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### Fast reduction of a copper center in laccase by nitric oxide and formation of a peroxide intermediate.

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**Abstract:** The rapid reduction of one of the copper atoms (type 2) of tree laccase by nitric oxide (NO) has been detected. Addition of NO to native laccase in the presence of oxygen leads to EPR changes consistent with fast reduction and slow reoxidation of this metal center. These events are paralelled by optical changes that are reminiscent of formation and decay of the peroxide intermediate in a fraction of the enzyme population. Although formation of this species is only possible if the trinuclear copper cluster (type 2 plus type 3) is fully reduced, this condition can be met if in a fraction of the enzyme the type 3 coppers are already reduced before addition of NO. Our data are consistent with this assumption. We have suggested recently that fast reduction of copper is the mechanism by which NO interacts with the dinuclear centre in cytochrome c oxidase. The present experiments using laccase strongly support this view, and suggest this reaction as a general mechanism by which copper proteins interact with NO. In addition, this provides an unexploited way to produce a stable peroxide intermediate in copper oxidases in which the full complement of copper atoms is present. This enables the O-O scission step in the catalytic cycle to be studied by electron addition to the peroxide derivative through the native electron entry site, type 1 copper.

Introduction. Nitric oxide is known to act as a signal transducer in neurotransmission1 and vasodilation2 by interacting with iron in the heme protein guanylyl cyclase. However, nitric oxide may also exert potent physiological effects through interaction with copper. We have recently reported very fast reactions between nitric oxide and cytochrome c oxidase3,4 and we have explained these through a mechanism in which NO acts as a one-electron reductant of the enzyme. We have proposed that reduction occurs at Cu<sub>B</sub><sup>2+</sup>, the oxidized copper in the dinuclear centre, and that, once reduced to Cu<sub>B</sub><sup>1+</sup> this rapidly equilibrates with other redox centers in the enzyme. This rapid interaction of NO with oxidized copper has been suggested to play a key role in the control of the activity of cytochrome c oxidase<sup>4</sup>, and is possibly relevant to the control of a number of other copper containing enzymes. However, the evidence for the reaction

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of NO with this metal is indirect, and only inferred from the observed electron transfer to other redox sites.

We have tested this model using laccase (pdiphenol:dioxygen oxidoreductase, EC 1.10.3.2), the simplest of the family of the multicopper oxidases, which catalyzes the four-electron reduction of dioxygen to water. Laccase contains four copper atoms, classified mainly according to their EPR features<sup>5</sup>: type 1 (T1) or blue, which has a small parallel hyperfine coupling ( $A_{\parallel} \sim 40\text{-}70 \text{ x } 10^{-4} \text{ cm}^{-1}$  or 43-75 G), type 2 (T2) with normal EPR features and a dinuclear EPRundetectable site termed type 3 (T3). Optically, T1 has a relatively strong band in the visible region ( $\epsilon_{615} \sim 5700 \text{ M}^{-1} \text{ cm}^{-1}$ ), T3 absorbs at 330 nm ( $\epsilon_{330} \sim 3,600 \text{ M}^{-1} \text{cm}^{-1}$ )<sup>6</sup>, whereas T2 is practically undetectable. Electron entry from the physiological substrate into the enzyme is at T1, which transfers electrons to a trinuclear copper center formed by T2 and T3<sup>7,8</sup>. It is at this center, analogous to the trinuclear centre found in ascorbate oxidase9 or ceruloplasmin10, where oxygen binds and is reduced.

The understanding of the function of this enzyme has been greatly improved by the use of two derivatives. One of these is type 2 depleted (T2D) laccase, in which one of the copper atoms (T2) is removed by a chelator<sup>11</sup>. Studies using this derivative showed that, in contrast to the oxygen carrier hemocyanin, the dinuclear T3 reduced site in laccase does not bind oxygen<sup>12</sup> in contrast with previous reports<sup>13</sup>. A second derivative<sup>14</sup> is T1Hg, in which the copper atom in the T1 center has been substituted by the redox-inactive Hg<sup>2+</sup>. Using this method, it is possible to obtain an enzyme reduced with only 3 electrons, all of them in the trinuclear center. The combination of fully reduced (3 electrons) T1Hg with oxygen results in the formation of a species which is described as a hydroperoxide, bridged between an oxidized T3 copper and reduced T2<sup>12,15</sup>. This intermediate has been suggested by these authors to precede the formation of the native intermediate<sup>8</sup> in the normal catalytic cycle.

The reaction of NO with tree laccase has been studied previously<sup>16</sup>. In these experiments, performed in the absence of oxygen and under a saturating NO concentration, the enzyme was fully reduced in a process that was reported to be very slow ( $t_{1/2} \sim 70$  min). These experiments, however, were made with a low temporal resolution (e.g., the first EPR measurement was collected 5 min after the addition of NO). On the other hand, the redox state of type 2 copper cannot be assessed optically and EPR measurements are complicated by changes in the redox potential of the metal centers upon freezing the samples<sup>16</sup>. Therefore, an accurate measurement of the level of reduction of type 2 copper in the first seconds was not given. In the present work, we present evidence that NO

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can reduce T2 copper in laccase in a very fast reaction ( $t_{1/2} < 1$  s).

**Experimental Section.** Laccase was obtained and purified according to the method of Reinhammar<sup>17</sup> from acetone powder of Rhus vernicifera (Saito & Co. Ltd. Tokyo, Japan). The concentration of laccase was determined using  $\epsilon_{614} = 5700 \, \text{M}^{-1} \, \text{cm}^{-1}$ .

Static spectra were collected with a Cary 5E UV-Vis-NIR spectrophotometer. The experiments were performed at room temperature in 0.1 M HEPES (N-[2-hydroxyethyl]piperazine-N'-[2-ethanesulfonic acid]) at pH 6 or pH 7.4. The pH was altered by passage through a Sephadex G-25 column equilibrated with buffer at the desired pH. Anaerobic laccase was prepared by degassing a solution in a cuvette pre-sealed with a rubber cap. This was purged with  $N_2\,$  gas, and after a few cycles the sample was immediately used.

Stopped-flow experiments were performed using a SX-18MV stopped-flow apparatus (Applied Photophysics, Leatherhead, UK). Laccase incubated anaerobically with NO was rapidly mixed with buffer containing oxygen at known concentrations. This was obtained by mixing anaerobically the degassed solution with the required volume of oxygen equilibrated buffer.

EPR spectra were measured on a Bruker EMX spectrometer with an ER 041XG microwave bridge (X-band). An ER 4122SP cavity was used. The temperature was controlled using an Oxford Instrument helium system. A Bruker WINEPR (v. 2.11) package was used for spectra analysis. Aliquots of the enzyme or reaction mixture were placed in 3mm ID EPR 'precision' tubes (Wilmad PQ) and frozen in ethanol termostated in dry ice. Once frozen, the tubes were transferred to liquid nitrogen and were normally stored there. The oxidation of the type 2 copper was monitored using the intensity of the low field component<sup>18</sup> at g=2.47, measured using conditions in which the signal-to-noise ratio was maximized (i.e., more scans and lower temperature (8K). After the addition of NO, the band shape of this component was slightly different from the control, although it was unchanged thereafter. Due to this fact, the average intensity corresponding to the samples after 1 hour was taken as 100 % type 2 copper oxidation.

Nitric oxide was obtained from a Kipps apparatus maintained in a certifed hood (due to the potential toxicity of NO gas). Sulfuric acid (1M) was mixed with sodium nitrite. The gaseous product(s) of this reaction were passed through a series of traps (NaOH, H<sub>2</sub>SO<sub>4</sub>, H<sub>2</sub>O, KI and dry ice)to remove nitrogen oxides other than NO. Finally NO was collected in a gas-tight syringe and injected into an anaerobic solution. The NO concentration was measured with an NO electrode (Iso-NO Mark II, World Precision Instruments). The electrode was pre-calibrated by the addition of a standard sodium nitrite solution to excess acidified potassium iodide; this generates NO stoichiometric to the added nitrite. Additions of NO were made using a gas-tight Hamilton syringe.

**Results.** Addition of NO to resting oxidized tree laccase (as prepared) in the presence of oxygen resulted in rapid spectral changes depicted in Fig 1 (spectrum A). This difference spectrum shows prominent positive features at 335 nm, 470 nm and a trough at 610 nm. These positive features are shared by a laccase intermediate that has been described as a

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peroxide, bridging oxidized T3 and reduced T2. This intermediate is obtained by mixing oxygen with the three electron-reduced derivative T1Hg<sup>20</sup>. Addition of NO to anaerobic lacease, however,

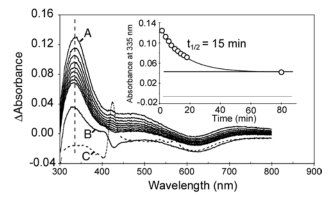


Figure 1. Spectral changes after addition of NO to laccase as prepared in the presence of oxygen. Difference spectra relative to the resting oxidized enzyme (as prepared) after the addition of an alliquot of 2 mM NO (final concentration  $\sim 100 \mu M$ ) to a solution containing  $\sim 70$ μM laccase in HEPES 100 mM pH 6.0. The first spectrum (A) was collected immediately after the addition of NO, and spectra were collected every 2 minutes thereafter. Spectrum B corresponds to the sample after 80 min. A vertical dotted line is plotted to indicate the blue shift of the band as the intermediate decays. Addition of NO to an anaerobic sample in the same conditions generates spectrum C. Insert: Time course followed at 335 nm (o) fitted to an exponential decay ( $t_{1/2} \sim 15 \text{ min}$ ) (———). The features of spectrum C at 400-425 nm and the small bands in the visible region (530 nm and 565 nm) correspond to the formation of a ferric-NO complex of lacquer peroxidase<sup>19</sup>, a contaminant (less than 0.5 % of the laccase concentration) that is common in these preparations. These features decayed rapidly at pH 6 ( $t_{1/2} \sim 30 \text{ s}$ ) and were not present at the time points shown (spectrum A and below). The trough at ~ 420 nm at 80 min (spectrum B) is caused by a further blue shift of the Soret band with respect to the oxidized enzyme.

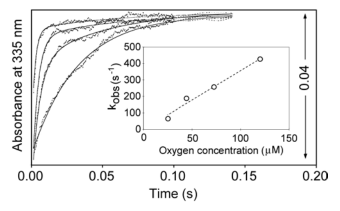
generates only minor optical changes (Fig. 1, spectrum C). As expected, after reaction of this sample with oxygen, spectrum A is generated (Fig. 1). Addition of oxygen to anaerobic laccase in the absence of NO did not produce any optical change. Similarly addition of sodium nitrite (up to 5mM) did not elicit the spectrum seen in Fig 1.

The amplitude of the change at 335 nm was the same at pH 7.4 or pH 6. At pH 6, the decay of this difference spectrum (Fig. 1 and insert) was slow ( $t_{1/2} \sim 15$  min), and  $\sim$  fivefold slower (not shown) when the experiment was performed at pH 7.4. Both the absolute rate of decay and its pH dependency are similar to those observed for the decay of the peroxide intermediate obtained using the T1Hg species<sup>20</sup>. Using the reported extinction coefficient for the intermediate, relative to the oxidized enzyme ( $\Delta \varepsilon_{335} \sim 5,000 \text{ M}^{-1} \text{ cm}^{-1}$ )<sup>20</sup>, the amplitude of the decay at 335 nm (Fig 1, insert) is consistent with a concentration of ~18 µM for this species (i.e., 25 % of the total laccase concentration used in the experiment). The final spectrum (Fig 1, spectrum B and insert), after the decay of the band at 335 nm, shows a remaining positive band at 330 nm which is consistent with the oxidation of ~ 25 % reduced dinuclear T3 centre ( $\Delta \varepsilon_{335} \sim 3,600 \text{ M}^{-1}\text{cm}^{-1}$ , see above), present in the native enzyme. A similar difference spectrum

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was observed (not shown) after addition of  $H_2O_2$  to laccase  $(H_2O_2/laccase molar ratio, 50:1)$ , as reported previously<sup>21</sup>. Addition of NO to the sample incubated with  $H_2O_2$  did not generate any optical change. Thus, the appearance of the features in spectrum A (Fig. 1) are accompanied by changes consistent with oxidation of a fraction of T3. This is also expected for the formation of the peroxide intermediate<sup>20</sup>. Appearance of spectrum A (Fig. 1) upon NO addition was also observed (not shown) when the enzyme had been previously incubated in 0.2 mM potassium ferricyanide for 1 h, in 1 mM fluoride for 30 min, or in 1 mM azide for 1 min.

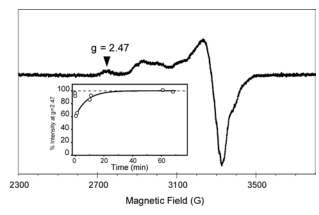
The rate constant for the formation of the band at 335 nm in Fig. 1 (spectrum A) was obtained by mixing oxygen containing buffer with an anaerobic solution containing the native enzyme plus NO (Fig 2). The time courses were fitted to a double exponential curve.



**Figure 2**. Time course of the changes generated following addition of oxygen to laccase incubated anaerobically with NO. Time courses followed at 330 nm in the stopped flow apparatus after mixing anaerobic laccase (65  $\mu$ M) incubated with ~100  $\mu$ M NO for 20 s with buffer at known oxygen concentrations. Oxygen concentration after mixing (bottom to top): 25, 45, 70 and 120  $\mu$ M. The traces were fitted to a double exponential increase. Insert: Plot of the observed pseudofirst order rate constant of the fast component versus the oxygen concentration.

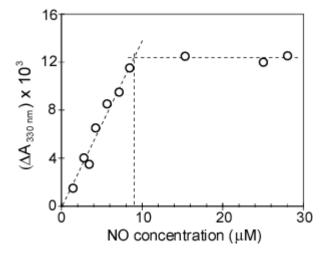
The fastest of these processes was found to be oxygen concentration dependent, yielding a value of 3.5 x  $10^6~M^{\text{-1}}~\text{s}^{\text{-1}}$  for the second order rate constant (Fig. 2, insert), similar to that found for the formation of the peroxide intermediate in T1Hgl5. The second process was slower (  $k_{\text{obs}} \sim 30~\text{s}^{\text{-1}}$  ) and independent on the oxygen concentration.

As the redox state of type 2 copper itself is difficult to detect optically, the changes occurring after addition of NO were monitored by EPR spectroscopy (Fig 3). This figure shows that the intensity of the low field component (g=2.47) corresponding to the oxidized type 2 copper (see Materials and Methods) drops  $\sim$  30-40 % after the addition of NO (Fig. 3, insert). This is followed by slow reoxidation.



**Figure 3**. Time course of the events described in Fig. 1 followed by EPR. EPR spectra of oxidized resting laccase at pH 6 (65 μM) measured at 77 K. Microwave power 3.2 mW, microwave frequency 9.4859 GHz, modulation frequency 100 kHz, modulation amplitude 1 G, time constant 0.041 s, sweep rate 9.15 G/s. The low field component of the type 2 copper signal (g = 2.47) is indicated. Inset: Intensity of this component changing during the experiment and normalized with respect to the intensity obtained after 1 h. The first two points (before addition of 100 μM NO) were taken as a control, and the following measurements were taken at different time points (0.5, 1, 10, 11, 60 and 68 min) after the addition of NO. The line represents an exponential fit to the data.

Note, however, that the first 2 measurements were obtained after 30 and 60 s after the addition of NO, and the percentage of T2 reduction could be even higher (see figure legend). This suggests that the reduction of the T2 coppers may not be restricted to the putative population of laccase having T3 reduced (23 %). In fact, when the enzyme in the presence of oxygen was titrated with NO and the absorbance at 335 nm corresponding to the putative peroxide intermediate was followed optically, it was observed (Fig. 4) that the



**Figure 4**. Effect of substoichiometric additions of NO to laccase in the presence of oxygen. Titration corresponding to the changes at 330 nm after the addition of alliquots of NO to 8.5  $\mu$ M laccase in the presence of oxygen. The time interval between NO addition and measurement was less than 5 s. The vertical dotted line indicates the enzyme concentration.

maximum amplitude of the absorbance change was obtained when the molar ratio NO/laccase was 1:1. This indicates that a

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complete formation of the putative peroxide intermediate is only obtained at equimolar concentration of NO and laccase. As the formation of the peroxide intermediate requires the reduction of T2, we conclude therefore that the affinity of T2 for NO may be similar, irrespective of the redox state of T3.

**Discussion.** On addition of NO to an aerobic solution of laccase or addition of oxygen to anaerobic solutions of laccase in the presence of NO rapid absorbance changes occur, as depicted in figures 1 and 2. The question arises as to whether these changes are due to the direct reaction of NO and O<sub>2</sub> with laccase or indirectly to the reaction of the products of the reaction between NO and O<sub>2</sub> with the enzyme.

In aerobic aqueous solutions nitrite is essentially the only product of the reaction between NO and  $O_2^{xx}$ . Not only does addition of nitrite to oxidised laccase not generate the spectra observed its formation is in any case much too slow to account for our observations. For example in Figure 2 an anaerobic solution of 65 µM laccase/100 µM NO was mixed with 25 μM oxygen and the spectrum described in Fig 1appeared at a rate of approximately 50 s<sup>-1</sup>. The rate equation for the reaction of NO with O<sub>2</sub> may be written as:

 $d[NO_2]/dt = 4 \times k_1[NO]^2[O_2],$ 

where  $k_1 = 2 \times 10^6 \text{M}^{-2} \text{s}^{-1}$ . Given the second order nature of this process in [NO], the chemical oxidation of NO is far too slow to produce species relevant to the laccase reactions observed here (the calculated k<sub>obs</sub> for nitrite formation in the above case is 2 x 10<sup>-6</sup> M.s<sup>-1</sup>, which indicates that under the concentration regime employed the half-time for the reaction is > 40s whereas the reaction with the laccase has a half time of some140ms). Although higher nitrogen oxides are formed as intermediates in the reaction of NO and O2, the calculated concentrations<sup>xx</sup> are far too small to be responsible for the observed stoichiometric changes in the laccase copper centers. The concentrations of N<sub>2</sub>O<sub>3</sub> and NO<sub>2</sub>may be calculated to be 1nM and 0.4 nM respectively versus a laccase concentration of 65 µM.. We conclude therefore that the spectra we have described are due directly to the action of NO and not to the products of its reaction with O<sub>2</sub>.

In fact the changes observed after addition of NO to laccase in the presence of oxygen can be interpreted as due to the formation of the peroxide intermediate in a fraction (25 %) of the molecules. However, to achieve this, a fully reduced trinuclear centre (T3 and T2) must be present<sup>6</sup>, which upon reaction with oxygen would produce the peroxide intermediate. Whereas we observe reduction of T2 by NO (see Fig. 2), it seems unlikely that NO also reduces T3. First, because reduction of T3 by NO has been reported to be very slow<sup>16</sup>, and second, because of the 1:1 stoichiometry observed in the appearance of the band at 335 nm (Fig. 4). The obvious alternative is that a fraction of T3 is already reduced before addition of NO, as suggested previously<sup>21</sup>. Indeed, Our data are consistent with the presence of 25 % T3 reduced in the native enzyme, becoming oxidized upon NO addition and further oxygen binding. Similar changes occurred after addition of hydrogen peroxide, an oxidant for T3 copper, consistent with oxidation of 25 % of T3, also reinforcing this hypothesis. Furthermore, addition of NO to this fully oxidized sample did not give rise to optical changes. Also, upon mixing oxygen with the enzyme anaerobically incubated with NO, the band at 335 nm is formed with a second order rate constant of  $3.5 \times 10^6 \, M^{-1} \, s^{-1}$ . This value is identical, within experimental error, to  $2 \times 10^6 \, M^{-1} \, s^{-1}$ , reported for the formation of the peroxide intermediate when mixing oxygen with fully reduced  $T1Hg^{15}$ , and also similar to 5 x  $10^6$  M<sup>-1</sup> s<sup>-1</sup>, obtained for the formation of the native intermediate when mixing oxygen with

the fully reduced enzyme<sup>22</sup>, In the last case, although the formation of the native intermediate is associated with oxidation of T1, the same kinetic process should be observed (e.g., oxidation of T3 coppers), as oxidation of T3 is the ratelimiting step <sup>20</sup>.

The appearance of the changes attributed to the peroxide intermediate upon NO addition, even after preincubation of laccase with ferricyanide or fluoride, can be explained by the fact that ferricyanide can not oxidize the dinuclear centre T3 <sup>21</sup>, therefore the population of molecules containing T3 reduced is still able to form the peroxide intermediate. Fluoride, on the other hand, although a strong ligand ( $K \sim 10^4$ M<sup>-1</sup>)<sup>23</sup> of oxidized type 2 copper, possibly precluding the formation of the intermediate, does not bind to this site when the dinuclear centre T3 is reduced<sup>23</sup>. The observation that the intermediate is formed also after incubation of laccase with azide is intriguing, as azide has been reported to bind preferentially to T2 when T3 is reduced7. The reaction may still be possible, however, if NO can reduce T2 and the peroxide group bridges between T2 and the T3 copper that is not bound to azide. This reaction possibly displays different kinetics.

The fact that the change in amplitude at 335 nm is complete at a stoichiometry NO/ laccase 1:1 suggests that all the T2 sites are reduced at the end of the titration, or at least bind NO. Thus, although the affinity of NO for T2 copper may be different depending on the redox state of T3, as for azide<sup>7</sup> or fluoride23, both fractions of the enzyme bind NO with an affinity > 10<sup>6</sup> M<sup>-1</sup>. Furthermore, the fact that other centres (T3 or T1) did not become reduced, suggests that only one NO is bound per molecule, at T2. The absence of bleaching of the bands at 614 or 330 nm after addition of NO to the anaerobic enzyme provides further evidence that binding of NO and reduction of Cu must occur at the optically undetectable T2 copper, confirming previous results16 which describe this center as the electron entry site in the reduction of laccase by NO. Although the EPR results (Fig. 3) suggest that only a fraction (30-40 %) of the T2 copper becomes reduced, it is possible that low temperature effects leading to electron redistribution<sup>16,24</sup> between T2 copper and NO may mask the true percentage of reduced T2 copper at room temperature. Formation of the native intermediate, as a product of the decay of the peroxide intermediate was not detected, either optically or by EPR. This intermediate, which displays a band at 360 nm relative to the oxidized enzyme<sup>22</sup>, is formed upon electron transfer from T1 to the peroxide intermediate, and has been associated to an oxygen radical which is detected only at low temperature<sup>25</sup>. Although such a species may be formed in our sample as the product of the decay of the peroxide intermediate, the fact that in our system the peroxide intermediate must be reduced by T2 (t<sub>1/2</sub> of minutes), and not by T1 (>1000 s<sup>-1</sup>), suggests that its formation is probably paralleled by its decay, and a significant population never accumulates.

In summary, we have observed rapid reduction of a type 2 copper in laccase upon NO addition. This has been observed directly by EPR, and also optically through a subpopulation of the enzyme (~ 25 %) which contains a dinuclear T3 centre

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already fully reduced (2 electrons). The kinetics of the formation of the peroxide intermediate are virtually identical to those reported for the formation of the native intermediate using fully reduced enzyme, or the formation of the peroxide intermediate using fully reduced T1Hg. The rate of decay of the species described is very slow, and similar to the decay of the peroxide intermediate, showing also a similar pH dependency. We conclude that the peroxide intermediate has been formed via reduction of T2 copper by NO in a fast reaction. We suggest that this reaction may not only hold implications for the regulation, by NO, of enzymes containing trinuclear copper centres (e.g. the major copper protein in blood plasma, ceruloplasmin), but also could be relevant to the regulation of copper enzymes in general. In addition, the method we report here for the formation of the peroxide intermediate in laccase molecules which posses the full

complement of copper atoms should prove useful to the study of the catalytic cycle of this enzyme. For example, it permits one to investigate the O-O scission step by the addition of a single electron through the native T1 copper. This reaction cannot be so easily isolated by other methods.

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