A Bioinspired Nonheme $Fe^{III}-(O_2^{2-})-Cu^{II}$ Complex with an $S_t = 1$ **Ground State**

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Cite This: *J. Am. Chem. Soc.* 2024, 146, [24808−24817](https://pubs.acs.org/action/showCitFormats?doi=10.1021/jacs.4c04492&ref=pdf) **Read [Online](https://pubs.acs.org/doi/10.1021/jacs.4c04492?ref=pdf)**

ACCESS | **ILLE** [Metrics](https://pubs.acs.org/doi/10.1021/jacs.4c04492?goto=articleMetrics&ref=pdf) & More | E Article [Recommendations](https://pubs.acs.org/doi/10.1021/jacs.4c04492?goto=recommendations&?ref=pdf) | **G** Supporting [Information](https://pubs.acs.org/doi/10.1021/jacs.4c04492?goto=supporting-info&ref=pdf) ABSTRACT: Cytochrome *c* oxidase (CcO) is a heme copper $O₂$ $o^{\leq o}$ oxidase (HCO) that catalyzes the natural reduction of oxygen to $Cu¹$ water. A profound understanding of some of the elementary steps o^{20} HO leading to the intricate $4e^-/4H^+$ reduction of O_2 is presently lacking. A total spin $S_t = 1 \text{ Fe}^{\text{III}} - (O_2^{\text{2}-}) - Cu^{\text{II}}$ (I_P) intermediate is \checkmark S_i = 1 end-on O_2^2

proposed to reduce the overpotentials associated with the reductive O−O bond rupture by allowing electron transfer from a tyrosine moiety without the necessity of any spin-surface crossing. Direct evidence of the involvement of I_P in the CcO catalytic cycle is, however, missing. A number of heme copper peroxido complexes have been prepared as synthetic models of I_p , but all of them possess the catalytically nonrelevant $S_t = 0$ ground state resulting from antiferromagnetic coupling between the $S = 1/2$ Fe^{III} and Cu^{II}

centers. In a complete nonheme approach, we now report the spectroscopic characterization and reactivity of the Fe^{III}−(O2^{2−})−Cu^{II} intermediates 1 and 2, which differ only by a single −CH3 versus −H substituent on the central amine of the tridentate ligands binding to copper. Complex 1 with an end-on peroxido core and ferromagnetically $(S_t = 1)$ coupled Fe^{III} and Cu^{II} centers performs H-bonding-mediated O−O bond cleavage in the presence of phenol to generate oxoiron(IV) and exchange-coupled copper(II) and PhO[•] moieties. In contrast, the μ - η^2 : η^1 peroxido complex 2, with a $S_t = 0$ ground state, is unreactive toward phenol. Thus, the implications for spin topology contributions to O–O bond cleavage, as proposed for the heme Fe^{III}−(O2^{2−})−Cu^{II} intermediate in CcO, can be extended to nonheme chemistry.

■ **INTRODUCTION**

Cytochrome *c* oxidase (CcO) is a multisubunit transmembrane protein, which catalyzes the 4e⁻/4H⁺ reduction of dioxygen to water during the last step of the electron-transport chain, thereby generating a transmembrane proton gradient respon-sible for driving ATP synthesis.^{[1](#page-8-0)−[3](#page-8-0)} CcO contains a unique heterobimetallic heme copper active site, the metal centers of which are separated by >4.0 Å.^{4,5} The four electrons required to fully reduce dioxygen to water are supplied by iron ($\bar{Fe}^{II} \rightarrow$ Fe^{IV}), copper (Cu^I → Cu^{II}), and a tyrosine residue (Tyr-OH → Tyr−O•), which is covalently tethered to one of the histidine moieties ligated to Cu. The consensus mechanism of CcO, shown in [Scheme](#page-1-0) 1, involves the initial binding of dioxygen to the reduced (R) form of the enzyme to form transient intermediate A, which is then rapidly converted to $P_M^{3,5}$ $P_M^{3,5}$ $P_M^{3,5}$ The electronic structure of P_M is unambiguously assigned as a ferryl-heme-cupric-hydroxide-tyrosyl-radical species based on various spectral and chemical evidence.^{[6](#page-8-0)} However, the identity of the species formed before the O−O cleaved intermediate P_M has proved controversial. For example, intermediate A, which is believed to formally be a ferric-superoxido species, exhibits a *ν*(Fe−O) stretch of 571

cm[−]¹ , [7](#page-8-0) which overlaps with end-on (hydro)peroxidos found in proteins $^{8-10}$ $^{8-10}$ $^{8-10}$ $^{8-10}$ $^{8-10}$ and various well-characterized synthetic low-spin heme copper peroxido complexes in organic media or on electrode surfaces.^{[5](#page-8-0),[11](#page-8-0)–[15](#page-8-0)} Accordingly, the lack of empirical vibrational data does not allow for a definite assessment of the degree of O_2 reduction for intermediate A. Furthermore, various theoretical and kinetic studies have hypothesized the involvement of a total spin $S_t = 1$ heme $Fe^{III} - O_2 - Cu^{II}$ peroxido intermediate (I_P) during the A to P_M conver- \sinh^{16-19} \sinh^{16-19} \sinh^{16-19} \sinh^{16-19} \sinh^{16-19} however, direct spectroscopic evidence of such intermediate in the CcO catalytic cycle has stayed elusive. Notably, the ferromagnetic coupling between the low-spin *S* = $1/2$ Fe^{III} and *S* = $1/2$ Cu^{II} centers in the *S*_t = 1 heme Fe^{III} – O2−CuII peroxido intermediate to form the exchange-coupled CuII−OH and Tyr• products in CcO is discussed as the

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prerequisite necessary for the efficient Tyr-mediated reductive O−O bond rupture without any spin-surface crossing.^{[18](#page-9-0),20,2}

Synthetic examples that demonstrate the structural feasibility of the proposed heme Fe^{III} – O_2 – Cu^{II} peroxido intermediate in CcO have been reported.^{5,11,12,[21](#page-9-0)–[28](#page-9-0)} Nevertheless, the acute Fe−O−O−Cu dihedral angle in all of these complexes have led to a catalytically nonrelevant antiferromagnetically coupled

 $S_t = 0$ ground state of the low-spin heme Fe^{III}–O₂–Cu^{II} adducts.^{[14,15](#page-8-0)} The presence of a bridging heme-peroxido-copper species has also been reported in the X-ray structures of the eracting state of CcO.^{[4,](#page-8-0)[29](#page-9-0)–[31](#page-9-0)} But the exact nature of the O₂reduced moiety and its formation in these "as isolated" structures have been disputed. In particular, the reported O−O and Fe/Cu−O bond lengths in these structures are inconsistent with the peroxido assignment.^{[5](#page-8-0)} One of the "as isolated" structures was reported to exhibit a UV−vis absorption feature at 650 nm, which when excited at this wavelength showed a resonance Raman (rRaman) active band at 755 cm⁻¹.^{[32](#page-9-0)} This band, which was initially thought to be a *ν*(O−O) stretching mode, was later assigned to a mode of the His-419 ligand based on rRaman spectroscopy.^{[33](#page-9-0)} This provided further doubt to the proposed binding of the peroxido ligand in the "as isolated" form of CcO.

Herein, we report the synthesis, spectroscopic characterization, and reactivity of a nonheme $[(CH₃CN)(trans$ cyclam)Fe^{III}(μ - η ¹· η ¹-O₂)Cu^{II}(AN)]³⁺ (1; AN = 3,3′-iminobis-(*N*,*N*-dimethylpropyl-amine)) complex, which may improve our understanding of the properties of the elusive catalytically relevant ferromagnetically coupled $(S_t = 1)$ low-spin heme Fe^{III} $-O_2-Cu$ ^{II} peroxido intermediate proposed during the A to P_M conversion in CcO. Notably, the corresponding complex $[(cis-cyclam)Fe^{III}(\mu-\eta^2:\eta^1-O_2)Cu^{II}(MeAN)]^{3+}$ (2; MeAN = 2,6,10-trimethyl-2,6,10-triazaundecane), where MeAN differs from AN in having one methyl group in the ligand structure, exhibits distinct $O₂$ binding mode and reactivity properties different from 1. The present study, therefore, highlights the importance of the interplay between metal centers and the local environment in governing key physical and chemical properties of biologically relevant dinuclear metal-dioxygen intermediates.

Figure 1. (A) Mößbauer spectra at variable magnetic field (parallel to the *γ*-beam) of ⁵⁷Fe-enriched 1 in a frozen solution of an acetone/CH₃CN 10:1 v/v mixture. The simulation (black lines) of the experimental data (dots) gave a best fit with two subspecies, sub1 (blue lines) with $S_t = 1$ corresponding to 1 (70%, δ = 0.26 mm s⁻¹, $\Delta E_{\rm Q}$ = -2.11 mm s⁻¹, $D_{\rm t}$ = 1.5 cm⁻¹, $A_{\rm xx}$ = -3.2 T, $A_{\rm yy}$ = -16.2 T, $A_{\rm zz}$ = 4.4 T, η = 0.43) and sub2 (green lines) corresponding to a partially formed (24%) Fe^{IV} (*S* = 1) species (see [Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S1 for further details); please note that the high-spin Fe^{III} signal observed in electron paramagnetic resonance (EPR) ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S3) is not detected in Mössbauer; (B) UV-vis absorption spectra of 1 (blue) and 3 (green) in a 10:1 acetone/CH₃CN v/v mixture at −90 °C. (C) rRaman spectra of 1 (pink) and ¹⁸O-labeled 1 (blue) measured in a 10:1 acetone/CH3CN v/v solution (407 nm excitation, 2 mW, −90 °C). Solvent features are marked by asterisks.

■ **RESULTS AND DISCUSSION**

Synthesis and UV−**Vis and Mössbauer Characterizations of 1.** In a previous study, we reported the formation and characterization of the $[(CH_3CN)(trans-cyclam)$ - $\mathrm{Fe}^{\mathrm{III}}(\mathrm{O_2}^{-\bullet})]^{2+}$ compound 3 by addition of $\mathrm{O_2}$ to a solution of $[(CH_3CN)_2$ (*trans-cyclam*) Fe^{II}](OTf)₂.^{[34](#page-9-0)} [Figure](#page-1-0) 1B shows the UV−vis spectrum after reaction of one equivalent of the well-characterized $[\text{Cu}^{\text{I}}(\text{AN})]\text{BF}_4$ complex^{[28](#page-9-0),[35](#page-9-0)} with 3, which led to the formation of a deep blue species, 1, with an absorption maximum at 615 nm (ϵ = 1.5 L mmol⁻¹ cm⁻¹) and a half-life time *t*1/2 = 10 min at −90 °C. The 14 K Mößbauer spectrum of 1 ([Figure](#page-1-0) 1A) shows a main species in ∼70% yield with parameters (isomer shift $\delta = 0.26$ mm s⁻¹, quadrupole splitting $|\Delta E_Q| = 2.11$ mm s⁻¹) that are typical for a low-spin (*S* $= 1/2$) Fe^{III} center. However, the low-temperature, low-field (5) K, 0.06 T) Mößbauer spectrum of 1 ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S1, top, blue lines) does not display a magnetic splitting but instead a broadened, asymmetric doublet, characteristic of an integer spin with a positive zero-field splitting (ZFS) parameter. This is confirmed by applied magnetic field Mößbauer studies of 1 ([Figure](#page-1-0) 1A), where the spectra are best fit in the slowrelaxation limit with a total spin $S_t = 1$ with a small axial zerofield splitting $D_t = 1.5$ cm⁻¹ and an anisotropic hyperfine coupling tensor. The asymmetry parameter $\eta = 0.43$ determined for 1 differs from the reported parameters for the axially symmetric $[{\rm Fe}^{\rm IV} (\rm O) (trans-cyclam) (\rm CH_3 CN)]\rm OTf_2$ $(\eta = 0)$ complex.^{[34](#page-9-0)} The Mössbauer spectra of the minor species [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S1, green lines) are completely compatible with an axial $Fe^{IV}=O$ complex. Altogether, the Mößbauer measurements propose a species with a low-spin Fe^{III} that is ferromagnetically coupled to a Cu^H center, giving rise to a S_t $= 1$ state in 1.

X-ray Absorption Near-Edge Structure (XANES) and EPR **Spectroscopy.** In addition to $\mathrm{Fe^{III}}$, the presence of a $\mathrm{Cu^{II}}$ center 1 is confirmed by the X-ray absorption near-edge structure (XANES) data [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S2). The XANES spectra at the Cu K-edge revealed an edge energy of 8986.2 eV, which is shifted by ca. 2.5 eV to higher energies relative to the $[Cu^{I}(AN)]BF_{4}$ starting compound (8983.6 eV), confirming a divalent oxidation state for copper in 1. Complex 1 is EPR silent in perpendicular mode at the X-band [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S3), consistent with its $S_t = 1$ ground state with considerable zerofield splitting; the signals shown in [Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S3A correspond to minor contributions (∼15% based on spin quantification; [Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S3B) arising from monomeric high- and low-spin Fe^{III} and Cu^{II} impurities.

rRaman Spectroscopy. Confirmation for the presence of a peroxido moiety in 1 comes from rRaman measurements (in acetone/CH3CN 10:1 v/v solution at −90 °C upon excitation at 407 nm) as shown in [Figure](#page-1-0) 1C. They exhibit a characteristic but very low-energy *ν*(O−O) vibrational mode at 739 cm⁻¹ (¹⁶O₂; 700 cm⁻¹ with ¹⁸O₂) and a diagnostic *ν*(Fe−O) mode at 635 cm⁻¹ (¹⁶O₂; 608 cm⁻¹ with ¹⁸O₂).^{[36](#page-9-0)}

Theoretical Studies. For a better understanding of the actual peroxido binding mode in 1, density functional theory (DFT) calculations were performed to identify optimized structures (Figure 2) and verify if they can theoretically reproduce the experimental results. Due to the flexibility of the mononucleating AN and cyclam ligands, a variety of different conformations and rotamers with and without an additional solvent molecule (CH₃CN) binding to the Fe^{III} center were needed to be considered. These contain the cyclam ligand to

Figure 2. DFT-optimized structures 1-*trans*-b (left) and 2-*cis*-b (right); (color code: Cu brown, Fe orange, O red, N blue, C gray, H white (C−H protons are omitted for clarity)). See [Tables](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S7 and S8, respectively, for their Cartesian coordinates.

Fe in either a folded (*cis*-V) conformation with alternating directions of the amino hydrogens (above vs below the ring) or a planar (*trans*-III) conformation with propylene-linked amino hydrogens pointing in the same and ethylene-linked ones in different directions. 37 One of the axial binding sites to Fe in *trans*-cyclam is occupied by a coordinating solvent molecule CH3CN, while solvent coordination at the *cis* binding sites is sterically hindered by the bound $\left[Cu(AN)(O_2) \right]$ moiety. The AN ligand can adopt either a bent (b) or a planar (p) conformation. $[(\text{cyclam})\text{Fe}^{\text{III}}(O_2)\text{Cu}^{\text{II}}(\text{AN})]^{3+}$ starting structures with all ligand conformers, different O−O binding modes (*μ*-*η*¹:*η*¹, *μ*-*η*¹:*η*², *μ*-*η*²:*η*¹, *μ*-*η*²:*η*²), and varying relative Fe− cyclam, O−O, and Cu−AN orientations were constructed and optimized. The same approach has been pursued for complex 2 (vide infra), in which the amino hydrogen of the AN ligand is replaced with a methyl group (MeAN). For each conformational combination, the lowest-energy structure, along with their relative energies, geometric parameters, Raman data of ¹⁸O-sensitive modes, and UV−vis absorption spectra are presented in [Table](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S1.

All geometry optimizations yielded Cu^H in a distorted tetrahedral coordination environment, ligated by only one of the peroxido oxygens. Independent of the conformation of the Cu ligand, starting structures with *trans*-III-cyclam converged to a *μ*-*η*¹ :*η*¹ (end-on) bridging motif with FeCu distances of 4.1−4.3 Å and a trend of short (1.8 and 1.9 Å, respectively) and long (2.6−2.8 Å) Fe···O and Cu···O distances, respectively. In contrast, those with *cis*-V-cyclam yielded a *μη*2 :*η*¹ motif with the peroxido unit binding side-on to Fe and end-on to Cu and shorter Fe···Cu distances of 3.7−3.9 Å, two shorter Fe \cdots O distances (1.8 and 2.0 Å), and one short and one long Cu···O distances (2.0 and 2.9 Å, respectively). For both binding modes, the O···O distances of 1.43−1.45 Å are very similar. Broken-symmetry (BS) DFT suggests triplet ground states and thus ferromagnetic Fe^{III}−Cu^{II} interactions for all of the structures.

Most interestingly, depending on the O−O binding mode, the conformers can be divided into two groups according to both the calculated UV−vis spectra and the vibrational frequencies of the peroxido bridge [\(Table](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S1 and [Figure](#page-3-0) 3). The Fe-cyclam conformation was found to be the controlling factor in determining the spectroscopic properties of the Fe^{III} O_2 −Cu^{II} core in 1. While all models show strong absorption below 450 nm, only the *trans-*III-cyclam, i.e., the end-on *μ*-

Figure 3. (A, B) Experimental UV−vis spectra of 1 and 2, respectively. (C, D) Time-dependent DFT (TD-DFT)-based UV−vis absorption spectra for 1-*trans*-b and 2-*cis*-b, respectively, calculated from the individual transitions (sticks) by Gaussian broadening with 80 nm full width at halfmaximum (fwhm). (E) Electron density difference maps (yellow: negative (loss) phase, green: positive (gain) phase, isovalue 0.0025 au) of the two main charge-transfer transitions at 610 and 649 nm in 1-*trans*-b.

*η*1 :*η*¹ peroxido structures exhibit an intense absorption band centered in the range λ_{LMCT} ^{cal} = 600−640 nm (vide infra), whereas the *cis*-V-cyclam/*μ*-*η*² :*η*¹ peroxido structures lack any strong absorption above 500 nm. At the same time, the end-on peroxido structures feature O−O stretching modes at 750− 767 cm^{-1} ($\Delta^{16/18}$ _O^{cal} = -37 to -40 cm⁻¹), while for the μ - η^2 : η^1 peroxido structures, they are in the range 852−875 cm[−]¹ $(\Delta^{16/18} \text{e}^{\text{cal}} = -43 \text{ to } -49 \text{ cm}^{-1})$. Thus, models with a $\mu \text{-} \eta^1 \text{-} \eta^1$ peroxido motif generally reproduce the experimental results for $1 (\lambda_{LMCT} = 615 \text{ nm}, \nu (O - \tilde{O}) = 739 \text{ cm}^{-1}, \Delta^{16/18}{}_{\tilde{O}} = -40 \text{ cm}^{-1}).$ In particular, considering the respective lowest-energy structures [\(Figure](#page-2-0) 2; see [Table](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S7), the calculated spectroscopic properties of 1*-trans-b* ($\lambda_{LMCT}^{cal} = 617$ nm, ν (O−O)^{cal} $= 737$ cm⁻¹, and $\Delta^{16/18}$ _O^{cal} = -37 cm⁻¹) are in excellent agreement with the experimental UV−vis and rRaman features of 1. Thus, although the corresponding *cis* and *trans* models cannot be compared energetically due to the presence of an additional axial CH₃CN ligand to the Fe^{III} ion in the *trans* structure, complex 1 can be safely assigned to an end-on *μη*1 :*η*¹ peroxido structure involving a [(CH3CN)(*trans*-IIIcyclam)FeIII−O2−CuII(*bent*-AN)]3+ (1-*trans*-b) structural motif as shown in [Figure](#page-2-0) 2. This is also supported by the identification of another 18 O-sensitive mode, involving the Fe $-$ O stretching vibration in the range of 608–627 cm⁻¹ (Δ ^{16/18}O^{cal} $= -28$ to -30 cm⁻¹) for the end-on peroxido structures, in close proximity to the experimentally observed 18O-sensitive band at 635 cm⁻¹ ($\Delta^{16/18}$ _O = −27 cm⁻¹) for 1. The corresponding Fe−O vibration is calculated at 565−592 cm⁻¹ $(\Delta^{16/18} \text{e}^{\text{cal}}) = -16$ to -22 cm^{-1}) for the $\mu \text{-} \eta^2 \text{:} \eta^1$ peroxido structures, which are significantly downshifted in energy relative to that of the experimentally observed value for 1.

The absorption spectrum of 1 was also analyzed in detail. For the absorption band peaking at 617 nm, we find two main contributions at 610 and 649 nm in the calculated spectrum (Figure 3C). From the TD-DFT difference density (Figure 3E) for the electronic transition at 610 nm, we readily can identify it as a ligand-to-metal charge-transfer (LMCT) band from the peroxido bridge to the Fe^{III} ion. Closer inspection by means of the corresponding natural transition orbitals (NTOs, [Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S6) shows that it is characterized by two orbital pairs: mainly an O_2^2 ⁻ π^* _v → Fe 3d_{xz} excitation (from the peroxido-Fe *π*-bonding MO into its *π*-antibonding counterpart, 65%), and a minor excitation (20%) from a mixed Fe 3d*xy*/Cu 3d*^z* 2 MO into a strongly delocalized *σ*-antibonding MO involving Fe $3d_z$ ², O_2 ² = π_{σ} , Cu $3d_{x^2-y^2}$ and orbitals on the N-donor atoms of both the AN and cyclam ligands. For the 649 nm band, the difference density and the NTOs indicate donor and acceptor orbitals delocalized mainly over the Cu ion, the peroxido ligand, and the nitrogens of the AN ligand. The donor orbital is composed of the Cu 3d*^z* ² orbital with substantial admixture from the peroxido oxygens (π^*) , while the acceptor orbital comprises the Cu $3d_{x,y}^2$ and the *σ*-antibonding ligand orbitals from the AN nitrogens and peroxido oxygens (*π***σ*). Thus, in the presented nonheme Fe^{III}−Cu^{II}-peroxido species 1, the specific O−O bridging motif is essential in defining both these transitions, such that a 600−650 nm band appears as a spectroscopic marker for this geometry.

Synthesis and Characterization of 2. An interesting structural aspect of 1-*trans*-b is the orientation of the amino hydrogen of the AN ligand in bent conformation, which is placed at distances of 3.92 and 4.11 Å from the peroxido O_{Cu} and O_{Fe} atoms, respectively, and possibly involved in a weak secondary interaction [\(Figure](#page-2-0) 2, [Scheme](#page-6-0) 3). Notably, the deuteration of the NH-group of AN resulted in an increased half-life of 1, further supporting an interaction of the −NH group with the peroxido moiety in 1 ([Figures](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) $S7-S8$). However, this interaction is too weak to effect any significant change in the Fe−O or O−O vibration modes, as corroborated by theoretical (the ${\Delta^{\scriptscriptstyle 1/2}}_{\rm H}^{\rm cal}$ isotope shift upon deuteration of the amino hydrogen is only 0.2 cm⁻¹ for the O-O stretch and 0.01 cm[−]¹ for the Fe−O stretch) and experimental rRaman (data not shown) studies. When the N−H group is replaced by a N−

Scheme 2. Overview of the Discussed Complexes and Intermediates and Their Reactivity

Figure 4. (A) UV−vis absorption spectral changes associated with the direct addition of DCHIm to 1 (dashed blue) and 2 (dashed pink) to form 1-DCHIm (dark blue) and 2-DCHIm (red), respectively (acetone/CH3CN 10:1 v/v mixture, −90 °C). (B) Zero-field Mößbauer spectra of 1- DCHIm (gray-dashed: experimental, dark blue: simulated main species) and 2-DCHIm (gray: experimental, red: simulated main species). Minor species are marked by asterisks, see Table S6 and [Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S12 for details.

Me group, a *μ*-*η*² :*η*¹ peroxido structure results in the corresponding $[(cis\text{-cyclam})\text{Fe}^{\text{III}}(\mu\text{-}\eta^2\text{:}\eta^1\text{-O}_2)\text{Cu}^{\text{II}}(\text{MeAN})]^{3+}$ complex 2. Experimentally, the iron center in 2 is also lowspin Fe^{III} as evident from zero-field Mößbauer studies (δ = 0.24 mm s⁻¹ and $\Delta E_{\rm Q}$ = -2.51 mm s⁻¹), but unlike the $S_{\rm t}$ = 1 ground state in 1, a $S_t = 0$ assignment is deduced for 2 based on the applied field Mößbauer, which is also consistent with EPR studies ([Figures](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S9 and S10). The different peroxido binding motifs and coupling situations in 1 and 2 are reflected in their different spectroscopic properties ([Figure](#page-3-0) 3A/B). Complex 2 lacks any UV−vis absorption feature above 400 nm and exhibits a higher peroxidic stretching frequency, *ν*(O−O) = 856 cm⁻¹ (¹⁶O₂; 803 cm⁻¹ for ¹⁸O₂, [Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S11), relative to that of 1 $(\nu(O-O) = 739 \text{ cm}^{-1})$. Our theoretical studies

predict a 2-*cis*-b structure with a *μ*-*η*² :*η*¹ peroxido binding motif ([Figure](#page-2-0) 2, [Table](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S8), which can reproduce the experimentally observed UV−vis absorption and the O−O stretching modes in 2 ([Figure](#page-3-0) 3 and [Table](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S1).

Extended X-ray Absorption Fine Structure (EXAFS) Analysis of 1 and 2. Analysis of the extended X-ray absorption fine structure (EXAFS) at the Fe and Cu K-edges of 1 and 2 [\(Figures](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S4, S5, Tables S2−S4) is in reasonable agreement with the bond lengths calculated by DFT for 1 *trans*-b and 2-*cis*-b. Satisfactory EXAFS simulations of the iron spectra of 1/2 were obtained with 4 N scatterers at a distance of 1.99 Å and additional O contributions at (mean) shorter $(1.79/1.73 \text{ Å})$ and longer $(2.68/2.12 \text{ Å})$ distances to the iron center that can be ascribed to the O atoms of the end-on

Figure 5. (A) UV−vis spectral changes associated with the conversion of 1 (blue) into 1-PhO• (purple) upon addition of 5 equiv of phenol (acetone/CH3CN 10:1 v/v mixture, −90 °C). (B) rRaman spectra of 1-PhO• (purple) and 1-Ph18O• (blue) measured with 568 nm excitation (2 mW) in an acetone- d_6 /CD₃CN 10:1 v/v solution. Solvent features are marked by asterisks.

peroxido unit in 1 (DFT: Fe−O1: 1.78 Å; Fe−O2: 2.72 Å) or the side-on peroxido unit in 2 (DFT: Fe−O1: 1.78 Å, Fe−O2: 1.99 Å). The apparent shortest Fe−O bonds in 1 and 2 from the EXAFS simulations may in part reflect contributions of $Fe^{IV}=O$ species in the samples (see below). Also, for the copper spectra, the simulations revealed two O atoms at 1.87/ 1.84 and 2.57/2.55 Å in 1/2, respectively (DFT: Cu−O2: 1.91/2.00 Å, Cu−O1: 2.62/2.93 Å) (see [Tables](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S2−S4 for details).^{[38](#page-9-0)} We note that in the presence of multiple species in the solution samples of the complexes, perfect agreement between interatomic distances from XAS and DFT is not expected, as a limited number of variable parameters (e.g., bond lengths) have to be used in EXAFS fit analysis to avoid data overinterpretation. Furthermore, low concentrations of 1 and 2 and the presence of scatterers from the ligand scaffold in similar distances did not allow for a reliable determination of Fe···Cu distances above 3.5 Å via EXAFS.

Axial Ligand Effect on the Peroxido Binding Motif. As suggested by DFT, the binding of an axial $CH₃CN$ ligand triggers the isomerization of the *cis*-cyclam moiety in 2 to *trans*-cyclam in 1, with the concomitant change of the peroxido binding motif at Fe from side-on to end-on ([Scheme](#page-4-0) 2 and [Figure](#page-2-0) 2). Consistent with this suggestion, the addition of 1,5 dicyclohexylimidazole (DCHIm) to a $CH₃CN/acetone$ solution of 2 at −90 °C led to the generation of a blue− green species 2-DCHIm ($t_{1/2}$ = 40 s at −90 °C) that shows the characteristic low-energy absorption feature at 640 nm [\(Figure](#page-4-0) [4](#page-4-0)A) associated with the μ - η ¹: η ¹ peroxido binding motif. The axial $CH₃CN$ ligand in 1 can also be replaced by DCHIm as evidenced by the red shift of the 615 nm band in 1 to 640 nm in 1-DCHIm ($t_{1/2}$ = 60 s at −90 °C). The Fe^{III} centers in 1-DCHIm and 2-DCHIm are electronically very similar as evidenced from their near identical Mößbauer parameters: for 1-DCHIm, $\delta = 0.28$ mm s⁻¹ and $|\Delta E_{\text{Q}}| = 1.87$ mm s⁻¹, and for **2**-DCHIm, $\delta = 0.29$ mm s⁻¹ and $|\Delta E_Q| = 1.81$ mm s⁻¹ [\(Figure](#page-4-0) [4](#page-4-0)B, [Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S12 and Table S6). Both are silent in conventional X-band EPR measurements in perpendicular mode indicating an integer total spin state [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S13). Although the metastable nature of 1-DCHIm and 2-DCHIm prevented us from measuring their *ν*(O−O) vibration modes, based on the similarity in their absorption and Mößbauer features, the presence of a $[(DCHIm)(trans-cyclam)Fe^{III}(\mu-\eta^1\cdot\eta^1-O_2)$ $Cu^{II}(AN/MeAN)³⁺$ motif can safely be concluded in both the complexes.

Reactivity Studies of 1 versus 2 with PhOH. The different electronic structures of 1 and 2 are also reflected in their reactivity properties. For example, complex 1, and not complex 2, exhibits reactions with phenol, possibly stressing the importance of the $S_t = 1$ ground state and the weak $-NH$ secondary interaction in the hydrogen atom transfer (HAT) reactivity of the peroxido complexes. Addition of phenol to 1 showed a fast reaction that could be monitored by UV−vis (Figure 5A). The typical absorption feature of 1 at 615 nm vanished, while a new purple species 1-PhO• is observed with an absorption maximum at 540 nm ($\varepsilon = 1.05$ L mmol⁻¹ cm⁻¹) and a broad feature between 600 and 1000 nm. When substituted phenols (*p*-kresol, *p*-methoxyphenol, and *p*chlorophenol) were added to 1, a shift of the absorption maximum $(1 - P^{\text{Me}}PhO^{\bullet})$: $\lambda_{\text{max}} = 525$ nm, $1 - P^{\text{OMe}}PhO^{\bullet}$: $\lambda_{\text{max}} =$ 580 nm, and $1^{p\text{-}\text{Cl}} \text{PhO}^*$: $\lambda_{\text{max}} = 570$ nm) can be observed, corroborating the binding of phenol-derived species [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) [S14](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf)). Gas chromatography-mass spectrometry (GC-MS) analyses of the reaction mixtures at room temperature revealed the formation of coupled phenol products (4,4′-bis(2,6-di*-tert*butylphenol) (19%), 3,3′,5,5′-tetra-*tert*-butyldiphenoquinone (10%), and 2,6-di-*tert*-butyl-1,4-benzoquinone (10%) for reaction with 2,6-di*-tert*-butylphenol and 5,5′-dimethoxy[1,1′ biphenyl]-2,2′-diol (∼20%) for reaction with 4-methoxyphenol) in moderate to high yields ([Figures](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S15, S16 and Table [S5](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf)). Further spectroscopic investigations show that the oxidative coupling of the phenol is accompanied by the reductive cleavage of the peroxido bridge in 1 and the formation of copper(II) and iron(III)-phenoxyl radical species. The initially formed oxoiron(IV) species presumably reacts with excess PhOH to generate the iron(III)-phenoxyl radical species ([Scheme](#page-6-0) 3). Indeed, the Mößbauer spectrum of 1-**PhO[•]** ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S17) shows the formation of a major Fe^{III} lowspin species in 70% yield ($\delta = 0.25$ mm s⁻¹, $|\Delta E_{Q}| = 2.57$ mm (s^{-1}) and a residual Fe^{IV} (δ = 0.07 mm s⁻¹, $|\Delta E_{\text{Q}}|$ = 1.92 mm s⁻¹) species in 25% yield. Notably, Mößbauer parameters of the Fe^{IV} species are comparable to the parameters for an authentic $[Fe^{IV}(O)(trans-cyclam)(CH₃CN)]²⁺$ reported previously ($\delta = 0.05$ mm s⁻¹, $\Delta E_{\text{Q}} = 2.49$ mm s⁻¹, [Table](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S6);³⁴ the differences in δ and $\Delta E_{\rm Q}$ values are plausibly attributed to additional interactions from the Cu^{II}, PhO[•], or OH moieties (see [Scheme](#page-6-0) 3). The EPR spectrum reveals the formation of multiple overlapping $S = 1/2$ species ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S18) arising from Cu^{II} centers in near-quantitative yields. rRaman studies were performed (Figures 5B and [S19\)](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) to confirm the $1 - XPhO^{\bullet}$ assignment. Two features were observed for 1-PhO[•] at 815 and 586 cm⁻¹, which showed no shift when ¹⁸O labeled 1 was used in the reaction, but shifted to 804 and 574 cm^{-1} , respectively, when ¹⁸O-labeled PhOH was used. The positions of these bands were also found to be sensitive to the nature of Scheme 3. Proposed Mechanism for the Reaction of 1 and 1-DCHIm with Phenol via a H-Bond-Assisted Mechanism

the *para*-substituents [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S19). Based on a recent study where we spectroscopically characterized a phenoxyl radical bound to an Fe^{II} center^{[39](#page-9-0)} and previous reports of metal bound phenoxyl radicals,^{[40,41](#page-9-0)} we assign the 586 and 815 cm⁻¹ bands to Fe−O and in-plane phenyl ring bending vibrations, respectively, for a metal bound PhO• radical.

The $S_t = 1$ [(CH₃CN)(*trans-cyclam*)Fe^{III}(μ - η ¹· η ¹-O₂)- $Cu^H(AN)³⁺$ core in 1 thus undergoes a reductive O-O bond cleavage step via a net H-atom transfer from phenol to generate [(AN)CuI I−OH]+, [FeIV(O)(*trans*-cyclam)- $\text{[CH}_{3}CN]^{2+}$ and PhO[•] (Scheme 3). A previous study on a heme $Fe^{III}-O_2-Cu^{II}$ species^{[39](#page-9-0)} has shown that the phenol can facilitate O−O cleavage via two possible mechanisms, which differ by the amount of proton transfer prior to the transition state. One mechanism involves nearly complete proton transfer from the phenol to the peroxo before the barrier. The second mechanism involves O−O homolysis by phenol H-bonding to peroxo, with the proton transfer occurring after the barrier. The two mechanisms can be distinguished by kinetic isotope effect (KIE) data for an H-atom-donating phenol inducing O− O cleavage. A KIE of 1.43 has been determined for the O−O bond cleavage step [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S20) in 1, consistent with the hydrogen-bonded mechanism (Scheme 3); 42 a much higher KIE (>5) is expected for complete proton transfer before the barrier. This H-bond assistance is further supported by an observed secondary KIE of 1.20 ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S20) in the phenol reaction upon deuteration of the −NH group of the AN ligand in 1. An increase of the reaction rate for the formation of $1 - XPhO^o$ with the pK_a of the corresponding substituted phenols ($k_2 = 0.38$ for 1^{-p} ^{Me}PhO[•], $k_2 = 0.83$ for 1-PhO[•], k_2 = 1.53 for 1-*^p***‑**ClPhO• , [Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S21) is also consistent with the mechanism shown in Scheme 3.

Effect of the Axial Ligand on PhOH Reactivity of 1. The complex $[Fe^{IV}(O)(trans-cyclam)(CH₃CN)²⁺ undergoes$ a side reaction with excess PhOH to generate [(*trans*- $\text{cyclam})\text{Fe}^{\text{III}}(\text{PhO}^{\bullet})(\text{OH})]^{2+}.$ This reaction is presumably triggered by the replacement of the axial $CH₃CN$ ligand in $[Fe^{IV}(O)(trans-cyclam)$ $(CH_3CN)]^{2+}$ by PhOH, followed by HAT by the Fe^{IV} = O center. Indeed, the side reaction is completely stopped in the reaction of 1-DCHIm with PhOH, where replacement of the stronger axial ligand DCHIm by

PhOH is not favored, and the Mößbauer analysis of the reaction mixture shows the formation of $[Fe^{IV}(O)(trans$ cyclam)(DCHIm)]²⁺ in near-quantitative yields (δ = 0.05 mm s^{-1} , $|\Delta E_Q| = 2.00$ mm s^{-1} , Table S6 and [Figures](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S22–S24). Consistent with our presumption, an authentic sample of $[(DCHIm)(trans-cyclam)Fe^{IV}=O]²⁺ obtained by replacement$ of the CH₃CN ligand in $[(CH_3CN)(trans-cyclam)Fe^{IV}=O]²⁺$ by DCHIm at −50 °C did not show any reactivity with PhOH ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S24). Notably, both 1-DCHIm and 2-DCHIm show increased yields of the coupled phenol products compared to 1 (Table S5 and [Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S25), reflecting the release of the phenoxyl radical instead of coupling to the iron center.

Reactions of 1 and 2 with PPh₃. Intermediate 1 also reacts at -90 °C with PPh₃ forming OPPh₃ (identified by GC-MS, [Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S26) and a new species 1-Cu with a UV−vis absorption feature at 675 nm ($\varepsilon = 0.19$ L mmol⁻¹ cm⁻¹) that further decays above -50 °C ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S27). Notably, both 2 and 3 also reacted with PPh_3 , albeit with rates, respectively, 1 and 3 orders of magnitude slower than 1 ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S28−S30). This confirms that the O−O bond in 1 is more activated relative to that in 2 and 3. In particular, the higher reaction rates of 1 and 2, relative to 3, demonstrate the positive influence of Cu in the reductive O−O bond cleavage reaction. The absorption spectrum and Mößbauer parameters [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) [S31,](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) $\delta = 0.07$ mm s⁻¹, $|\Delta E_{Q}| = 2.08$ mm s⁻¹) of 1-Cu are similar compared to that reported for $[Fe^{IV}(O)(trans-cyclam)$ - $(CH₃CN)$]OTf₂ ([Table](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S6).^{[34](#page-9-0)} The XANES spectra at the Fe K-edge of 1-Cu and 2-Cu ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S32) show an intense preedge peak at about 7115 eV that is typical for oxoiron complexes. EXAFS analysis additionally revealed a short iron− oxygen bond of 1.62 Å in ca. 50% of the complexes [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S33, [Table](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S3), which is indicative of an $Fe^{IV}=O$ species. The ESI-MS spectrum of 1-Cu shows a signal at $m/z = 706.1$ with mass and isotope distribution pattern consistent with a $[Fe(O)(cyclam)(OTf)Cu(AN)(Cl)]^+$ assignment, which is shifted by two and one mass units upon ¹⁸O- and ⁵⁷Felabeling, respectively ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S33). 1-Cu is EPR silent in perpendicular mode X-band EPR ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S34). All of these point to the formation of $\text{Fe}^{\text{IV}}=$ O, Cu^I, and PPh₃O products in the reaction of the $S_t = 1$ $[(CH_3CN)(trans-cyclam)Fe^{III}(\mu \eta$ ¹: η ¹-O₂)Cu^{II}(AN)]³⁺ (1) with PPh₃.

■ **CONCLUSIONS**
In summary, a nonheme iron(III)-superoxo complex 3 reacts with $[Cu^{I}(AN)]$ generating a low-spin iron(III)-peroxidocopper(II) complex 1 with a $S_t = 1$ ground state. 1 undergoes efficient O−O bond cleavage in the presence of H-bonded PhOH and a strong axial ligand DCHIm leading to the stoichiometric formation of an $oxoiron(IV)$ species. This process has some relevance to the CcO catalytic cycle. A key matter that remains unsolved in CcO is the reaction coordinate connecting intermediates A and P_M . While it has not been observed experimentally, the involvement of a peroxido level intermediate $(I_p$ in [Scheme](#page-1-0) 1) in CcO has been suggested in many studies, whereby a $S_t = 1$ $[Fe^{III}(\mu - \eta^1 \cdot \eta^1 - O_2)Cu^{II}]$ core undergoes a fast low-barrier (<12.4 kcal mol[−]¹) O−O bond cleavage via H-atom abstraction from Tyr-OH that is involved in a hydrogen-bonding network with water.^{5,[43](#page-9-0)–[45](#page-9-0)} In the present study, complex 1 not only reproduces the proposed S_t $= 1$ [Fe^{III}(μ - η ¹: η ¹-O₂)Cu^{II}] motif, suggested for I_P, but the −NH group of the AN ligand also provides the platform for HAT from a H-bonded phenol that is required for the efficient O−O bond cleavage step. The fast and efficient O−O cleavage

occurring in 1 can be attributed to the observed ferromagnetic coupling between the low-spin Fe^{III} and Cu^{II} centers in 1, which, as previously proposed based on theoretical stud-ies^{[16](#page-8-0)−[19](#page-9-0)} in \tilde{I}_p ensures efficient reductive O−O rupture to form $S = 1$ Fe^{IV}=O and antiferromagnetically coupled Cu^{II} and PhO• via electron transfer from PhO[−] without the necessity of any spin-surface crossing. However, the exchange-coupled Cu^{II}–PhO[•] species is presumably transient in our case and decays to mononuclear Cu^{II} (detected in EPR, data not shown) and coupled phenol products. Consistent with this proposition, a related low-spin Fe^{III}-peroxido-Cu^{II} species 2 in otherwise similar Fe- and Cu-coordination environments, but with an $S_t = 0$ ground state due to antiferromagnetic coupling between the $\mathrm{Fe}^{\mathrm{III}}$ and $\mathrm{Cu}^{\mathrm{II}}$ centers, did not show any reactivity with PhOH.

Complexes 1 and 2 differ in their peroxido binding mode to Fe^{III} (end-on vs μ - η^2 : η^1 in 1 and 2, respectively), which is also reflected in their *ν*(O−O) modes and the transitions of the O_2^2 ⁻ to Fe^{III} charge transfer in the UV−vis absorption spectrum. Considering the substantial differences in structural parameters for the two peroxido bridging motifs and of the Fe coordination geometries, it is apparent that metal−ligand orbital interactions and MO energies will be largely different in 1 and 2. The fact that in 2, the Fe^{III} ion is directly ligated by both instead of only one O donor leads to a considerably different interaction of the peroxo *π** with the Fe 3d orbitals. For example, the $\pi^*_{\,\,v}$ orbital exhibits stronger interactions with the Fe 3d*xy* and 3d*yz* orbitals at the expense of the interaction with its 3d_{*xz*} orbital compared to the situation in complex 1. It can thus be understood that the dominant excitation within the strongest LMCT transition (610 nm) in 1 does not exist as such in 2 due to a mitigated *π*-bonding interaction between the O_2^2 ⁻ π^* _v and the Fe 3d_{xz} orbital.

Comparison of 1 to the very well characterized heme-based iron-peroxo-copper adduct $[(DCHIm)(F_8)Fe-O_2-Cu(AN)]^+$ by Karlin and Solomon, $12,28,42$ $12,28,42$ $12,28,42$ $12,28,42$ which they have identified to possess a *μ*-*η*¹ :*η*1 /end-on binding mode, shows that they resemble each other in that they both feature similar LMCT excitations from the O_2^2 ⁻ π^* _v orbital into the Fe 3d_{xz} and Cu 3d*^x 2* −*y ²* orbitals, albeit at significantly lower energies (789 and 951 nm, respectively) and of lower intensity. As orbitals from the equatorial ligands to the Fe^{III} ions contribute only insignificantly in both cases, comparison between the heme and nonheme complexes is legitimate in this respect. The higher energy of the transition from the *π*-bonding $O_2^2 - \pi^*$ _v/ Fe $3d_{xz}$ MO into its π -antibonding counterpart in 1 than in $[(DCHIm)(F_8)Fe-O_2-Cu(AN)]^+$ indicates a stronger Fe–O bond, in line with the somewhat shorter calculated Fe···O distance (1.79 in 1 vs 1.82 Å in $[(DCHIm)(F_8)Fe-O_2-$ Cu(AN)]+). At the same time, the considerably lower *ν*(O− O) stretching vibrational energy of 739 cm⁻¹ in 1 compared to 794 cm^{-1} in the heme complex (Table 1) demonstrates concomitant weakening of the O−O bond, here consistent with the longer O···O distance (1.45 in 1 vs 1.40 Å in $[(DCHIm)(F_8)Fe-O_2-Cu(AN)]^+$).

In heme-copper model chemistry, a lower value for *ν*(O−O) is in general correlated with a side-on binding mode to one or both metal centers. Table 1 shows selected examples of hemecopper model complexes (note that the $[(F_8)Fe-O_2-$ Cu(AN)] system also incorporates the AN ligand at the copper site) with a linear correlation between the observed O− O stretching frequency and the Fe···Cu-distance, correspond-ing to the assignment from side-on^{[11,13](#page-8-0),[24](#page-9-0)} to end-on^{[11,](#page-8-0)[27](#page-9-0),[28](#page-9-0)} via a

Table 1. Comparison of the Frequencies of the O−O Stretching Modes and Selected Bond Lengths of Fe−O₂−Cu Intermediates

compound	ν (O-O) $\left[\Delta^{\scriptscriptstyle{16/18}}\right]$ (cm^{-1})	$O \cdot O$ (Å)	Fe \cdots Cu (Å)
1	739 $[-40]$	nd	nd
1-trans-b (DFT; end-on)	735 [-39]	1.45	4.21
2	856 [-50]	nd	nd
2-cis-b (DFT; $\mu - \eta^2$: η^1 -O ₂)	862 [-49]	1.43	3.71
$[F_8Fe-O_2-CuAN]^+$ (side- on ²⁸	756 [-48] (DFT 821)	1.46 (DFT)	3.63 (XAS) 3.73 (DFT)
$[(TMP)Fe-O,-]$ Cu(5MeTPA) ⁺ $(\mu - n^2 : n^1 -$ Q_2 ²²	790 $[-44]$	1.46 (XRD)	3.92 (XRD)
$[(DCHIm)F_8Fe-O_2-$ CuAN ^{\vert+} (end-on) ²⁸	796 $[-42]$ (DFT 840)	1.40 (DFT)	4.01 (XAS) 4.01 (DFT)
resting oxidized state of CcO	750 ^a	$1.49 - 1.70$	$4.6 - 4.8$

^a Alternatively assigned as an O-sensitive histidine breathing mode.^{[33](#page-9-0)}

 μ - η ²: η ¹- $(O_2)^{22,27}$ $(O_2)^{22,27}$ $(O_2)^{22,27}$ binding mode.^{5,12} Similar observations and assignments were also made for other heme-based models. Studies on the resting oxidized states of CcOs show a more complex situation.^{1[,29](#page-9-0)–[33](#page-9-0)} While the investigated structures and trapped intermediate states are not directly involved in the catalytic cycle of CcO, they are thought to provide valuable insights into the character of actual intermediate I_p. All studies agree on a long Fe···Cu distance of about 4.6−4.8 Å, which would only allow for an end-on bound peroxido unit, but the O−O-bond length could not be determined precisely.^{[5](#page-8-0)} The O−O frequency that is of course correlated with the bond length is also a matter of discussion. In initial studies, a rRaman feature at 750 cm[−]¹ was observed in one resting oxidized CcO example and assigned to the O−O stretching mode,^{[7](#page-8-0),[32](#page-9-0)} but a recent study discussed this feature as the wagging of the axial histidine that is sensitive to the peroxido unit.^{[33](#page-9-0)} One reason why the assignment of the 750 cm[−]¹ band to the O−O mode was questioned was the discrepancy between its low energy, indicating a side-on binding mode in heme chemistry while the environment in the active center would presuppose an end-on bound O_2^2 ⁻. The present report of the $S_t = 1$ [(CH₃CN)- $(trans-cyclam)Fe^{III}(μ - η ¹: η ¹-O₂)Cu^{II}(AN)³⁺ complex 1, which$ can reproduce both the UV−vis absorption and rRaman features associated with the as-isolated CcO, corroborates that a bridging peroxido moiety in an end-on bound O_2^2 ⁻ structure and with a significantly weak O−O bond can be accommodated within the Fe \cdots Cu distance of >4.0 Å that is established in CcO. Whether the 750 $\rm cm^{-1}$ signal in the resting state of CcO can be possibly assigned as the O−O-vibration mode associated with an end-on $Fe^{III}-O_2-Cu^{II}$ peroxido binding motif is now an intriguing question, which needs further investigation.

■ **ASSOCIATED CONTENT**

\bullet Supporting Information

The Supporting Information is available free of charge at [https://pubs.acs.org/doi/10.1021/jacs.4c04492.](https://pubs.acs.org/doi/10.1021/jacs.4c04492?goto=supporting-info)

> Instrumental and physical methods, synthesis of compounds and intermediates, reactivity and kinetic data, theoretical studies, Figures S1−S34 (further characterization and reactivity data), and Tables S1− S8 (reactivity and DFT data) [\(PDF](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf))

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Notes

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(36) Notably, the *ν*(O−O) and *ν*(Fe−O) vibrational modes are both very intense in 1 and appear with nearly equal intensities [\(Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) [S2C](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf)). The decay of 1 leads to a species also with a vibration mode at 635 cm[−]¹ ([Figure](https://pubs.acs.org/doi/suppl/10.1021/jacs.4c04492/suppl_file/ja4c04492_si_001.pdf) S2D). However, this mode is significantly weaker relative to the Fe−O vibration mode in 1.

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