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Mechanism of phenol oxidation by heterodinuclear Ni-Cu bis(µ-oxo) complexes involving nucleophilic oxo groups

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Received (in XXX, XXX) Xth XXXXXXXXX 20XX, Accepted Xth XXXXXXXXX 20XX
DOI: 10.1039/b000000x

Oxidation of phenols by heterodinuclear CuIII(µ-O)xNiIII complexes containing nucleophilic oxo groups occurs by both proton coupled electron transfer (PCET) and hydrogen atom transfer (HAT) mechanisms; the exact mechanism depends on the nature of the phenol as well as the substitution pattern of the ligand bound to Cu.

The conversion of phenols to phenoxyl radicals plays a vital role in a number of biological systems. Photosystem II is the most prominent example,1,2 but evidence exists that similar processes are involved in the function of several other biochemical systems.3 The oxidative chemistry of phenols has also other potential biological application, notably their antioxidant properties.4,5 Additionally, oxidative dehydrodimerization of phenols is an important class of reactions, being involved in the first stages of natural processes such as lignin formation.6-10 Last but not least, phenol oxidation has important synthetic applications.11-13 Metal mediated oxidation of phenols by a PT-ET mechanism is unknown in the literature, although this mechanism has been previously invoked for the oxidation of phenols by organic radicals.16

Scheme 1. Possible reaction pathways for the oxidation of phenols.

Very recently we reported the isolation and spectroscopic characterization of a novel mixed-metal NiIII-CuIII bis(µ-oxo) complex (Scheme 2), [MeAN]CuIII(µ-O)xNiIII[bis(µ-oxo)] (1, MeAN = tert-butylbenzenesulfonamide).17,18 Moreover, the formation of a phenoxyl radical from a neutral phenol can occur via direct hydrogen atom transfer (HAT) or via a proton-coupled electron transfer (PCET) (Scheme 1) process. For the HAT and PCET mechanisms, homolytic O-H bond cleavage constitutes the rate-determining step of the reaction. In the HAT mechanism, the proton and electron of the H· radical both come from the same orbital. Conversely, proton and electron transfers are both rate determining for the PCET process, but occur from different orbitals in a concerted mechanism.12 Alternatively the proton and electron transfers can be uncoupled (PT-ET) with either proton transfer (PT) or the electron transfer (ET) being the rate determining step (Scheme 1). A HAT mechanism has been established for the phenol oxidation mediated by the terminal MnV-oxo (TBP8Cz)MnV0 (1) (TBP8Cz = octakis(para-tert-butylphenyl)corrolazinato3−)13 and CrIII-superoxo, [CrIII(TMC)(O2)(Cl)]+ (TMC = 1,4,8,11-tetramethyl-1,4,8,11-tetraazacyclotetradecane) complexes.14 In contrast, the reactions of phenols with two distinct homodinuclear dicopper-dioxygen complexes having bis (µ-oxo)dicopper(III) and (µ-η2:η2-peroxo)dicopper(II) cores exhibit a PCET mechanism.15 Metal mediated oxidation of phenols by a PT-ET mechanism is unknown in the literature, although this mechanism has been previously invoked for the oxidation of phenols by organic radicals.16

Scheme 2. Scheme showing the reactions of 1 and 2 with different substituted phenols.

Electronic Supplementary Information (ESI) available: [Synthesis and characterization of 2; additional kinetic data]. See DOI: 10.1039/b000000x/
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N,N,N',N'-pentamethyldipropylentriamine and L = [HC(CMeNC6H3)(P3)2]]. Unlike typical homodinuclear bis(µ-oxo) complexes, this species contains nucleophilic oxo groups that can carry out deoxygenation of aldehydes. We initiated the present study to investigate whether the PCET mechanism established for phenol oxidation by homodinuclear Cu2O2 cores having electrophilic oxo groups also persists for the Cu(µ-O)2Ni core of 1, which has nucleophilic oxo groups. We report a systematic study of the oxidation of neutral phenols by 1 and provide deeper insights in the O-H bond activation mechanism.

Figure 1. A) Changes in the absorption spectra associated with the reaction of 1 (0.2 mM in CH2Cl2) with 2,4-di-tert-butylphenol (100 equiv) at -90 °C. The inset shows the time trace of the decay of the 895 nm band upon addition of 2,4-di-tert-butylphenol, which can be fitted to a pseudo first-order kinetic model to obtain a first-order rate constant (kobs). Figures 1B and 1C depict the linear dependence of kobs on the concentrations of 2,4-di-tert-butylphenol and the mono-deuterated analogue for the reactions with 1 and 2, respectively.

The Cu(µ-O)2Ni complex 1 was generated by the reaction of equimolar amounts of [MeANCu][BF4] and [L(Ni)O2] complexes at -90 °C in CH2Cl2 as previously reported. Treatment of a preformed solution of 1 with excess 2,4-di-tert-butylphenol led to pseudo-first order decay of the characteristic absorption feature of 1 at 895 nm (Figure 1A). The rate constant increases proportionally with the substrate concentration (Figure 1B), affording a second-order rate constant, k2, of 2.55 M⁻¹s⁻¹ at -90 °C. These kinetic behaviors indicate that the reaction between 2,4-di-tert-butylphenol and 1 is a simple bimolecular process.

Table 1. Oxidation potential (E°0) of ArOH and the second order rate constants (k2) and the yields (based on ArOH) of the phenol oxidation products for reactions of ArOH with 1 and 2. C-C coupled products are observed in all cases except for 2,4,6-tri-tert-butylphenol, where formation of 2,4,6-tri-tert-butylphenoxynyl radical occurs.

<table>
<thead>
<tr>
<th>Phenol</th>
<th>E°0 (V)</th>
<th>k2 (M⁻¹s⁻¹)</th>
<th>T/K</th>
<th>(RT/F) ln k²</th>
<th>% yield</th>
</tr>
</thead>
<tbody>
<tr>
<td>2,4-DTBP</td>
<td>1.46</td>
<td>2.55</td>
<td>183.16</td>
<td>0.015</td>
<td>45</td>
</tr>
<tr>
<td>2,6-DTBP</td>
<td>1.62</td>
<td>0.03</td>
<td>223.16</td>
<td>-0.067</td>
<td>38</td>
</tr>
<tr>
<td>2,4,6-TTBP</td>
<td>1.58</td>
<td>0.15</td>
<td>223.16</td>
<td>-0.037</td>
<td>95</td>
</tr>
<tr>
<td>4-phenylphenol</td>
<td>1.52</td>
<td>4.23</td>
<td>183.16</td>
<td>0.023</td>
<td>42</td>
</tr>
<tr>
<td>4-phenoxylphenol</td>
<td>1.49</td>
<td>10.92</td>
<td>183.16</td>
<td>0.038</td>
<td>42</td>
</tr>
</tbody>
</table>

DTBP = di-tert-butylphenol, TTBP = tri-tert-butylphenol, E°0 (V) vs SCE values are taken from ref 15 except for 4-Br-2,6-DTBP, which has been determined by cyclic voltammetry (see Electronic Supplementary Information for details). Reaction of complex 1 with 4-Br-2,6-DTBP is very slow, and hence, no kinetic measurements could be performed.

The k2 values (Table 1) obtained for the reaction of 1 with different 4-substituted phenols (ArOH) were found to be dependent on the E°0 values of the phenols; in general, k2 increased with decreasing E°0 for all investigated phenols with the exception of 2,4-di-tert-butylphenol. Plots of (RT/F) ln k² versus E°0 afford a good linear correlation with a slope of -0.81±0.05 (Figure 2A); the corresponding value for 2,4-di-tert-butylphenol is again significantly below the trend line, which may be suggestive of a change in mechanism for this substrate. If electron transfer from phenols to 1 is rate-determining, and is followed by fast proton transfer, then according to Ram and Hupp19 the slope of the (RT/F)lnk² versus E°0 plot should be -0.5, as one would expect from Marcus theory22 for a pure electron transfer reaction. On the other hand, if proton transfer is rate-determining and the electron transfer is in equilibrium then the slope should be -1.0. If the rates of electron transfer and proton transfer are comparable and thereby coupled to each other (PCET mechanism), a value between -0.5 and -1.0 would be expected.
In contrast, the $k_2$ values for a HAT mechanism are expected to be constant irrespective of the $E'^0$ values, as has been reported previously for the HAT reactions with N,N-dimethylanilines and for the oxidation of phenols by cumylperoxyl radical. The slope of -0.81 in Figure 2A therefore establishes a PCET mechanism for the oxidation of all investigated phenols by 2,4-di-tert-butylphenol, for which a different mechanism may be involved.

**Figure 2.** Plots of (RT/F)ln($k_2$) against the oxidation potential ($E'^0$) of ArOH for the reactions of ArOH with 1 (Figure 2A) and 2 (Figure 2B) in CH$_2$Cl$_2$.

In order to obtain additional mechanistic insights, deuterium kinetic isotope effects (KIE) on the second-order rate constant $k_2(O-H)/k_2(O-D)$ were measured for 4-phenoxynaphthalene and 2,4-di-tert-butylphenol. A deuterium KIE of 1.66 (Figure S4) at -90 °C was determined for 4-phenoxynaphthalene, indicating that proton transfer is involved in the rate determining step. This value is smaller than the KIE values obtained for most metal-oxo mediated HAT reactions (3 – 25). It is nearly the same as the KIE reported for the PCET reaction of guanidine with Ru(bpy)$_3^{2+}$, as well as those of 4-substituted phenols and dicopper-dioxygen complexes. Interestingly, 2,4-di-tert-butylphenol exhibited a significantly larger KIE of 4.72 (Figure 1B), which is in the typical range for a HAT process. The KIE values determined for the oxidations of 4-phenoxynaphthalene and 2,4-di-tert-butylphenol by 1 may, therefore, point to the involvement of a PCET mechanism for 4-phenoxynaphthalene and a HAT mechanism for 2,4-di-tert-butylphenol, which would be also consistent with the (RT/F)ln($k_2$) versus $E'^0$ plot shown in Figure 2A.

Karlin et al. have previously reported that the electronic structure of the dicopper-dioxygen cores supported by the N,N-dimethylpropylamine) forms a Cu$_{II}$(μ-O)$_2$ core in the presence of dioxygen, the corresponding (MeAN)Cu$_{II}$ complex affords a Cu$_{II}$(μ-peroxo) structure. In an effort to understand the role of the ligand structure of the copper precursor on the structure and reactivity of heterodinuclear bis-μ-oxo intermediates, we treated (AN)Cu$_{II}$ with the [L(Ni)O$_2$] complex at -90 °C in CH$_2$Cl$_2$ to generate 2 in near-quantitative yields (from EPR quantification studies). Complex 2 possesses a half-life of 9000 seconds at -50 °C, an order of magnitude higher than 1 ($t_{1/2} = 900$ s at -50 °C). However, the accumulated spectroscopic (absorption, X-ray absorption near-edge, resonance Raman, and electron paramagnetic resonance) properties (Figures S5-S6) of 2 are almost indistinguishable from those of 1, indicating that 2 also possesses a heterodinuclear Cu$_{II}$(μ-O)Ni$_{II}$ core similar to 1 (Scheme 2). Thus, unlike homodinuclear dicopper-dioxygen chemistry, where MeAN stabilized a μ-peroxo core, and AN stabilized a bis (μ-oxo) core, both of the MeAN and AN ligands lead to formation of a heterodinuclear Cu-Ni bis (μ-oxo) complex.

The reactivity of 2 was also investigated with various 4-substituted phenols and the second-order rate constants (Table 1, Figure S7) were compared with those obtained for 1. A plot of (RT/F)ln($k_2$) versus $E'^0$ for the reaction of 2 affords a good linear correlation with a slope of -0.61±0.03 (Figure 2B), which supports a PCET mechanism for the oxidation of phenols. In particular, reaction of 2,4-di-tert-butylphenol with 2 follows the linear correlation, in contrast to 1, where it appeared outside the linear trend. The KIEs for the second-order rate constant $k_2(O-H)/k_2(O-D)$ was also determined, with values of 1.71 (Figure 1C) and 1.77 (Figure S8) for the reactions of 2 with 2,4-di-tert-butylphenol and 4-phenoxynaphthalene, respectively. These KIEs are similar to the KIE value of 1.66 obtained for the reaction with 1 with 4-phenoxynaphthalene, but significantly lower than the value of 4.72 obtained for the corresponding reaction of 1 with 2,4-di-tert-butylphenol.

In summary, the oxidation of phenols mediated by the heterodinuclear Cu$_{II}$(μ-O)$_2$Ni$_{II}$ complex 2 is shown to proceed via a proton coupled electron transfer mechanism based on the rate dependence of the reaction on the one-electron oxidation potentials of the phenol substrates, as well as deuterium kinetic isotope effects of magnitude less than 2. Thus, while the oxygen atoms of the Cu$_{II}$(μ-O)$_2$Ni$_{II}$ core in 2 are nucleophilic, they prefer to oxidize phenol by a concerted PCET mechanism similar to what has been observed before for the corresponding Cu$_{II}$(μ-O)$_2$Cu$_{II}$ species involving electrophilic oxygen atoms. In contrast, for complex 1, which differs from 2 with respect to the substitution pattern of the ligand attached to the Cu center, but possesses identical spectroscopic properties, both HAT and PCET mechanisms may be feasible for the oxidation of phenols. Specifically, the oxidation of 2,4-di-tert-butylphenol proceeds by a HAT mechanism, while oxidations of 2,6-di-tert-butylphenol, 2,4,6-tri-tert-butylphenol, 4-phenoxynaphthalene and 4-phenoxynaphthalene proceed by a PCET mechanism. The different mechanisms observed for the oxidation of 2,4-di-tert-butylphenol by 1 and 2, therefore, highlights the importance of subtle electronic changes in modulating the reactivity of biologically relevant metal-dioxygen intermediates.

We gratefully acknowledge financial support of this work from the Cluster of Excellence 'Unifying Concepts in Catalysis' (EXC 314/1), Berlin. XAS data were obtained on beamline X3B of the National Synchrotron Light Source (Brookhaven National Laboratory, Upton, NY, USA), which is operated by the Case Western Reserve University and Brookhaven National Laboratory under contract number DE-AC02-98CH10886.
Western Reserve University Center for Synchrotron Biosciences, supported by NIH Grant P30–EB–009998. NSLS is supported by the United States Department of Energy, Office of Science, Office of Basic Energy Sciences, under Contract DE–AC02–98CH10886. We also thank Prof. Dr. Peter Hildebrandt and Dr. Uwe Kuhlmann for the resonance Raman measurement of 2 and Prof. Dr. Matthias Driess and Dr. Shenglai Yao for the supply of the nickel superoxide precursor.

Notes and references

21 The formation of the bis(hydroxo)CuIINiII product in scheme 2 is suggested on the basis of the analysis of the reaction mixture of 1 and 2,4-di-tert-butylphenol by ESI-MS (Figure S2) and EPR (Figure S3) spectroscopic methods. We emphasize, however, that the formation of other metal-containing products in addition to the bis(hydroxo) species cannot be excluded at this point.
Both PCET and HAT mechanisms can be feasible for the oxidation of phenols by heterodinuclear Cu(II)(µ-O)(µ-N) complexes.