4$] with Cl$^-$ as a terminal ligand were unsuccessful, even when the μ-oxo-bridged diiron [Fe$_2$OCl$_4$]$_2$−complex was used as starting material. The mononuclear complex 2 was the only product obtained. Complexes 1 and 2 are stable in air and over a large temperature range, and they are soluble in methanol and acetonitrile.

Description of the Structures. [Fe$_2$O(L)$_2$(NO$_3$)$_2$(CH$_3$OH)$_2$]- (NO$_3$)$_2$ (1). The structure of complex 1 consists of the [Fe$_2$O$_2$-(L)$_2$(NO$_3$)$_2$(CH$_3$OH)$_2$]$^{2+}$ cation and NO$_3$ counterions. A view of the centrosymmetric cation is shown in Figure 1. Selected bond lengths and angles are summarized in Table 2. The bridging oxo group (O(1)) lies on the crystallographic center of symmetry that relates both halves of the molecule. Each iron atom sits in a N$_3$O$_3$ donor set and has a distorted octahedral geometry. Three N atoms come from the tridentate planar ligand L and three O atoms come from the bridging oxo group, the nitrate, and the methanol. The Fe−N(benzimidazole) bond lengths (2.113(5) and 2.118(5) Å) are longer than those found in [N$_2$FeFeCl$_3$]$^+$ (where N$_5$ = N,N′,N″).

(22) Sheldrick, G. M. SHELXTL, System for Crystal Structure Solution, Revision 5.1; University of Göttingen: Göttingen, Germany, 1986.
(23) Carlin, R. L. Magntochemistry; Springer-Verlag: Berlin, 1986; pp 33 and 64.
The Fe–O–Fe bond angle is 180°, and the Fe⋯Fe separation is 3.546 Å. The two planar ligands, with maximum deviation of 0.2 Å from the least-squares planes in I, are transoid to the oxo bridge and parallel to each other. This causes significant π–π stacking interaction between the two parallel planar rings of the ligands (Figure 2). The π–π stacking distance between the two planes is 3.546 Å, which is close to the 3.35 Å known for graphite sheets and the 3.4 Å for stacked nucleotide residues in DNA. Interligand π–π stacking interactions are becoming increasingly common in coordination complexes involving aromatic ligands and have been used to control binding of aromatic substrates in host–guest systems. The theoretical estimation of the magnitude of these interactions is still sparse, but recently it has been shown that face-to-face (rather than edge-to-edge) stacking is favored when the size of the aromatic system is increased. This interligand π-stacking interaction participates in the stability of singly oxo-bridged diiron complexes and may explain the short Fe–O value in I. To the best of our knowledge, this is the first structurally characterized (µ-oxo)diiron complex with a tridentate planar ligand and methanol coordination. Reiff and co-workers have reported a (µ-oxo)diiron complex with a planar terpyridine ligand, formulated as [Fe2O(terpy)]2+[NO3]−4H2O, which was only characterized by spectroscopic methods. It is reasonable to assume that this complex has a structure similar to that of I.

Fe(L)Cl2·CH3OH (2·CH3OH). A perspective view of the molecule with the atomic numbering scheme is shown in Figure 3. Selected bond lengths and angles are summarized in Table 3. The molecule is a neutral Fe(III) complex in which Fe is hexacoordinated with a distorted octahedral geometry. The tridentate N3 planar ligand (L) coordinates the iron(III) center in a meridional fashion. The three monodentate chloride ligands occupy the remaining meridional sites. The Fe–N(benzimidazole) distances (2.111(3), 2.124(3) Å) are shorter than the Fe–N(pyridine) distance (2.168(5) Å) as observed in a related manganese complex.

The Fe–Cl(equatorial) distance (2.262(2) Å) is in the range found for other Fe–Cl ligand bond lengths. The Fe–O(oxo) bond (1.773(1) Å) is in the range found for other Fe–O values observed in methymethylen and methymethylenimethyl azide, which average 1.78 Å.

The tridentate N3 planar ligand (L) coordinates the iron(III) center in a meridional fashion. The theoretical estimation of the magnitude of these interactions is still sparse, but recently it has been shown that face-to-face (rather than edge-to-edge) stacking is favored when the size of the aromatic system is increased. This interligand π-stacking interaction participates in the stability of singly oxo-bridged diiron complexes and may explain the short Fe–O value in I. To the best of our knowledge, this is the first structurally characterized (µ-oxo)diiron complex with a tridentate planar ligand and methanol coordination. Reiff and co-workers have reported a (µ-oxo)diiron complex with a planar terpyridine ligand, formulated as [Fe2O(terpy)]2+[NO3]−4H2O, which was only characterized by spectroscopic methods. It is reasonable to assume that this complex has a structure similar to that of I.
significantly shorter than the Fe–Cl(axial) distances (2.326(1), 2.427(1) Å). The angle Cl(2)–Fe–Cl(1) (173.58°) is close to the ideal value of 180°, while bond angles in the equatorial plane show large distortions from the ideal value of 90° (73.6, 73.9, 106.13, and 88.14° for N(1)–Fe–N(3), N(3)–Fe–N(5), N(5)–Fe–Cl(3), and Cl(3)–Fe–N(1), respectively) due to the stereo requirement of L. The crystallographic study shows that the solvated molecule methanol in Fe(L)Cl3•CH3OH does not interact with any atom of Fe–(L)Cl3. The molecular packing is mainly determined by van der Waals interactions.

**Electronic Spectroscopy.** The electronic spectra of 1 and 2 (10 × 10⁻⁴ M) recorded in methanol show the splitting of intraligand π→π* transition at ~355 and ~293 nm, indicating that ligand L is coordinated to iron(III). Complex 1 also exhibits two bands at 542 and 705 nm, which are tentatively assigned to 6A1 → (4A1, 4E) and 6A1 → 4T2, respectively.42

**Vibration Spectroscopy.** In the IR spectrum of complex 1, the Fe–O–Fe asymmetric stretching mode is assigned to the peak at 842 cm⁻¹ on the basis of the literature reports.7,40 The nitrate groups in 1 yield three bands at 1482, 1351, and 820 cm⁻¹. The magnitude for the splitting of the two higher energy bands is 131 cm⁻¹, suggesting that nitrate groups are present as monodentate ligands.43 Meanwhile, we observed the ionic nitrate band at 1385 cm⁻¹. These results show that complex 1 in methanol has both ionic and coordinated nitrate groups, which is consistent with the X-ray crystal structure.

**Mössbauer Spectroscopy.** The Mössbauer spectrum of complex 1 was recorded at room temperature and shows an asymmetric quadruple doublet (Figure 4). The isomer shift

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and the quadruple splitting (\(\Delta E_0\)) for 1 are 0.46 and 1.27 mm s\(^{-1}\), respectively. Since the isomer shifts for high-spin mononuclear and \(\mu\)-oxo-bridged iron(III) complexes generally fall in the 0.3–0.6 mm s\(^{-1}\) range,\(^{45}\) the 0.46 mm s\(^{-1}\) isomer shift obtained is consistent with the presence of high-spin iron(III) in 1. The 1.27 mm s\(^{-1}\) quadruple splitting for 1 lies at the lower end of the 1.27–1.85 mm s\(^{-1}\) range found in other \(\mu\)-oxo-bridged diiron(III) complexes, indicating a near symmetric charge distribution around the iron nucleus in 1.\(^{46}\)

**Magnetic Properties.** The solution magnetic susceptibility measurement was carried out for 1 in acetonitrile solution using the Evans method.\(^{47}\) An effective magnetic moment of 1.68 \(\mu_B/\text{Fe}\) was obtained. This is much smaller than the 5.9 \(\mu_B\) for high-spin mononuclear iron(III), indicating that the (\(\mu\)-oxo)diiron(III) core of complex 1 is retained in solution. Solid-state variable-temperature magnetic susceptibility studies have been performed for 1. A nonlinear least-squares fitting program\(^{48}\) was used to fit the observed data. With \(g\) fixed at 2, optimization of the fit for 1 gave \(J = -107.7\) cm\(^{-1}\), \(P\% = 8.82 \times 10^{-2}\), \(t\text{ip} = 4.43 \times 10^{-4}\), and \(R = 9.76 \times 10^{-3}\). A plot of molar magnetic susceptibility (\(\chi_m\)) as a function of temperature is shown in Figure 5. The solid line represents the calculated data. The \(J\) value determined for 1 lies in the \(-95\) to \(-191\) cm\(^{-1}\) range reported for singly oxo-bridged diiron complexes,\(^{5}\) but it is significantly smaller than the values \((-187\), \(-191\) cm\(^{-1}\)) reported for linear (\(\mu\)-oxo)diiron porphyrin complexes.\(^{49a,b}\) In the latter case, the difference may be attributed to the slightly shorter Fe–O\(_{\text{oxo}}\) bond length in the linear (\(\mu\)-oxo)diiron porphyrin complexes, because the \(J\) value for (\(\mu\)-oxo)diiron complexes increases as the distance of the iron–oxo bond decreases.\(^{5,49c,d}\)

![Figure 4](image-url)  
Figure 4. Mössbauer spectrum of complex 1 at room temperature.

![Figure 5](image-url)  
Figure 5. Molar susceptibility (\(\Delta\)) for [Fe\(_2\)O\(_2\)(MeOH)\(_2\)(NO\(_3\))\(_2\)](NO\(_3\))\(_2\) (1) as a function of temperature. The solid line is the theoretical fit.

**Catalytic Activity of Complexes 1 and 2.** The catalytic activity of complexes 1 and 2 was tested in the oxidation of cyclohexane, phenol, adamantane, and \(cis\)-1,2-dimethylcyclohexane by tert-butylhydroperoxide (t-BuOOH) and H\(_2\)O\(_2\) in acetonitrile solutions at 35°C under N\(_2\) atmosphere. The product distributions are given in Table 4. Under our standard reaction conditions (complex 1 or 2; substrate:oxidant ratio equal to 1:2250:2500), complex 1 catalyzes the oxidation of cyclohexane to cyclohexanol and cyclohexanone (\(\sim 1\):1 ratio) with 51% conversion of the substrate. Under the same conditions, substrate (cyclohexane) conversion decreased to 37% when H\(_2\)O\(_2\) was used. However, the ratio of cyclohexanol to cyclohexanone (\(\sim 1\):1) was the same as with t-BuOOH. The addition of the tetrabutylammonium bromide (TBAB) phase-transfer catalyst to the H\(_2\)O\(_2\) reaction mixture increased the conversion to 48%. Neither t-BuOOH nor H\(_2\)O\(_2\) with complex 2 shows catalytic activity in the oxidation of cyclohexane. Both complexes 1 and 2, under our reaction conditions, efficiently catalyze the oxidation of phenol to hydroquinone, with 61% and 23% conversions, respectively. Interestingly, a higher selectivity of the catalytic oxidizing system for the tertiary C–H bonds of adamantane with t-BuOOH by complex 1 has been observed. The C \(3\)°\(c\) ratio is 15.4 (Table 4, entry 8), \(cis\)-1,2-Dimethylcyclohexane is converted to \(cis\)- and \(trans\)-1,2-dimethylcyclohexanol with a \(cis\)/\(trans\) ratio of 1.1 (Table 4, entry 9). The catalytic activities of complex 1 for the oxidation of several alkenes (stylene, 2-methyl-2-butenne, cyclohexene, 1-methylcyclohexene, and \(cis\)/\(trans\)-2-heptenes) were determined. The product distributions are summarized in Table 5. Typical reactions with a ratio of catalyst 1:substrate/t-BuOOH equal to 1:2250:2500 were run in acetonitrile solution for 2.5–3.0 h at 35°C. Complex 1 catalyzes the oxidation of alkenes with 63–85% conversion to epoxides and ketones. However, the selectivity for epoxide formation is high. The product distributions (Table 5) for epoxides range from 77 to 99% and 0.3–7.9% for ketones or aldehydes. This high stereo-epoxidation selectivity is in sharp contrast with the iron complex containing a tetradentate macrocyclic cyclamide...
and t-BuOOH oxidant, where only the allylic product was observed.\(^\text{50}\) The cis/trans-2-heptenes were converted to their corresponding epoxides in 72–76% yields with 93–97% retention of configuration (Table 5, entries 5 and 6). In the case of cumene, complex 1 gives only the 2-phenyl-2-propanol product. To the best of our knowledge, complex 1 is one of a few non-heme iron catalysts that are capable of catalyzing the oxidation of alkenes with epoxides as a main product.\(^\text{51}\) Complex 2 is not catalytic in the oxidation of all alkenes. Control experiments performed in the absence of catalyst 1 or in the presence of the ligand alone showed little or no oxidation. Autoxidation by Fe(NO\(\text{3}\))\(_4\)-9H\(_2\)O and FeCl\(_3\)-6H\(_2\)O was much slower and gave very low yields and selectivity. This indicates that the metal complex is required for the reactions to occur and is indeed acting as a catalyst.

Complex 1 efficiently catalyzes both the oxidation of cyclohexane and epoxidation of alkenes with t-BuOOH or H\(_2\)O\(_2\) as the oxidant in good yields (36–85%) under mild conditions. Complex 2 has no catalytic activity toward the oxidation of cyclohexane and alkenes under the same conditions. The difference in catalytic activity between complexes 1 and 2 may be associated with the presence of the exchangeable ligands, methanol molecules and nitrate groups, in the diiron core of 1. The methanol molecules trans to the bridged oxygen atom can be readily replaced by the oxidant or the substrate. This observation indicates that the presence of exchangeable ligands in non-heme iron catalysts provides the best catalytic activities, as reported in the literature.\(^\text{8b,11}\) The Cl\(^-\) in Fe–Cl bonds in 2 is not as labile, which makes it a poor catalyst.\(^\text{11c}\) This is further supported by the efficient catalysis of the oxidation of phenol by both 1 and 2. Apparently, phenol has strong coordinating ability and can replace Cl\(^-\) in complex 2.\(^\text{52}\) A significant decrease was observed in the oxidation of cyclohexane by H\(_2\)O\(_2\) compared to t-BuOOH, because complex 1 is less soluble in aqueous H\(_2\)O\(_2\) than in the acetonitrile reaction mixture. This is underscored by the fact that the addition of the tetrabutylammonium bromide phase-transfer catalyst to the reaction mixture increases the oxidation.

It is worth pointing out that complex 1 slowly decomposes after 5 h and some precipitate was obtained in acetonitrile under reaction conditions. It is probable that the dinuclear oxo-bridged iron(III) core cleaved and two mononuclear iron species were formed during the reaction. This is further supported by the disappearance of the intense absorption band at ~542 nm after 5 h, which is assigned to \(^6\)A\(_1\) \(\rightarrow \) \(^4\)A\(_1\), \(^4\)E) for a dinuclear \(\mu\)-oxo-bridged motif (Figure 6). The characteristic absorption bands assigned to the \(\pi-\pi^*\) transition at 293 and 355 nm for the coordinated ligand are unchanged.

**Mechanistic Probes of Alkane Oxidation.** The alcohol/ketone (A/K) ratio is a simple test that probes the mechanism of alkane oxidation. Complex 1’s A/K ratio is 1.2 with t-BuOOH and 1.1 with H\(_2\)O\(_2\) (as shown in Table 4). Both values are close to 1, indicating that cyclohexane oxidation catalyzed by complex 1 involves either alkyl radicals with long lifetimes or \(\cdot\)OH radicals.\(^\text{8b,53}\)

The intermolecular kinetic isotope effect (KIE) for the formation of cyclohexanol was determined in competition

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**Table 4. Product Distributions for the Oxidation of Alkanes and Phenol with t-BuOOH and H\(_2\)O\(_2\), Catalyzed by Complexes 1 and 2**

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<td>1</td>
<td>1</td>
<td>t-BuOOH</td>
<td>cyclohexane</td>
<td>51.3</td>
<td>cyclohexanol/cyclohexanone</td>
<td>28.1</td>
<td>A/K(^f) = 1.2</td>
</tr>
<tr>
<td>2</td>
<td>2</td>
<td>t-BuOOH</td>
<td>cyclohexane</td>
<td>0</td>
<td>cyclohexanol/cyclohexanone</td>
<td>23.2</td>
<td>KIE(^g) = 1.8</td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>30% H(_2)O(_2)</td>
<td>cyclohexane</td>
<td>36.8</td>
<td>cyclohexanol/cyclohexanone</td>
<td>19.5</td>
<td>A/K = 1.1</td>
</tr>
<tr>
<td>4</td>
<td>2</td>
<td>30% H(_2)O(_2)</td>
<td>cyclohexane</td>
<td>0</td>
<td>cyclohexanol/cyclohexanone</td>
<td>17.3</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>1</td>
<td>30% H(_2)O(_2) + TBAB(^h)</td>
<td>cyclohexane</td>
<td>47.6</td>
<td>cyclohexanol/cyclohexanone</td>
<td>25.4</td>
<td>A/K = 1.1</td>
</tr>
<tr>
<td>6</td>
<td>1</td>
<td>t-BuOOH</td>
<td>phenol</td>
<td>60.7</td>
<td>cyclohexanol/hydroquinone</td>
<td>59.8</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>2</td>
<td>t-BuOOH</td>
<td>phenol</td>
<td>23.4</td>
<td>hydroquinone/cyclohexanol</td>
<td>0.9</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>1</td>
<td>t-BuOOH</td>
<td>adamantane</td>
<td>50.7</td>
<td>cis-1,2-dimethyl-cyclohexanol isomers</td>
<td>47.6</td>
<td>cis/cis = 1.1</td>
</tr>
<tr>
<td>9</td>
<td>1</td>
<td>t-BuOOH</td>
<td>cis-1,2-dimethyl-cyclohexane</td>
<td>38.8</td>
<td>trans-1,2-dimethyl-cyclohexanol isomers</td>
<td>20.2</td>
<td></td>
</tr>
</tbody>
</table>

\(^a\) Reactions were performed in acetonitrile solution under N\(_2\) atmosphere. Reaction temperature: 35 °C. Reaction time: 2 h. t-BuOOH: tert-butylhydroperoxide. \(^b\) TBAB: tetrabutylammonium bromide. \(^c\) Yield based on the substrate. \(^d\) A/K = cyclohexanol/cyclohexanone. \(^e\) KIE = intermolecular kinetic isotope effect of cyclohexanol formation. \(^f\) A/K = (1-adamantanol)/(2-adamantanol + 2-adamantanone). \(^g\) cis/trans = cis-1,2-dimethylcyclohexanol isomers/trans-1,2-dimethylcyclohexanol isomers = [(\(1\R,2\R\))-1,2-dimethylcyclohexanol + (\(1\R,2\S\))-1,2-dimethylcyclohexanol + (\(1\S,2\R\))-1,2-dimethylcyclohexanol].

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experiments between cyclohexane and [D\textsubscript{12}]cyclohexane. A value of 1.8 (Table 4, entry 1) was obtained in acetonitrile, which is in the range of 1–2 for radical-type oxidations.\(^{(54)}\)

With adamantane as a substrate, a 3\(^{9}/2\textsuperscript{o}\) ratio of 15.4 (Table 4, entry 8) was observed. The selectivity for the oxidation at the tertiary position is higher than the average value of 2.7 found for Gif-type oxidations, about 2 for the oxidation of alkanes by \(\cdot\text{OH}\), and 9.5–10 for the oxidation with \([\text{Fe}^{II}(\text{bpy})_4(\text{H}_2\text{O})_2](\text{ClO}_4)_4/\text{t-BuOOH}\).\(^{(55)}\) It is in the range of 15–33 for oxidation with \([\text{Fe}^{III}(\text{TPA})(\text{CH}_3\text{CN})_2]/\text{H}_2\text{O}_2\).\(^{(56)}\) These observations indicate that an oxidant more selective than the hydroxyl radical is also involved in the oxidation.

The stereoselectivity of the alkane hydroxylation reaction was examined with \(\text{cis-1,2-dimethylcyclohexane}\) as a substrate, and the results are shown in Table 4 (entry 9). Both isomeric \(\text{cis-1,2-dimethylcyclohexanol}\) and \(\text{trans-1,2-dimethylcyclohexanol}\) were formed with a cis/trans ratio of 1.1, which is in the range of 1.1–1.3 found for a catalytic autoxidation reaction.\(^{(57)}\) This indicates the formation of alkyl radicals with a lifetime sufficient to allow epimerization at the radical site.

The data imply that alkyl radicals are involved in the oxidation of the alkanes studied here, although it does not exclude the possibility that a metal-based oxidant is also

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**Table 5.** Product Distributions for the Oxidations of Alkenes and Cumene with t-BuOOH, Catalyzed by Complex 1

<table>
<thead>
<tr>
<th>Entry</th>
<th>Substrate</th>
<th>Conversion of Substrate (%)</th>
<th>Product</th>
<th>Yield(^{b})</th>
<th>Selectivity for Epoxide(^{c})</th>
<th>RC(^{d})</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>styrene</td>
<td>79.6</td>
<td></td>
<td>79.3</td>
<td>99.2%</td>
<td>0.3</td>
</tr>
<tr>
<td>2</td>
<td>2-methyl-2-butene</td>
<td>84.5</td>
<td></td>
<td>83.9</td>
<td>98.6%</td>
<td>0.6</td>
</tr>
<tr>
<td>3</td>
<td>cyclohexene</td>
<td>72.0</td>
<td></td>
<td>71.1</td>
<td>97.5%</td>
<td>0.9</td>
</tr>
<tr>
<td>4</td>
<td>1-methylcyclohexene(^{e})</td>
<td>57.9</td>
<td></td>
<td>50.7</td>
<td>75.1%</td>
<td>7.2</td>
</tr>
<tr>
<td>5</td>
<td>(\text{cis-2-heptene})</td>
<td>81.2</td>
<td></td>
<td>76.9</td>
<td>96.7%</td>
<td>1.3</td>
</tr>
<tr>
<td>6</td>
<td>(\text{trans-2-heptene})</td>
<td>79.8</td>
<td></td>
<td>72.5</td>
<td>93.9%</td>
<td>2.3</td>
</tr>
<tr>
<td>7</td>
<td>Cumene(^{e})</td>
<td>63.4</td>
<td></td>
<td>63.4</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

\(^{a}\) Reactions were performed in acetonitrile solution under N\(_2\) atmosphere. Reaction temperature: 35 °C. Reaction time: 3 h. \(^{b}\) Yield based on the substrate. \(^{c}\) Selectivity for epoxidation = epoxide product/(epoxide product + ketone or aldehyde product). \(^{d}\) RC = 100((cis – trans)epoxide/(cis + trans)epoxide for epoxidation of \(\text{cis-2-heptene}\) or 100((trans – cis)epoxide/(cis + trans)epoxide for epoxidation of \(\text{trans-2-heptene}\). \(^{e}\) Reaction time: 2.5 h.

involved in the oxidation because of a higher 3°/2° ratio of 15.4 for the oxidation of adamantane.

As far as the oxidation of alkenes is concerned, complex 1 exhibits highly stereoselective epoxidation of the alkenes studied, indicating a type of two-electron oxidation process\(^\scriptscriptstyle{55}\) and involvement of a metal-based oxidant.\(^\text{(58)}\) With styrene as a substrate, complex 1 afforded almost all epoxide product with very little benzaldehyde, which is derived from the O\(_2\) trapping of a radical cation species,\(^\text{(59)}\) and points to the involvement of a nonradical oxidant. The oxidation of 1-methylcyclohexene with \(\text{t-BuOOH}\) gave 75% epoxide product and 25% ketone product. With cyclohexene, which is susceptible to allylic oxidation, 99% epoxide product was found and little ketone product was observed. These results indicate that a nonradical oxidant is involved in the oxidations. In the oxidations of \(\text{cis}-\text{trans}-2\)-heptenes both compounds were converted into their corresponding \(\text{cis}-\text{trans}\)-epoxides with 93–98% retention of configuration. This high degree of retention of configuration is consistent with a nonradical mechanism\(^\text{(51b)}\) or, less likely, with a radical mechanism having very short-lived radical intermediates. Complex 1 also gave only 2-phenyl-2-propanol product in the oxidation of cumene under our standard reaction conditions, indicating involvement of a metal-based oxidant intermediate in the oxidation.\(^\text{(60)}\) These results indicate that nonradical species, probably metal-based intermediate oxidants, are involved in the oxidations of the alkenes studied and cumene.

On the basis of the data above, we suggest that two different oxidants (RO\(^\text{•}\) and iron–O oxo) might be involved in the oxidations\(^\text{(55,61)}\) and we propose the following possible mechanism. In the first step of the reaction, the binding of the oxidant to the dinuclear complex results in the cleavage of the dinuclear unit into monomers, forming an \(\text{LF}_{\text{Fe}}\text{O}_{\text{O}} \) intermediate. This unstable peroxo complex would either generate an \(\text{Fe}^{\text{IV}}\text{O} \) and the highly reactive RO\(^\text{•}\) by O–O bond homolysis or generate a hypervalent oxo–iron intermediate \(\text{Fe}^{\text{V}}\text{O} \) by bond O–O heterolysis, or it could simultaneously generate all reactive species both by O–O bond homolysis and heterolysis. A transient species \(\text{Fe}^{\text{V}}\text{O} \) with a non-heme macrocyclic ligand was recently isolated and fully characterized by X-ray diffraction,\(^\text{(62)}\) showing that non-heme ligands can stabilize the \(\text{Fe}^{\text{V}}\text{O} \) transient species. In the case of the oxidation of alkenes, the O–O bond heterolysis predominates, which results in \(\text{Fe}^{\text{V}}\text{O} \) active oxidizing intermediates. This is followed by the intermediate \(\text{Fe}^{\text{V}}\text{O} \) transfer of oxygen atoms to alkenes, giving epoxides as major products. It is possible that the nuclophilic nature of the C–C bonds in alkenes favors the formation of \(\text{Fe}^{\text{V}}\text{O} \) transient species, resulting in the \(\text{Fe}^{\text{V}}\text{O} \) intermediates domination of the oxidations of alkenes. With the oxidations of alkenes, O–O bond homolysis dominates and generates reactive alkoxyl radicals RO\(^\text{•}\) that are involved in the oxidation of alkenes. This is in good agreement with the 1:1 ratio of cyclohexanol/cyclohexanone during the oxidation of cyclohexene. However, more experiments are needed to confirm the validity of this proposed mechanism.

Conclusions

To prepare functional models that can mimic the alkane functionalization chemistry of methane monoxygenase, the dinuclear oxo-bridged complex 1 was synthesized with the planar tridentate 2,6-bis(N-methylbenzimidazol-2-yl)pyridine, \(\text{L}\), and \(\text{Fe(NO}_3\text{)}_3\cdot 9\text{H}_2\text{O}\) or \(\text{Fe(OAc)}_3\cdot 2\text{H}_2\text{O}\) in methanol solution. Mononuclear complex 2 was also obtained by reaction of \(\text{L}\) with \((\text{N}_{\text{Et}})_2\text{Fe(OCl)}_3\). Crystallographic studies revealed that each \(\text{Fe(III)}\) in \([\text{Fe}_2\text{OL}_2\text{H}_2\text{O}(\text{NO}_3)_{2}\text{]}\text{NO}_3\) (1) has a distorted octahedral geometry containing a \(\text{N}_2\text{O}_3\) donor set with three \(\text{N}\) atoms from \(\text{L}\) and three O atoms from nitrate, methanol, and a bridged oxo unit. In \(\text{FeLCl}_3\), \(\text{Fe(III)}\) is coordinated to three \(\text{N}\) atoms from \(\text{L}\) and three Cl\textsuperscript{–} anions. Complex 1 is the first example of a structurally characterized linear, dinuclear, oxo-bridged complex with a planar tridentate ligand containing a methanol coordinated trans to the \(\mu\)-oxo bridge. In catalytic studies, complex 1 displays high catalytic activity in the oxidation of alkanes and is an excellent catalyst for stereoepoxidation of the alkenes studied, while complex 2 is inactive. The difference in catalytic activity between complexes 1 and 2 shows that the presence of a labile coordinated group is essential to

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\*Figure 6. UV/vis spectra of catalyst 1 at various reaction times: (a) at time zero; (b) after 1 h; (c) after 5 h.*
permit the oxidant or substrate to bind to the Fe for an efficient catalyst. Mechanistic studies show that the mechanism of catalytic oxidation by complex 1 depends on the substrates used: a A/K ratio (close to 1) and a low KIE value (1.8) for cyclohexanol formation as well as the lack of stereoselectivity in the oxidation of cis-dimethylcyclohexane suggest that alkyl radicals are involved in the oxidation. Stereospecific epoxidation of the alkenes studied, stereoselectivity of the oxidation of cumene, and the high degree of retention of configuration in the oxidation of cis/trans-2-heptenes indicate that nonradical species, probably metal-based intermediate oxidants, are involved in the oxidations of the alkenes and cumene.

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Supporting Information Available: Tables of crystal data and structure solution, refinement details, atomic coordinates, bond lengths and angles, and anisotropic thermal parameters for compounds 1 and 2. This material is available free of charge via the Internet at http://pubs.acs.org.

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