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Catalytic Cycle of Multicopper Oxidases Studied by Combined Quantum- and Molecular-Mechanical Free-Energy Perturbation Methods

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Abstract

We have used combined quantum mechanical and molecular mechanical free-energy perturbation methods in combination with explicit solvent simulations to study the reaction mechanism of the multicopper oxidases, in particular the regeneration of the reduced state from the native intermediate. For 52 putative states of the trinuclear copper cluster, differing in the oxidation states of the copper ions and the protonation states of water- and O₂-derived ligands, we have studied redox potentials, acidity constants, isomerisation reactions, as well as water- and O₂ binding reactions. Thereby, we can propose a full reaction mechanism of the multicopper oxidases with atomic detail. We also show that the two copper sites in the protein communicate so that redox potentials and acidity constants of one site are affected by up to 0.2 V or 3 pK_a units by a change in the oxidation state of the other site.

Keywords: multicopper oxidases, CueO, redox potential, acidity constant, quantum-mechanical calculations, free-energy perturbation, QM/MM

Introduction

The multicopper oxidases (MCOs) are a group of enzymes that couple the oxidation of various substrates with the reduction of O₂.¹



They occur in all types of organisms and typical examples are ceruloplasmin, laccase, ascorbate oxidase, bilirubin oxidase, and the bacterial copper-resistance protein CueO.^{2,3} They can oxidise a great range of organic substrates, as well as a number of metal ions^{1,3} and have therefore found use both in biotechnological and bioelectrochemical (biofuel cells) applications.^{4,5}

As the name implies, the MCOs contain at least two copper sites. The substrate oxidation takes place close to a type 1 (blue) copper site, in which a single copper ion is bound to a cysteine (Cys) and two histidine (His) residues, and in some proteins also to a methionine residue. The O₂ molecule binds to a trinuclear copper cluster (TNC), in which one copper ion (the type 2 copper ion, Cu_{T2}) is bound to two His residues and to one solvent molecule (water or OH⁻), whereas the other two Cu ions (the type 3 copper ions, Cu_{T3}) bind to three His ligands each, and possibly to additional solvent or O₂-derived molecules. The eight His ligands of the TNC occur pair-wise in the amino-acid sequence (four His-X-His motifs) and in one case, the X residue is the Cys ligand of the type 1 Cu ion (Cu_{T1}), providing a favourable path for the transfer of electrons between the two sites, which are at a Cu-Cu distance of ~13 Å.

The MCOs have been extensively studied by both experimental^{1,3,6-11} and theoretical¹⁰⁻¹³ methods. Crystal structures are available from ~20 different organisms.^{1,6-8,13-15} In the oxidised resting state of the enzyme, all three copper ions of the TNC are in the oxidised Cu^{II} state and the two Cu_{T3} ions are bridged by a OH⁻ molecule. In the reduced state, the three TNC copper ions are in the reduced Cu^I state and there is no ligand bridging the two Cu_{T3} ions. This is the state that binds the O₂ molecule.

By ingenious spectroscopic studies, Solomon and coworkers have characterised two intermediates in the catalytic mechanism of the MCOs.^{1,3,11} Directly after the binding of O₂ to the TNC, a peroxide intermediate (PI) is formed, in which two Cu ions are oxidised to Cu^{II} and O₂ is reduced to the peroxide level.¹⁶ Theoretical calculations suggested that the peroxide binds in the centre of the TNC in a diagonal manner (μ₃-1,1,2), interacting with all three Cu ions^{12,17-19} and this was later confirmed by combined experimental and computational studies^{20,21}. Rotation of the O₂²⁻ ion in the TNC is facile both in isolated clusters and in the protein.²²

A second state, the native intermediate (NI), has also been identified experimentally and was shown to contain four oxidised Cu ions and the O₂ substrate fully reduced to the level of two water molecules.²³ Theoretical calculations indicated that it consists of an O²⁻ ion bound symmetrically in the centre of the TNC (μ₃),^{12,17,19} which was later confirmed by other studies^{10,24}. Interestingly, NI is the only fully oxidised state in the catalytic cycle of the enzyme, whereas the oxidised resting state seems to be outside this cycle, as the conversion of the NI to the oxidised resting state has been shown to be too slow to comply with the overall catalytic turnover.^{10,11,25}

Based on these findings, a reaction mechanism of the MCOs was postulated, shown in Figure 1.^{1,10-13} It starts with O₂ binding to the reduced state, accompanied with the transfer of two electrons from two Cu^{II} ions to O₂, directly leading to the PI. Next, an electron is transferred from Cu_{T1} to the TNC, resulting in the transient NI' species in the Cu^{II}Cu^I₂ state. This triggers the cleavage of the O-O bond and the formation of the NI. The latter step has been studied in detail with quantum-mechanical (QM) and QM/MM methods,^{20,26} showing a relatively facile reaction (with estimated barriers of 20-60 kJ/mol). Finally, the reduced state should be regenerated by the transfer of three more electrons from Cu_{T1} to the TNC. These electron transfers should be accompanied also by proton transfers (ultimately from the surrounding solvent). However, relatively little information is available for the detailed structure of the states between NI and the reduced state, or on the exact protonation of the water- and O₂- derived ligands of the TNC. An alternative resting state has been observed and characterized in bilirubin oxidase by spectroscopy and crystallography, indicating a TNC in the Cu(I)₂Cu(II) state, with the two Cu_{T3} ions bridged by an oxygen ligand with a long Cu-Cu distance (5.0 Å).²⁷ Similar crystal structures have also been published by other groups, in which the two Cu_{T3} ions are bridged by either one or two oxygen

atoms,^{28, 29} but it is likely that at least some of them are affected by partial photoreduction of the TNC during data collection (i.e. showing a mixture of several different oxidation states).¹³ Solomon and coworkers have studied the one-electron reduction of the native intermediate, giving rise to the Cu(I)Cu(II)₂ state.²⁵ Very recently, they have also published a combined kinetic, electron paramagnetic resonance (EPR) and QM study of all states involved in the conversion of NI to the fully reduced state.³⁰

As already discussed, the reduction of O₂ involves the uptake of four electrons and four protons. The electrons are transported to the TNC via the Cu_{T1} site and they are taken from the substrates, which are oxidised at the Cu_{T1} site. The protons are taken from solution and are most likely transported to the TNC via two conserved residues, Asp-112 (throughout this article, we use the residue numbering of CueO¹⁵), which is connected to Cu_{T2} via two water molecules, and Glu-506, which is connected to the ligand bridging the two Cu_{T3} ions via a water molecule. Differences involving electrons and protons are very hard to discern in crystal structures. Unfortunately, reactions involving a change in the number of electrons or protons are also hard to study by theoretical methods, because the net charge of the system changes. This gives rise to large and long-ranged electrostatic energies and huge solvation effects.³¹ For example, the Coulombic interaction between two groups with a unit charge at a distance of 15 Å is 93 kJ/mol (0.96 eV; although it is probably screened by a dielectric constant of 4–80) and the Born solvation energy in water of a unit charge is 23 kJ/mol (0.24 V) even for a spherical system with a radius 30 Å (i.e. similar to the size of a MCO). This makes the calculation of absolute redox potentials and acidity constants of protein sites a formidable task. Further complications arise from the fact that there are large differences in the electron-correlation energies when the number of electrons in a metal change, leading to large differences in reduction potentials calculated with different density-functional theory (DFT) methods.^{32, 33}

Many methods to calculate the redox potentials of metal sites in proteins have been developed. A first estimate can be obtained by simply calculating the QM energies of the active-site clusters in a continuum solvent. However, to obtain information on the influence of the surrounding protein, more sophisticated methods are needed.^{30, 32, 34-38} In particular, several groups have estimated and rationalised the redox potentials of Cu_{T1} sites in proteins^{36, 39-42} and the TNC in MCOs has also been studied.⁴³

Likewise, many methods have been developed to calculate acidity constants, both of small molecules in solution and of various groups in proteins.^{33, 34, 44-49} The former are typically based on QM calculations, to obtain the intrinsic proton affinity of the molecules of interest, combined with continuum-solvation methods to estimate the effect of transferring the reactants from vacuum to solvent. In proteins, the intrinsic proton affinity is often of less interest; instead, the main issue is how the protein alters the acidity constant of a group relative to water solution (for which the acidity constant typically is known). Many approaches have been suggested and tested for the calculation of the differential solvation effect in the protein, based on e.g. free-energy perturbations (FEP) or continuum-solvation models, using the Poisson–Boltzmann (PB), the Langevin dipole (LD) approach, or the generalised Born (GB) methods.^{34, 47, 50, 51} In general, an accuracy of one or a few pK_a units is claimed.^{33, 44, 47} Fewer groups have tried to estimate the acidity constants of metal-bound ligands, mainly because of the large effect of the charged metal and the fact that it is difficult to model metals by molecular-mechanical (MM) methods. However, several attempts have been made, especially for metal-bound water molecules, using the LD and PB methods.^{32, 34, 52-54}

In this work, we study electron- and proton transfer reactions in the reaction cycle of the MCOs with the help of FEP calculations at the combined QM and MM (QM/MM) level. By estimating redox potentials, acidity constants, isomerisation free energies, as well as O₂ and water-binding free energies of putative intermediates, we obtain information about the most likely steps in the reaction mechanism, especially for the regeneration of the reduced state from the NI.

Methods

The protein

The calculations on the MCOs are based on the 1.4-Å crystal structure of CueO (PDB code 1KV7).¹⁵ CueO is a bacterial copper homeostasis factor and this structure was selected because

of its high resolution and that it lacks the extended glycosylation found for most eukaryotic MCOs. In addition, we have already performed extensive theoretical studies on this protein.^{12, 13, 17-19, 26} The whole protein was included in all calculations, as well as a large number of surrounding water molecules. Initially, all Glu and Asp residues were assumed to be deprotonated and all Lys and Arg residues were assumed to be protonated. The protonation state of the His residues were assigned based on their solvent exposure, the hydrogen-bond network, and the local surroundings.¹² Eight His residues on the protein surface were assumed to be protonated on both N atoms (His-145, 224, 314, 405, 406, 465, 488, and 494), whereas the remaining ten His residues, which are ligands of the Cu ions, were assumed to be protonated on the N atom not coordinating to the metal. The Cys ligand of Cu_{T1} was assumed to be deprotonated. This gave a net charge of the full simulated system of -1 to +4 e, depending on the studied state of the two copper sites. No counter-ions were used in the simulations.

In some simulations, we also tested to neutralise all solvent-exposed charged groups, as has often been done in calculations of reduction potentials or acidity constants.^{36, 44, 52} The solvent exposure was determined from the contact of each residue with water molecules during molecular dynamics (MD) simulations³² and it was also checked by the number of protein atoms surrounding the groups²⁰ and by visual inspection. We decided to keep the charge on 17 buried residues: Arg-232, 234, 242, 280, and 466, Lys-188 and 470, Asp-112, 180, 187, 260, 439, and 471, as well as Glu-279, 282, 457, and 506. Of these, Asp-112, 439, 471, form hydrogen bonds to His ligands of the copper sites and Glu-506 interacts with the TNC bridging ligand via a water molecule. All the other residues form ionic pairs with each other, except Lys-188, which is deeply buried in the protein, but the closest carboxylate group is four hydrogen bonds away. All the other (exposed) groups were neutralised by adding or removing a proton, using the AMBER neutralised residues ASH, GLH, and LYN and a neutralised Arg model.⁵⁵ All sites were visually inspected so that protons were added or removed on proper positions to optimise the hydrogen-bond networks.

The QM/MM-2QM approach

For the QM/MM geometry optimisations, the QM/MM-2QM approach was employed,^{37, 43} in which the two copper sites were both treated by QM, but in separate QM calculations. This is possible because the two copper centres are ~13 Å apart. The QM/MM-2QM implementation was based on the COMQOM program^{56, 57} which is an interface between the Turbomole⁵⁸ and Amber⁵⁹ softwares.

In this QM/MM-2QM approach, the protein and solvent are divided into three subsystems: the two QM systems (systems 1a and 1b) and the MM system (system 2). In the QM calculations, systems 1a and 1b are represented by a wavefunction, whereas all the other atoms are represented by an array of partial point charges, one for each atom, taken from MM libraries. Thereby, the polarisation of the QM system by the surroundings is included in a self-consistent manner (electrostatic embedding, EE). In the QM/MM calculations, the MM system was kept fixed at the original (crystallographic) coordinates (but it was relaxed in the subsequent QTCP calculations, described in the next section).

When there is a bond between systems 2 and 1a or 1b (a junction), the hydrogen link-atom approach is employed: The QM region is truncated by hydrogen atoms (called hydrogen link atoms, HL), the positions of which are linearly related to the corresponding carbon atoms in the full system (called carbon link atoms, CL).⁵⁶ In the point-charge model of the surroundings, all atoms were included, except the CL atoms.⁶⁰ For our QM systems, the CB atom of the His and Met Cu ligands and the CA atom of the Cys ligand were replaced by HL atoms. In the proton-transfer reactions, the CA atom of Asp-373 was replaced by a HL atom.

The total QM/MM-2QM energy was calculated in the following way:

$$E_{QM/MM-2QM}^{EE} = E_{QM1a+ptch1b2}^{HL} - E_{MM1a,q=0}^{HL} + E_{QM1b+ptch1a2}^{HL} - E_{MM1b,q=0}^{HL} + E_{MM1ab2,q_{1a}=q_{1b}=0}^{CL} \quad (2)$$

Here, $E_{QM1a+ptch1b2}^{HL}$ is the QM energy of system 1a, truncated with HL atoms, and including a point-charge model of the surrounding protein (also of system 1b) in the one-electron Hamiltonian. $E_{QM1b+ptch1a2}^{HL}$ is the corresponding QM energy for system 1b. $E_{MM1a,q=0}^{HL}$ and $E_{MM1b,q=0}^{HL}$ are the MM

energies of systems 1a and 1b (still with HL atoms), but with all charges zeroed. Finally, $E_{MM1ab2,q1a=q1b=0}^{CL}$ is the MM energy of all systems with CL atoms, but with the charges of the two QM systems zeroed (because all electrostatics interactions within the QM systems and between the QM and MM systems are considered in the two QM terms. However, the electrostatic interactions between the two QM systems is still accounted for in both $E_{QM1a+ptch1b2}^{HL}$ and $E_{QM1b+ptch1a2}^{HL}$. This problem was solved by scaling down the point charges of the other QM system by a factor of 2 in both terms.⁴³

The point-charge model of each QM system was obtained by a fit to the electrostatic potential (ESP) calculated for a wavefunction polarised by a point-charge model of the surroundings, but without the point charges when the ESP was calculated.^{43, 61} The ESP points were sampled with the Merz–Kollman approach⁶² as implemented in Turbomole⁵⁸. The charges were updated in each step of the geometry optimisation. The QM/MM-2QM calculations were performed on spherical systems in which the whole CueO protein was solvated in a 38-Å sphere of explicit water molecules (~21 000 atoms in total).¹³

QTCP-2QM calculations

Based on the QM/MM-2QM optimised structures, we used the QTCP (QM/MM thermodynamic cycle perturbation) approach^{63, 64} to calculate the various reaction free energies. This is a method to calculate free energies with a high-level QM/MM method, using sampling only at the computationally cheaper MM level. It calculates the free energy from three separate terms:

$$\Delta G_{QTCP}(R \rightarrow P) = -\Delta G_{MM \rightarrow QM/MM}(R) + \Delta G_{MM}(R \rightarrow P) + \Delta G_{MM \rightarrow QM/MM}(P) \quad (3)$$

where R and P are the reactant and product states. Thus, the free energy difference between P and R is calculated with a normal FEP calculation at the MM level. Then, additional FEP calculations are performed from the MM to the QM/MM levels, one for the R state and one for the P state. To obtain stable MM→QM/MM free energies, the QM systems are kept fixed in the MD simulations. Test calculations at the semiempirical level have shown that this does not affect the final free energies significantly.⁶⁵

The QTCP-2QM calculations were performed as previously described.^{32, 43, 63, 64, 66} First, each state was optimised by QM/MM-2QM, as described in the previous section. Second, the total system was moved into a periodic octahedral box and was further solvated with water molecules extending at least 9 Å from the original system, giving a total of ~12300 water molecules or ~43000 atoms. For the reactant state, the system was first subjected to a 1000-step minimisation, keeping the QM systems fixed and all heavy atoms except water molecules restrained towards their positions in the crystal structure with a force constant of 418 kJ/mol/Å². Then, a 20-ps constant-volume MD simulation was run with only the heavy atoms restrained, but with a force constant of 2092 kJ/mol/Å². Next, a 20-ps constant-pressure MD simulation was run with the same constraints but a force constant of 418 kJ/mol/Å². Finally, the box size was equilibrated by a 100-ps MD simulation with a constant pressure and only the heavy atoms of the two QM systems restrained to the QM/MM-2QM structure with a force constant of 418 kJ/mol/Å². The final structure of this simulation was then used as the starting structure for the simulations of the other state. It is not possible to keep the QM atoms exactly fixed in the constant-pressure simulations. Therefore, the QM atoms were moved back to the QM/MM structure before an equilibration of 200 ps and a production simulation of 400 ps were run with a constant volume and with the two QM systems fixed. During the production run, 200 snapshots were collected every 2 ps. Finally, free energies were estimated from these snapshots, using FEP or thermodynamic integration (TI). FEP free energies and their uncertainties were estimated by cumulant expressions.^{65, 67}

Several methods were used to estimate long-range electrostatic effects outside the simulated system. The results presented in the text and the tables were obtained with periodic systems and Ewald summation (as in the MD simulations). However, non-periodic calculations with an infinite cut-off and solvation energies outside the simulated system calculated by the Onsager or Born models^{32, 68} give results that differ by only a few kJ/mol (isomerisation free energies coincide within 1 kJ/mol, O–O bond cleavage reaction free energies differ by 1–3 kJ/mol,

the reduction potentials are typically 0.04 V more positive, and the acidity constants differ by 1.4 pK_a units on average). However, if the long-range solvation effects were instead estimated with the generalised Born method,⁶⁹ much larger differences were often encountered (results not shown).

The perturbations were divided into several steps, depending on the type of energies studied, as is illustrated in Figure 2. Reduction potentials were studied by transferring an electron from the Cu_{T1} site to the TNC (i.e. between the two QM systems). Such a reaction does not change the net charge of the protein, thereby making the calculated energies more stable. We performed nine simulations with the charges of the QM system linearly transformed from the reactant state to the product state by a coupling parameter, λ , assuming the values of 0, 0.125, 0.25, 0.375, 0.5, 0.625, 0.75, 0.875, and 1. This was accomplished by simply changing the charges in the parameter file, whereas the coordinates of the QM system were kept to those of the reactant state. Free energies were estimated by TI. Then, the coordinates of the QM system were perturbed from that of the reactant state to that of the product state, using charges of the product state. This was accomplished in five FEP simulations using another coupling parameter attaining the values of 0, 0.25, 0.5, 0.75, and 1. For the two end states (reactant and product states), we also performed a perturbation from MM to QM/MM-2QM energies, as has been detailed before.^{32, 43, 64} These calculations were performed on the same octahedral systems as all the other QTCP-2QM calculations, but without any periodicity or Ewald summation and instead with an infinite cut-off. We studied the O–O cleavage reaction in the same way (but without transferring any electron, so that the Cu_{T1} site was not affected by the reaction).

For the acidity calculations, we instead transferred a proton from the TNC to a carboxylate group on the surface of the protein (which now was the other QM system, whereas the Cu_{T1} site was part of the MM system). After a set of test calculations (described in Table S1 in the supporting information), we decided to use Asp-373 as the proton acceptor. It is located on the surface of the protein, without any interactions with other protein residues and it is ~24 Å from Cu_{T1} and 34–37 Å from the Cu ions in the TNC. When the residue was protonated, only charges in the QM system (an acetate ion) were modified, whereas the charges on the protein back-bone (N, H, HA, C, and O atoms) were the same as for a normal negatively charged Asp residue.

The movement of a proton is a more complicated reaction, involving disappearance and appearance of atoms.³² We started from a protonated TNC and a deprotonated Asp-373 residue. Then, we introduced a dummy atom on Asp-373, calculating its contributions to the free energy analytically from local configurational integrals.^{32, 70, 71} Next, the van der Waals parameters of this appearing proton were perturbed from those of a dummy atom with a zeroed a theoretical depth to polar hydrogen by a single-step perturbation. After that, the charges of the two QM sites were perturbed from those of the starting state to those of the final state (and zero charges for the disappearing protons) in nine steps, in the same way as for the electron-transfer reactions (keeping the geometry and van der Waals parameters fixed). Subsequently, the van der Waals parameters of the disappearing proton of the TNC were perturbed from those of a polar hydrogen to a dummy atom with a zeroed well-depth in a single step. Next, the resulting non-interacting dummy atom was deleted, again calculating its contributions to the free energy analytically. In fact, the two analytical contributions will cancel in the net result. Finally, the coordinates of the two QM systems were transformed from those of the starting state (without the dissociating proton) to those of the final state in five steps, as described above. The same approach was used for isomerisation reactions, for which a proton was moved within the TNC cluster (a Cu_{T1} site was included as the other QM system, although it was not explicitly involved in the reaction).

All calculations were automatized and performed by four Linux shell scripts and a number of Fortran programs that are available from the authors upon request. Some further details of the calculations can be found in <http://www.teokem.lu.se/~ulf/Methods/qtcp.html>.

QM/MM-PBSA-2QM calculations

The QM/MM-PBSA method^{43, 61} can be viewed as a post-processing of QM/MM calculations to obtain more stable energies, including a solvation energy obtained by continuum solvation methods. It is an adaptation of the widely used MM/PBSA approach⁷² for QM/MM calculations. In this approach, an approximation to the total free energy for each state is obtained from:

$$G = E_{QM/MM-2QM}^{ME} + G_{Solv} + G_{np} - TS_{QM/MM} \quad (4)$$

Here

$$E_{QM/MM-2QM}^{ME} = E_{QM1a}^{HL} - E_{MM1a}^{CL} + E_{QM1b}^{HL} - E_{MM1b}^{CL} + E_{MM1ab2}^{CL} \quad (5)$$

where E_{QM1a}^{HL} and E_{QM1b}^{HL} are the QM energy of the two QM systems, truncated with HL atoms, but without any point-charge model. E_{MM1a}^{CL} and E_{MM1b}^{CL} are the corresponding MM energies of the two QM systems, but now with CL atoms and with full charges. Finally, E_{MM1ab2}^{CL} is the MM energy of the total system with CL atoms and full charges. This energy is similar to the QM/MM-2QM energy in Eqn. (1), but it instead employs mechanical embedding (ME), i.e. the electrostatic interactions between the QM and MM systems are calculated at the MM level. G_{Solv} is the polar solvation energy, estimated by a continuum approach, obtained either by the generalised Born (GB) approach developed by Onufriev, Bashford and Case (model II, i.e. igb=5)⁶⁹ or by solving the Poisson–Boltzmann (PB) equation using the PB solver in Amber.⁵⁹ G_{np} is the non-polar solvation energy, estimated from the solvent-accessible surface area (SASA), through the linear relation $G_{np} = 0.0227 \text{ SASA } (\text{\AA}^2) + 3.85 \text{ kJ/mol}$.⁷³ The entropy term was ignored, because entropy effects are expected to be quite small in these electron- and proton-transfer reactions.⁴³

Two variants of QM/MM-PBSA-2QM were tested. They differ in how the QM energy and the charges on the QM atoms are calculated.^{32, 43, 66} In the first approach, the QM energy and the charges are calculated from a vacuum wavefunction. In the second approach, the QM wavefunction is optimised with a point-charge model of the other systems. However, the final energy and the QM charges are calculated without this point-charge model and without re-optimising the wavefunction (obtained by setting the number of SCF iterations to 1 and using the final DFT integration grid in this single step). Thereby, the QM system is polarised by the MM surroundings and the cost of the polarisation is included in the calculations.

The QM/MM-PBSA-2QM calculations were based on the spherical QM/MM-2QM structures. The calculations were automatized and performed by Linux shell scripts, which are available from the authors on request. Further details of the calculations can be found in http://www.teokem.lu.se/~ulf/Methods/qmmm_pbsa.html.

Throughout this article, the presented results are from the QTCP method if available, which are the most accurate. However, O₂ and H₂O binding energies are harder to estimate with QM/MM-FEP methods (owing to the dissociation ligand); therefore, the reported results are from the QM/MM-PB(GB)SA-2QM calculations and the same applies to the more approximate absolute redox potentials. Naturally, the latter methods also give results for the other types of calculations. Our results show that QM/MM-PBSA-2QM typically gives a somewhat larger variation in the results than QTCP, whereas QM/MM-GBSA-2QM often gives a smaller variation. However, for the most of the redox, isomerisation, and O–O bond- cleavage reactions, all methods indicate the same direction of the reaction, increasing the credibility of the calculated results. For the proton-transfer reactions, the QM/MM-GBSA results are consistently more negative than the QTCP results (by 14–21 pK_a units on average), whereas the QM/MM-PBSA results are more positive (by 10–12 pK_a unit on average). This shows that there is a difference in how the three methods estimate the change in the solvation energy when the charge of a site changes by one unit when one site is solvent exposed and the other is buried in the protein. QM/MM-2QM results are also available for all reactions, but they in general overestimate variations by a factor of about two, owing to the missing dynamic and long-range solvation effects, and they are therefore not discussed.

To examine the effect of the neutralisation of the surface charges (employed for the proton-transfer and isomerisation reactions, but not for the redox potentials), we repeated one of the QTCP redox calculations with neutralised surface charges. Quite encouragingly, the predicted redox potential changed by only 0.18 V when the surface charges were neutralised. The effect is similar for the QM/MM-GBSA calculations, in which the calculated redox potentials changed by 0.18 V on average for all 21 systems. The effect is even smaller for the proton-transfer reaction, less than 0.8 pK_a units. Consequently, we think that the present results can be trusted to within 30

kJ/mol (0.3 eV or 5 pK_a units), except for states for which there is a large difference in the results obtained with the different DFT functionals (in particular the O₂ binding free energies).

QM calculations

QM/MM-2QM optimisations and the QTCP-2QM calculations were performed with the Perdew–Burke–Ernzerhof (PBE) density functional⁷⁴ and the def2-SVP basis set.⁷⁵ These energies were then improved by single-point energy calculations with the hybrid B3LYP method^{76,77} and the larger def2-TZVPD basis set with diffuse functions on all atoms.^{75,78} All QM calculations were sped up by expanding the Coulomb interactions in an auxiliary basis set, the resolution-of-identity (RI) approximation.^{79,80} Test calculations (cf. Table S2 in the supplementary material) showed that the def2-TZVPD basis set gives reaction energies (isomerisation, redox potentials, acidity constants, and O–O cleavage reaction energies) that are converged to within 3 kJ/mol compared to calculations with the even larger def2-QZVPD basis set. Diffuse functions are needed to give such accurate energies for the acidity constants and redox potentials – the def2-TZVP basis set gives errors of up to 14 kJ/mol. All QM calculations were performed with the Turbomole 6.5 software⁵⁸.

Dispersion effects were included by the DFT-D3 approach⁸¹ with Becke–Johnson damping⁸² and third-order terms, calculated with the dftd3 program.⁸³ Results presented in the text and the tables were obtained at the B3LYP-D3/def2-TZVPD level of theory. The dispersion correction is in generally modest for isomerisation, electron- and proton-transfer reactions (3–5 kJ/mol absolute average with a varying sign). However, for the O–O cleavage, O₂ binding, and the H₂O-dissociation reactions it is large and significant (–24, –35 and 22 kJ/mol on average, respectively).

We used an acetate model for Asp-373, a Cu(imidazole)₂(CH₃S)(CH₃SCH₃) model for the Cu_{T1} site, and a Cu₃(imidazole)₈ model for the TNC cluster with a variable number of water- and O₂-derived ligands. A water molecule hydrogen-bonded to the ligand bridging the two Cu_{T3} atoms was included in all systems.¹² Typical examples of both sites are shown in Figure 3. The Cu_{T1} site was studied in both the oxidised and reduced states, with a net charge of either zero or one, and with no or one unpaired electron, respectively. For the TNC, all possible oxidation states between Cu(I)₃ and Cu(II)₃ were studied with unrestricted formalism. With more than one Cu(II) ion, the antiferromagnetically coupled state with no or one unpaired electron¹³ was studied using the broken-symmetry approach⁸⁴ for all calculations, both geometry optimisations and single-point energy calculations.

MM and MD calculations

All MM and MD calculations were carried out with the sander module in the Amber 10 software⁵⁹ using the Amber 99SB force field.⁸⁵ The bond lengths involving hydrogen atoms were constrained using the SHAKE algorithm.⁸⁶ The electrostatics were treated using the particle-mesh Ewald method^{87,88} with a grid size of 80³, a fourth-order B-spline interpolation, a tolerance of 10^{–5}, and a real-space cut-off of 8 Å. The temperature was kept constant at 300 K and the pressure was kept constant at 1 atm using the Berendsen weak-coupling algorithm⁸⁹ with a time constants of 1 ps in both cases. The MD time step was 2 fs and the non-bonded pair list was updated every 50 fs. All MD simulations were performed with explicit water molecules, using the TIP3P water model.⁹⁰

Result and Discussion

In this paper, we study the full catalytic cycle of the MCOs using theoretical methods. We have optimised 52 putative intermediates in the reaction mechanism using the QM/MM-2QM method and have subsequently used the QTCP-2QM approach to obtain accurate energies for the various steps. The QTCP-2QM results were compared to the corresponding QM/MM-2QM and QM/MM-PBSA-2QM energies to confirm that reliability and consistency of the results. Our focus has been to study possible electronic and protonation states, in particular during the reformation of the reduced state of the TNC from the native intermediate.

According to the current consensus mechanism,^{1, 13} the resting oxidised state (Ox) involves a Cu(II)_3 cluster for the TNC with a hydroxide ion coordinated to $\text{Cu}_{\text{T}2}$ and another hydroxide ion bridging between the two $\text{Cu}_{\text{T}3}$ ions. Likewise, the reduced state (Red) contains a Cu(I)_3 cluster with a water molecule or hydroxide ion coordinated to $\text{Cu}_{\text{T}2}$ and no bridging ligand between the two $\text{Cu}_{\text{T}3}$ ions.¹⁴ The PI is formed directly from the Red state and it contains a O_2^{2-} ion bound in the centre of the TNC in a diagonal manner. Finally, in the NI, the O–O is cleaved and one of the O^{2-} ions is bound in the centre of the TNC, whereas the other O atom is probably protonated and bridges the two $\text{Cu}_{\text{T}3}$ ions. These four states constitute the starting point of the present investigation and they are shown in Figure 3. Except for the Ox state, the protonation state of the $\text{Cu}_{\text{T}2}$ ligand is uncertain (for simplicity, we have used a hydroxide ligand for the other states in the figure).

In this study, we have tried to connect these states by adding electrons or protons and letting water molecules dissociate. In principle, the TNC in the NI state should be connected to the Red state by the addition of three electrons and three protons (the fourth electron and proton are transferred to the TNC during the conversion of the PI to the NI). Apparently, there are two intermediate oxidation states of the TNC between NI and Red (note that NI is at the same oxidation level as the Ox state): IOx with $\text{Cu(II)}_2\text{Cu(I)}$ and IRed with Cu(II)Cu(I)_2 . To simplify the discussion, we have used the following nomenclature to describe the various states: First the oxidation state of the TNC is specified by giving the state (Red, IRed, IOx, NI, PI, or Ox; we use NI' for the one-electron reduced PI, i.e. the NI state before the O–O bond is cleaved, sometimes denoted $\text{PI}+\text{e}$). Then, the ligands (without formal charges) are given within brackets in the order $\text{Cu}_{\text{T}2}$ ligand, (i.e. the ligand binding only to $\text{Cu}_{\text{T}2}$), the ligand bridging the two $\text{Cu}_{\text{T}3}$ ions, and possibly a central ligand interacting with all three TNC Cu ions. If only two ligands are given, the central ligand is missing for IOx, IRed, and Red, but the bridging ligand for PI and NI'. With this nomenclature, the four states shown in Figure 3 are Ox(OH,OH) , $\text{Red(H}_2\text{O,-)}$, $\text{PI(H}_2\text{O,O}_2)$, and $\text{NI(H}_2\text{O,OH,O)}$.

In total, 52 different states of the TNC were studied (7 Red states, 4 PI states, 4 NI' states, 8 NI states, 13 IOx states, 14 IRed states, and 2 Ox states). They are connected by addition of protons and electrons or dissociation of water molecules in the way illustrated in Figure S1 in the supporting information. In fact, there are additional possible states: We have considered only states with water or OH^- as the $\text{Cu}_{\text{T}2}$ ligand and only states in which the central ligand has the same number of protons or less protons than the bridging $\text{Cu}_{\text{T}3}$ ligand (because the additional interactions with the copper ions of the former ligand are expected to make it more acidic). After identifying an initial tentative reaction pathway, we have ensured to include all states that differ by one electron, one proton, or by isomerisation, within the above rules. This led to the states included in Figure S1. For all states of the TNC, we have considered structures with both a reduced and an oxidised $\text{Cu}_{\text{T}1}$ site. Moreover, structures have been obtained both with and without neutralised solvent-exposed charges, as well as with Asp-373 as the second QM system (instead of the $\text{Cu}_{\text{T}1}$ cluster) in the protonated or deprotonated state.

In the following, we will discuss the electronic structure, the geometries, redox potentials, acidity constants, isomerisation reactions, the chemical reaction of O–O cleavage, and O_2 and water-binding free energies in separate sections. Finally, we discuss the implications of the present results on the reaction mechanism of the MCOs and how the two copper sites influence each other.

Electronic structure

We will start by discussing the electronic structure of the various complexes. All Red structures are closed-shell systems, with a well-defined Cu(I)_3 state for the TNC. However, for the other states, one (IRed and NI'), two (IOx and PI), or all three (NI and Ox) copper ions are formally oxidised, but the unpaired spin on the copper ions are typically delocalised also onto the ligands, especially on those that have a formal negative charge. The electronic structure is further complicated by the fact that all these states, except IRed and NI' are expected to be antiferromagnetically coupled so that the net number of unpaired electrons (in the broken-symmetry approximation) is zero (IOx and PI) or one (NI and Ox). Consequently, the spin density on the oxidised copper ions is typically 0.4–0.7 e (and never above 0.80 e). The observed Mulliken

spin population on the three Cu ions and the O ligands are collected in Table S3 for the various complexes (obtained at the B3LYP/def2-TZVPD level).

The spin density on the Cu ions is quite varying. For the Ox states, it is high on all Cu ions, but slightly lower on Cu_{T2} than on the two Cu_{T3} ions (0.43–0.58 and 0.65–0.76 e, respectively), although one is negative, reflecting the antiferromagnetic coupling and that all three Cu ions are oxidised. The NI states mostly have a similar electronic structure, with rather large spins (0.29–0.56 on Cu_{T2} and 0.47–0.80 e on Cu_{T3}, but with varying signs). The only exception is the NI(H₂O,O,O) complex, for which the spin is small (<0.07 e) on Cu_{T2}. The spin on the copper ions is slightly higher when calculated with the def2-TZVPD basis set than with the def2-SV(P) basis set and typically ~0.25 e higher when calculated with B3LYP than with PBE and the spin on the ligands is typically also higher.

For eight of the IRed states, one of the copper ions has a larger spin population, indicating that it is oxidised. However, for the other five states, at least two Cu ions have spin densities of similar magnitude, making it hard to identify the oxidised ion. Likewise, only for nine of the IOx states there is a pair of Cu ions with a large spin population with opposite spin and a minor population on the third Cu ion, as expected for a Cu(II)₂Cu(I) state. For the other states, all Cu ions have significant spin or one Cu ion has appreciably more spin than the other two ions.

In the PI state, the spin is typically similar (but with varying signs) on all three copper ions (0.05–0.18 e with O₂²⁻ and 0.18–0.43 e with HO₂⁻), making it hard to identify which two copper ions are oxidised. However, the electronic structure seems to be flexible and for some calculations, one Cu ion has either a larger or lower spin population than the other two. For the NI' states with a OH⁻ Cu_{T2} ligand, the Cu_{T2} ion is oxidised (0.33–0.48 e spin) and the two Cu_{T3} ions reduced. However, in NI'(H₂O,HO₂), both Cu_{T3} ions have a high spin (0.24–0.29 e) and in NI'(H₂O,O₂), all three Cu ions have a low spin (0.00–0.15 e).

All the PI and NI' states have a O₂-derived ligand in the centre of the cluster. For the PI states, the spin on this ligand is fairly low (0.09–0.34 e), indicating a rather pure peroxide level, as has been observed before.¹² However, sometimes larger spin populations are observed for the HO₂⁻ ligand (up to 0.49 e), connected to increased Cu populations. In the NI' states, the spin on the O₂ ligand increases to 0.40–0.74 e, indicating an electronic structure more between the peroxide and superoxide levels, except for the NI(OH,HO₂) state for which the spin population decreases from 0.30–0.40 e to 0.12–0.16 e.

Water ligands always have only a minimal spin, typically 0.00–0.03 e, but occasionally up to 0.10 e for the Cu_{T3} ligands in a few complexes.

O²⁻ ligands have a spin that varies with the oxidation state of the TNC. For the IOx states, the spin is typically rather low 0.00–0.34 e and often negative, reflecting that it is always a bridging ligand and that the net spin of the cluster is vanishing. For the IRed clusters, it has a higher spin, 0.35–0.53 e when the Cu_{T2} ligand is H₂O, and 0.10–0.21 e when it is OH⁻, i.e. similar or even larger than for the three Cu ions (up to 0.61 e). This mainly reflects that these complexes are not antiferromagnetically coupled. For the NI structures, the spin on O²⁻ is similar, 0.51–0.70 e when the Cu_{T2} ligand is H₂O (but 0.33–0.49 e for the two O²⁻ groups in Nat(H₂O,O,O)), and 0.17–0.31 e when it is OH⁻, reflecting that these complexes are antiferromagnetically coupled but have a single unpaired electron.

Hydroxide ligands also have a varying spin. For most states, it is rather low, 0.00–0.25 e. However, for three complexes with two OH⁻ ions, Ox(OH,OH) and IOx(OH,OH), it is much higher, 0.36–0.43 e, on the Cu_{T2} ligand. It is also high on the Cu_{T2} ligand in the IRed(OH,H₂O) and PI(OH,HO₂) complexes, 0.26–0.39 e.

Geometries

Cu–ligand distances of all optimised structures are collected in Table S4. The Cu_{T1} site consists of a Cu ion bound to one Cys, one Met, and two His residues. When it is reduced, all Cu–ligand distances are elongated. The bond to Cys-500 increases from 2.18–2.20 to 2.22–2.23 Å, with a variation of only 0.01 Å over the 52 studied TNC states. The bonds to the two His residues increase from 1.98–2.00 Å to 2.00–2.02 Å for His-505 but to 2.03–2.07 Å for His-443. The former bond is 0.01–0.02 Å longer in the optimisations with the neutralised protein, whereas the latter

bond is shorter by a similar amount. Finally, the bond to Met-510 is appreciably weaker and it shows a larger variation (2.93–3.36 Å), but it always increases by 0.11–0.22 Å when the site is reduced.

The Cu–Cu distances in the TNC primarily depend on the nature of the central and bridging ligands. In particular, a central O^{2-} ligand always gives rise to short Cu–Cu distances (average distance < 3.45 Å), whereas all other states have longer Cu–Cu distances. The oxidation state and the bridging ligand also has some influence, but in a less systematic manner. For example, shortest Cu–Cu distances are observed for the IRed(H_2O, OH, O), IOx(H_2O, O, O), and Red(H_2O, O) states (average Cu–Cu distances of 2.84, 2.90, and 2.97 Å, respectively), whereas the other states with a central O^{2-} ligand have average Cu–Cu distance between 3.13 and 3.45 Å. States with other ligands show less systematic trends, but states without any central or bridging ligands give the longest Cu–Cu distances (4.54–4.72 Å), followed by complexes with only a bridging water molecule (4.38–4.51 Å). States with a HO_2^- ligand also give longer average Cu–Cu bonds (4.07–4.40 Å) than states with a O_2^{2-} ligand (3.92–4.11 Å).

The Cu_{T2} ion binds to two His residues with distances of 1.86–2.22 Å. The longer distances (>2.0 Å) are observed only for the IOx(OH, OH, O), IRed(H_2O, OH, O), and Red(H_2O, O) states. If the Cu_{T2} ligand is OH^- , it binds at a distance of 1.85–1.93 Å in most cases, except in the IOx/IRed(OH, O) and NI($OH/(H)O_2$) states, in which it is 1.95–2.02 Å. If it is instead H_2O , it binds at a distance that increases from ~2.12 Å for the Ox states to ~2.35 Å for the Red states. If there is a central ligand, or even a bridging Cu_{T3} ligand, the distance can increase even more, especially for the IRed and Red states, up to 4.00 Å for Red(H_2O, O). A central O^{2-} ligand always coordinate also to the Cu_{T2} ion at a distance of 1.87–2.03 Å. A central OH^- ion binds at a similar distance (1.92–2.03 Å).

A central (H) O_2 ligand is normally close to the Cu_{T2} ion 2.08–2.35 Å, except in the PI(H_2O, HO_2) state (and also in the PI(OH, HO_2) and NI'(H_2O, HO_2) with the neutralised protein), up to 3.31 Å. The distance is often longer in the PI than the NI' states, and longer for HO_2^- than for O_2^{2-} . The second O atom is further away from the Cu_{T2} ion (2.92 Å or larger), owing to the diagonal binding of the peroxide. Thus, most NI' complexes have a significant Cu_{T2} –O interaction with the (hydro)peroxide ligand in contrast to what was reported by Solomon and coworkers. It should be noted that for each complex, many different structures can be found, often with quite different Cu–O bond lengths (and of course also with the diagonal (hydro)peroxide directed either towards the first or the second Cu_{T3} ion). These structures are typically close in energy (within ~10 kJ/mol). We have spent quite some effort to obtain the lowest energy structure for each complex (presented in the Tables), although this selection does not significantly affect the reported energies.

The Cu–N distances of the two Cu_{T3} ions are typically 1.94–2.15 Å, i.e. somewhat longer than for the Cu_{T2} ion, which reflects the higher coordination number of the Cu_{T3} ions. The distance for His-499 is longer than the other five distances (>1.99 Å). Quite frequently longer distances are encountered when there is a central O^{2-} or OH^- ion. In the IOx/IRed(OH, H_2O, O) states, one of the His ligands has dissociated and instead forms a hydrogen bond to the Cu_{T3} water ligand (which is not bridging).

A central O^{2-} ion typically binds with nearly symmetric Cu_{T3} distances of 1.87–1.99 Å. However, the presence of another bridging Cu_{T3} ligand often makes the distances longer (up to 2.08 Å) and less symmetric.

Eight states have a central OH^- ion. Those with a bridging OH^- ion have quite symmetric distances to this ion (1.93–2.07 Å), whereas the distances to the central OH^- ion are less symmetric, 1.92–2.11 and 2.28–3.10 Å (depending on the direction of the proton). However, the two complexes with a bridging water molecule are much more distorted because one of the protons on the water molecule does not have any natural hydrogen-bond partner. Consequently, there is one short and one long Cu–O bond for both the OH^- ion (1.97–2.09 and 3.25–3.32 Å) and the water molecule (2.04–2.14 and 2.99–3.18 Å).

A (H) O_2 ligand binds in a diagonal fashion with one short and one long bond to each Cu_{T3} ion. The former bond is slightly shorter for O_2^{2-} (1.93–2.06 Å) than for HO_2^- (1.99–2.12 Å), but there is no consistent difference between the PI and NI' oxidation states. The longer bond is 2.62–3.22 Å. In all cases, the O–O bond is intact, 1.32–1.50 Å. This is intermediate between the bond length

of a free superoxide ion, calculated at the same level (1.35 Å for O_2^- ; 1.32 Å for HO_2) and a free peroxide ion (1.62 Å for O_2^{2-} ; 1.53 Å for HO_2). The O–O bond length is 0.03–0.08 Å longer for the NI' states than for the PI states, indicating a stronger superoxide character in the latter cases (which is also supported by the spin densities). It is also 0.02–0.07 Å longer for HO_2^- than for O_2^{2-} and it is 0.03–0.08 Å longer if the Cu_{T2} ligand is a hydroxide than if it is water. Consequently, the PI/NI'(H₂O,O₂) states have the shortest O–O bond and the PI/NI'(OH,HO₂) states have the longest bond, whereas the PI/NI'(OH,O₂) and PI/NI'(H₂O,HO₂) states have similar O–O bond lengths.

A bridging OH[−] group binds with Cu–O distances of 1.91–2.09 Å with an average difference between the two Cu ions of 0.05 Å (the distorted Red(H₂O,OH,O) state has one Cu–O distance of 2.10–2.17 Å). A bridging water ligand has longer Cu–O distances 2.04–2.77 Å and a larger asymmetry (in many cases with central ligands, the water molecules binds effectively only to one of the two Cu_{T3} ions). The bond lengths decrease when the oxidation state of the Cu ions increases.

A change in the protonation state of Asp-373 has a very small effect of the geometry of the TNC: All the Cu–ligand distances change by less than 0.02 Å, showing that the geometry optimisations give stable results. Likewise, the Cu–ligand distances of the TNC are not much influenced by the oxidation state of the Cu_{T1} site, with differences typically below 0.05 Å.

Redox potentials

We have calculated redox potentials between 21 different states of the TNC. To make the calculations more stable, we do not calculate the redox potential of the TNC directly, but instead study the transfer of an electron from the Cu_{T1} site to the TNC. Thereby, the net charge of the total system does not change and long-range electrostatic effects are reduced. The absolute redox potential of the TNC can be estimated using the experimental redox potential of the Cu_{T1} site, 0.46 V at pH 7.0⁹¹ (throughout this article, all redox potentials are versus the standard hydrogen electrode).

The redox potentials of the various TNC states relative to the Cu_{T1} site, calculated with QTCP approach are listed in Table 1. It can be seen that the statistical precision of all calculated potentials is reasonable, 0.01–0.03 V (1–3 kJ/mol). A positive relative redox potential indicates that the TNC can be reduced by the Cu_{T1} site, i.e. that the forward reaction in the MCO reaction cycle is favourable. The results show that most of the reactions are predicted to be favourable in this direction. Moreover, the relative redox potential of the TNC decreases (becomes more negative) as the number of protons decrease, i.e. the net charge of the TNC decreases: The relative redox potential is positive for TNC states with a net charge of +3, +4, and +5 (0–1.4, 1.6–2.5, and 4.7 V, respectively) whereas it is negative (−0.4 to −1.5 V) for states with a net charge of +2. This is a combined effect of the electrostatic solvation energy, which always is negative, but decreasing with the charge (around −7.4, −5.6, −3.8, and −2.1 V for TNC charges of +5, +4, +3, and +2) and the intrinsic QM redox potential, which is always positive, but decreasing even more with the net charge.

Acidity constants

We have also calculated acidity constants for 40 states of the TNC (and for the Cu_{T1} site in both the reduced and oxidised states). To make the results more stable by conserving the total charge of the system, the proton was moved to a solvent-exposed carboxylate group on the other side of the protein, rather than simply being deleted. After some testing (discussed in the SI), we selected Asp-373 as the proton acceptor, because it gave results close to the average among the tested groups and it does not form any hydrogen bond with other protein groups, neither in the protonated nor the deprotonated state. Since the pK_a value of solvent-exposed carboxylate groups on the surface of a protein seldom change by more than one unit,^{34, 44} we can obtain the acidity constant of the TNC by using the intrinsic pK_a of an Asp group, ~4.0.^{34, 70} Consequently, protonation of the TNC is predicted to be favourable at pH 7 if the calculated relative pK_a value in Table 2 is larger than 3.

As shown in Table 2, it can be seen that the results predict that the Ox(OH,OH) state is more stable than the Ox(H₂O,OH) state throughout the physiological pH range, in accordance with

experimental observations,^{10, 92} giving some credence to the calculations. As expected, the relative pK_a value always increases (becomes more positive) if a TNC site with the same ligands is reduced by one electron (by 10–40 pK_a units; average 30) and it decreases as the number of protons (and therefore the net charge) of the TNC in the same oxidation state is increased (pK_a values of –59 to –63, –24 to –46, –25 to 19, and 8–38 for TNC sites with a net charge of +5, +4, +3, and +2, respectively). Consequently, the calculations always predict that the TNC states with a net charge of +2 or +3 are most stable.

The statistical precision of the calculated results is reasonable, 0.6–0.9 pK_a units, except for the three reactions involving the IOx(H₂O,H₂O,O) state (1–3 pK_a units) and also one reaction involving the corresponding IRed(H₂O,H₂O,O) state (0.9–1.1 pK_a units), which come from steric clashes in one of the coordinate perturbations.

Isomerisation free energies

We have also studied 19 isomerisation reactions between states that differ in the position of a single proton (in all cases with the Cu_{T1} site in both the reduced and oxidised states and using a neutralised protein). The calculated isomerisation free energies are collected in Table 3, using the convention that a negative sign indicates that the product is more stable. The precision is 1–3 kJ/mol, again except for four reactions involving the IOx and IRed(H₂O,H₂O,O) states.

The results show that it is more favourable to protonate the Cu_{T2} ligand than a peroxide ligand (i.e. (H₂O,O₂) is more favourable than (OH,H₂O₂)). States with two O²⁻ ions (X,O,O) are not favourable, but states with two OH⁻ ions (X,OH,OH) are energetically preferred over (X,H₂O,O) states, independent on the oxidation state. On the other hand, the proton transfer from a Cu_{T2} water molecule to a central O²⁻ ion (i.e. from (H₂O,X,O) to (OH,X,OH)) can be either favourable or unfavourable, depending on the oxidation state of the TNC and the presence of a bridging Cu_{T3} ligand.

We have also studied the reaction free energies for the cleavage of the O–O bond in three NI' states. These results are also included at the end of Table 3 and they show that the reaction is slightly uphill if the peroxide is not protonated, whereas it is strongly downhill if it is protonated. The reaction is further favoured if the Cu_{T2} ligand is a OH⁻ ion. These results are similar to our previous QM/MM results of the same reactions,²⁶ although there are differences of 15–34 kJ/mol in the energies, owing to differences in the methods and the basis sets (the DFT-D3 dispersion correction, which was not included in the previous study, contributes by 23–25 kJ/mol), as well as the rather large uncertainty of the QTCP results, 5–10 kJ/mol.

Yoon and Solomon have studied the O–O bond cleavage with QM-cluster calculations.²⁰ They started from a NI'(H₂O,O₂) state, but with Glu-506 protonated and moved so that it formed a direct hydrogen bond to the lower O₂ atom, and also with several bond lengths, angles, and dihedrals fixed to the values in the crystal structure to avoid artificial changes in the structures. They observed that the proton is transferred to the peroxide ion, either before or during the reaction and obtain a much more exergonic reaction free energy (204 kJ/mol) than in any of our calculations (although we do not study exactly the same reactions). They assigned the different modes of proton transfer to the experimental observation of different kinetic isotope effects at low and high pH, but this is unlikely, as a change of the external pH would change the total number of protons in the modelled reaction (e.g. by removing the extra proton on the Glu-506 model at high pH). They reported that the state in which the proton has moved from Glu-506 to O₂ is 9 kJ/mol lower than the other state and that the barrier for this proton transfer is low. This means that the state with Glu-506 protonated would instantly be converted to the other state and should not have any influence on the reaction rate, independent of the pH. In addition, they reported an activation barrier of only ~12 kJ/mol, which would correspond to a rate of ~10¹⁰ s⁻¹, (using a standard prefactor in the Arrhenius equation, 6·10¹² s⁻¹⁹³), much larger than the observed rate of > 350 s⁻¹. We strongly believe that our calculations including the full protein without any constraints should be more realistic.

Binding free energies

Our results also let us to estimate the binding free energy of O₂ to the Red complexes, forming the PI, and the dissociation of water from the various IOx, IRed, and Red complexes. These free energies are more approximate than the other energies in this article, because they are based on the QM/MM-PBSA calculations (because it is difficult to estimate binding free energies with the QTCP approach). Moreover, they include a contribution from the dissociated molecule that can be estimated in several different ways. Fortunately, it is only a constant factor that contributes equally to all reactions with the same binding molecule. For simplicity, we decided to estimate this contribution from a B3LYP/def2-TZVPD calculation in the COSMO continuum solvent with a dielectric constant of 80 (water). The dissociated species is also favoured by an entropy term from the enhanced translational and rotational freedom. The size of this term has been much discussed⁹⁴⁻⁹⁶ and we have modelled it by simply adding 30 kJ/mol to the dissociated species.

The resulting free energies are shown in Table 4. It can be seen that the dissociation of H₂O from all complexes is favourable (we have only studied the dissociation of a water molecule from a bridging position between the two Cu_{T3} ions). The reaction seems to be more favourable for the IOx and IRed complexes than for the Red complexes.

On the other hand, binding of O₂ is favourable for the Red(OH,-) complex, but not for the Red(H₂O,-) complex. However, these free energies are uncertain, because there is a very large difference between the results obtained by the PBE and the B3LYP methods: With PBE, both binding reactions are favourable by 49–91 kJ/mol. The reason for this is the change in the oxidation state of two copper ions and the O₂ ligand during binding: Large differences between pure (e.g. PBE) and hybrid DFT functionals (e.g. B3LYP) are often observed when the oxidation state of metal complexes is changed.^{32,97,98} For the dissociation of H₂O (which does not involve any change in the oxidation state of the TNC), the results with the two DFT functionals agree within 17 kJ/mol, except for two complexes (23–31 kJ/mol).

Communication between redox sites

The present calculations allow us to study how the two copper sites communicates, i.e. how changes in the redox or protonation state affects the redox potential and acidity of the other site. From Table 2, it can be seen that the acidity constants of the TNC in general decrease by ~3 pK_a units when the Cu_{T1} site is oxidised, reflecting that the Cu_{T1} site is much closer to the TNC (~12.5 Å Cu–Cu distance) than to Asp-373 (~24 Å Cu–CG distance; the TNC is always positively charged and this charge is one unit higher in the protonated state than in the deprotonated state; the Cu_{T1} site is neutral in the reduced state but has a single positive charge in the oxidised state).

For the isomerisation and O–O cleavage reactions in Table 3, it can be seen that the effect of the Cu_{T1} oxidation state is more varying (because the reactions conserve the net charge of the TNC), so it is no longer obvious which state will be favoured. For most reactions, the effect is restricted, 0–9 kJ/mol, but for seven reactions, it is larger, up to 28 kJ/mol, although this mainly reflects the larger computational uncertainty of these energies (especially those involving the IOx and IRed(H₂O,H₂O,O) states).

Further information about the communication between the two Cu sites in the MCOs can be obtained by using the QM/MM-PB(GB)SA results to calculate absolute reduction potentials for either that Cu_{T1} site or for the TNC in the various states of the other site. Concentrating first on the Cu_{T1} site, Figure 4 shows how the reduction potential of the Cu_{T1} site varies when the oxidation and protonation state of the TNC is varied. It can be seen that the Cu_{T1} potentials are predicted to vary by over 0.4 V, depending on the state of the TNC. However, this is primarily a charge effect, reflecting the repulsion between the oxidised state of the Cu_{T1} site (with a net charge of +1 e) and the TNC (with a net charge of 1–5 e), giving the highest potentials (Cu_{T1} reduced state most stable) for the highest charge. For the TNC states with a net charge of 2–3 (which in general seem to be preferred), the variation is only 0.18 V (0.35–0.53 V). Still, the results clearly show that the state of the TNC need to be specified if it should be meaningful to discuss the redox potential of the Cu_{T1} site to an accuracy better than 0.2 V and that the Cu_{T1} potential obtained for the resting Ox state probably is not representative for turn-over conditions.

Conversely, our calculations show that the Cu_{T1} site has a somewhat smaller effect on the redox potential of the TNC. An oxidised Cu_{T1} site gives a 0.06–0.13 V higher potential (0.08 V on average) than the reduced Cu_{T1} site (results in Table S4 in the SI). The reason is again the electrostatic interaction that is more unfavourable for the oxidised Cu_{T1} site (net charge +1) and the oxidised TNC, which always has one unit more positive charge than the reduced TNC.

Implication on reaction mechanism

We can now use these results to study the full reaction mechanisms of the MCOs. We start from the Red state with no bridging ligand. It can have either a water or OH[−] ion as the Cu_{T2} ligand, Red(H₂O,−) or Red(OH,−). The results in Table 2 indicate that these two states are close in energy at pH 7. They should bind O₂ and according to Table 4, this reaction is possible only for Red(OH,−). However, this suggestion is uncertain because the PBE calculations indicate that the O₂ binding to Red(H₂O,−) is also favourable by 49 kJ/mol.

When O₂ is bound, a PI state is formed, either PI(H₂O,O₂) or PI(OH,O₂). The results in Table 2 indicate that the PI(OH,O₂) state is most stable at pH 7. Moreover, it can be seen that the peroxide ion is unlikely to be protonated, in accordance with experimental studies.¹⁰ From Table 1, it can be seen that only the PI(OH,O₂) state is unlikely to be reduced by the Cu_{T1} site. This is a general problem for all the considered states: The relative acid constants indicate that the states with a net charge of the TNC of +2 or +3 (i.e. quite deprotonated) are most stable, whereas the relative redox potentials indicate that these states are unlikely to be reduced, but the more protonated states are readily reduced by Cu_{T1}. A possible solution to this dilemma is that a proton transfer accompanies the electron transfer, i.e. a coupled electron–proton transfer (EPT), as has also been suggested by Solomon and coworkers for some states.³⁰ Therefore, we have in Table 5 combined the results from Tables 1 and 2. The electron is taken from Cu_{T1} and the proton from Asp-373 and the results are expressed in pK_a units, so that a forward electron and proton transfer is predicted to be favourable at pH 7 if the result is more positive than 3 (as in Table 2). These results are somewhat more uncertain than the results in Table 1 and 2, because they are obtained from two different QTCP calculations, using different optimised structures. It can be seen from Table 5 that the EPT is still slightly unfavourable (by 1–3 pK_a units or 9–17 kJ/mol), which is within the error limits of the present method (we seem underestimate the stability of the reduced TNC by ~25 kJ/mol, as we will see also for the other reactions).

The EPT from PI(OH,O₂) gives either the NI'(H₂O,O₂) or NI'(OH,HO₂) states, which are the most stable protonation states according to Table 2. However, the isomerisation energies in Table 3 indicate that the NI'(H₂O,O₂) state is the most stable NI' state, by 30–34 kJ/mol. This is somewhat unexpected, because the cleavage of the O–O bond in this state is slightly uphill (by 23–25 kJ/mol), whereas it is strongly downhill for the NI'(OH,HO₂). The favourable O–O cleavage in NI'(OH,HO₂) gives rise to NI(OH,OH,O), which is the most stable NI state. Alternatively, NI'(H₂O,O₂) may give rise to the NI(H₂O,O,O) state, which is predicted to isomerise to the NI(OH,OH,O) state, which is 63–84 kJ/mol more stable. This strongly favourable isomerisation may drive a slightly uphill O–O bond cleavage.

Again, NI(OH,OH,O) cannot be directly reduced by the Cu_{T1} site, but the EPT to IOx(OH,OH,OH) is only slightly unfavourable (by 4 pK_a units or 25 kJ/mol). The protonation and isomerisation energies in Tables 2 and 3 show that IOx(OH,OH,OH) is the most stable IOx state.

The direct reduction of IOx(OH,OH,OH) is unfavourable, but the EPT to IRed(OH,H₂O,OH) is favoured by 4 pK_a units (24 kJ/mol). From the latter state, a water molecule can readily dissociate (Table 4), giving rise to the IRed(OH,OH) state.

Again, the reduction of the IRed(OH,OH) state is unfavourable, but the EPT to Red(H₂O,OH) or Red(OH,H₂O) is favourable or nearly favourable at pH 7. The latter state is predicted to lose the water to form the Red(OH,−) state. IRed(H₂O,OH) seems to be the more stable, but it needs to be protonated to Red(H₂O,H₂O) before a water is predicted to dissociate (which could drive the protonation). Thereby, we are back to the reduced state without any bridging ligand, which is the starting point of our catalytic cycle. Structures of the key intermediates in this suggested mechanism are shown in Figure 5.

Conclusions

In this study we have used advanced QM/MM-FEP methods to study the complete reaction mechanism of the multicopper oxidases, including the re-reduction of the TNC after the formation of the NI. This mechanism involves four reduction and protonation reactions, which are quite challenging to study with theoretical methods, owing to the change in the net charge of the protein and the correlation energy. We have reduced these problems by studying only the internal electron transfer between the Cu_{T1} ion and the TNC or the proton transfer between the TNC and a carboxylate group on the protein surface, thereby keeping the net charge of the protein constant. Moreover, we have tested to neutralise all charged groups on the surface of the protein, which are known not to affect the reduction potentials or pK_a values, thereby making the calculated energies more stable. Finally, we have used several different methods, employing both explicit water simulations and continuum solvation methods, to ensure that the calculated energies are stable and consistent. Fortunately, the values of most of the calculated energies are quite large so that we can predict the direction of most of the reactions even if the accuracy of the calculations is not better than ~30 kJ/mol.

Our calculations give rise to a tentative reaction mechanism, depicted in Figure 5. It can be seen that the calculations predict that the net charge of the TNC (i.e. the copper ions and their direct ligands) is 1–3 e throughout the whole catalytic cycle (and the most stable state at each oxidation-state has a charge of 2 or 3 e), as could be expected from the presence of two conserved neutralising carboxylate groups close to TNC in the protein, Asp-112 and Glu-506. The TNC is predicted to be reduced by the Cu_{T1} site in the PI(OH,O₂), NI(OH,OH,O), IOx(OH,OH,OH), and IRed(OH,OH) states by coupled electron–proton transfer reaction. Moreover, water molecules are predicted to dissociate rather late in the mechanism, probably from the IRed(OH,H₂O,OH) and Red(OH,H₂O) or Red(H₂O,H₂O) states. However, it is less clear whether the O–O cleavage takes place for the NI'(H₂O,O₂) state (for which the reaction may be slightly uphill) or in the NI'(OH,HO₂) state, for which the reaction is strongly favourable.

In a recent QM-cluster study, Solomon and coworkers studied the re-reduction of NI to the Red state.³⁰ They also used B3LYP calculations, but with smaller basis sets and treating the surrounding protein only as a continuum solvent with a dielectric constant of 4, thereby being forced to fix two atoms in each imidazole ligand. They consider only the isolated TNC and in several cases show protons on OH[−] and H₂O groups pointing in unexpected directions (e.g. towards the Cu ions). Our QM/MM-FEP calculations should give a much more detailed account of the surrounding protein and dynamic effects. Solomon et al. considered only water as the Cu_{T2} ligand, whereas our results (cf. Table 2) indicate that most states are more stable with OH[−] as the Cu_{T2} ligand, giving a less positive net charge of the TNC, and therefore much less favourable electron-transfer energies. Moreover, they suggest that both water molecules dissociate in the Red state (actually the second water molecule never dissociates in their calculations, in contrast to the crystal structure of the Red state¹⁴), whereas the first water ligand dissociates in the IRed state in our mechanism. In fact, our results indicate that IRed(H₂O,H₂O,OH), involved in their mechanism has the most favourable water dissociation energy of all studied complexes in Table 4. This seems to be connected with a desire to reproduce kinetically estimated driving forces of the three electron-transfer reactions between NI and Red (IRed(H₂O,OH) would give a too low driving force for the last electron transfer, as in our calculations). We are not fully convinced that the driving forces suggested in ref. 30 are the only that can fit the kinetic data (a complicated mechanism with at least 14 kinetic parameters) nor that the QM-cluster or QTCP calculations are accurate enough to allow a quantitative comparison with the kinetic data. Still, our mechanisms agree that the most stable NI state is NI(X,OH,O) (where X is H₂O in their calculations and OH[−] in our calculations), that the most stable IOx state is IOx(X,OH,OH), and that the IRed(X,H₂O,OH) state is involved in the mechanism.

Our calculations show that there is a significant communication between the two Cu sites in the MCO, through the net charge (+1 or 0 for the Cu_{T1} site and +1–3 for the TNC): The redox potential of the Cu_{T1} is predicted to vary by up to 0.18 V depending on the state of the TNC. Likewise, the oxidation state of Cu_{T1} site is predicted to affect the redox potential of the TNC by ~0.08 V and modify acidity constants by ~3 pK_a units. This shows that it is rather meaningless to

discuss reduction potentials and acidity constants of one site without specifying the state of the other site.

In conclusion, we have shown how advanced multi-scale modelling can be used to gain atomistic information of the structures and energies of the various states in the reaction mechanism of the MCOs. We think that the presented methodology represents a robust computational treatment of challenging problems in the bioinorganic chemistry, establishing a link between computations and experimental data.

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Supporting Information Available

Test of various proton-accepting sites, influence of the basis sets on the calculated energies, spin states and geometries of the various states, and calculated absolute redox potentials. This information is available free of charge via the Internet at <http://pubs.acs.org>.

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Table 1. Redox potentials for the various TNC states relative to the Cu_{T1} site (ΔE° in V), calculated with QTCP. The protonation state of all ligands are the same in both oxidation states so the symbols in brackets for the former state are omitted. The more stable TNC state is marked in bold face, i.e. the reduced TNC state if $\Delta E^\circ > 0$ and the oxidised state otherwise.

TNC states	ΔE°
PI→ NI' (H ₂ O,HO ₂)	2.36 ±0.02
PI→ NI' (H ₂ O,O ₂)	0.35 ±0.02
PI→ NI' (OH,HO ₂)	1.08 ±0.02
PI →NI'(OH,O ₂)	-1.42 ±0.02
NI→ IOx (H ₂ O,H ₂ O,O)	1.61 ±0.02
NI→ IOx (H ₂ O,OH,O)	0.03 ±0.02
NI →IOx(OH,OH,O)	-1.45 ±0.02
IOx→ IRed (H ₂ O,H ₂ O,O)	0.35 ±0.02
IOx →IRed(H ₂ O,OH,O)	-1.68 ±0.02
IOx →IRed(OH,OH,OH)	-1.37 ±0.02
IOx→ IRed (H ₂ O,OH)	2.28 ±0.02
IOx→ IRed (H ₂ O,O)	0.59 ±0.02
IOx→ IRed (OH,OH)	1.35 ±0.02
IOx →IRed(OH,O)	-1.04 ±0.02
IRed→ Red (H ₂ O,H ₂ O)	2.42 ±0.02
IRed→ Red (H ₂ O,OH)	0.16 ±0.02
IRed →Red(H ₂ O,O)	-1.30 ±0.03
IRed →Red(OH,OH)	-0.38 ±0.02
IRed→ Red (H ₂ O,-)	2.72 ±0.02
Ox→ IOx (H ₂ O,OH)	4.72 ±0.01
Ox→ IOx (OH,OH)	2.52 ±0.02

Table 2. Calculated pK_a values for the various states of TNC relative to Asp-373, calculated with QTCP. Results are given both with the Cu_{T1} site reduced and oxidised. The more stable TNC state at pH 7 is marked in bold face, i.e. the protonated state if the pK_a value is larger than 3 and the deprotonated state otherwise.

TNC states	Cu_{T1} oxidised	Cu_{T1} reduced
$PI(H_2O, HO_2) \rightarrow (H_2O, O_2)$	-45.7 ± 0.6	-43.0 ± 0.6
$PI(H_2O, HO_2) \rightarrow (OH, HO_2)$	-31.2 ± 0.7	-28.7 ± 0.7
$PI(H_2O, O_2) \rightarrow (OH, O_2)$	-8.6 ± 0.7	-4.7 ± 0.7
$PI(OH, HO_2) \rightarrow (OH, O_2)$	-22.3 ± 0.6	-18.1 ± 0.6
$NI'(H_2O, HO_2) \rightarrow (H_2O, O_2)$	-10.4 ± 0.6	-6.1 ± 0.6
$NI'(H_2O, HO_2) \rightarrow (OH, HO_2)$	-2.7 ± 0.8	0.8 ± 0.8
$NI'(H_2O, O_2) \rightarrow (OH, O_2)$	25.9 ± 0.8	28.4 ± 0.8
$NI'(OH, HO_2) \rightarrow (OH, O_2)$	17.7 ± 0.6	20.7 ± 0.6
$NI(H_2O, H_2O, O) \rightarrow (H_2O, OH, O)$	-46.4 ± 0.6	-44.3 ± 0.6
$NI(H_2O, OH, OH) \rightarrow (H_2O, OH, O)$	-35.1 ± 0.6	-34.2 ± 0.6
$NI(H_2O, OH, O) \rightarrow (H_2O, O, O)$	15.5 ± 0.6	18.7 ± 0.6
$NI(H_2O, OH, O) \rightarrow (OH, OH, O)$	-11.2 ± 0.7	-10.7 ± 0.7
$IOx(H_2O, H_2O, OH) \rightarrow (H_2O, H_2O, O)$	-24.9 ± 2.1	-18.6 ± 3.3
$IOx(H_2O, H_2O, O) \rightarrow (H_2O, OH, O)$	-9.5 ± 2.4	-7.7 ± 3.1
$IOx(H_2O, H_2O, O) \rightarrow (OH, H_2O, O)$	-6.7 ± 1.9	-2.8 ± 1.4
$IOx(H_2O, OH, OH) \rightarrow (OH, OH, OH)$	-12.6 ± 0.7	-9.9 ± 0.7
$IOx(H_2O, OH, O) \rightarrow (OH, OH, O)$	14.1 ± 0.7	15.8 ± 0.7
$IOx(H_2O, OH, O) \rightarrow (H_2O, O, O)$	34.4 ± 0.6	38.1 ± 0.6
$IOx(OH, H_2O, O) \rightarrow (OH, OH, O)$	12.4 ± 0.6	15.5 ± 0.6
$IOx(OH, OH, OH) \rightarrow (OH, OH, O)$	22.9 ± 0.7	26.1 ± 0.7
$IOx(H_2O, OH) \rightarrow (H_2O, O)$	-28.0 ± 0.6	-26.7 ± 0.6
$IOx(H_2O, OH) \rightarrow (OH, OH)$	-24.4 ± 0.6	-23.9 ± 0.6
$IOx(H_2O, O) \rightarrow (OH, O)$	-11.1 ± 0.7	-8.0 ± 0.8
$IOx(OH, OH) \rightarrow (OH, O)$	-14.9 ± 0.6	-12.1 ± 0.7
$IRed(H_2O, H_2O, OH) \rightarrow (H_2O, H_2O, O)$	-14.5 ± 0.9	-8.2 ± 1.1
$IRed(H_2O, H_2O, OH) \rightarrow (OH, H_2O, OH)$	-18.5 ± 0.7	-18.1 ± 0.7
$IRed(H_2O, H_2O, O) \rightarrow (H_2O, OH, O)$	26.9 ± 0.7	27.3 ± 0.7
$IRed(OH, H_2O, OH) \rightarrow (OH, OH, OH)$	30.3 ± 0.8	26.1 ± 0.8
$IRed(H_2O, H_2O) \rightarrow (H_2O, OH)$	-45.0 ± 0.6	-42.3 ± 0.6
$IRed(H_2O, OH) \rightarrow (OH, OH)$	1.3 ± 0.7	5.0 ± 0.7
$IRed(H_2O, OH) \rightarrow (H_2O, O)$	5.7 ± 0.6	9.3 ± 0.6
$IRed(H_2O, O) \rightarrow (OH, O)$	17.0 ± 0.9	18.0 ± 0.9
$IRed(OH, OH) \rightarrow (OH, O)$	15.5 ± 0.6	16.7 ± 0.7
$Red(H_2O, H_2O) \rightarrow (H_2O, OH)$	-6.4 ± 0.6	-2.9 ± 0.6
$Red(H_2O, H_2O) \rightarrow (OH, H_2O)$	3.0 ± 0.7	5.4 ± 0.7
$Red(H_2O, OH) \rightarrow (H_2O, O)$	28.2 ± 0.7	37.9 ± 0.6
$Red(H_2O, OH) \rightarrow (OH, OH)$	17.4 ± 0.7	21.2 ± 0.7
$Red(OH, H_2O) \rightarrow (OH, OH)$	8.1 ± 0.6	11.9 ± 0.6
$Red(H_2O, -) \rightarrow (OH, -)$	2.6 ± 0.7	3.6 ± 0.7
$Ox(H_2O, OH) \rightarrow (OH, OH)$	-59.4 ± 0.6	-58.6 ± 0.7

Table 3. Calculated isomerisation and reaction free energies (in kJ/mol) for the various states of TNC, calculated with QTCP. Results are given both with the Cu_{T1} site reduced and oxidised. The more stable state is marked in bold face, i.e. the reactant if the isomerisation energy is positive and the product state otherwise.

Reactant	Product	Cu _{T1} oxidised	Cu _{T1} reduced
PI(H₂O,O₂)	PI(OH,HO ₂)	78.8 ±1.6	78.3 ±1.5
NI'(H₂O,O₂)	NI'(OH,HO ₂)	34.5 ±3.0	30.1 ±2.4
NI(H ₂ O,H ₂ O,O)	NI(H₂O,OH,OH)	-55.5 ±0.7	-53.7 ±0.6
NI(H₂O,OH,O)	NI(OH,H ₂ O,O)	121.9 ±1.7	112.7 ±1.7
NI(H₂O,OH,O)	NI(OH,OH,OH)	30.5 ±1.4	24.6 ±1.2
NI(H ₂ O,O,O)	NI(OH,OH,O)	-62.6 ±1.7	-83.5 ±1.8
IOx(H ₂ O,H ₂ O,O)	IOx(H₂O,OH,OH)	-1.3 ±19.5	-17.0 ±6.0
IOx(H ₂ O,H ₂ O,O)	IOx(OH,H ₂ O,OH)	10.1 ±11.6	-18.4 ±16.2
IOx(H₂O,OH,O)	IOx(OH,H ₂ O,O)	15.5 ±2.1	8.4 ±2.4
IOx(H ₂ O,OH,O)	IOx(OH,OH,OH)	-41.7 ±2.8	-41.3 ±2.4
IOx(H₂O,O)	IOx(OH,OH)	7.9 ±1.3	19.5 ±1.2
IRed(H₂O,H₂O,O)	IRed(H ₂ O,OH,OH)	35.7 ±18.8	17.0 ±8.4
IRed(H ₂ O,H ₂ O,O)	IRed(OH,H ₂ O,OH)	-13.7 ±10.7	36.9 ±7.1
IRed(H ₂ O,OH,O)	IRed(OH,OH,OH)	-53.5 ±2.6	-51.9 ±2.7
IRed(H ₂ O,OH,O)	IRed(OH,H₂O,O)	-38.2 ±2.3	-39.3 ±2.5
IRed(H₂O,OH)	IRed(OH,H ₂ O)	112.6 ±1.4	115.5 ±1.3
IRed(H ₂ O,O)	IRed(OH,OH)	-6.3 ±1.8	-15.4 ±1.6
Red(H₂O,OH)	Red(OH,H ₂ O)	50.7 ±1.9	48.2 ±1.8
Red(H ₂ O,O)	Red(OH,OH)	-61.0 ±2.9	-36.0 ±3.0
NI'(H₂O,O₂)	NI(H ₂ O,O,O)	22.8 ±8.7	25.3 ±8.3
NI'(H ₂ O,HO ₂)	NI(H₂O,OH,O)	-92.3 ±9.8	-86.8 ±9.4
NI'(OH,HO ₂)	NI(OH,OH,O)	-117.7 ±4.8	-116.4 ±4.8

Table 4. O₂ binding free energies and H₂O dissociation free energies for the various TNC states. The energies were calculated with the QM/MM-PBSA method with surface charges neutralised. An entropy contribution of 30 kJ/mol has been added to the dissociated O₂ or H₂O molecule and the energy of the unbound molecule was estimated by a B3LYP/def2-TZVPD calculation in the COSMO continuum solvent with a dielectric constant of 80 (water). The more stable state is marked in bold face, i.e. the reactant state if the energy is positive and the product state otherwise.

	Cu _{T1} oxidised Cu _{T1} reduced	
Red(H₂O,-) + O₂ → PI(H₂O,O₂)	31.3	31.3
Red(OH,-) + O ₂ → PI(OH,O₂)	-44.2	-38.9
IOx(H ₂ O,H ₂ O,OH) → IOx(H₂O,OH) + H₂O	-43.8	-47.7
IOx(H ₂ O,H ₂ O,O) → IOx(H₂O,O) + H₂O	-49.2	-51.2
IOx(OH,OH) + H₂O	-17.3	-15.2
IOx(OH,H ₂ O,O) → IOx(OH,O) + H₂O	-86.8	-81.6
IRed(H ₂ O,H ₂ O,OH) → IRed(H₂O,OH) + H₂O	-122.6	-121.0
IRed(H ₂ O,H ₂ O,O) → IRed(H₂O,O) + H₂O	-46.8	-40.5
IRed(OH,H ₂ O,OH) → IRed(OH,OH) + H₂O	-10.8	-11.4
IRed(OH,H ₂ O,O) → IRed(OH,O) + H₂O	-98.5	-97.6
IRed(H ₂ O,H ₂ O) → IRed(H₂O,-) + H₂O	-63.3	-63.6
Red(H ₂ O,H ₂ O) → Red(H₂O,-) + H₂O	-5.6	-4.5
Red(OH,H ₂ O) → Red(OH,-) + H₂O	-8.9	-10.0

Table 5. Coupled electron–proton transfer reaction energies for the various TNC states (ΔE_{EPT}), obtained from the QTCP energies in Tables 1 and 2. Thus, the electron comes from the Cu_{T1} site and the proton from Asp-373, and the results are give in pK_a units relative to that of an Asp residue (pK_a $\approx$ 4). Consequently, the coupled electron–proton transfer reaction is predicted to be favourable in the forward direction at pH 7 if ΔE_{EPT} is larger than 3 and the more stable states have been marked in bold face.

TNC states	ΔE_{EPT}
PI(H₂O,O₂) $\rightarrow$ NI'(H ₂ O,HO ₂)	-3.9 $\pm$ 0.7
PI(OH,HO ₂) $\rightarrow$ NI'(H₂O,HO₂)	13.2 $\pm$ 0.8
PI(OH,O₂) $\rightarrow$ NI'(H ₂ O,O ₂)	1.5 $\pm$ 0.9
PI(OH,O₂) $\rightarrow$ NI'(OH,HO ₂)	0.1 $\pm$ 0.7
NI(H₂O,H₂O,O) $\rightarrow$ IOx(H ₂ O,H ₂ O,OH)	2.2 $\pm$ 2.1
NI(H₂O,OH,O) $\rightarrow$ IOx(H ₂ O,H ₂ O,O)	-9.0 $\pm$ 2.4
NI(H ₂ O,O,O) $\rightarrow$ IOx(H₂O,OH,O)	19.3 $\pm$ 0.7
NI(OH,OH,O) $\rightarrow$ IOx(H ₂ O,OH,O)	-10.2 $\pm$ 0.7
NI(OH,OH,O) $\rightarrow$ IOx(OH,H ₂ O,O)	-12.0 $\pm$ 0.7
NI(OH,OH,O) $\rightarrow$ IOx(OH,OH,OH)	-1.4 $\pm$ 0.8
IOx(H₂O,H₂O,O) $\rightarrow$ IRed(H ₂ O,H ₂ O,OH)	-8.5 $\pm$ 0.9
IOx(H₂O,OH,O) $\rightarrow$ IRed(H ₂ O,H ₂ O,O)	-1.8 $\pm$ 3.1
IOx(H ₂ O,O,O) $\rightarrow$ IRed(H₂O,OH,O)	10.0 $\pm$ 0.6
IOx(OH,H ₂ O,O) $\rightarrow$ IRed(H₂O,H₂O,O)	3.2 $\pm$ 1.5
IOx(OH,OH,OH) $\rightarrow$ IRed(OH,H₂O,OH)	7.2 $\pm$ 0.9
IOx(OH,OH,O) $\rightarrow$ IRed(H ₂ O,OH,O)	-12.4 $\pm$ 0.7
IOx(OH,OH,O) $\rightarrow$ IRed(OH,OH,OH)	3.0 $\pm$ 0.8
IOx(H₂O,OH) $\rightarrow$ IRed(H ₂ O,H ₂ O)	-6.8 $\pm$ 0.7
IOx(H ₂ O,O) $\rightarrow$ IRed(H₂O,OH)	13.6 $\pm$ 0.7
IOx(OH,OH) $\rightarrow$ IRed(H₂O,OH)	19.1 $\pm$ 0.7
IOx(OH,O) $\rightarrow$ IRed(H ₂ O,O)	0.7 $\pm$ 0.9
IOx(OH,O) $\rightarrow$ IRed(OH,OH)	4.2 $\pm$ 0.7
IRed(H₂O,OH) $\rightarrow$ Red(H ₂ O,H ₂ O)	-2.7 $\pm$ 0.7
IRed(H ₂ O,O) $\rightarrow$ Red(H₂O,OH)	9.2 $\pm$ 0.8
IRed(OH,OH) $\rightarrow$ Red(H₂O,OH)	7.6 $\pm$ 0.7
IRed(OH,OH) $\rightarrow$ Red(OH,H ₂ O)	1.8 $\pm$ 0.7
IRed(OH,O) $\rightarrow$ Red(H₂O,O)	27.9 $\pm$ 0.9
IRed(OH,O) $\rightarrow$ Red(OH,OH)	39.3 $\pm$ 0.7
Ox(OH,OH) $\rightarrow$ IOx(H₂O,OH)	19.3 $\pm$ 0.7

Figure 1. The consensus reaction scheme of the MCOs, considering only the oxidation state of the TNC.^{1, 10, 13}

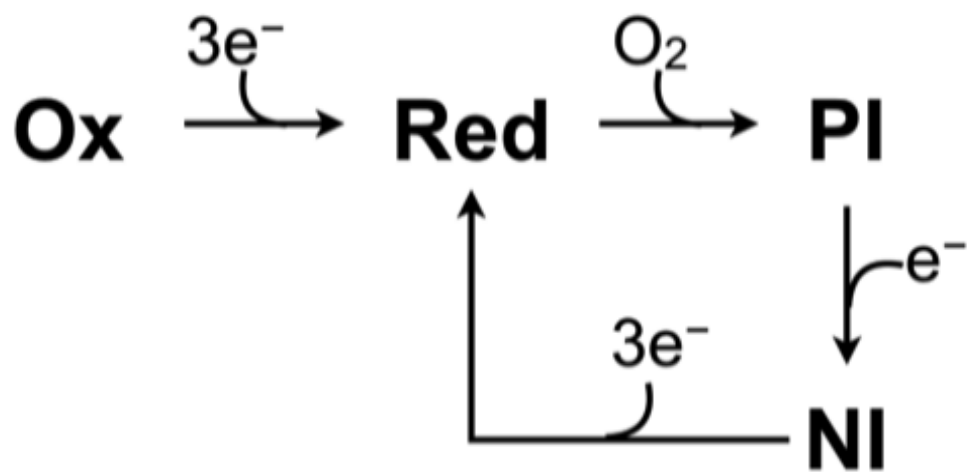


Figure 2. Schematic picture of the various perturbations performed for the electron- and proton-transfer reactions. In the former case the two letters for each system represent the charges and the coordinates, respectively, which may be either those for the starting state (A, in which the TNC is oxidised and Cu_{T1} is reduced) or the final state (B, in which the TNC is reduced and Cu_{T1} is oxidised). For the proton-transfer reaction, the four letters represent the proton on the TNC site, charges, the coordinates, and the proton on Asp-373. The proton can be either modelled as a full proton, with zeroed dummy parameters (D), or be deleted (Ø). The charges and coordinates can be either those of the starting state (A, in which the TNC is protonated and Asp-373 is deprotonated) or the final state (B, in which the TNC is deprotonated and Asp-373 is protonated). The label for each free energy and the number of perturbation steps are also indicated.

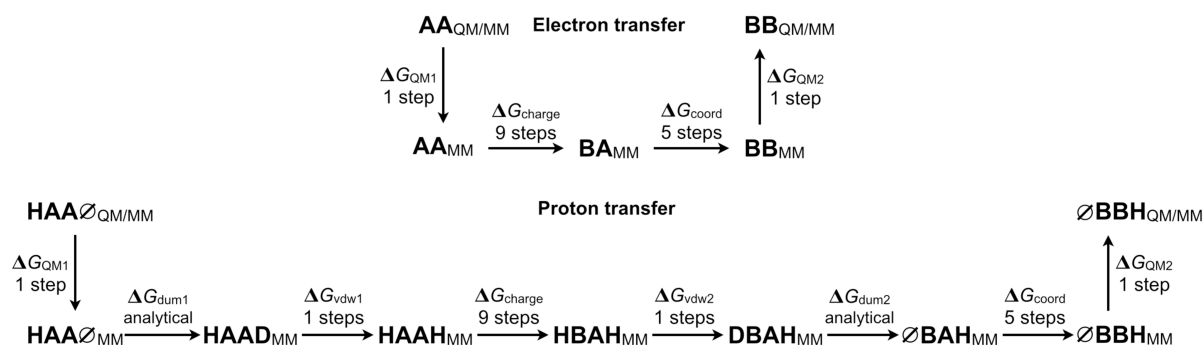


Figure 3. Structures of the QM systems for a) the Cu_{T1} site and for the TNC in the b) Ox(OH,OH), c) Red(OH), d) PI(OH,O₂), and e) NI(OH,OH,O) states. Cu ions are orange, S atoms yellow, C atoms grey, N atoms blue, O atoms red, and H atoms white.

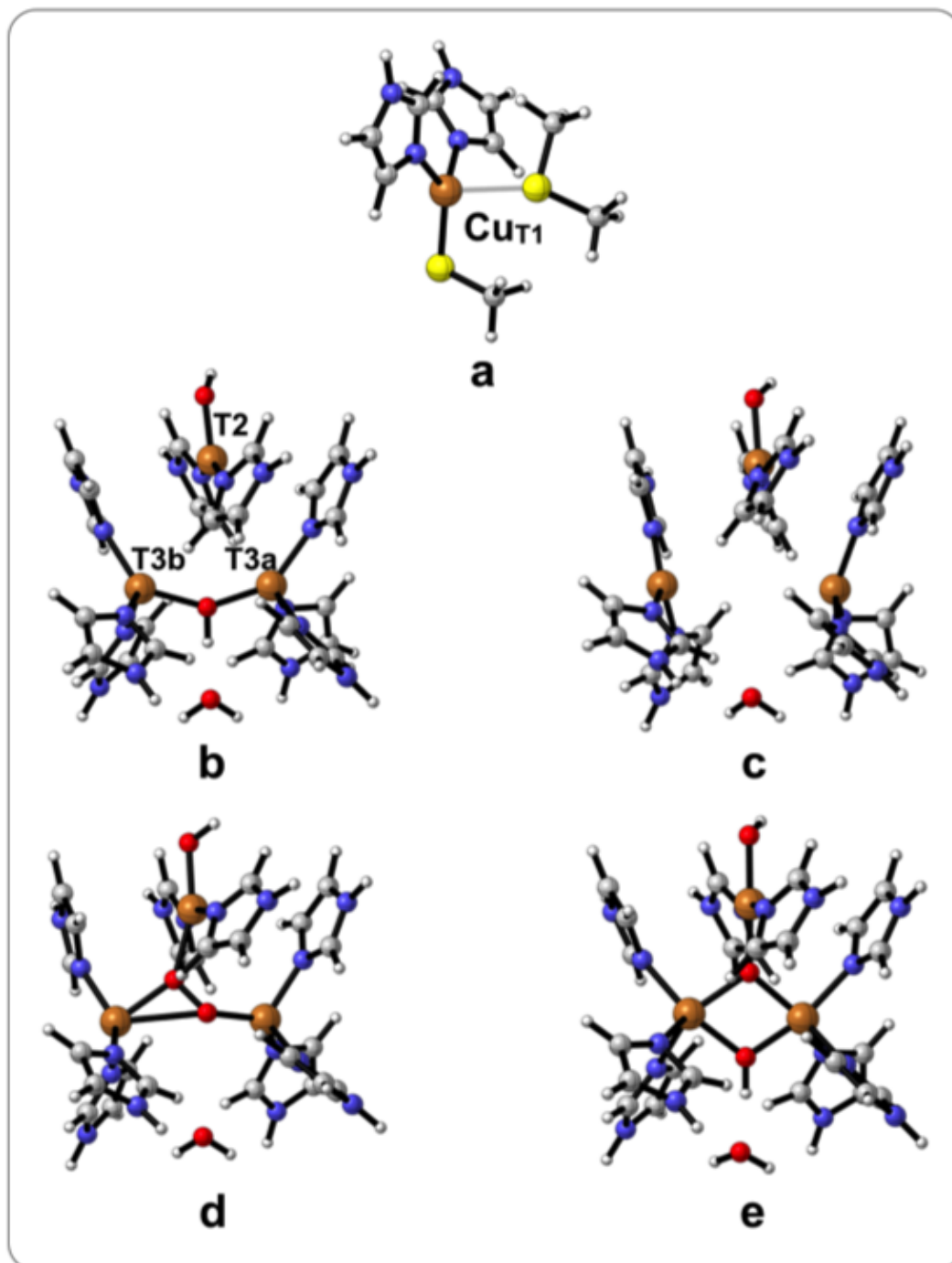


Figure 4. Dependence of the estimated redox potential of the Cu_{T1} site on the state and charge of the TNC, colour-coded after the charge of the TNC. The results are based on the QM/MM-GBSA results (which give the smallest influence on the redox potentials among methods used) for the proteins with neutralised charges, sorted after the net charge (legend) and oxidation state of the TNC site.

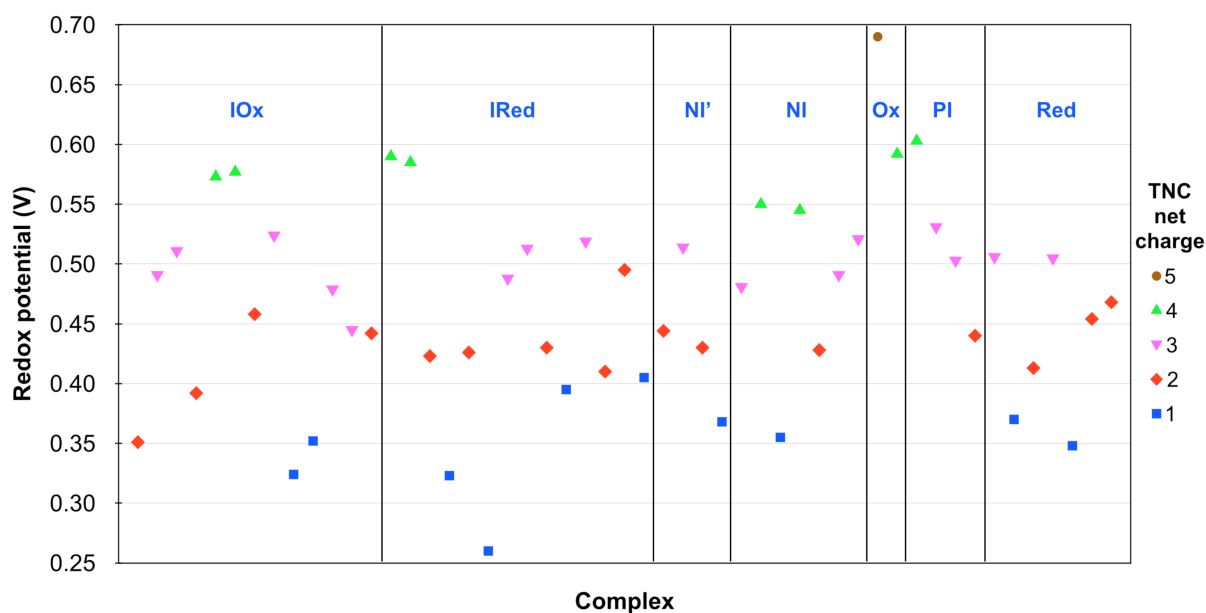
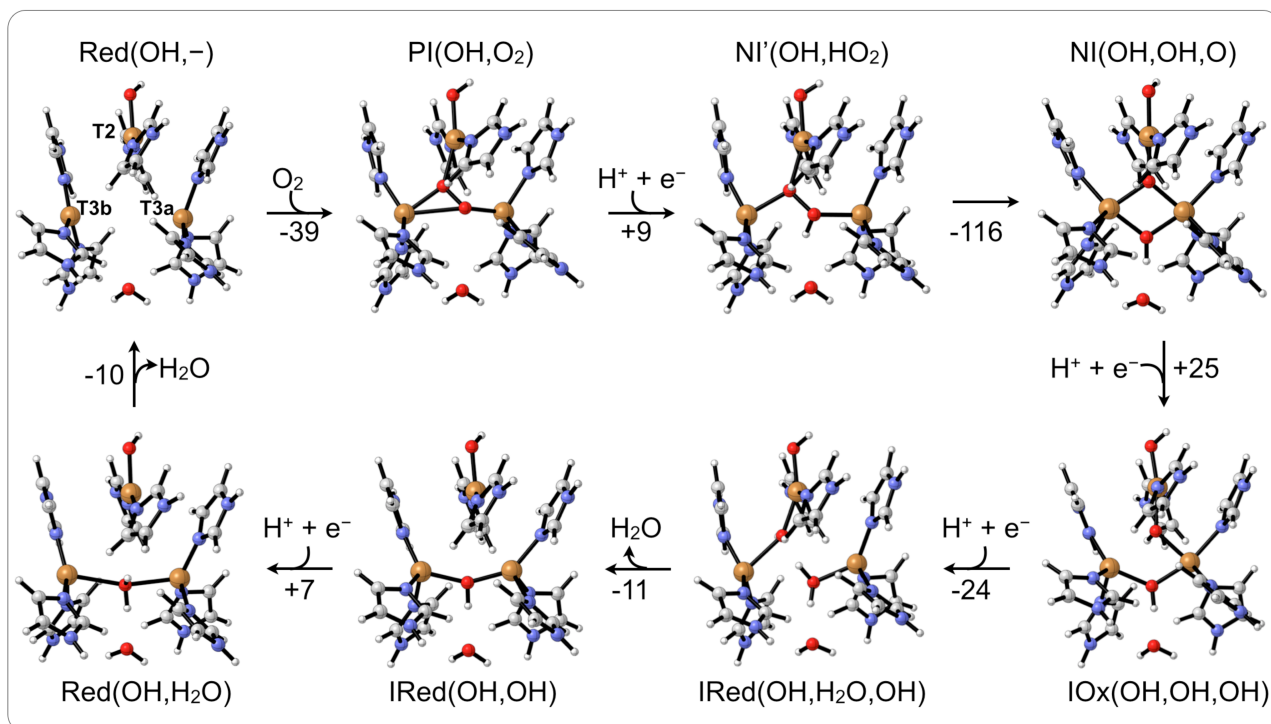


Figure 5. The suggested catalytic cycle of the MCOs. Calculated reaction free energies (in kJ/mol from Tables 3–5) are given for each step, assuming a pH of 7, that the electrons come from the Cu_{T1} site, and that the Cu_{T1} site is reduced in the steps not involving electron transfer.



TOC graphics

