Mechanism of hydrogen peroxide formation by lytic polysaccharide monooxygenase†

Octav Caldararu, a Esko Oksanen, b,c Ulf Ryde a and Erik D. Hedegård a,*

Lytic polysaccharide monooxygenases (LPMOs) are copper-containing metalloenzymes that can cleave the glycosidic link in polysaccharides. This could become crucial for production of energy-efficient biofuels from recalcitrant polysaccharides. Although LPMOs are considered oxygenases, recent investigations have shown that H$_2$O$_2$ can also act as a co-substrate for LPMOs. Intriguingly, LPMOs generate H$_2$O$_2$ in the absence of a polysaccharide substrate. Here, we elucidate a new mechanism for H$_2$O$_2$ generation starting from an AA10-LPMO crystal structure with an oxygen species bound, using QM/MM calculations. The reduction level and protonation state of this oxygen-bound intermediate has been unclear. However, this information is crucial to the mechanism. We therefore investigate the oxygen-bound intermediate with quantum refinement (crystallographic refinement enhanced with QM calculations), against both X-ray and neutron data. Quantum refinement calculations suggest a Cu(i)–O$_2$ system in the active site of the AA10-LPMO and a neutral protonated –NH$_2$ state for the terminal nitrogen atom, the latter in contrast to the original interpretation. Our QM/MM calculations show that H$_2$O$_2$ generation is possible only from a Cu(i) center and that the most favourable reaction pathway is to involve a nearby glutamate residue, adding two electrons and two protons to the Cu(i)–O$_2$ system, followed by dissociation of H$_2$O$_2$.

1 Introduction

Lytic polysaccharide monooxygenases (LPMOs) are metalloenzymes capable of activating molecular oxygen, and thereby inserting a single oxygen atom into the C–H bonds that comprise the glycoside link in polysaccharides. This oxidation leads to cleavage of the glycoside link, which may disrupt the surface of crystalline polysaccharides sufficiently to boost polysaccharide decomposition. Hitherto, this decomposition has been a major obstacle for energy-efficient production of biofuels from recalcitrant polysaccharides such as cellulose, which is one of the most abundant polysaccharides on earth.

A number of different LPMOs have been categorized, belonging to four distinct classes, AA9, AA10, AA11, and AA13. The various LPMOs have remarkably varying amino-acid sequences (no residues besides the histidine brace are strictly conserved between AA9, AA10, AA11 and AA13 LPMO classes) and target a wide range of different polysaccharide substrates. However, all LPMOs contain a copper ion coordinated by two histidine residues in what has become known as the histidine brace (cf. Fig. 1). In the histidine brace, one histidine is the amino-terminal residue that coordinates bidentately with both the N-terminus and the imidazole side chain.

It is believed that the mechanism of the LPMOs is initiated by reduction of Cu(II) to Cu(I), followed by binding of O$_2$ to Cu(I) to form a superoxide complex, [CuO$_2$]$^+$.

\[
\text{Cu}^{2+} + e^- \rightarrow \text{Cu}^+ \quad (1)
\]

\[
\text{Cu}^+ + \text{O}_2 \rightarrow [\text{CuO}_2]^+ \quad (2)
\]

![Fig. 1](image-url) The active site of AA10-LPMOs with the histidine brace and the nearby Phe164 residue, numbering according to PDB 5VG0. The figure also shows to the employed quantum system in the quantum-refinement calculations. (A) and (B) refer to chain A and B, respectively.
This complex has been suggested to be the reactive intermediate. However, theoretical work has shown that the superoxide is not sufficiently reactive to abstract a hydrogen from the glycoside C–H bond and points rather to either a copper-oxyl, or a copper-hydroxy, species. These more reactive copper–oxygen species can be generated by successive reductions and protonations of the superoxide species.

Another issue concerns the co-substrate. Bissaro et al. recently suggested that H$_2$O$_2$, rather than O$_2$, is the actual co-substrate, whereas Hangasky et al. proposed that both O$_2$ and H$_2$O$_2$ can be utilized by LPMOs as co-substrates. Intriguingly, LPMOs in fact generate hydrogen peroxide in absence of a polysaccharide substrate.

In a landmark study, combining theoretical and experimental methods, Kjærsgaard et al. investigated the initial reduction and reaction with O$_2$ (eqn (1) and (2)). Their data showed that Cu(II) and O$_2$ form a superoxide [CuO$_2$]$^+$ species, which is rapidly displaced by water to generate the resting state. Based on DFT calculations, they further suggested a mechanism for superoxide release by concerted O$_2$ dissociation and protonation of O$_2$ to HO$_2$ by an axially coordinated water molecule. This is a feasible mechanism for H$_2$O$_2$ formation, as HO$_2$ is known to form H$_2$O$_2$ through disproportionation in aqueous solution. However, no second-sphere residues were taken into account as possible proton donors.

Regarding the oxygen species, the paper by Kjærsgaard et al. is today supported by the X-ray and neutron diffraction study by O’Dell et al., which shows an O$_2$-bound intermediate from an AA9 LPMO, although they interpreted the species as a peroxide-bound (O$_2^\cdot$) intermediate. More recently, a similar intermediate was trapped in a crystal structure of an AA10 LPMO, investigated by both X-ray and neutron diffraction. This study has further attracted attention since the protein crystallizes as a mixture of [CuO$_2$]$^+$ species, or a mixture of [CuOH]$^+$ species. These more reactive copper–oxygen species can be generated by successive reductions and protonations of the superoxide species.

The nature and coordination of this species in more detail. This is particularly pertinent, given that both the metal ion and the coordinated oxygen species are susceptible to radiation damage, making conclusions from X-ray structures dubious. Indeed, this is a known issue even in high resolution X-ray structures and has also previously been a problem in crystal structures for LPMOs. The problem can be partly remedied by a combination of quantum chemical methods and crystallographic refinement in what is known as quantum refinement. This method has been successfully used within X-ray crystallography to resolve issues with poor density around metal atoms. The N-terminal protonation state suggested from the neutron diffraction data of ref. 28 is also not entirely unambiguous. Interpreting nuclear scattering length density maps from refinements against only neutron data have proven quite difficult for several reasons. First, neutron data usually has a lower resolution than X-ray data, since including hydrogen atoms introduces many more parameters, so the data-to-parameter ratio is low. Second, if hydrogen and deuterium atoms are close to each other, the deuterium and hydrogen signals can cancel out, as they have opposite signs. Similar to X-ray crystallography, the local structure around critical sites can be improved by quantum refinement. Quantum refinement for neutron crystallography is still in its infancy and only a proof-of-principle study has been undertaken so far. Here we employ for the first time quantum mechanics to facilitate both the refinement of X-ray and neutron diffraction. The result is the most probable interpretation of the LPMO copper–oxygen species, and from these results we suggest a new mechanism for generation of hydrogen peroxide.

2 Methods

Joint X-ray–neutron refinement

The low data-to-parameter ratio in neutron crystallography can often be significantly improved by refining against both X-ray and neutron data at the same time (joint X-ray–neutron refinement). Unfortunately, such a procedure can be complicated when there are differences between the reported neutron and X-ray structures as is the case in the structure reported by Back et al., where the backbones of the two protein models do not perfectly superpose. This is presumably why no joint refinement was performed for the deposited neutron structure. However, the space groups of the X-ray and neutron crystals are identical and the unit cell parameters are very similar, suggesting that the two datasets are suited for joint X-ray–neutron refinement. We therefore decided to redo the refinement as a joint refinement. X-ray structure factors, coordinates, occupation numbers, B factors, as well as neutron structure factors were obtained from the Brookhaven Protein Data Bank (PDB). Joint X-ray–neutron refinement was performed in...
using the phenix.refinement module and the coordinates from the X-ray model. Deuterium atoms were added at exchangeable sites and H atoms at the rest of the sites with the phenix.ready_set module. Deuterium atoms at the N-terminus were added manually.

Quantum refinement

Crystallographic refinement is a global pseudo-energy minimization using an energy function of the form

$$E_{\text{cyst}} = w_X E_{\text{Xray}} + E_{\text{MM}}$$  

(3)

or

$$E_{\text{cyst}} = w_X E_{\text{Xray}} + w_N E_{\text{neutron}} + E_{\text{MM}}$$  

(4)

for pure X-ray refinement and joint X-ray and neutron refinement, respectively. Here $E_{\text{MM}}$ is an MM (or another empirical or statistical) energy function of the protein model, whereas $E_{\text{Xray}}$ and $E_{\text{neutron}}$ are target functions describing how close the current model reproduces the experimental X-ray and neutron data, respectively. We have used a joint maximum-likelihood target function. The two weight factors $w_X$ and $w_N$ are needed because $E_{\text{MM}}$ typically is in energy units, whereas the other two terms are unit-less pseudo-energies.

For quantum refinement, $E_{\text{QM/MM}}$ replaces $E_{\text{MM}}$, meaning that QM replaces MM for a restricted, but interesting, part of the protein (system 1). In this work we employ

$$E_{\text{QM/MM}} = E_{\text{QM}} + E_{\text{MM}} - E_{\text{MM}}$$  

(5)

where the $E_{\text{QM}}$ is the energy of the system described by QM, which we here denote system 1. $E_{\text{MM}}$ is the MM energy of the same system. The remaining system is denoted system 2. Technical parameters associated with the QM/MM calculations are described in the next subsection.

The quantum refinement method is implemented in the ComQum-X software, which combines Turbomole with the Crystallography and NMR system (CNS), replacing $E_{\text{MM}}$ in eqn (1) with $E_{\text{QM/MM}}$ in eqn (2) to give our combined energy function

$$E_{\text{ComQum-X}} = w_X E_{\text{Xray}} + w_N E_{\text{neutron}} + E_{\text{QM}} + w_{\text{MM}} (E_{\text{MM}} - E_{\text{MM}})$$  

(6)

In eqn (6), an additional scaling factor, $w_{\text{MM}}$, was introduced because the statistics-based force field in CNS typically gives energies that are ~3 times larger than an energy-based force field (i.e. $w_{\text{MM}}$ is set to 1/3).

The quantum-refinement calculations were performed starting from the joint X-ray–neutron refined coordinates, $B$ factors and occupancies. However, CNS does not support anisotropic $B$ factors, so those were ignored. The quantum system used for all refinements is shown in Fig. 1. It consisted of the copper ion, the imidazole ring of His-109, the phenyl ring of Phe-164, the full His-32 residue, which coordinates to Cu through the terminal amino group and a crystal water molecule that coordinates to the copper atom. For the investigation of the protonation of the N-terminus, we ran quantum refinement calculations including the neutron data. We can include neutron data in eqn (6) simply by adding a $E_{\text{neutron}}$ term and the corresponding weight factor, the equation thus becoming

$$E_{\text{ComQum-U}} = w_X E_{\text{Xray}} + w_N E_{\text{neutron}} + E_{\text{QM}} + w_{\text{MM}} (E_{\text{MM}} - E_{\text{MM}})$$  

(7)

This has recently been implemented in the ComQum-U software, combining Turbomole with the joint X-ray–neutron refinement version of CNS, nCNS. We assumed that the total charge of the quantum system was 0, i.e. corresponding to a Cu(I)-peroxide system (CuO2) as in ref. 27. In a separate set of calculations, we exclusively employed quantum refinement of the X-ray diffraction data in ref. 28, and in this case we investigated both CuO2 and [CuO2]− forms. The full protein was used in all calculations, including all crystal water molecules. In each cycle of the geometry optimization, the surrounding protein was allowed to relax by one cycle of crystallographic minimization and one cycle of individual $B$ factor refinement. However, the new coordinates and $B$ factors were accepted only if the $R$ factor was reduced. After quantum-refinement, anisotropic $B$ factor and occupancy refinement was performed using phenix.refine. All refinements were done at the TPSS/def2-SV(P) level of theory. Separate refinements were run with the quantum system in subunit A and subunit B respectively.

Model quality metrics

The quality of the models was compared using the real-space difference density Z-score (RSZD) per atom, calculated by EDSTATS in the CCP4 software suite, which measures the local accuracy of the model. The maximum absolute value of RSZD is typically <3.0 for a good model. RSZD− shows the maximum negative RSZD value, whereas RSZD+ shows the maximum positive RSZD value.

QM/MM calculations

QM/MM calculations were performed with the ComQum interface, which combines the QM program Turbomole and the MM program AMBER. Here we employed Turbomole 7.1 (ref. 46) and AMBER 14, respectively. When there is a bond between systems 1 and 2 (a junction), the hydrogen link-atom approach is employed: the QM region is capped with hydrogen atoms (hydrogen link atoms), the positions of which are linearly related to those of the corresponding carbon atoms (carbon link atoms) in the full system.

The chosen QM method was exclusively density functional theory using the dispersion-corrected TPSS-D3 functional. We performed two sets of QM/MM calculations. The first was carried out as comparison to the quantum refinement. Here the QM systems (system 1) were designed to be identical (i.e. the one in Fig. 1), and we employed the def2-SV(P) basis set. We again employed total charges of the quantum systems of either +1 ([CuO2]−) or 0 (CuO2). In the former model, this involves Cu(II) coupled to a superoxide anion, O2−, whereas the latter model involves either Cu(i) and O2− or Cu(i) and peroxide O2− (a detailed analysis of the wave function is required to determine which of the two formulations is the most appropriate). Note
that while CuO$_2$ was exclusively considered in the doublet spin-state, the [CuO$_2$]$^+$ moiety can, from previous experience,\textsuperscript{17,21,52} result in a triplet or an (open-shell) singlet state that are nearly degenerate in energy, and we therefore considered both cases. Separate calculations were performed with the quantum system in subunit A or B, respectively.

In a second set of calculations, we studied the mechanism of H$_2$O$_2$ generation. Here we extended the QM system with the side chain of the Glu-201 residue, which is a possible proton donor in the vicinity of the active site. In a some of the calculations, six additional water molecules close to the active site were added to the quantum system. These studies were performed only on subunit B, as this showed less variation in the binding of the oxygen species. The studies of proton donation and dissociation of the resulting species exclusively employed the def2-TZVPD basis sets.\textsuperscript{51,53}

A more detailed account of the computational and protein setup (including the alternate configurations and protonation states of individual amino acids) are provided in the ESI.\textsuperscript{†}

3 Results and discussion

In this paper, we discuss two important aspects of the LPMO copper center. First, we re-investigate the nature of the Cu–oxygen species observed in crystal structures, both in terms of oxidation state and protonation state of the terminal amino group. This is done by carrying out a joint refinement of the X-ray and neutron diffraction data reported in ref. 28. Moreover, we employ quantum refinement against both neutron and X-ray data. Second, we suggest a new mechanism of H$_2$O$_2$ formation, based on QM/MM calculations starting from the oxygenated species we deem most probable from the quantum re

Joint X-ray–neutron refinement

The joint X-ray–neutron crystallographic refinement employed the X-ray coordinates (entry 5VG0) as starting coordinates and we first considered the N-terminus in both subunits as ND$_2$. The model was refined to X-ray $R_{\text{free}}$ and $R_{\text{work}}$ values of 14.4% and 13.9% respectively, and neutron $R_{\text{free}}$ and $R_{\text{work}}$ values of 24.8% and 19.2%. These values are comparable to those for the models deposited in the PDB (X-ray $R_{\text{free}}$ of 12.8% and neutron $R_{\text{free}}$ of 26.5%), showing an improvement of the model quality for the neutron structure but a slightly worse X-ray model. This is generally expected from a joint refinement.\textsuperscript{39}

The structures of the active site resulting from the joint refinement are shown in Fig. 2A and B for subunits A and B, respectively, together with the nuclear density. The coordination environment is the same as in the original crystal structure for both subunits.\textsuperscript{39} Selected structural parameters for the active site are shown in Table 1. The coordination of the dioxygen species is side-on in subunit A (Cu–O distances of 2.18 and 2.48 Å) and end-on in subunit B (Cu–O distances of 1.84 and 2.91 Å), as in the original structure. However, in monomer A, the Cu–O coordination distances are $\approx$0.3 Å longer than in the original X-ray model (Table 1). This difference can be attributed to the weak nuclear density of the O$_2$ unit from the neutron data, especially in subunit A (cf. Fig. 2). As oxygen atoms contribute less to the neutron data and our model is refined against both X-ray and neutron data, it is expected that the O$_2$ entity will have a different position in joint refinement compared to the neutron-only crystallographic refinement done in ref. 28.

One of the deuterium atoms (D1) in the terminal amino group forms a hydrogen bond to the backbone carbonyl of Ala-107 with an O–D distance of 2.41–2.60 Å. As can be seen in Fig. 2A and B, no nuclear difference-density peaks are observed on the N-terminus at a 3.0$\sigma$ level, neither on subunit A nor on subunit B, indicating that the model with two deuterium atoms is consistent with the crystallographic data. Similarly, no difference density peaks can be seen in the X-ray $mF_o - DF_c$ maps (Fig. S2$^\dagger$). This can also be seen in the RSZD $- mF_o$–$DF_c$ maps of the N-terminus and deuterium atoms, which are $<$0.1 in both subunits and for both types of maps (X-ray and neutron).

For comparison, we performed also joint refinement of the protein with only one deuterium atom on the N-terminus (i.e. a deprotonated amino terminal), both in subunit A and in subunit B. The resulting structures of the active site are shown in Fig. 2C and D, and selected bond distances are given in Table 1. The structures are nearly identical to those in the model with two deuterium atoms, the only atom showing a significant movement is D1, which moves to an intermediate position compared to the two atoms in the previous model (Fig. 2A and B). The remaining deuterium atom still forms a hydrogen bond with the backbone carbonyl of Ala-107. The neutron $mF_o - DF_c$ map of the active site of subunit B, which was previously modeled as ND$_2$, shows a positive difference density peak at a 3.0–3.2$\sigma$ level, indicating that the ND$_2$ state is a better interpretation of the crystal structure, based on the joint refinement. These findings are consistent with the previous joint X-ray–neutron structure of an AA9 LPMO by O’Dell et al. (Nc PMO-2,
X-ray A 2.00 2.01 2.12 1.84 2.14 1.48
X-ray B 2.01 2.01 2.14 1.83 2.69 1.46
Neutron ND2 A 2.55 2.56 3.40 c c
Neutron ND2 B 2.48 2.49 2.94 1.84 2.49 3.37 b
JXN-R ND2 A 1.94 1.97 2.12 2.18 2.48 1.45 2.60
JXN-R ND2 B 1.92 1.95 2.11 1.84 2.91 1.45 2.41
JXN-R ND2 A 1.95 1.97 2.10 2.22 2.52 1.45 2.55
JXN-R ND2 B 1.96 1.99 2.10 1.98 2.90 1.45 2.38
CQU ND2 A 1.90 1.88 2.19 2.21 2.72 1.35 2.50 0.01 0.48 0.53
CQU ND2 B 1.98 1.98 2.18 2.01 2.75 1.38 2.33 0.11 0.53 0.35
CQU ND2 A 1.94 1.90 2.17 2.23 2.80 1.37 2.49 0.18 0.40 0.42
CQU ND2 B 2.01 1.98 2.15 2.00 2.74 1.39 2.32 0.20 0.46 0.35

* The first column shows the type of calculation (X-ray/neutron for the deposited structures 5VG0 and 5VG1 respectively, JXN-R for joint refinement, CQU for quantum refinement), the second column shows the protonation state of the N-terminus. b No deuterium atoms were reported in the deposited X-ray and neutron structure. c Subunit A in the deposited neutron structure does not contain any oxygen species bound to copper.

PDB entry 5TK1,27 which shows no indication of a deprotonated N-terminus.

### Joint X-ray-neutron quantum refinement

Next, we investigated whether the ND2 state of the N-terminus agrees both with the crystallographic data and with quantum chemical calculations by quantum refinement. In this approach, we use the X-ray and neutron crystal structure factors and replace the MM potential used in the joint X-ray–neutron refinement by an QM/MM potential. The atoms modeled by a QM potential are shown in Fig. 1.

The quantum-refined structures are shown in Fig. 3A and B, and selected bond lengths are given in Table 1. The structures are generally in close agreement with the structures obtained by traditional joint X-ray–neutron refinement in both subunits. The D1 atom still forms a hydrogen bond with the backbone carbonyl of Ala-107. The quantum refinement slightly improves the hydrogen bonding geometry, the O–D1 bond length being 0.1 Å shorter in the structure obtained with traditional joint refinement, 2.50 Å in subunit A and 2.33 Å in subunit B.

Although the density maps calculated from nCNS structures are noisier, the nuclear difference density maps calculated from the quantum-refined structure show no negative density around the D2 atom in either of the subunits. This is also reflected in the RSZD− scores of the N-terminus and the two deuterium atoms, of 1.3–2.2, 0.1–0.5 and 0.8–1.3 respectively, which are all lower than 3.0, albeit higher than in the traditional joint X-ray–neutron refinement. We also performed the quantum refinement with the N-terminus in the ND− state; the resulting structures are shown in Fig. 3C and D. The structures with ND− show no change in the geometry of the active site in either subunit, but a positive nuclear difference density at the 2.4–3.0σ level is observed at the position of the D2 atom in both subunits. Thus, the ND2 state of the N-terminus agrees better than the ND− state both with the crystallographic data and with quantum-chemical calculations.

Interestingly, the geometry of the oxygen species changes somewhat when a QM potential is introduced. In subunit A, the position of the proximal oxygen atom is identical to the one from joint refinement, whereas the distal oxygen atom moves away from the copper atom, resulting in an end-on coordination with Cu−Odist = 2.72 Å (in contrast to the side-on coordination both in the joint refinement, as well as in the original crystal structure). In subunit B, the dioxygen geometry changes less. The coordination stays end-on, and the Cu−Oprox distance is shorter by 0.2 Å.

### Nature and coordination of the oxygen species

While our joint crystallographic and quantum refinement calculations both agree on the protonation state of the N-terminus, the two methods give slightly diverging results concerning the coordination of the dioxygen species. We therefore decided to investigate this issue further, focusing on the structures with two hydrogen atoms on the N-terminus. To this end, we performed QM/MM structure optimisations quantum-
refinement calculations, as well as vacuum QM calculations (to calculate strain energies), testing both the CuO₂⁻ and [CuO₂⁺] forms. Considering the low nuclear density for the oxygen atoms, we chose to perform quantum refinement only against the X-ray data.

Key bonding parameters for Cu and the oxygen species are listed in Table 2, whereas strain energies and RSZD values are summarized in Table 3. The different QM/MM and quantum-refined structures are compared in Fig. 4. A more detailed discussion of the QM/MM and quantum refinement structures is provided in the ESL† Here we only summarise the main findings.

From the vacuum calculations (cf. Table 3) we see that the quantum-refined CuO₂⁻ system is 30 kJ mol⁻¹ more strained than the [CuO₂⁺] system (for both subunits). Both QM/MM and quantum refinement show that in subunit B, the geometry of the oxygen species in the crystal structure is closer to that of a superoxide ion, as the [CuO₂⁺] system has a lower RSZD score and is less strained than the corresponding [CuO₂⁻] system. In contrast, in subunit A none of the calculations yield geometries that are close to the side-on coordination in the crystal structure and all optimised geometries fit to the data rather poorly. The observed coordination is thus either an artifact or resulting from a superposition of different configurations of the Cu–O₂ bond. This would also explain why different Cu–O₂ binding modes are obtained for subunits A and B, and why the density of the oxygen species is so weak in the crystallographic data (see further discussion in the ESI†).

To investigate this option, we carried out a potential energy scan of the Cu–O–O angle of the [CuO₂⁻] system in vacuum. This investigation reveals that the energy difference between various binding modes of the superoxide is only 15 kJ mol⁻¹ (Table S1†). In subunit A, we also obtain two different end-on conformations of [CuO₂⁻] (for the peroxide, we always obtained the same conformer). The two conformers are nearly degenerate (within 8 kJ mol⁻¹). Therefore, these two conformers may co-exist in the crystal structure (cf. Fig. S4†).

**Formation of H₂O₂**

Knowing the protonation state of the N-terminus and the nature of the oxygen species, we proceeded to study the mechanism of H₂O₂ formation.

### Table 2

<table>
<thead>
<tr>
<th>Structure</th>
<th>Subunit</th>
<th>State</th>
<th>Cu–Oprog</th>
<th>Cu–Odiss</th>
<th>O–O</th>
<th>Cu–Oprog–Odiss</th>
<th>q⁻⁻β Cu</th>
<th>q⁻⁻β Oprog</th>
<th>q⁻⁻β Odiss</th>
<th>Figure</th>
</tr>
</thead>
<tbody>
<tr>
<td>5VG0</td>
<td>A</td>
<td></td>
<td>1.84</td>
<td>2.14</td>
<td>1.48</td>
<td>79.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>5VG0</td>
<td>B</td>
<td></td>
<td>1.83</td>
<td>2.69</td>
<td>1.46</td>
<td>109.5</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>QM/MM</td>
<td>A</td>
<td>[CuO₂⁻]</td>
<td>1.99</td>
<td>2.72</td>
<td>1.40</td>
<td>105.8</td>
<td>0.30</td>
<td>0.36</td>
<td>0.24</td>
<td>Fig. 4A</td>
</tr>
<tr>
<td>QM/MM conf 1</td>
<td>A</td>
<td>[CuO₂⁻]</td>
<td>2.08</td>
<td>2.82</td>
<td>1.28</td>
<td>112.0</td>
<td>0.38</td>
<td>0.36</td>
<td>0.16</td>
<td>Fig. 4A</td>
</tr>
<tr>
<td>QM/MM conf 2</td>
<td>A</td>
<td>[CuO₂⁻]</td>
<td>2.02</td>
<td>2.79</td>
<td>1.29</td>
<td>112.9</td>
<td>0.38</td>
<td>0.33</td>
<td>0.20</td>
<td>Fig. 4A</td>
</tr>
<tr>
<td>QM/MM</td>
<td>B</td>
<td>[CuO₂⁻]</td>
<td>2.01</td>
<td>2.15</td>
<td>1.40</td>
<td>75.6</td>
<td>0.26</td>
<td>0.44</td>
<td>0.32</td>
<td>Fig. 4B</td>
</tr>
<tr>
<td>ComQum-X</td>
<td>A</td>
<td>[CuO₂⁻]</td>
<td>2.06</td>
<td>2.92</td>
<td>1.27</td>
<td>120.2</td>
<td>0.34</td>
<td>0.34</td>
<td>0.15</td>
<td>Fig. 4B</td>
</tr>
<tr>
<td>ComQum-X</td>
<td>A</td>
<td>[CuO₂⁻]</td>
<td>2.21</td>
<td>2.68</td>
<td>1.34</td>
<td>94.8</td>
<td>0.24</td>
<td>0.43</td>
<td>0.37</td>
<td>Fig. 4C</td>
</tr>
<tr>
<td>ComQum-X</td>
<td>B</td>
<td>[CuO₂⁻]</td>
<td>2.31</td>
<td>2.96</td>
<td>1.25</td>
<td>108.6</td>
<td>0.32</td>
<td>0.37</td>
<td>0.21</td>
<td>Fig. 4C</td>
</tr>
<tr>
<td>ComQum-X</td>
<td>B</td>
<td>[CuO₂⁻]</td>
<td>2.20</td>
<td>2.54</td>
<td>1.34</td>
<td>87.8</td>
<td>0.22</td>
<td>0.44</td>
<td>0.37</td>
<td>Fig. 4D</td>
</tr>
<tr>
<td>Vacuum</td>
<td>A</td>
<td>[CuO₂⁻]</td>
<td>2.27</td>
<td>3.00</td>
<td>1.25</td>
<td>113.6</td>
<td>0.31</td>
<td>0.35</td>
<td>0.22</td>
<td>Fig. 4D</td>
</tr>
<tr>
<td>Vacuum</td>
<td>B</td>
<td>[CuO₂⁻]</td>
<td>2.03</td>
<td>2.82</td>
<td>1.33</td>
<td>112.5</td>
<td>0.04</td>
<td>0.44</td>
<td>0.53</td>
<td>Fig. 4S</td>
</tr>
<tr>
<td>Vacuum</td>
<td>B</td>
<td>[CuO₂⁻]</td>
<td>2.11</td>
<td>2.88</td>
<td>1.26</td>
<td>115.2</td>
<td>0.29</td>
<td>0.37</td>
<td>0.24</td>
<td>Fig. 4S</td>
</tr>
</tbody>
</table>

Fig. 4 Active sites of the original crystal structure (entry 5VG0) (blue), calculated/quantum-refined structures with peroxide oxygen species (red) or superoxide oxygen species (green). (A) – QM/MM structure, subunit A; (B) – QM/MM structure, subunit B; (C) – quantum-refined structure, subunit A; (D) – quantum-refined structure, subunit B.

In a set of initial calculations, we performed QM-cluster calculations of the [CuO₂⁻]⁻ system and the dissociation products, Cu(II) and O₂ or Cu(I) and O₂, respectively. These calculations were carried out at the TPSS/def2-TZVPD level of theory in either gas phase or with a COSMO solvation model (with ε = 80). The QM-cluster calculations allow us to obtain zero-point energies (ZPE) and entropies. We have collected the results in Table 4. For both dissociation reactions, the thermochemical corrections...
(ZPE and entropy) favour dissociation, but their contributions are roughly equal (around 45–55 kJ mol\(^{-1}\)). The lack of solvation makes dissociation of the oxygen species as superoxide highly disfavoured in gas phase, while dissociation of \(O_2\) is favourable, mainly owing to the entropy contribution. As expected, solvation has a large stabilizing effect on dissociation of \(O_2\), but the dissociation is still unfavourable by \(\sim 80\) kJ mol\(^{-1}\), whereas dissociation of dioxygen from Cu(i) is favourable in COSMO solvent with a free energy of \(\sim 20\) kJ mol\(^{-1}\).

The above result indicates that dissociation of \(O_2^–\) is not feasible. In a series of more elaborate QM/MM calculations, we also attempted to dissociate \(O_2\), following the recipe of ref. 26. To this end, we performed several linear transit QM/MM calculations, displacing the O\(_{\text{prox}}\) atom of the [CuO\(_2\)]\(^+\) species from the copper atom (up to 3.4 Å), optimising all atoms in the QM system. Similar to the QM-cluster calculations, these calculations were carried out with the def2-TZVPD basis set. During the dissociation, the triplet is always more stable than the (open-shell) singlet. In the equilibrium structure, the singlet-triplet splitting is 15 kJ mol\(^{-1}\) and this magnitude is kept at larger distances, although at larger distances the spin density is concentrated on the oxygen atoms, suggesting a Cu(i) state and triplet molecular oxygen.

As also noted in ref. 26, the use of a triple-zeta basis set has some influence and results in an equilibrium structure where the axial water molecule no longer binds to the copper (the Cu–OH distance is 2.83 Å), in contrast to the calculations in the previous section, which were done with the def2-SVP(P) basis set. At a Cu–O\(_{\text{prox}}\) distance of 3.0 Å the water molecule binds back to the copper atom in the axial position, as shown in Fig. 57.† This is again consistent with the findings of Kjaergaard et al. 36. However, unlike ref. 26, we did not observe any transfer of a proton from the water ligand to the oxygen ligand, even if we changed the orientation of the water molecule to be optimal for proton transfer. Moreover, if the restraints on the Cu–O\(_{\text{prox}}\) distance were removed, the O\(_2\) entity binds back to the copper atom, forming the [CuO\(_2\)]\(^+\) species again. This is perhaps understandable given the results in vacuum where dissociation is strongly disfavoured, even with inclusion of solvent (cf. Table 3).

In an attempt to keep the oxygen fully dissociated, we included six additional water molecules in the QM system and reoptimised from a structure with the O\(_2\) already dissociated, at a Cu–O\(_{\text{prox}}\) distance of 4.6 Å, without any restraints (Fig. S8†). However, the optimisation also led to a structure with the superoxide bound to the copper atom, suggesting that the release of the unprotonated superoxide does not occur.

Prompted by our previous study on an A99 LPMO\(^{38}\) we next investigated a mechanism in which [CuO\(_2\)]\(^+\) is successively protonated and reduced. We investigated the following six reactions:

\[
\begin{align*}
[CuO_2]^+ + H^+ &\rightarrow [Cu(O_2H)]^2+ + HO_2 \rightarrow Cu^2+ \quad (8) \\
[Cu(O_2H)]^2+ + H^+ &\rightarrow [Cu(O_2H_2)]^3+ \\
[Cu(O_2H_2)]^{1+} + e^- &\rightarrow [Cu(O_2H_2)]^2+ + HO_2 \rightarrow Cu^2+ \\
[Cu(O_2H)^+ + H^+ &\rightarrow [Cu(O_2H_2)]^{2+} \\
[Cu(O_2H_2)]^{2+} + e^- &\rightarrow [Cu(O_2H_2)^+] + HO_2 \rightarrow Cu^+ \\
[Cu(O_2H)^{2+} + e^- &\rightarrow [Cu(O_2H)^+] + HO_2 \rightarrow Cu^+ \\
\end{align*}
\]

where dissociation of the dioxygen species, HO\(_2\) or H\(_2\)O\(_2\), may occur from either Cu(ii) in [Cu(O\(_2\)H)]\(^{2+}\) or [Cu(O\(_2\)H\(_2\))]\(^{2+}\) or from Cu(i) in [Cu(O\(_2\)H)]\(^+\) or [Cu(O\(_2\)H\(_2\))]\(^+\).

The first protonation of the superoxide species (eqn (8)) was started from the previously optimised equilibrium structure of [CuO\(_2\)]\(^+\), with a proton added to the Glu-201 residue (cf. Fig. 5A). In the triplet spin state, the proton from Glu-201 does not transfer to \(O_2\) and all attempts to start from the product (with Cu–O\(_2\)) leads back to the reactant. In contrast, for the open-shell singlet the Glu-201 proton transferred spontaneously to the distal oxygen of the superoxide, resulting in a Cu–O\(_H\) species (Fig. 5B), with a Cu–O\(_{\text{prox}}\) distance of 1.88 Å, 0.05 Å shorter than with the unprotonated \(O_2\). The reaction energy of this transfer of the proton from Glu-201 to the superoxide moiety is \(-8\) kJ mol\(^{-1}\).

Starting from the [Cu(O\(_2\)H)]\(^{2+}\) species, we performed a second protonation, resulting in the [Cu(O\(_2\)H\(_2\))]\(^{3+}\) species (eqn (9)). The proton was moved in 0.1 Å steps from the OE1 atom of the glutamate to the proximal oxygen atom of the superoxide. The reaction energy for this proton transfer was 36 kJ mol\(^{-1}\), with a barrier of 41 kJ mol\(^{-1}\). Interestingly, this second protonation occurs with an un- and re-binding of HO\(_2\), the superoxide species moving as far as 2.4 Å away from the copper ion in the transition state, as shown in Fig. 6A. However, if we attempt the second protonation after first reducing the copper center to Cu(i) (first part of eqn (11) and (13)), the proton transfers practically at the same place as in the first protonation (Fig. 5B). Moreover, if the restraints on the Cu–O\(_{\text{prox}}\) distance were removed, the O\(_2\) entity binds back to the copper atom, forming the [CuO\(_2\)]\(^+\) species again. This is perhaps understandable given the results in vacuum where dissociation is strongly disfavoured, even with inclusion of solvent (cf. Table 3).

In an attempt to keep the oxygen fully dissociated, we included six additional water molecules in the QM system and reoptimised from a structure with the O\(_2\) already dissociated, at a Cu–O\(_{\text{prox}}\) distance of 4.6 Å, without any restraints (Fig. S8A†). However, the optimisation also led to a structure with the superoxide bound to the copper atom, suggesting that the release of the unprotonated superoxide does not occur.

Prompted by our previous study on an A99 LPMO\(^{38}\) we next investigated a mechanism in which [CuO\(_2\)]\(^+\) is successively protonated and reduced. We investigated the following six reactions:

\[
\begin{align*}
[CuO_2]^+ + H^+ &\rightarrow [Cu(O_2H)]^2+ + HO_2 \rightarrow Cu^2+ \quad (8) \\
[Cu(O_2H)]^2+ + H^+ &\rightarrow [Cu(O_2H_2)]^3+ \\
[Cu(O_2H_2)]^{1+} + e^- &\rightarrow [Cu(O_2H_2)]^2+ + HO_2 \rightarrow Cu^2+ \\
[Cu(O_2H)^+ + H^+ &\rightarrow [Cu(O_2H_2)]^{2+} \\
[Cu(O_2H_2)]^{2+} + e^- &\rightarrow [Cu(O_2H_2)^+] + HO_2 \rightarrow Cu^+ \\
[Cu(O_2H)^{2+} + e^- &\rightarrow [Cu(O_2H)^+] + HO_2 \rightarrow Cu^+ \\
\end{align*}
\]

Table 4: Energy components for the dissociation of [CuO\(_2\)]\(^+\) from QM-cluster calculations in gas phase and COSMO solvent, \(\nu = 80\), in kJ mol\(^{-1}\). Note that the frequencies are from gas phase calculations and hence ZPVE and entropy effects are estimated from the gas phase.

<table>
<thead>
<tr>
<th>Reaction</th>
<th>Electronic</th>
<th>Solvation</th>
<th>ZPE</th>
<th>(-T\Delta S)</th>
<th>(\Delta G\text{vac})</th>
<th>(\Delta G\text{COSMO})</th>
</tr>
</thead>
<tbody>
<tr>
<td>[CuO(_2)](^+) → Cu(i) + O(_2)</td>
<td>920.7</td>
<td>-795.1</td>
<td>-3.9</td>
<td>-42.6</td>
<td>874.3</td>
<td>79.2</td>
</tr>
<tr>
<td>[CuO(_2)](^+) → Cu(i) + O(_2)</td>
<td>26.0</td>
<td>9.3</td>
<td>-4.8</td>
<td>-50.9</td>
<td>-29.6</td>
<td>-20.3</td>
</tr>
</tbody>
</table>
optimisation starting from any of the investigated Cu–O

system) resulted in stable structures with the dioxygen moieties released from the active site, at Cu–O distances of 3.13 and 3.15 Å, respectively (Fig. 7). The spin density on the dissociated HO₂ is ~0.8, showing that the system indeed contains Cu(i) and HO₂, even at larger Cu–HO₂ distances. Interestingly, the resulting Cu(i) coordination sphere is different in the HO₂ calculation compared to the H₂O₂ calculation. In the former, the Cu(i) adopts a trigonal pyramid geometry, with a water molecule in the axial position (Cu–Owat distance of 2.2 Å), whereas in the latter, the copper ion is only coordinated by the three histidine nitrogen atoms, in a trigonal geometry, the Cu–Owat distance becoming 2.26 Å.

For Cu(i), linear-transit calculations between the equilibrium structures and the structures with the dioxygen species dissociated show that dissociation is much easier for H₂O₂ than for HO₂. The dissociation of HO₂ has a barrier of 60 kJ mol⁻¹, whereas H₂O₂ dissociates from Cu(i) with a barrier of only 4 kJ mol⁻¹. Moreover, the reaction energy for the dissociation of HO₂ is higher than that for the dissociation of H₂O₂, 28 kJ mol⁻¹ compared to 3 kJ mol⁻¹. However, assuming entropy and ZPVE contributions of similar size as for the dissociation of O₂ or O₂⁻, HO₂ dissociation would be roughly thermoneutral.

Therefore, we conclude that the formation of the hydrogen peroxide co-substrate in AA10-LPMO is possible only after the reduction of the active site by one electron. Hydrogen peroxide could be formed from the release and disproportionation of HO₂, but the dissociation energy of HO₂ is higher than that of H₂O₂. Since protonation of HO₂ (after reduction of the active site) is practically spontaneous, and dissociation of H₂O₂ has

Fig. 5 Protonation of the superoxide moiety. (A) – reactant, proton on Glu-201 (triplet); (B) – product with the superoxide protonated (open-shell singlet).

Fig. 6 Second protonation of the superoxide moiety starting from (A) – Cu(i)–HO₂ or (B) – Cu(i)–O₂H. Relative energies of the reactants, products and transition state (for Cu(i)–O₂H) are depicted above each structure.
a very low activation energy this seems to be the preferred pathway. Our favoured overall pathway involves Cu(i) and consumes two protons and two electrons leading to H₂O₂ and regeneration of Cu(i), corresponding to eqn (2), first parts of eqn (8) and (13) and eqn (11) and a total catalytic reaction

\[
O_2 + 2H^+ + 2e^- \xrightarrow{\text{LPMO-Cu(I)}} H_2O_2
\]  

(14)

In all cases the proton donor is Glu-201. The steps are either spontaneous or involve barriers lower than 10 kJ mol⁻¹. We note that while this pathway gives the lowest activation energies, the pathways involving eqn (9) and last part of 13 have barriers of 41 and 60 kJ mol⁻¹ (where the latter should be considered an upper limit as it does not include thermochemical corrections). These are both feasible, considering that the reported lower-limit rate constant is 0.15 s⁻¹, corresponding to an activation energy of ~80 kJ mol⁻¹, as calculated from transition state theory. In summary, we cannot find support for dissociation of neither O₂, HO₂ nor H₂O₂ from Cu(n). However, alternate protonations and reductions starting from [CuO₂]⁺ with Glu-201 as proton donor, followed by dissociation of either HO₂ or (more likely) H₂O₂ from Cu(i) are feasible. This is different than the hitherto accepted molecular mechanism involving direct release of O₂. In light of our results, we speculate that the formation of H₂O₂ from release of O₂ only becomes relevant if the proton donors in the second coordination sphere are perturbed, e.g., by mutations. Such a study was recently performed by Span et al. (albeit on a different LPMO, see further below). Their study indeed indicated H₂O₂ formation from O₂ in solution increased by such mutations. While we here have only discussed one specific AA10 LPMO, it is naturally interesting to assess how general our suggested mechanism is for other LPMO classes. The large sequence variation among the LPMOs makes such an assessment difficult. However, a general idea can be obtained by comparing the second coordination sphere for a few different LPMOs. Among the AA10 LPMOs, there is not only variation compared to AA9, but also considerable variation within the group, depending on their substrate specificity (chitin or cellulose). The target LPMO in this study is JdLPMO10A, whose substrate is chitin; cellulose-active AA10 LPMOs also have conserved hydrogen-bonding motifs in the second coordination sphere. For example, the structure with PDB entry 4OY7 (ref. 57) has both an Arg and a Glu residue (Arg-212 and Glu-217) close to the Cu ion. These residues could play the same role as Glu-201 in our calculations. In AA9 LPMOs, a histidine and a glutamine (His-147 and Gln-162 in 5ACP31) replace the Arg and Glu residues from cellulose-active AA10 LPMOs and these two residues are generally conserved among AA9 LPMOs. Proton transfer from His147 to [CuO₂]⁺ has been found to be feasible, making it likely that the current mechanism is also valid for AA9 LPMOs. Yet, further investigations are required before we can truly comment on how general our current mechanism is.

4 Conclusions

In this article, we have studied the mechanism of hydrogen peroxyde generation from a recent AA10 LPMO structure with a bound dioxygen species. Since the reduction level and protonation state of this species is ambiguous, we have first employed quantum refinement of the crystal structure against both X-ray and neutron data to obtain a proper description of the active site. These calculations show conclusively that the N-terminus, which coordinates to the copper atom, is not deprotonated in the crystal structure. Furthermore, both quantum-refinement and QM/MM calculations indicate that the dioxygen species bound in the crystal is a superoxide, O₂⁻, which is in accordance to the study by Kjaergaard et al. on an AA9 LPMO. The mechanism of H₂O₂ generation was studied using the QM/MM approach, at the TPSS/def2-TZVPD level. The results show that unprotonated superoxide does not dissociate after protonation from a water molecule, as seen in the AA9 LPMO studied by Kjaergaard et al. Moreover, we were unable to dissociate any dioxygen species from Cu(n), no matter the protonation and reduction state. However, dissociation of both HO₂ and H₂O₂ is possible from Cu(i); the former can then generate hydrogen peroxyde in bulk through disproportionation. The QM/MM calculated energies show that dissociation of H₂O₂ from Cu(i) is more favourable than that of HO₂. Thus, our proposed mechanism consists of adding two protons and two electrons to dissociate hydrogen peroxyde from Cu(i). The first protonation is spontaneous, whereas the second protonation has a very low activation barrier, provided that the first electron has been added to the system. Then, hydrogen peroxyde dissociates from Cu(i) with a barrier of only 4 kJ mol⁻¹, in contrast to hydrogen superoxide, which needs to overcome a dissociation energy barrier of 60 kJ mol⁻¹. In summary, our calculations support that hydrogen peroxyde needs to be formed at a Cu metal center through successive protonations and reductions before being dissociated into the bulk to act as a cosubstrate for the monooxygenase reaction.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors thank grants from the Swedish research council (project 2014-5540) and from COST through Action CM1305.
Notes and references


