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Recent structural insights into the function of copper nitrite reductases

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Copper nitrite reductases (CuNiR) carry out the first committed step of the denitrification pathway of the global nitrogen cycle, the reduction of nitrite (NO_2^-) to nitric oxide (NO). As such, they are of major agronomic and environmental importance. CuNiRs occur primarily in denitrifying soil bacteria which carry out the overall reduction of nitrate to dinitrogen. In this article, we review the insights gained into copper nitrite reductase (CuNiR) function from three dimensional structures. We particularly focus on developments over the last decade, including insights from serial femtosecond crystallography using X-ray free electron lasers (XFELs) and from the recently discovered 3-domain CuNiRs.

Introduction

Copper nitrite reductases (CuNiRs) are key enzymes in the nitrogen cycle, occurring in a wide range of denitrifying bacteria and fungi.¹ The denitrification pathway in these organisms couples ATP synthesis with the reduction of nitrate (NO_3^-) or nitrite (NO_2^-) to dinitrogen (N_2), *via* nitric oxide (NO) and nitrous oxide (N_2O) intermediates.^{2,3} This pathway is critical in the return of fixed nitrogen to the atmosphere and in controlling the level of biologically available nitrogen in the soil. The extensive use of nitrogen containing fertilisers from the Haber–Bosch process can cause major environmental problems if the denitrification pathway becomes overloaded.² Moreover, the N_2O intermediate in the denitrification pathway is a potent greenhouse gas.⁴ The roles of different metalloenzymes in denitrification and their environmental and agricultural impacts have been recently reviewed.⁵

The first committed step (*i.e.* the first effectively irreversible reaction in the pathway that commits the organism to generate the final product) in denitrification is the reduction of nitrite to NO (eqn (1)) and this is carried out either by Fe-containing cytochrome *cd1* nitrite reductases, penta- or octa-haem nitrite reductases or by CuNiRs, encoded by the *nirk* gene.



Many detailed biochemical, spectroscopic, kinetic and structural studies have investigated the catalytic mechanisms of the CuNiRs and we describe only the key points in this review, focusing on structural insights. We describe the three-dimensional structures of CuNiRs and how these have led to profound insights into enzyme function and catalytic mechanism. We particularly focus

on developments in the last decade including the discovery of extended CuNiRs, the application of serial femtosecond crystallography (SFX) and approaches to characterise the enzyme mechanism within protein crystals.

Overall fold and oligomeric state of CuNiRs

The majority of CuNiRs are trimeric, with each ~ 37 kDa monomer comprising two cupredoxin (β -sandwich) domains, Fig. 1A and B.⁶ Each of these domains has a similar architecture to that of monomeric cupredoxin proteins, such as azurin, pseudoazurin and rusticyanin, that act as electron donor partner proteins to CuNiRs.⁷ The C-terminal region of each cupredoxin-like monomer forms an extended loop that wraps around an adjacent monomer contributing to the stability of the trimer.⁶

These well-characterised two-domain CuNiRs are assigned as classes I and II, depending on the colour arising from their type 1 Cu (T1Cu) centre, with class I being blue and class II being green. Class III has an additional T1Cu domain with a hexameric oligomer and is greenish-blue due to the presence of two Cu centres, one each of class I and II.⁸ This class III CuNiR is an example of a recently discovered group of extended (three-domain) CuNiRs,⁹ that exist with additional fused N- or C-terminal domains containing a T1Cu or cytochrome haem redox centre, respectively (Fig. 1).¹⁰ Regardless of such variations, the core structure of T1Cu and type 2 Cu (T2Cu) sites in the trimeric core is largely conserved. We will first discuss the structure and function of the trimeric CuNiRs before highlighting variations in the more recently characterised extended CuNiRs.

The copper centres of CuNiRs in the resting state

All CuNiRs characterised to date contain an electron-providing T1Cu centre and a catalytic T2Cu centre. At the T1Cu site, the Cu is coordinated by two His, one Cys and an axial Met forming a tetrahedral geometry, Fig. 2.⁶ The structural changes within

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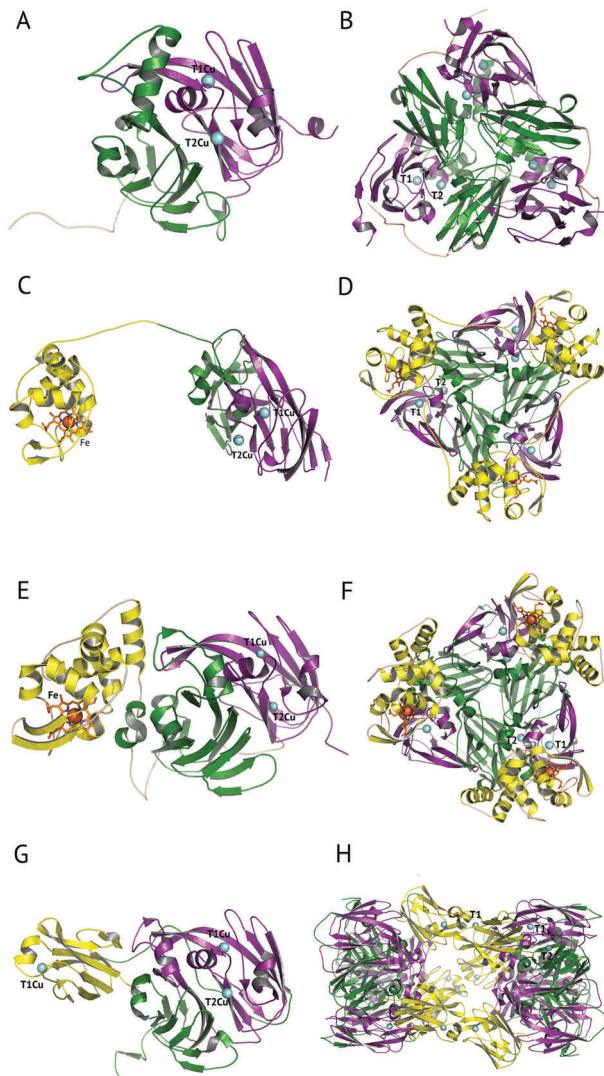


Fig. 1 The domain and oligomeric structures of CuNiRs. In class I and II proteins, two cupredoxin domains are present per monomer, here coloured green and purple. Class III CuNiRs have an additional C- or N-terminal domain (yellow) containing T1Cu or haem. The two-domain proteins form a trimeric oligomer, shown in panels A and B. In the three-domain proteins from *PhNiR* and *RpNiR* the additional C-terminal cytochrome domain, panels C and E, forms trimeric oligomers, panels D and F, respectively. In the three-domain *HdNiR* an additional T1Cu N-terminal domain is present, panel G, and the biological unit appears to be a hexameric oligomer, panel H.

T1Cu sites between the Cu(II) and Cu(I) states are small (consistent with an entactic/rack state), and close to the level of precision at the typical resolutions of crystal structures.

Atomic resolution structures provide more accuracy for bond length parameters but usually at the expense of higher X-ray dose which can readily reduce the T1Cu to the Cu(I) state. EXAFS studies suggest an increase in the Cu–Cys bond length on reduction of the blue copper protein azurin of some 0.07 Å.¹¹

As the main chromophore of CuNiR, the T1Cu site is responsible for the colour of the protein in the cupric state. In general, T1Cu sites in copper proteins can vary widely in colour, including blue, green, red and purple.^{12,13} The relationship

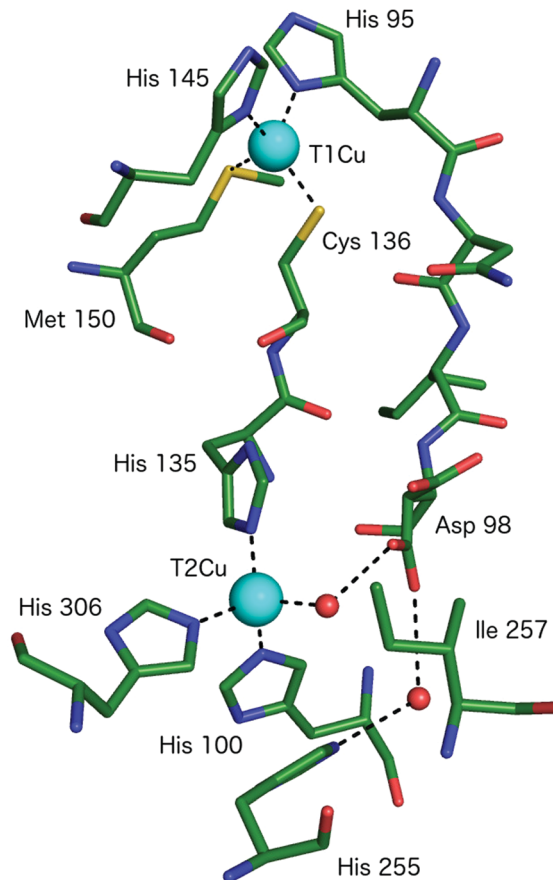


Fig. 2 The T1Cu and T2Cu centres of AcNiR and the protein links between them. The T1Cu atom has (His)₂CysMet ligation and is linked to the catalytic T2Cu centre by a Cys–His bridge. A second link is made between the T1Cu ligand His95 and T2Cu site residue Asp98, proposed to be a sensor loop, enabling intramolecular T1–T2 electron transfer. The T2Cu coordination is completed by three more His residues, one of which originates from an adjacent monomer of the trimer and a water molecule. Image generated from the structure of resting state AcNiR at 0.9 Å resolution (PDB 2BW4).¹⁷

between T1Cu colour and geometry has been systematically studied. CuNiRs from different species appear blue, green or 'bluish-green' despite possessing an identical set of ligand residues. Insights into the relationship between T1Cu structure, colour and redox potential have been obtained by comparison of structures of blue and green CuNiRs and by mutagenesis of first and second coordination shell residues. Blue CuNiRs typically exhibit a long/weak Cu–Met bond ~ 2.9 Å while in green CuNiRs this bond is much shorter.

Work from Solomon and co-workers revealed that in green CuNiRs, two spectral forms were present, with the ratio between these being strongly temperature dependent.¹⁴ At low temperatures, a green form with a strong Met–Cu interaction is dominant, while at higher temperatures, a blue form becomes more populated, with a weaker Cu–Met interaction. The T1Cu geometry, colour and redox potential are profoundly affected by mutation of ligand residues with consequent effects on enzyme activity.

The catalytic T2Cu is separated from the T1Cu by ~ 12.6 Å and the centres are connected by a Cys–His bridge (for rapid



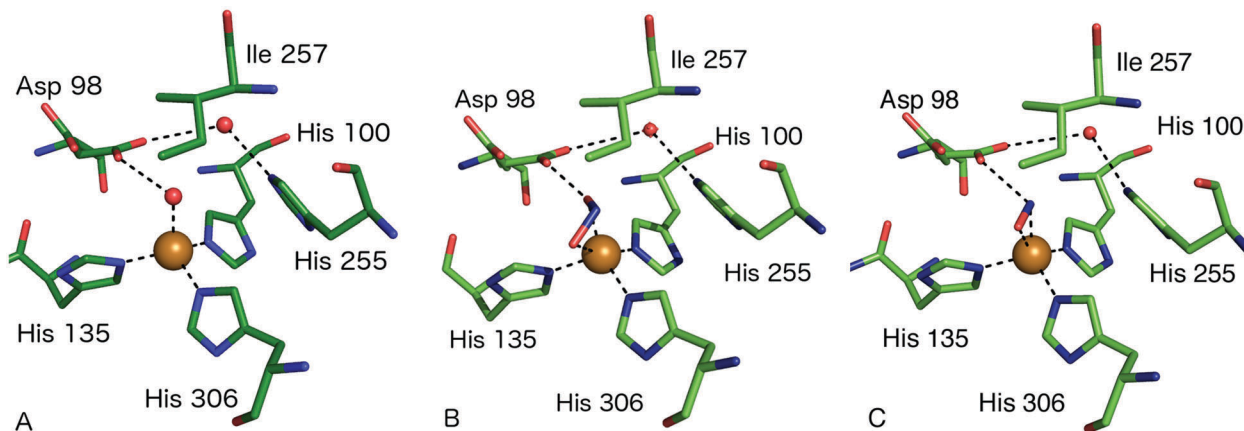


Fig. 3 Type 2 Cu site structures (A) resting state AcNiR with a water molecule bound to the T2Cu (PDB 2BW4); (B) nitrite bound to the T2Cu site in a side-on bidentate O manner (PDB 5AKR); (C) the T2Cu site of AcNiR bound to NO at 1.22 Å resolution (PDB 5I6N). The NO ligand is bound in a side-on manner with near-equidistant Cu–N and Cu–O bonds. Residue Asp98 forms a H-bond to the N atom of NO and is oriented toward the proton delivery channel. Panels A and B are from the structures described in ref. 18 while panel C is from ref. 55.

electron transfer) comprised of the T1Cu ligand Cys and a T2Cu ligating His.⁶ The T2Cu has (His)₃–H₂O ligation in the resting state with a tetrahedral geometry, Fig. 2 and 3A. One of the three His ligands is provided from an adjacent monomer of the CuNiR trimer. Upon reduction (in the absence of substrate), the T2Cu loses the coordinated H₂O resulting in Cu–(His)₃ ligation.¹⁵ The T2Cu centre possesses only a weak UV-visible absorption spectrum that is obscured by the much stronger absorbance of the T1Cu centre.¹⁶ The T2Cu ligand binding pocket contains several conserved residues with proposed functional roles related to substrate access, specificity, activity, electron transfer & proton transfer.¹⁶ The Cu-coordinated water molecule is H-bonded to an aspartic acid residue (Asp_{CAT}) and, *via* a water bridge, to a conserved His (His_{CAT}).⁶ Asp_{CAT} has been proposed to play roles in substrate guidance/stabilisation and to act as a proton donor,¹⁷ while His_{CAT} acts as a second proton donor. Crystal structures show that Asp_{CAT} can adopt two different conformations, named ‘gatekeeper’ – pointed away from T2Cu and ‘proximal’ – oriented towards T2Cu and H-bonding to the bound water molecule in the resting state.¹⁸ Mutation of Asp_{CAT} to asparagine (Asn) in AxNiR eliminated catalytic activity providing strong evidence for its role in proton transfer.¹⁹ The properties of Asp_{CAT} and His_{CAT} may provide the rationale for the observed pH dependence of CuNiR activity. Solomon and coworkers have reported a pK_a of 6.4 for Asp_{CAT} (in AxNiR) when NO₂[–] is bound.²⁰ Finally, an isoleucine (Ile_{CAT}) residue that lies above the T2Cu binding site has been shown by structural studies²¹ to profoundly influence substrate orientation and consequently the catalytic activity.

Enzymatic mechanism of nitrite reduction: structural insights

The consensus mechanism for two domain CuNiRs begins with the resting state enzyme, with a water bound to the T2Cu centre, Fig. 3A. On the basis of X-ray crystallography and EXAFS studies on native and site-selected AxNiR mutants, supported by kinetics measurements, a catalytic mechanism was initially proposed in which NO₂[–] coordinates to the T2Cu(II), displacing

the water molecule in the native enzyme, Fig. 3B. In the absence of nitrite at the T2Cu there is a limited potential for electron transfer (ET) from the T1Cu, but following nitrite binding ET from T1Cu(I) to the T2Cu site occurs,^{15,22,23} which reduces the T2Cu. Two protons required for the catalytic reaction are thought to be provided by residues His₂₅₄ (His_{CAT}) and Asp₉₇ (Asp_{CAT}) and the proton-coupled electron-transfer (PCET) from T1Cu to T2Cu, involving the His_{CAT} and Asp_{CAT} residues, is thought to be the rate-determining step.^{24,25} In this PCET process, the two protons are provided to the bound NO₂[–] along with an ET from the reduced T2Cu, to yield the NO product, Fig. 3C. This product then dissociates from the T2Cu, allowing water to rebind to reform the oxidised resting T2Cu state, Fig. 3A. Structural and kinetic studies on a number of other CuNiRs gave apparently conflicting evidence on whether the intramolecular ET occurred before or after NO₂[–] binding [*e.g.* ref. 15, 26 and 27]. These conflicts have been mostly reconciled by a kinetic scheme for catalytic turnover in which two alternative mechanistic pathways are possible, varying in the sequence of ET and binding of nitrite, depending on the pH and nitrite concentration.²⁸ The catalytic mechanism at the T2Cu site of CuNiRs has also been extensively explored using theoretical methods based on the crystal structures, though as yet no clear consensus has been reached on the precise nature and order of the proposed PCET reaction.^{29–34}

Key features relevant to this mechanism that are informed by structures include the modes of substrate and product binding, the likely electron and proton transfer pathways and steps and the conserved water structure within the active site pocket.

Substrate binding to the T2Cu centre and specificity

The first structure of a nitrite-bound CuNiR complex adduct was determined in 1991.⁶ Since then, structures have been determined of NO₂[–] and NO-bound complexes at a variety of resolutions and pH of native and mutant CuNiRs from different organisms.^{18,19,21–23,27,35–56} Nitrite has typically been observed



to bind either in an asymmetric bidentate binding geometry with its O1 and O2 atoms ligating to T2Cu(II) or in a 'side-on' geometry with similar distances from T2Cu to the three atoms of nitrite. One of the highest resolution structures of a CuNiR-nitrite complex is that of *AcNiR*, to 1.10 Å resolution,¹⁸ where nitrite was observed with bidentate binding to T2Cu and dual conformations of Asp98_{CAT} were present, with the proposal that the gatekeeper conformation of Asp98 (Asp_{CAT}) may be important for guiding NO₂⁻ to its binding site at the T2Cu. The proximal conformation, on the contrary, is significant for proton transfer, bond-cleavage and formation and release of the product, NO. The resting state has water bound to oxidised T2Cu and forms a H-bond to the negatively charged Asp_{CAT} residue. Nitrite is observed to bind to the T2Cu ion *via* a O-coordinate bidentate mode, displacing the bound water molecule and forming H-bonds between the O^{δ2} atom of Asp_{CAT} and the O2 atom of nitrite. Proton transfer occurs from the Asp_{CAT} to the nitrite to form the intermediate T2Cu⁺NOOH on reduction of the T2Cu site. His255 (His_{CAT}) facilitates bond cleavage of the N–ON bond to form the product, NO, by supplying the second proton in the widely-accepted mechanism. The NO product dissociates and leaves the resting water-bound oxidised T2Cu ion. Structures have been determined of ascorbate-reduced and reduced nitrite-soaked *AfNiR* at 2.0 [PDB 1AQ8] and 1.85 Å [PDB 1AS8] resolutions, respectively, with nitrite bound to reduced T2Cu, revealing few structural changes relative to the oxidised form.⁴¹

In the different structures of nitrite complexes, bidentate nitrite has been modelled in both 'top-hat' and 'side-on' positions. XFEL SFX⁵⁴ data have revealed that the 'top-hat' mode is the binding geometry initially present in the T2Cu(II) form of the enzyme and that the side-on may be the result of T2Cu reduction. This highlights that great care must be taken in crystallographic studies of nitrite-bound CuNiRs due to the propensity for X-ray generated photoelectrons to initiate catalysis within the crystal, leading to the conversion of substrate to NO at the T2Cu site. Structures determined from CuNiR nitrite complexes using high X-ray doses in many cases reveal a significant population of the T2Cu-NO product.²⁷ Most recently this phenomenon has been exploited to drive *in crystallo* reactions to generate 'structural movies' of enzyme catalysis in green *AcNiR*^{55,57} and thermostable *Geobacillus thermodenitrificans* (*GtNiR*).⁵⁸

T2Cu site mutations in *AfNiR* significantly altered the binding mode of nitrite²¹ and these were related to changes in enzyme activity. Mutation of this Ile_{CAT} to Val or Leu led to asymmetric bidentate NO₂⁻ binding while mutation to Met, Thr, Gly or Ala resulted in monodentate coordination, allowing a water molecule to also bind to T2Cu in a position close to that usually occupied by the O2 atom of nitrite. Monodentate nitrite binding was associated with greatly reduced catalytic activity, pointing to a key role for Ile_{CAT} in promoting a nitrite binding geometry conducive during catalysis. Crystal structures were determined of the oxidised and chemically-reduced Asp_{CAT} to Asn mutant in *AfNiR*.³⁵ In these structures, nitrite was bound in a side-on manner with an increased level of disorder, ascribed to a shift of Asn relative to Asp producing an unfavourable geometry for H-bonding with the bound protonated nitrite. However, this

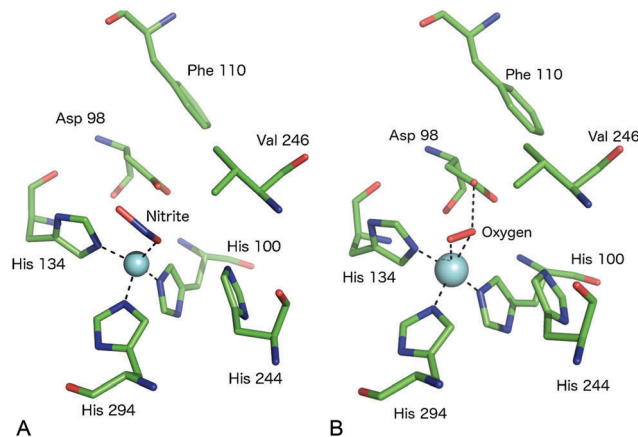


Fig. 4 The T2Cu site environment of *GtNiR*. (A) The nitrite-complex (PDB code 3WKP),⁵² note the monodentate binding of nitrite *via* a single O atom, the substitution of Ile_{CAT} with a Val residue and the steric restriction of the gatekeeper Asp_{CAT} (Asp98) conformation (seen in other CuNiRs) by Phe110, resulting in only the proximal conformation being observed; (B) oxygen binding to *GtNiR* (PDB code 3WNJ).⁶¹ Note that partially occupied water molecules are omitted for clarity. Cu atoms are shown as cyan spheres.

disorder could simply represent a mixture of top-hat and side-on nitrite molecules within the crystal. The His_{CAT} to Asn mutant of *AfNiR* showed no displacement of the water molecule and the nitrite is bound to the T2Cu in a monodentate manner *via* an O atom. Nitrite is oriented away from Ile257 and towards an Ala residue and the pocket H-bonding network is drastically altered in the H255N mutant, indicating an essential role for nitrite binding and function.³⁵

A somewhat different binding mode for nitrite was observed in a structure of the C135A mutant (which eliminates T1Cu binding) of the thermophile *GtNiR*,⁵² a close homologue of *GkNiR*. The 1.90 Å structure showed nitrite bound in a monodentate manner *via* its O2 atom with this atom being positioned to form a H-bond to Asp_{CAT}. His_{CAT} adopted a rotated conformation likely due to the formation of a H-bond to a Gln residue rather than the Thr which interacts with His_{CAT} in other CuNiRs. Interestingly, in this enzyme, the gatekeeper conformation of Asp_{CAT} and the 'vertical' conformer of His_{CAT} are sterically restricted by the unusual position of a Phe residue near to the catalytic pocket which provides additional rigidity to the active site in this enzyme from a thermophile, Fig. 4A. Intriguingly, when the structure of the nitrite complex of *GtNiR* was determined at 320 K, the observed substrate binding mode was side-on, similar to that observed in other CuNiRs,⁵³ a result ascribed to increased dynamic flexibility of the pocket at the higher temperature.

Substrate and copper access to the T2Cu site

The T2Cu site, as mentioned previously, makes up the active substrate-binding site and is coordinated by three His and one H₂O/OH⁻ ligand.³⁸ In the resting state nitrite has access to the T2Cu centre *via* a channel some ~12 Å deep from the protein surface and displaces the H₂O ligand to initiate catalysis. Intriguingly, mutation of the surface-exposed Phe306 to a Cys



residue some 12 Å away from T2Cu in A_xNiR resulted in a change to the second coordination sphere of the Cu with consequent effects on the rate-limiting step of the enzyme reaction.⁵⁹ In this F306C mutant, the substrate access channel is more open and contains an additional water bound to the T2Cu-coordinating water molecule. The specific activity was remarkably increased four-fold while the apparent nitrite affinity and the rate of intramolecular ET were lower than in native A_xNiR. In this mutant, intramolecular ET became the rate-limiting step.

Binding of small molecule ligands to the T2Cu centre

Native CuNiR can bind small molecules including acetate, nitrate, formate, azide and N₂O, of which complex structures have all been determined crystallographically. The binding mode of such ligands⁶⁰ is strongly influenced by the pocket residues Ile_{CAT} and Asp_{CAT} and these, together with the dimensions of the substrate access channel, act to prevent the binding of larger ligands. Monoatomic species such as chloride have also been shown to bind to T2Cu.^{42,54}

A recent study of GtNiR at 1.20 Å resolution⁶¹ revealed electron density consistent with the interpretation that an oxygen ligand was bound at the T2Cu site, in the absence of nitrite or NO, Fig. 4B. CuNiRs have the ability to catalyse the two-electron reduction of O₂ to hydrogen peroxide (H₂O₂) but no structural data for this molecular oxygen ligand has previously been observed. In the GtNiR structure, the O₂ ligand was modelled in a side-on manner with one short (2.1 Å) and one longer (2.4 Å) Cu–O bond length. Oxygen forms H-bonds with Asp_{CAT} and a water molecule and also makes van der Waals contacts with Val_{CAT}. This latter hydrophobic interaction appears to be related to the side on binding mode as previously proposed for NO binding. Oxygen binding has not been observed in other CuNiRs and was proposed to be observable in GtNiR due to the presence of a Val residue in place of Ile_{CAT}, which creates an unusually open substrate binding pocket and/or the slower O₂-reduction rate in the CuNiR from this species. Interestingly, the structure of a CuNiR from *Nitrosomonas europaea*⁵⁰ revealed a much more restricted substrate channel than in other CuNiRs, proposed to be the basis of the unusual O₂ tolerance of this enzyme enabling it to function in an aerobic environment. In addition, a second water channel was observed and assigned as a water exit channel in this enzyme.

It is a notable feature of CuNiRs that they possess a significant superoxide dismutase activity,^{15,62} for example A_xNiR at a remarkable ~56% activity compared to bovine Cu, Zn superoxide dismutase (CuZnSOD), one of the most efficient enzymes known. The structural similarity between the Cu(I) form of the T2Cu site and that of CuZnSOD has been noted from both EXAFS and crystallographic data.¹⁵ Interestingly, CuZnSOD does not display NiR activity.

Identification of intramolecular electron and proton transfer pathways

The CuNiR enzyme reaction requires that one electron and two protons are supplied to the catalytic T2Cu centre and it has been demonstrated that this occurs in a coupled (PCET) manner.²⁵

Spectroscopic and mutational studies showed that electrons are transferred from physiological redox partner proteins such as azurin, pseudoazurin or cytochrome *c*₅₅₁ to reduce the T1Cu centre. Intramolecular ET occurs between the Cu sites to deliver the electron to the T2Cu centre, in the process reoxidising the T1Cu. The pathway for intramolecular ET was identified as a Cys–His bridge between ligating residues of the two Cu centres, providing a facile through-bond pathway.

T1Cu site mutations have been characterised revealing effects upon enzyme colour, redox potential and ET. A M150E mutation in A_fNiR resulted in a protein binding Zn to the T1Cu site³⁹ with a 1000-fold loss of activity, while a C130A mutation in A_xNiR led to a vacant T1Cu site²³ and abolition of activity.

Analysis of high resolution crystal structures led to proposals for proton channels connecting the protein surface (and bulk solvent) with the T2Cu site. A ‘primary’ proton channel in A_xNiR linked the T2Cu site residue Asp_{CAT} with the surface *via* a well-ordered network of water molecules.⁴² A ‘secondary’ channel was later identified in a structure at high pH where residues Asn90, Asn107 and Ala131 together with water molecules also provide a H-bonding network to the surface.⁴³ Supporting evidence was provided from activity and structural work on mutated enzymes. For example, a H254F mutation abolished the ‘primary’ channel yet retained near-native levels of activity. In contrast the N90S variant targeting the secondary channel lowered activity by 70%. These data led to the proposal that this secondary channel was in fact the most important route for protons.^{22,25} Similarly, the ET pathway has been investigated by structural and related studies on mutants. As well as the C130A A_xNiR mutation described above, a H129V mutation in A_xNiR resulted in an unprecedented bis-His T2Cu binding site⁶³ while disrupting the Cys–His bridge leading to a complete loss of enzyme activity.

The structure of the T2Cu-NO species

All crystal structures of CuNiR–NO complexes have revealed a side-on binding mode to T2Cu, Fig. 3C.^{18,45,55} Structures have been determined of NO complexes by several approaches: (i) NO-soaking into A_fNiR crystals,⁴⁵ (ii) endogenously present NO, in crystals grown from protein isolated from *Achromobacter cycloclastes* cells grown under denitrifying conditions¹⁸ and (iii) produced from nitrite *in situ* by X-ray driven catalysis using ‘multiple-structure one crystal’ (MSOX) serial crystallography.⁵⁵ The structures determined by each method consistently show side-on binding with near-equidistant bond lengths to Cu for the N and O atoms and reveal that bound NO interacts with active site residues. Several issues regarding the NO complex remain challenging to address from the electron density alone. The electron density for the N and O atoms is very similar and structures have been determined with different assignments of the density peaks modelled.^{18,45,55} Consideration of bonding further suggests that the NO molecule interacts with Asp_{CAT} which adopts its ‘proximal’ position in the product complex. Similarly, crystal structures are not yet able to give a definitive answer to the identification of the side-on NO complex as Cu(I)–NO• or Cu(II)–NO[−].



An outstanding mystery is the suggestion that solution spectroscopic data supports an end-on mode of NO binding [Cu–N–O] rather than side-on binding. In particular, ENDOR data⁶⁴ of native and the I289V and I257A mutants of *RsNiR* suggests a Cu–N–O angle of 160° in native *RsNiR* with magnetic interaction of O with the C^{δ1} atom of Ile257. Simulations of the complex show 3–10 kcal mol⁻¹ energetic preference for the end-on form.⁶⁵ A subsequent EPR study of model compound analogues found a clear dependence of g_z upon the Cu–N–O angle.⁶⁶ Taken together with the previous EPR data of *RsNiR*,⁶⁴ this work suggested that a strongly bent (~136°) Cu–N–O binding geometry is feasible in CuNiRs. One proposal from computational simulation was that Ile257 dictates the side-on binding mode observed in structures with the proposal that a small structural change when the protein is in solution would allow the end-on binding mode to be sterically feasible.⁶⁷

Surface properties and intermolecular electron transfer

The electrostatic surfaces of CuNiRs are variable, with distinct differences between the blue and green classes of 2 domain enzymes. This, together with comparable analysis of the electrostatic surfaces of the partner proteins⁶⁸ provides a rationale for the observed specificity of CuNiRs from different species for their cognate electron donors.⁶⁸ Azurins reacted rapidly with blue CuNiRs but slowly with green CuNiRs. In contrast, pseudoazurin reacted effectively with either colour of CuNiR. Notably *AfNiR* (green) has a strong surface negative charge complementing the positive charge of the pseudoazurin complementary surface.⁶⁸ Moreover, a common structural feature of CuNiRs is a hydrophobic patch on the enzyme surface near to the location of the T1Cu centre, which has been proposed to form part of the charged protein–protein interface in the ET complex.^{69,70}

Identifying the precise route(s) of intermolecular ET between the redox centre of the electron donor protein (T1Cu or haem Fe) and the CuNiR T1Cu centre is more challenging, in part due to the difficulty of obtaining crystal structures of transient ET complexes. Mutational analysis of putative pathway residues have provided insights. For example, a W138H mutation in *AxNiR* produced a large drop in activity with the physiological redox partner azurin but retained full activity with the artificial electron donor methyl viologen.³⁶ This supported a role for Trp138 in intermolecular ET, consistent with a docking model of the azurin–*AxNiR* complex which positioned Tyr197 close to the azurin T1Cu ligand His117. Tyr197 is close to Trp138 which is adjacent in sequence to a T1Cu His of *AxNiR*.³⁶

Structural characterisation of complexes between CuNiRs and redox partners

Structural characterisation of the ET complexes between CuNiRs and their partner proteins have remained elusive due to the transient nature of the complexes. Computational docking and mutagenesis have attempted to identify interface regions between CuNiRs and their specific azurin, pseudoazurin or *c*-type cytochrome partners *e.g.*⁶⁷ Paramagnetic nuclear magnetic resonance (NMR) data have been used to characterise the interacting region in the complex between *AfNiR* and its partner

pseudoazurin [Paz].⁷¹ In this case, distance restraints derived from NMR data of gadolinium added to engineered surface cysteines were used to restrain computational docking simulations. The resulting structural model positions the T1Cu atoms of Paz and *AfNiR* some 15.5 (±0.5) Å apart.

One crystal structure available for such an ET complex is that between *AxNiR* and a functional partner protein, Cyt_{c551}, at 1.7 Å resolution⁴⁹ (PDB 2ZON). This CuNiR has been shown to accept electrons either from azurin, pseudoazurin or a cytochrome electron donor. In the binary *AxNiR*–Cyt_{c551} complex, the edge of the haem moiety lies some 10.5 Å from T1Cu in an arrangement likely to promote efficient intermolecular ET, Fig. 5A. The protein–protein interface involves the previously proposed hydrophobic patch surrounded by a semicircle of interface water molecules.⁴⁹ Analysis of the structure suggested the main ET route to be from the haem CBC methyl carbon through space to Pro88 of the CuNiR. Through-bond ET then proceeds to the adjacent T1Cu ligand His89.

Recently, a heterogeneous complex between *AxNiR* and a pseudoazurin from *H. denitrificans* as the electron donor was solved at 3.0 Å resolution (PDB 5B1J). The intermolecular contact at the hydrophobic patches adjacent to the T1Cu sites positions the Cu centres of the two proteins to within ~16 Å. The solvent-exposed His81 Cu-ligand of the pseudoazurin lies ~2.6 Å from the Ala86 oxygen atom of *AxNiR*, an arrangement that would allow through-space ET between the proteins,⁷² Fig. 5B.

3-Domain fused CuNiRs

A recent development has been the discovery of CuNiRs containing additional domains. The first example was the structure of the hexameric CuNiR from *Hyphomicrobium denitrificans* (*HdNiR*),⁷³ determined at 2.2 Å resolution,⁷⁴ Fig. 1. *HdNiR* has high sequence homology to *AniA*, while the structure revealed a novel additional N-terminal cupredoxin domain, containing an additional T1Cu(T1Cu_N) centre. The extended enzyme forms a 340 kDa hexameric prism-shaped oligomer. Interestingly, the additional T1Cu_N cupredoxin domain has low sequence homology to the other cupredoxin domains of *HdNiR* but high homology with those of *AfNiR*. The separation of the T1Cu_N centres is 14 Å in a head to head arrangement, suggesting that ET proceeds by initial reduction of T1Cu_N by Cyt_{c550} followed by ET between T1Cu_N and T1Cu_C, Fig. 6A. The additional cupredoxin domain in *HdNiR* may not be involved in the catalytic mechanism⁷⁴ due to the large distance separating it from the T2Cu centre where catalysis occurs. Subsequent bioinformatics analysis led to the identification of other N- and C-terminal 3-domain CuNiRs.⁹ The C-terminally-extended enzymes possess an additional haem-cytochrome domain. A crystal structure of a C-terminally extended CuNiR from *Pseudoalteromonas haloplanktis* (*PhNiR*) was deposited in the Protein Data Bank in 2008 (PDB 2ZOO) but remains unpublished. The first detailed and published structural analysis of a C-terminally extended CuNiR, came from the atomic resolution (1.01 Å) structure of the 499 residue CuNiR from *Ralstonia pickettii* (*RpNiR*) by Hasnain and co-workers.⁷⁵ The 1.01 Å resolution structure of *RpNiR*⁷⁵ revealed a domain binding motif that positioned the T1Cu and Fe atoms only 10.6 Å apart, notably



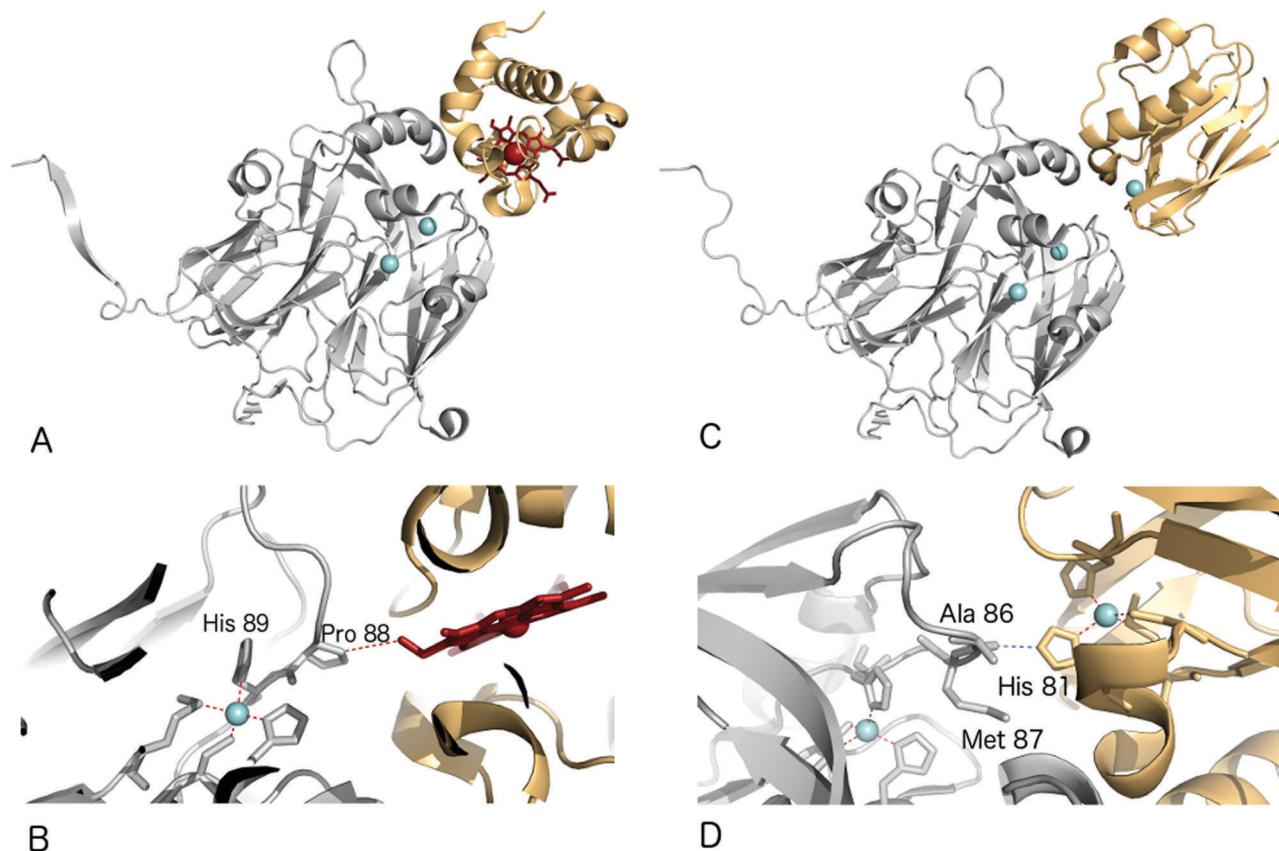


Fig. 5 Structures of electron transfer complexes between CuNiRs and electron transfer proteins. Cu atoms are shown as cyan spheres, while the haem group of cyt. c_{551} is represented using red sticks (A) structure of the proposed electron transfer complex between AxNiR (gray) and cyt. c_{551} (orange); (B) the interface region of the complex shown in (a), note the close contact between Pro88 of AxNiR and the haem edge of cyt. c_{551} . (C) Structure of the proposed electron transfer complex between AxNiR (gray) and pseudoazurin from *H. denitrificans* (PAz, orange); (D) interface region of the complex shown in (C) here note the close contact between Ala86 of AxNiR and the T1Cu ligand His81 in Paz.

closer than the separation of centres in the AxNiR–Cyt c_{551} complex, Fig. 5 and 6. A series of water molecules were present between the cytochrome domain and the surface above the T1Cu site, suggesting a role in electronic coupling across the interface. This hypothesis was tested by structural and functional analysis of site-directed mutants which confirmed the roles of Met92 and Pro93 in interface ET. As well as the differences in domain structure, notable differences are apparent between RpNiR and the 2-domain NiRs in the vicinity of the catalytic T2Cu centre. In contrast to 2-domain NiRs where the T2Cu-bound water is H-bonded to Asp_{CAT} and His_{CAT}, in RpNiR this water instead forms an H-bond to Asp_{CAT} and a second water molecule. In PhNiR, the water is only H-bonded to Asp_{CAT}.

Notably, in the substrate binding channel of RpNiR, a Tyr residue acts to ‘plug’ the substrate binding channel, excluding water molecules.⁷⁵ The atomic resolution structure of RpNiR provided clear evidence for the observed preference for nitrite binding to reduced T2Cu in this enzyme, rather than the preferred oxidised state in 2-domain CuNiRs.

Membrane associated CuNiRs

The crystal structure of the soluble domain of the outer membrane AniA protein from *Neisseria gonorrhoeae* was determined in 2002.⁴⁴

The AniA protein has around half the specific activity of AfNiR, is essential for growth in low-oxygen conditions with nitrite present and protects against human-sera killing of the pathogen. The soluble domain has a similar fold to the typical CuNiRs but has two shortened loops, with one loop possibly related to interactions with a membrane anchored azurin electron donor and the other likely to interact with the membrane itself. The Cu centres in AniA are similar to those of 2-domain CuNiRs, but with a visible colour intermediate between that of green and blue sites. While bioinformatics identified related proteins to AniA across a range of species, further structures have not yet been forthcoming.

Variations on the classical CuNiR structure and further classification of CuNiRs

In the vicinity of the metal centres, CuNiRs have been divided into classes based on the length of the two loops, the linker loop between the cupredoxin domains forming a monomer and the ‘tower loop’ adjacent to the T1Cu site.⁴⁴ The majority of structures are of class I enzymes, while class II includes AniA.⁴⁴ Some recently discovered CuNiRs fall between these two classes with intermediate length loops, e.g. GkNiR. Several CuNiRs from thermophilic bacteria have been structurally characterised. The structure of *Geobacillus kaustophilus* HTA426 CuNiR (GkNiR)⁵¹



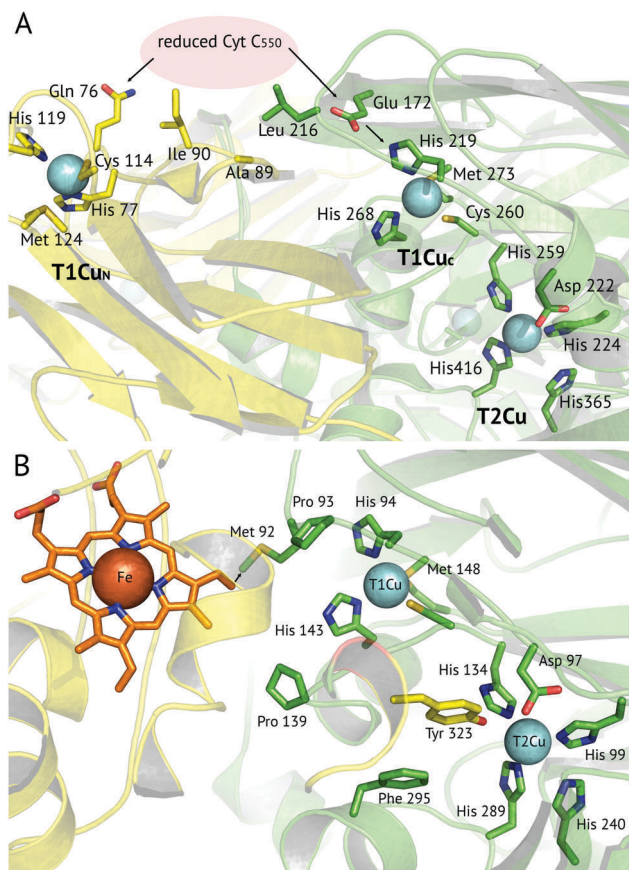


Fig. 6 Proposed electron transfer routes between the redox centres of 3-domain CuNiRs. In (A) *HdNiR* the external electron donor partner protein cyt. C_{550} is proposed to bind to a hydrophobic patch spanning the N-terminal (near Ala89, Ile90) and cupredoxin domains (near Leu216), where it is positioned for electron transfer to both the T1Cu_N and T1Cu_C sites (indicated by arrows).⁷⁴ In (B) *RpNiR*, the proposed internal electron transfer route is from the haem (CBC atom) to Met92 and to the T1Cu site.⁷⁵ Interface and T2Cu water molecules have been omitted from the figures for clarity.

revealed a novel 28 residue additional α -helical region close to the N-terminus (residues 41–68). Other notable features of *GkNiR* include the presence of a Val residue in place of Ile_{CAT} and differences in the hydrogen bonding network around the T2Cu centre.

Much of the structural data presented above has come from single, static crystal structures. Recent developments in X-ray sources, rapid detectors and data handling have allowed the determination of both ‘damage free’ single structures from large numbers of microcrystals and also multiple (potentially hundreds of) sequential datasets from just one protein crystal, making it possible to follow reactions *in crystallo*. Both approaches show promise for determining structures of relevant catalytic intermediates.

Insights from multiple-structures one crystal (MSOX) serial crystallography

Multiple Structures from One Crystal (MSOX) is a data collection technique which uses solvated electrons generated during X-ray

diffraction experiments to drive reactions requiring electron transfer.⁷⁶ By taking sequential diffraction datasets over the same volume of one crystal it is possible to observe a reaction, occurring through a series of high resolution crystal structures with progressively increasing X-ray dose, Fig. 7. For NiRs, this approach was used to obtain 3 (low, medium and high-dose) datasets from tobacco assimilatory NiR, a Fe-S cluster and sirohaem containing protein that converts nitrite to ammonium.⁷⁷ A series of this sort for a CuNiR, comprising 45 consecutive datasets collected in 19 s each for *AcNiR*, between 1.07 Å and 1.62 Å resolutions, clearly showed conversion of bound NO_2^- to side-on NO^{55} within the crystal. At the end of the 45 structure series, a Cu(II) species with bound water is observed after loss of the NO ligand. Crystal structures are not yet able to give a definitive answer to the identification of the observed side-on NO complex as Cu(I)- NO^\bullet or Cu(II)- $\text{NO}^{\bullet-}$.

Little to no change to the overall fold of the enzyme or in the vicinity of the T1Cu site were seen throughout the MSOX series. The low dose atomic resolution starting structure exhibited a previously unseen dual conformation of NO_2^- , in ‘top-hat’ and ‘side-on’ bidentate orientations, of which the top-hat conformation matches the vertical substrate coordination from ‘radiation-damage-free’ XFEL structures,^{54,58} corroborating the idea that this vertical binding mode represents the initial nitrite binding position. Overall, the 45-dataset MSOX series demonstrated several structural changes at the T2Cu site, Fig. 7. Starting with the dual conformation nitrite and both gatekeeper and proximal Asp_{CAT} conformations, by dataset four the series showed nitrite adopting a single side-on conformation and increasing proximal Asp_{CAT} occupancy, lending credence to the operation of the sensor loop in response to

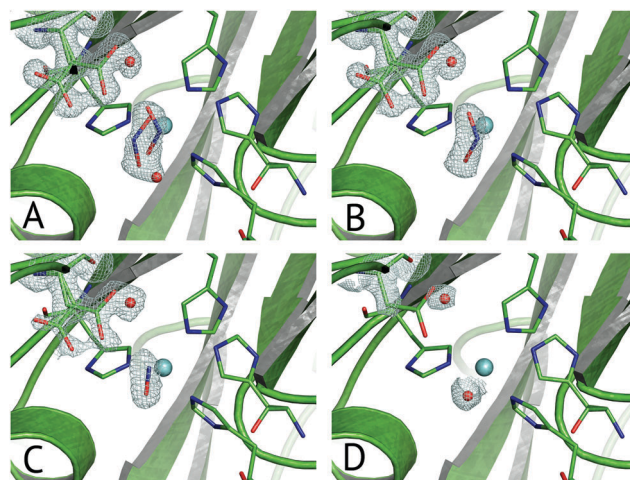


Fig. 7 Selected frames from the 45-frame MSOX structural movie of *AcNiR* catalysis at 100 K.⁵⁵ (A) The starting nitrite-bound state with partial occupancies in the top-hat and side-on modes (dataset 1, PDB 5i6k); (B) fully side-on nitrite (dataset 4, PDB 5i6l); (C) side-on nitric oxide product complex (dataset 17, PDB 5i6n); (D) water rebinding to the T2Cu following dissociation of NO (dataset 40, PDB 5i6k). Electron density is shown, contoured at $0.53-0.36 \text{ e}^- \text{ \AA}^{-3}$, for the ligands and the Asp_{CAT} residue, present in its ‘gatekeeper’ and ‘proximal’ orientations. The T2Cu atom is depicted by the cyan sphere and water molecules by red spheres.



the copper oxidation states during catalysis.²³ By dataset eleven a second previously unseen conformation was observed, with either NO_2^- , NO or a H_2O molecule at the T2Cu, indicative of catalytic turnover in the crystal. The changes in gatekeeper and proximal positions of Asp_{CAT} provide some structural evidence for the role of Asp_{CAT} in substrate delivery and product release.

XFEL structures of intact redox states of CuNiRs by serial femtosecond crystallography

Reduction of metalloproteins by X-ray generated solvated electrons has been well-characterised, presents a major challenge for structural biology of such proteins,^{78,79} and has been demonstrated to occur in CuNiRs.²⁷ Various online spectroscopic techniques have been developed to monitor the redox state of metals in crystals.^{80–82} Strategies to mitigate such reduction include merging of composite, low dose partial datasets⁸³ or helical data collection approaches.⁸⁴ Use of rapid X-ray detectors and cryogenic temperatures – typically 100 K – minimises but does not eliminate the effects of photoreduction or radiation damage to crystals. However, intact fully-oxidised structures may require the use of femtosecond X-ray pulses that can only be provided with sufficient brilliance by an X-ray free electron laser (XFEL).⁸⁵ The first CuNiR XFEL structures at room temperature were recently obtained using the Spring-8 Compact Free Electron Laser (SACLA).^{54,58}

Comparison of the 1.43 Å resolution serial femtosecond crystallography (SFX) structure of resting state *GtNiR*⁵⁸ with conventional synchrotron radiation (SRX) structures revealed a 10° rotation of the imidazole ring of His_{CAT} , in the SRX data in relation to the SFX structure. This led to the suggestion that His_{CAT} may function as a proton-relay switch, whereby T2Cu reduction causes His_{CAT} rotation thus altering the H-bonding network of His_{CAT} with Thr268 and Glu267 to destabilise the positive charge of His_{CAT} and facilitate proton transfer to the bridging water.

Subsequently, SFX structures were determined of *Alcaligenes faecalis* NiR (*AfNiR*) in its resting and nitrite bound (PDB 5D4I) states at 2.03 Å and 1.60 Å resolutions, respectively.⁵⁴ The SFX structure of intact nitrite bound *AfNiR* revealed a single, vertical ‘top-hat’ binding mode of NO_2^- compared to the side on conformation typically observed in SRX structures,¹⁸ suggesting the conformational change to side on NO_2^- results from photoreduction of the metal centres. *AfNiR*, as in *GtNiR*, shows a conformational change in His_{CAT} in photoreduced SRX structures compared to oxidised SFX structures. However, unlike *GtNiR* where His_{CAT} rotates 10° pivoting on $\text{N}^{\delta 1}$ and C^{β} with a 0.3 Å shift in C^{β} , in *AfNiR*, His_{CAT} rotates 20° about C^{γ} with no change in C^{β} , switching the H-bonding partner of His_{CAT} $\text{N}^{\delta 2}$ from the Glu279 carbonyl to the hydroxyl of Thr280 which is both a longer and weaker H-bond, Fig. 8. Based on this conformational change between SFX and SRX structure⁵⁴ an updated reaction mechanism for CuNiRs was suggested in which nitrite binds to the T2Cu and is protonated by Asp_{CAT} . This is followed by intramolecular ET to give side-on nitrite and rotation of His_{CAT} , where weaker H-bonds facilitate proton transfer from His_{CAT} to the bridging water by destabilising the positive charge of His_{CAT} . With this the necessary

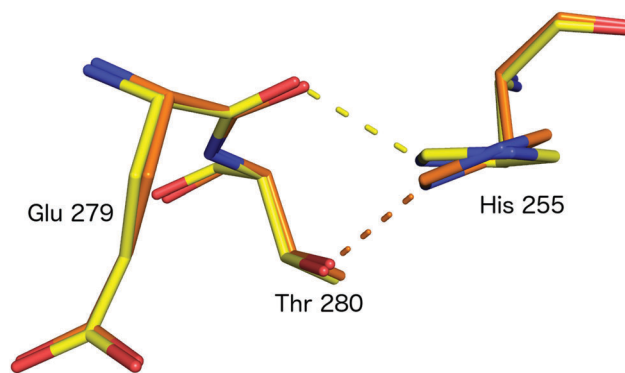


Fig. 8 His_{CAT} rotation in the proposed redox coupled proton transfer mechanism derived from comparison of XFEL (SFX; PDB 4YSC) and synchrotron radiation (SRX; PDB 4YSE) structures of *AfNiR*.⁵⁴ Note the H-bond switching of His_{CAT} (His255) between Glu279 O atom (damage free oxidised XFEL structure, yellow) and Thr280 (reduced SRX structure, orange).

second proton can be delivered to form NO in either end-on or side-on binding modes. However, the recently published MSOX series⁵⁵ showing X-ray induced ligand turnover from NO_2^- to side-on NO to water shows no change in the H-bond distances between His_{CAT} and Glu279 (2.7 Å) between low and high X-ray doses and a gradual increase in the bond length between His_{CAT} and Thr280 with increasing dose (2.8 Å at 0.69 MGy and 3.1 Å at 27.60 MGy).

Outlook and remaining questions

Several exciting new discoveries of CuNiR enzymes together with methodological developments have taken place in recent years. The advent of rapid mixing technologies at XFEL sources may allow time-resolved structural analysis of the CuNiR reaction with damage-free structures for reaction intermediates. The ever-increasing number of sequenced genomes promises the discovery of new variations of CuNiRs with differing domain structures. However, several outstanding questions remain that structural analysis will play a key role in addressing. For example, the protonation states of the important ‘catalytic residues’ have yet to be unequivocally determined for all stages of the reaction mechanism. Neutron diffraction – another way to obtain room temperature radiation-damage free structures – may help provide this information in the future and its potential for studying CuNiRs, in combination with high resolution X-ray structures, has already been shown.⁸⁶ Establishing the precise chemical nature of the side-on NO intermediate also remains challenging as do the routes of substrate access and product escape. While significant insights have been gained from the current structures of protein–protein electron transfer complexes, improved structural resolution is required to clarify details of the emerging picture. The dynamic behaviour of CuNiRs is clearly of vital importance and can be addressed by combining structural data, measured at different temperatures, with molecular dynamics and QM/MM simulations.

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