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Crystal structures from the *Plasmodium* peroxiredoxins: new insights into oligomerization and product binding

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Abstract

Background: *Plasmodium falciparum* is the protozoan parasite primarily responsible for more than one million malarial deaths, annually, and is developing resistance to current therapies. Throughout its lifespan, the parasite is subjected to oxidative attack, so *Plasmodium* antioxidant defences are essential for its survival and are targets for disease control.

Results: To further understand the molecular aspects of the *Plasmodium* redox system, we solved 4 structures of *Plasmodium* peroxiredoxins (Prx). Our study has confirmed PvTrx-Px1 to be a hydrogen peroxide (H₂O₂)-sensitive peroxiredoxin. We have identified and characterized the novel toroid octameric oligomer of PyTrx-Px1, which may be attributed to the interplay of several factors including: (1) the orientation of the conserved surface/buried arginine of the NNLA(I/L)GRS-loop; and (2) the C-terminal tail positioning (also associated with the aforementioned conserved loop) which facilitates the intermolecular hydrogen bond between dimers (in an A-C fashion). In addition, a notable feature of the disulfide bonds in some of the Prx crystal structures is discussed. Finally, insight into the latter stages of the peroxiredoxin reaction coordinate is gained. Our structure of PyPrx6 is not only in the sulfinic acid (RSO₂H) form, but it is also with glycerol bound in a way (not previously observed) indicative of product binding.

Conclusions: The structural characterization of *Plasmodium* peroxiredoxins provided herein provides insight into their oligomerization and product binding which may facilitate the targeting of these antioxidant defences. Although the structural basis for the octameric oligomerization is further understood, the results yield more questions about the biological implications of the peroxiredoxin oligomerization, as multiple toroid configurations are now known. The crystal structure depicting the product bound active site gives insight into the overoxidation of the active site and allows further characterization of the leaving group chemistry.

Background

There are at least 500 million clinical episodes of malaria annually with more than a million Africans dying each year, most of whom are children under 5 years of age [1]. The causative agent for the most lethal form of malaria is a protozoan parasite, *Plasmodium falciparum*, while *P. vivax* causes a less severe form, *P. knowlesi* is responsible for macaque malaria (but it can also infect humans [2,3]), and *P. yoelii* and *berghei* infect

rodents. *Plasmodium* parasites are frequently subject to oxidative attack, for example, in the erythrocyte from H₂O₂ release during heme metabolism and from NO and reactive oxygen species (ROS) generation during the host immune response [4,5]. In addition, oxidative stress is sustained during the sexual maturation of the parasite within the *Anopheles* mosquito midgut and salivary gland prior to transmission [6,7]. As such, *Plasmodium* antioxidant defences are essential to its survival, and thus are expected to be targets for the effective control of the disease [8,9].

Interestingly, neither the *Plasmodium* parasites nor the trypanosomes contain a catalase or a selenocysteine-

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containing glutathione peroxidase (GPx), which are enzymes notably efficient for the detoxification of hydroperoxides [10,11]. *Plasmodium* does however possess 2 superoxide dismutases, 6 proteins homologous to thiol-dependent peroxidases, and a glutathione-S-transferase (GST). The GST has only weak GSH peroxidase activity, but it might contribute significantly to the antioxidant capacity of the parasite due to its high concentration [12]. Of those homologous to the thiol-dependent peroxidases, there is the GPx-like thioredoxin peroxidase, which is a non-selenocysteine GPx known to be significantly less active than its selenium homologue [13]. The 5 remaining thiol-dependent peroxidase homologues identified in *Plasmodium* include thioredoxin peroxidase 1 and 2 (Trx-Px1 and Trx-Px2) from the peroxidase subfamily Prx1, a 1-Cys peroxidase (1-Cys Prx) from the Prx6 subfamily, antioxidant protein (AOP) from subfamily Prx5, and a very recently characterized nuclear peroxidase (*PfnPRx*) [14] (Table 1). Interestingly, peroxidases have recently been implicated in a different role, namely as a non-transcriptional rhythmic marker, indicative of the

circadian clock [19]. Various strategies have been used to classify the members of the peroxidase family including a phylogenetic tree analysis that categorizes them into 6 subfamilies (Prx1, Prx6, Prx5, Trx-Px, BCP, and AhpE), each of which may include the mechanistically distinct 1-Cys and 2-Cys peroxidases [20,21].

All peroxidases contain a conserved cysteine residue at the N-terminus that is referred to as the peroxidatic cysteine (C_P). During catalysis, it is oxidized by the ROS substrate (generally H₂O₂ or an alkyl hydroperoxide) to sulfenic acid (Cys-S-OH). Typical 2-Cys Prx contain 2 conserved cysteines, including the C_P and a C-terminal cysteine (termed the resolving Cys (C_R)). During catalysis, the C_P sulfenic acid reacts with the C_R of the adjacent monomer to form the intermolecular disulfide of the homodimer that is subsequently reduced by another (undetermined) thiol. In 1-Cys Prx, the C_P sulfenic acid is directly reduced by an unidentified redox partner.

Building on the dimer formation, the 2-Cys Prx enzymes organize themselves into higher order oligomers, such as decamers, which have higher peroxidase

Table 1 *Plasmodium* peroxidase orthologues and corresponding PDB codes for solved structures

| Subfamily | Name | Mechanistic Classification | Cellular Location | PlasmoDB ID | PDB ID | Reference |
|-----------|-------------------|----------------------------|-------------------|---------------|--------------|-----------|
| Prx1 | <i>Pf</i> Trx-Px1 | 2-Cys (C50, C170) | cytosolic | PF14_0368 | | [15] |
| | <i>Pv</i> Trx-Px1 | 2-Cys (C50, C170) | cytosolic | PVX_118545 | (2H66, 2I81) | this work |
| | <i>Py</i> Trx-Px1 | 2-Cys (C50, C170) | cytosolic | PY00414 | (2H01) | this work |
| | <i>Pb</i> Trx-Px1 | 2-Cys (C50, C169) | cytosolic | PB000037.01.0 | | |
| | <i>Pk</i> Trx-Px1 | 2-Cys (C50, C170) | cytosolic | PKH_126740 | | |
| | <i>Pf</i> Trx-Px2 | 2-Cys (C67, C187) | mitochondrial | PFL0725w | (2C0D) | [15,16] |
| | <i>Pv</i> Trx-Px2 | 2-Cys (C67, C187) | mitochondrial | PVX_123435 | | |
| | <i>Py</i> Trx-Px2 | 2-Cys (C67, C187) | mitochondrial | PY02747 | | |
| | <i>Pb</i> Trx-Px2 | 2-Cys (C59, C179) | mitochondrial | PB001545.02.0 | | |
| | <i>Pk</i> Trx-Px2 | 2-Cys (C67, C187) | mitochondrial | PKH_143220 | | |
| Prx6 | <i>Pf</i> Trx-Px1 | 1-Cys (C47) | cytosolic | PF08_0131 | | |
| | <i>Pv</i> Trx-Px1 | 1-Cys (C47) | cytosolic | PVX_093630 | | |
| | <i>Py</i> Trx-Px1 | 1-Cys (C47) | cytosolic | PY04285 | (3TB2) | this work |
| | <i>Pb</i> Trx-Px1 | 1-Cys (C47) | cytosolic | PB000600.02.0 | | |
| | <i>Pk</i> Trx-Px1 | 1-Cys (C47) | cytosolic | PKH_011610 | | |
| Prx5 | <i>Pf</i> Trx-Px1 | 1-Cys (C117) | apicoplast | MAL7P1.159 | (1XIY) | [17] |
| | <i>Pv</i> Trx-Px1 | 1-Cys (C114) | apicoplast | PVX_081760 | | |
| | <i>Py</i> Trx-Px1 | 1-Cys (C122) | apicoplast | PY01475 | | |
| | <i>Pb</i> Trx-Px1 | 1-Cys (C28) | apicoplast | PB000177.01.0 | | |
| | <i>Pk</i> Trx-Px1 | 1-Cys (C114) | apicoplast | PKH_021360 | | |
| BCP | <i>Pfn</i> Prx | 1-Cys (C56) | nuclear | PF10_0268 | | [14,18] |
| | <i>Pvn</i> Prx | 1-Cys (C52) | nuclear | PVX_111355 | | |
| | <i>Pyn</i> Prx | 1-Cys (C52) | nuclear | PY03834 | | |
| | <i>Pbn</i> Prx | 1-Cys (C61) | nuclear | PBANKA_051140 | | |
| | <i>Pkn</i> Prx | 1-Cys (C52) | nuclear | PKH_061160 | | |

Abbreviations include: *Pf*, *P. falciparum*; *Pv*, *P. vivax*; *Py*, *P. yoelii*; *Pb*, *P. berghei*; and *Pk*, *P. knowlesi*. Cellular location from experimental result, from result of orthologue, or from predictive targeting sequences. Mechanistic classification from experiment or based on experimental result and sequence alignment of orthologue

activity. Formation of the higher order oligomers is dependent on the redox state of C_P (and C_R), as well as other factors [22,23]. Both Trx-Px1 and Trx-Px2 have been identified as typical 2-Cys Prx enzymes; and a crystal structure of *P. falciparum* Trx-Px2 (PDB ID: 2C0D) has been published [16]. Distinguishing these two *Plasmodium* thioredoxin peroxidases is their cellular location, as Trx-Px1 is predicted to be cytosolic and Trx-Px2 has a mitochondrial targeting sequence [15]. Both features were recently confirmed [24]. Like the *Plasmodium* 1-Cys Prx from subfamily Prx6, AOP is also a 1-Cys Prx; and the *P. falciparum* AOP structure has been solved (PDB ID: 1XIY) [17]. AOP is thought to be an apicoplast enzyme due to its N-terminal signal motif, while the other *Plasmodium* 1-Cys Prx is cytosolic. Both predictions were recently confirmed experimentally [24]. Prx enzymes are highly expressed in *Plasmodium* (0.5% of cellular protein) and have been predicted from competitive kinetic analysis with human cells to be responsible for the reduction of 90% of mitochondrial H₂O₂ and nearly 100% of cytoplasmic H₂O₂ [25].

In order to further the understanding of the molecular details of the *Plasmodium* redox system, we solved the crystal structures of Trx-Px1 from *P. vivax* (PvTrx-Px1) in the reduced and oxidized forms, Trx-Px1 from *P. yoelii* (PyTrx-Px1) in the oxidized form, and a 1-Cys Prx from *P. yoelii* (termed PyPrx6, herein) with C_P oxidized to the sulfinic acid and with glycerol bound within the active site pocket. In addition, we have structurally confirmed and characterized the PvTrx-Px1 as a H₂O₂-sensitive peroxiredoxin; PyTrx-Px1 as forming an octamer (instead of the typical decamer and dodecamer arrangements); and PyPrx6 as a product bound complex revealing some interesting features of these enzymes.

Results

Expression and initial characterization of Trx-Px1 from *P. falciparum*, *P. vivax*, *P. yoelii*, and *P. knowlesi* and Prx6 from *P. yoelii*

Constructs of the Trx-Px1 enzymes from *P. falciparum*, *P. vivax*, *P. yoelii*, and *P. knowlesi* were expressed and purified as described previously [26]. PyPrx6 was also expressed in Studier auto-induction media [27]. All were full length constructs except PyTrx-Px1 which was also expressed for crystallization with a 6-residue truncation at the N-terminus. According to our mass spectroscopic analysis, all of our purified Trx-Px1 enzymes were disulfide-linked dimers (Table 2). As verified by mass spectroscopy, each of the 4 purified Trx-Px1 enzymes could be completely reduced using 20 mM dithiothreitol (Table 2).

By analytical gel filtration, all of the purified Trx-Px1 constructs (oxidized form and at high μM concentrations) eluted primarily as the higher order oligomer (*i.e.* octamer or decamer or dodecamer) (data not shown). Oligomerization has been shown to be dependent on numerous factors including ionic strength, pH, concentration of divalent metals, and most importantly redox state [28]. In our gel filtration, a small amount of presumably aggregated protein was followed by the oligomeric protein around (corresponding to a calculated mass of 272 or 314 kDa). In the case of PfTrx-Px1, some presumably dimeric protein was also observed, which was not noticed in previous gel filtration analyses of PfTrx-Px1 by Akerman and coworkers. [29]. These authors only observed higher order oligomers corresponding to 400 and 250 kDa for PfTrx-Px-1 and further detected the (α₂)₅ quaternary form by electron microscopy.

Table 2 Mass spectroscopy of reduced and oxidized Plasmodium Trx-Px1 orthologues

| | Expected MW of Reduced Monomer | Expected MW of Oxidized Dimer | Purified Enzyme (Da) | Purified Enzyme + 20 mM DTT (Da) |
|--------------------|--------------------------------|-------------------------------|--|----------------------------------|
| PfTrx-Px1 | 23702.08 | 47400.12 | 47456.56 (99%+) 23729.32 (trace) ^a | 23730.96 |
| PvTrx-Px1 | 23567.94 | 47131.84 | 47133.20 (major) 23568.94 (minor) | 23568.77 |
| PyTrx-Px1 | 23534.86 | 47065.68 | 47066.49 (99%+) 23533.70 (trace) | 23533.70 |
| PyTrx-Px1: Q7-L195 | 23234.40 | 46468.76 | 46470.60(99%+) 23234.49 (trace) | 23235.18 |
| PKTrx-Px1 | 24225.58 | 48447.12 | 48449.43 (99%+) 24226.10 (trace) | 24226.38 |

The calculated MW in Da is determined from the amino acid sequence and includes the His₆-tag incorporated into our constructs. The expected monomer MW is determined from the calculated monomer MW and subtracts the known *E. coli* post translational modification, namely clipping of the N-terminal Met (-131.19) of the His₆-tag [26]. The expected MW of the oxidized dimer is calculated by doubling the expected MW weight of the monomer (clipping of the N-terminal Met accounted for) and subtracting 4 H (-4.04). ^aThe MW of the purified monomeric PfTrx-Px-1 is off by 28 Da (and the dimer is off by twice this) which may be accounted for by one of the following: addition of ethyl addition, N, N-dimethylation of Arg or Lys, 2,4 bis-Trp-6,7-dione formation, or addition of formaldehyde (CHO)

PyPrx6 was expressed in two ways: PyPrx6 expressed from our typical protocol was purified in the reduced form according to mass spectroscopy (27189.1 Da) without addition of exogenous reducing agents (with the expected molecular weight after cleavage of the *N*-terminal Met weight being 27188.2 Da), while PyPrx6 expressed from Studier auto-induction media was purified in the sulfinic acid form (see below). From a calibrated gel filtration column, the enzyme eluted at 220 mL (corresponding to a calculated gel filtration mass of 62 kDa) which is consistent with the expected behaviour of a dimer.

Crystal structures of Trx-Px1 from *P. vivax* and *P. yoelii*

The crystal structures of *P. vivax* Trx-Px1 were solved in both the reduced (*PvTrx-Px1_red*, PDB ID: 2I81) and oxidized forms (*PvTrx-Px1_ox*, PDB ID: 2H66) at 2.45 Å and 2.5 Å, respectively (Figure 1). Aside from these 2 structures, to date only rat and *Salmonella typhimurium* Prx1 subfamily structures have been solved in both fully reduced and oxidized (as the disulfide) redox states [21]. In comparison to other solved structures, *PvTrx-Px1* is most similar (52% sequence identity) to *PfTrx-Px2* [16]. It has 85% sequence identity with *PfTrx-Px1* and 47% identity to its closest human orthologues. Herein, the oxidized form of *PyTrx-Px1* has also been solved to 2.3 Å (PDB ID: 2H01) (Figure 1C).

The structures of each (*PvTrx-Px1_red*, *PvTrx-Px1_ox*, and *PyTrx-Px1_ox*) contain the typical thioredoxin-fold found in known peroxiredoxins. There is a central 7-stranded β -sheet comprised of $\beta 2$ - $\beta 1$ - $\beta 5$ - $\beta 4$ - $\beta 3$ - $\beta 6$ - $\beta 7$ with $\beta 1$ and $\beta 6$ running anti-parallel relative to the other strands. This β -sheet is sandwiched by $\alpha 1$ and $\alpha 4$ on one side and $\alpha 2$, $\alpha 3$, and $\alpha 5$ on the other side. The root mean square deviation (rmsd) between *PvTrx-Px1_ox*, and *PyTrx-Px1_ox* is 0.91 Å when superimposing over the monomer encompassing $C\alpha$'s from residues 2 to 177. The remaining residues of the *C*-terminus are disordered (from 178 to 195) in the *PvTrx-Px1_ox* structure.

In *PvTrx-Px1_red* structure, the C_P residue (Cys50 for *P. vivax*) is located at the first turn of helix $\alpha 2$ at the end of a narrow accessible channel formed by a loop-helix motif and surrounded by 3 conserved residues Pro43, Thr47, and Arg125 (Figure 2A, red/dark grey). The pyrrolidine ring of Pro43 limits solvent accessibility and protects the reactive cysteinyl sulfenic acid from further oxidation during catalysis. The distance between Cys50 and the corresponding C_R thiol from its dimeric partner is 13.5 Å in the reduced form. In both the oxidized structures of *PvTrx-Px1_ox* and *PyTrx-Px1_ox*, the $\alpha 2$ helix is locally unfolded (LU) around the C_P , such that the $\alpha 2$ helix begins after the C_P with Ser52 (coloured in orange in Figure 2A for *Pv* and *Py*: *PvTrx-*

Px1_ox) or Ser46 (Figure 2B for *PyTrx-Px1_ox*). Some conserved residues that form the active site in the reduced structure are in the same positions and orientations in both the oxidized structures (Pro43, Thr47, and Arg125 for *PvTrx-Px1*, Figure 2A, orange/light grey and Pro37, Thr41, and Arg119 for *PyTrx-Px1*, Figure 2B). The C_P has rotated to the surface as part of a highly exposed loop; and S_P (sulfur of C_P) is engaged in a disulfide bond with S_R (sulfur of C_R) of the enzyme's dimeric partner forming a domain swapped homodimer.

PvTrx-Px1 and *PyTrx-Px1* are H₂O₂-sensitive peroxiredoxins

The 3 sequence motifs that define the H₂O₂-sensitive peroxiredoxins include: (1) the conserved loop-helix from Pro43 to Glu53 surrounding the C_P ; (2) a 3_{10} helix-loop from Pro89 to Ile98; and (3) the 29 *C*-terminal residues from Gly167 including C_R and conserved bulky residues [30]. The loop-helix is completely conserved among human, rat, and *Plasmodium* peroxiredoxins. However, the *Plasmodium* enzymes are a slight variation of 3_{10} helix-loop and *C*-terminal tail motifs with sequences including $^{93}GGIG^{96}$ and $^{191}YL^{192}$, respectively, instead of the typical GGLG and YF (Figure 3A). Upon addition of 500 μ M to 5 mM H₂O₂ to reduced (by DTT) *PfTrx-Px1*, *PvTrx-Px1*, *PkTrx-Px1*, or *PyTrx-Px1* enzymes, additions of 2 and 3 oxygen atoms were observed by mass spectroscopy, confirming that these enzymes are H₂O₂-sensitive. Structurally, the first loop-helix and the *C*-terminal arm of *P. falciparum* were predicted by modelling to undergo the same structural rearrangement during catalysis as the mammalian peroxiredoxins [31]. Figure 3B, 3C illustrate the structural changes that *PvTrx-Px1* does indeed undergo during catalysis further supporting its characterization as a H₂O₂-sensitive peroxiredoxin. Although predicted by Kawazu [31], to be fluid (*i.e.* structurally disordered) from Pro171 immediately following C_P , the *C*-terminal tail is an ordered loop from Pro171 to Gly177 in the *PvTrx-Px1_ox* structure. The *PyTrx-Px1_ox* structure also shows a similar arrangement of conserved residues, a 3_{10} helix-loop motif at $^{90}PLSQGGIGNI^{98}$, and a *C*-terminal tail bearing a $^{191}YL^{192}$ motif (nearly identical to *PvTrx-Px1* that folds upon reduction into a loop followed by an α -helix), so it is also expected to be a H₂O₂-sensitive peroxiredoxin (Figure 2B and 3A). The conserved $^{191}YL^{192}$ motif that is located on the α -helix close to the surface stabilizes the full-folded conformation. This motif therefore slows the resolution reaction and allows overoxidation by reaction with a second equivalent of peroxide. In contrast robust peroxiredoxins do not have residues protecting the C_P and are quickly oxidized to the disulfide [30].

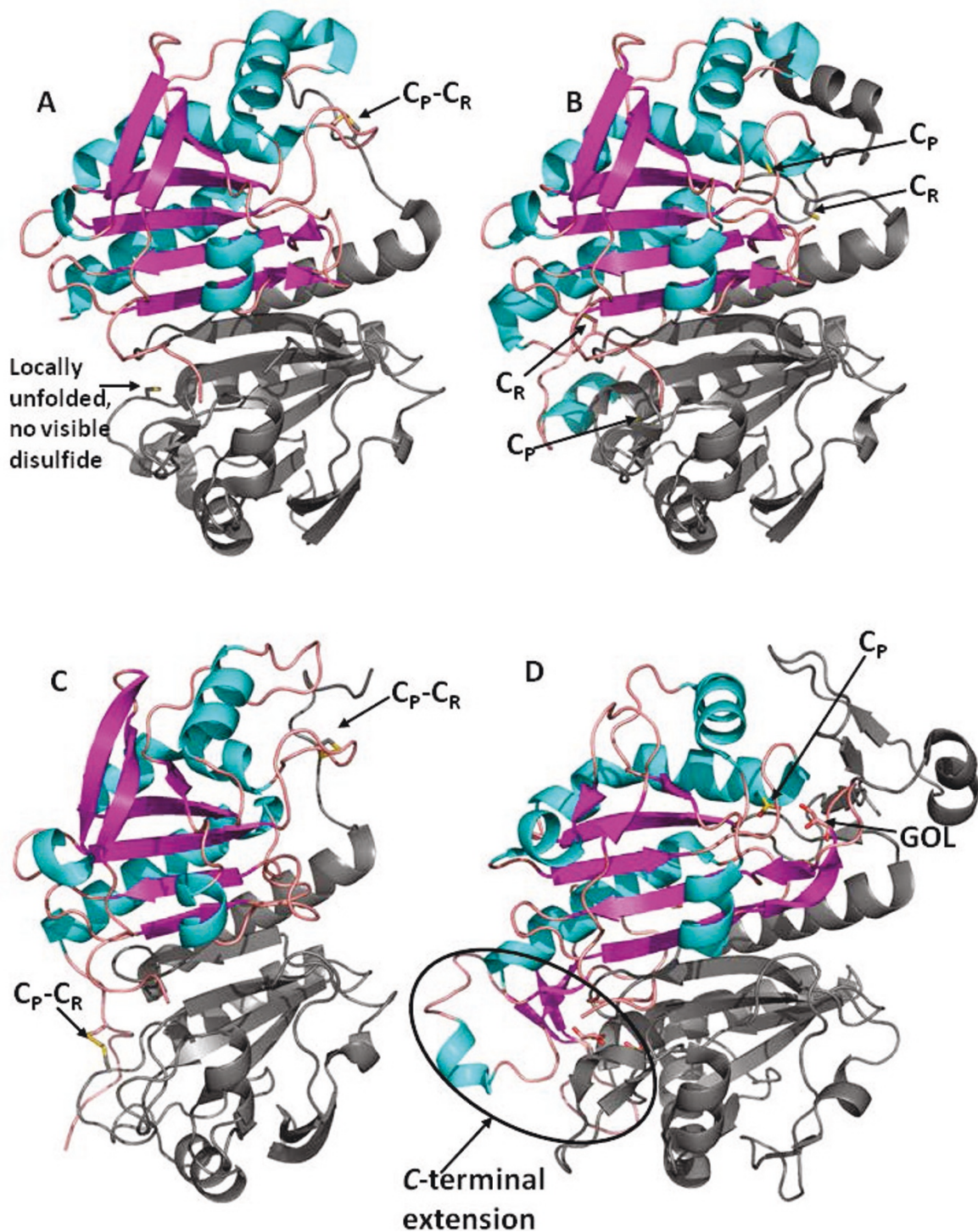
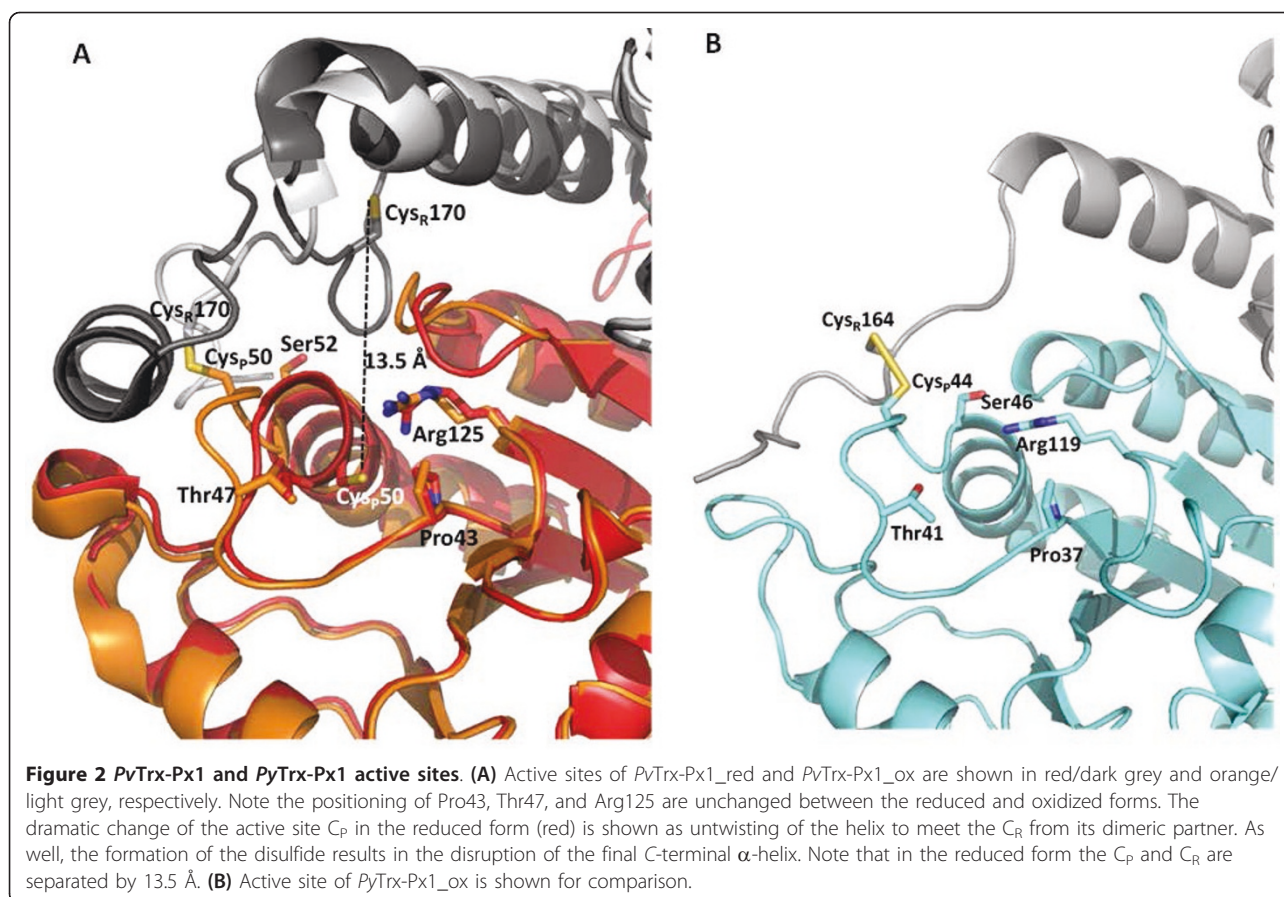


Figure 1 Structures of the dimeric units of the *Plasmodium* peroxiredoxins. The structures of the dimeric units of the peroxiredoxins with the C_P and C_R thiol side chains are shown to display the secondary structure in one of the monomers with α -helices shown in blue, β -sheets shown in pink, and sulfur and oxygen atoms displayed in yellow and red, respectively. **(A)** *PvTrx-Px1_ox* is shown with only one disulfide visible, as the second one is not visible due to the lack of structure at the C-terminus (thus C_P) of the monomer shown with secondary structure colours. **(B)** *PvTrx-Px1_red* with all 4 reduced thiols clearly visible. **(C)** *PvTrx-Px1_ox* with the 2 disulfides clearly displayed; and **(D)** *PvPrx6* is shown with its sulfenic acid active site cysteine and with glycerol bound.



Disulfide bonds in the *PvTrx-Px1*_{ox} and *PyTrx-Px1*_{ox} structures

Upon examination of each dimer of *PvTrx-Px1*_{ox} decamer ($(\alpha_2)_5$ oligomer) (Figures 1A and 4A), only 4 of a possible 10 disulfide bonds are clearly defined in the crystal structure. There is inadequate density to define the C-terminal tail from around the C_R for the remaining residues, so that for several of the monomers only the side chain of C_P but not that of C_R is visible. In the case of the *PyTrx-Px1*_{ox} octamer ($(\alpha_2)_4$ oligomer) (Figures 1C and 4B) (for which the data was collected at a home source) the Cys-S_P to Cys-S_R distances measure ~3 Å (notably, bond lengths at the resolution of these structures are derived from a combination of x-ray data and chemical constraints). There are no reports indicating that disulfide bonds are labile under the conditions used in our *PyTrx-Px1* experiments. As expected, the C-terminal tails of *PvTrx-Px1* and *PyTrx-Px1* have higher B-factors than the other parts of the molecule, indicative of a more fluid region and also of the apparent absence of detectable disulfide bonds in portions of the *PvTrx-Px1* structure and the distortion of the disulfide bond length in the *PyTrx-Px1* structure. In previously published crystal structures of oxidized 2-Cys Prx, the Cys-

S_P to Cys-S_R distance is also longer than expected for a disulfide bond (as the typical bond length is 2.05 for a disulfide). For example in the structure of a 2-Cys Prx from *Helicobacter pylori* (*HpaHpc*) (PDB ID: 1ZOF), the Cys-S_P to Cys-S_R distances measurements range from 2.0 to 3.2 Å and in a *P. falciparum* 2-Cys Prx (*PfTrx-Px2*) (PDB ID: 2C0D) one of the disulfides is shown in two different orientations (2.0-2.2 Å) indicative of the structural flexibility of these structures, while the other measures at 2.6 Å. Our analysis using mass spectroscopy (already discussed) and the overall structural configurations (*i.e.* local unfolding about C_P and at the C-terminus) both support *PvTrx-Px1*_{ox} and *PyTrx-Px1*_{ox} being in the oxidized form making these long Cys-S_P to Cys-S_R distances not easily accounted for, yet prevalent in the Prx1 subfamily.

Oligomeric organization of *PvTrx-Px1* and *PyTrx-Px1*

Oligomeric peroxiredoxins are formed via at least 2 types of interactions: (1) B-type interactions where edge to edge associations of $\beta 7$ strands of the central sheet meet to extend it into a 14-stranded β -sheet; and (2) A-type where the interface is a tip to tip association centred on helices $\alpha 4$ and $\alpha 5$ packing against helices $\alpha 4$

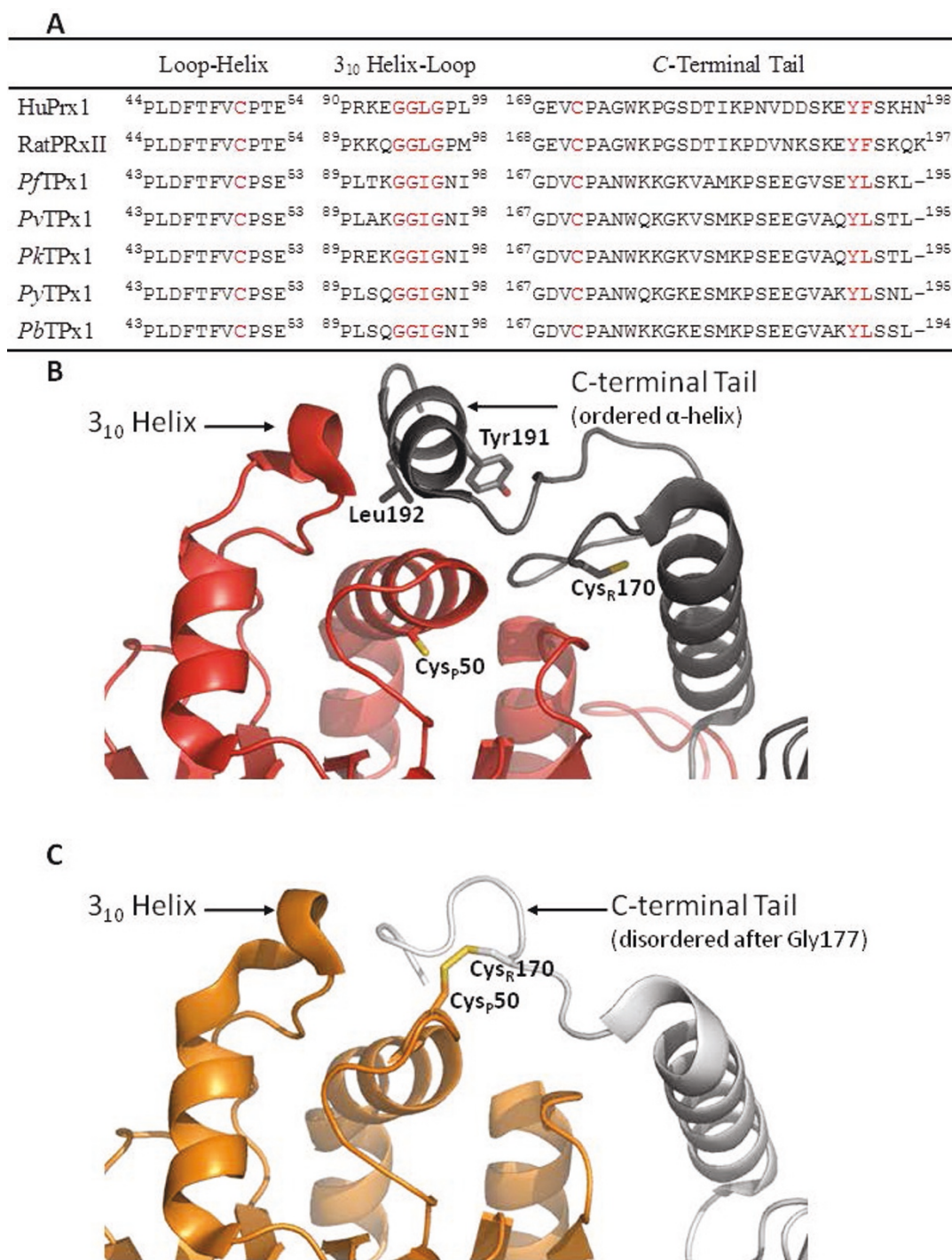


Figure 3 *PvTrx-Px1* is H₂O₂-sensitive. (A) Comparison of the sequences of the H₂O₂-sensitive mammalian peroxidoredoxins to the *Plasmodium* peroxidoredoxins discussed herein. (B) & (C) Active sites of the reduced and oxidized forms of *PvTrx-Px1* exemplifying the features described in the text of the H₂O₂-sensitive peroxidoredoxins.

and $\alpha 5$ of the other chain [17]. The dimer interface of the *PvTrx-Px1_{ox}* and *PyTrx-Px1_{ox}* (i.e. where the interactions resulting at the surfaces formed by the disulfide bond) is termed the B-type interaction face.

Accordingly, the same dimeric unit is expected in the reduced structures. These dimers then associate via A-type interactions to form the higher order oligomers, which are typically decameric and dodecameric. *PvTrx-*

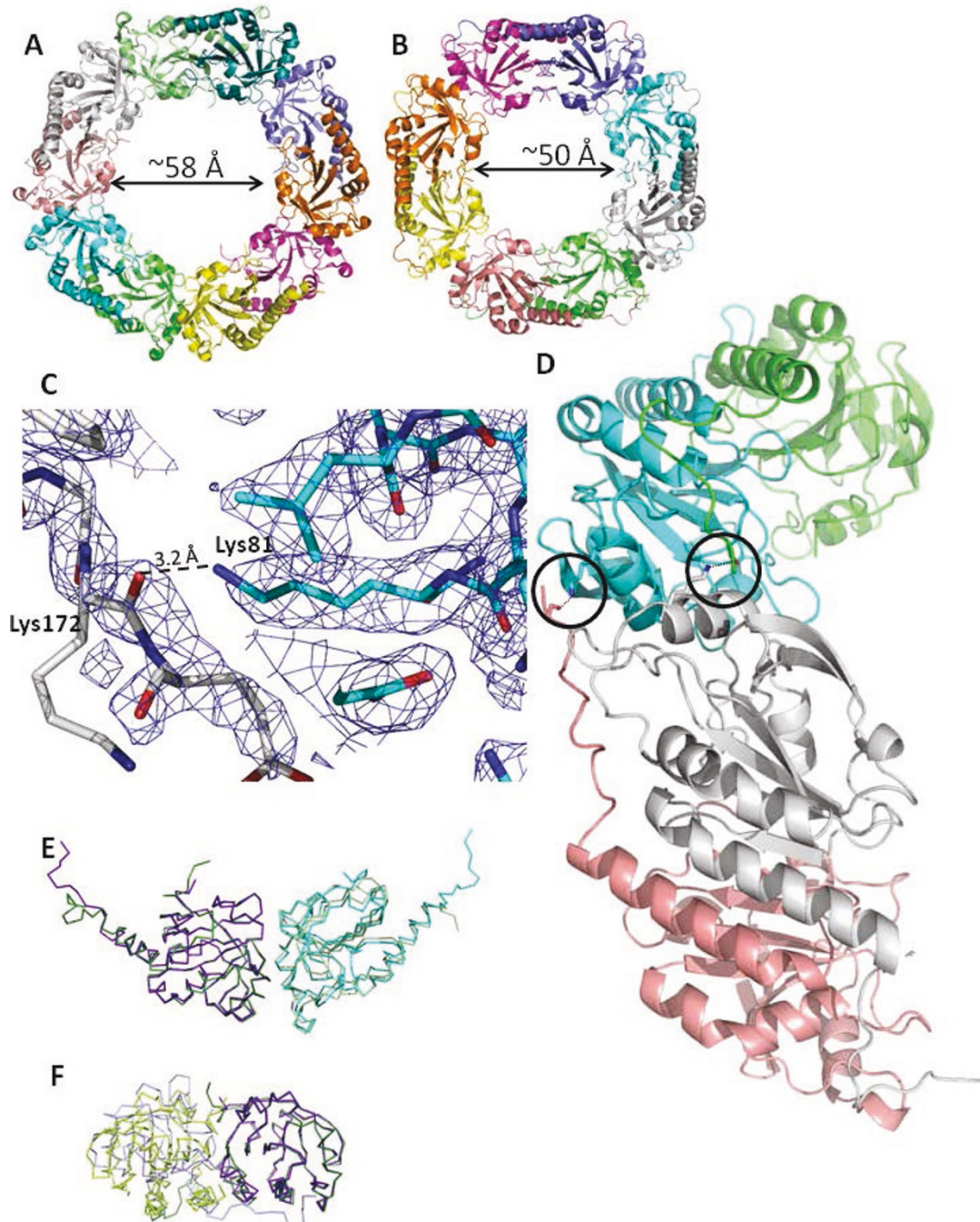
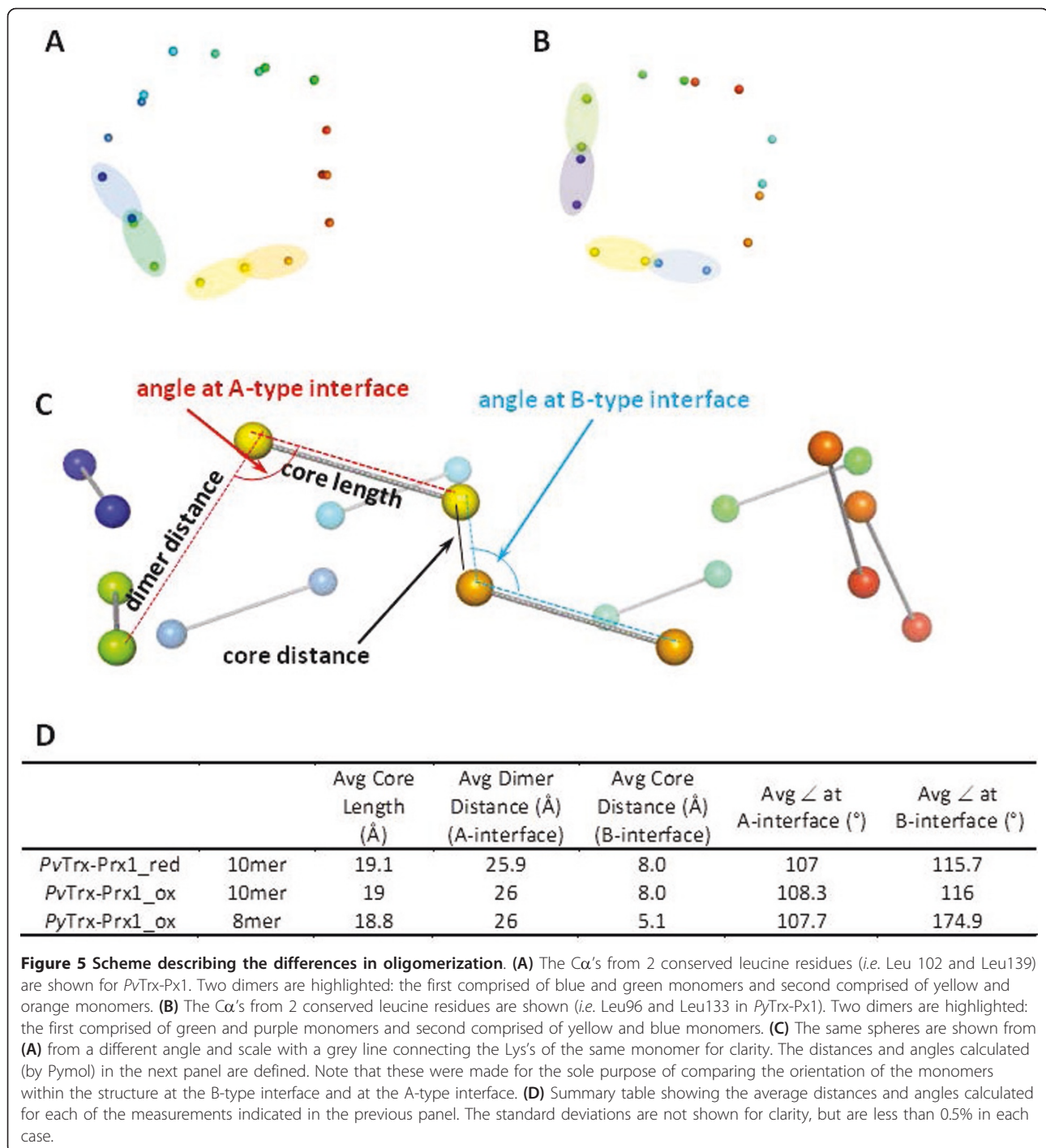


Figure 4 Plasmodium peroxiredoxin oligomeric structures. Oligomeric structures of (A) *PvTrx-Px_ox* and (B) *PyTrx-Px1_ox* showing the internal dimer. (C) The close-up view of a hydrogen bond at the A-type interface between the side chain of Lys81 from one monomer (cyan) and the main chain carbonyl of Lys172 from an adjacent molecule (grey) with the 2mFo-dFc electron map contoured at 1.0 σ (blue mesh). (D) Examples of the novel hydrogen bonding interactions of the *PyTrx-Px_ox* are circled in black. The panel is simplified to show only a tetramer for clarity, but indeed there are hydrogen bonding interactions across all of the A-type interfaces. Although the hydrogen bonding interaction is at the A-type interface, it is formed between distant chains in an A-C fashion (where A, B, C, and D chains are pink, grey, cyan, and green, respectively). For example, in the pink chain Lys172 carbonyl backbone is shown hydrogen bonding with the side chain of Lys81 from the cyan chain (A-C fashion) and in the grey chain Lys81 is hydrogen bonding with the main chain carbonyl of green Lys172 (B-D fashion). A comparison of the A-type interactions (E) and B-type interactions (F) between *PvTrx-Px1* and *PyTrx-Px1* is shown. In (E) and (F) the monomer structures (dark green and purple) are structurally aligned (rmsd = 0.573 Å), so that upon comparison of the corresponding dimeric partners the difference in the interfaces are shown (*PvTrx-Px1* shown in light greens and *PyTrx-Px1* shown in light blue/purple). A full alignment of the A-type dimers gives a rmsd = 1.065 Å, while a full alignment of the B-type dimers gives a rmsd = 3.137 Å.

Px1_ox and PvTrx-Px1_red arrange in a decameric fashion (according to our crystal structure), which from a survey of peroxiredoxin structures deposited to the PDB is the most common oligomerization. According to our results (namely a crystal structure and the gel filtration elution times characteristic of the higher order oligomer and not a dimer), the toroid PyTrx-Px1 structure is unique in that it is octameric (Figure 4B). PvTrx-Px1_ox and PvTrx-Px1_red decamers have internal diameters of ~58 Å, while PyTrx-Px1_ox octameric diameter is correspondingly smaller at ~50 Å. Li et al. reportedly solved the structure of another octameric peroxiredoxin (*Mycobacterium tuberculosis* AhpE), but this has been disputed as simply a crystallographic artifact, as predominantly dimers were observed in gel filtration and the octameric interface is not extensive and does not involve the typical interfaces [32,33]. It could be argued that the octameric arrangement of PyTrx-Px1 is a result of the crystal packing of PyTrx-Px1_ox dimers; however, no dimers were observed in the gel filtration of the PyTrx-Px1_ox sample which was done at a high μM concentration comparable to crystallization experiments. Interestingly, there is a hydrogen bonding interaction in the PyTrx-Px1 structure between distant monomers and more specifically between adjacent dimers, where the C-terminal tail of one monomer crosses its dimeric partner to hydrogen bond to next monomer. Contributing to the stability of the octamer, the side chain of Lys81 is within hydrogen bonding distance (3.2 Å) of the backbone carbonyl of Lys172 (Figure 4C). Indeed, there are interactions between each of the pairs A-C, B-D, C-E, D-F, E-G, F-H, G-A, and H-B, as one would expect from symmetry (Figure 4D). Although these 2 residues are conserved in PvTrx-Px1, there is no similar interaction in the PvTrx-Px1_ox or PvTrx-Px1_red structures (where the corresponding residues are Lys87 and Gly177). There is 83% sequence identity between the *P. yoelii* and *P. vivax* Trx-Px1 enzymes, so the differences in oligomeric state were not predictable. Previous reports have identified alterations of the B-type interface as conferring the different orders of oligomeric state, while the A-type interface remains constant [33]. An alignment of the Ca's of a monomer from the decameric PvTrx-Px1_ox with a monomer from octameric PyTrx-Px1_ox (rmsd = 0.573 Å) is shown in Figure 4E, F to illustrate the overlap of each accompanying monomer in the A-type interface and B-type interface, respectively. In Figure 4E, the corresponding A-type interface dimer is shown for the octamer and decamer which visually appears to preserve the overlap for their respective dimeric partners. Indeed, a structural alignment of the A-type dimers between the octamer and decamer gives a rmsd of 1.065 Å. Figure 4F shows the same monomers aligned, but in this case, their respective B-type interface

with the dimeric partner from the opposite side shown. Here, one can see that the overlap is poor which is also reflected as a much larger rmsd of 3.137 Å for the alignment of octameric and decameric B-type dimeric partners. In order to more directly compare the interfaces, each monomer is simplified to the 2 Ca's of conserved leucine residues at its core, for example, Leu 102 and Leu139 from PvTrx-Px1 and corresponding Leu96 and Leu133 from PyTrx-Px1 (Figure 5A, B, respectively). An analysis of the changes between the PyTrx-Px1 octamer and the PvTrx-Px1 decamer was undertaken using these conserved core residues as a representation of each monomer. Indeed, the model is validated, as the distance between the selected leucine residues (termed core length) is conserved throughout each of the structures and is similar between the two structures (Figure 5D). The orientation (termed angle between the dimers) and distance between the dimers within the oligomer is also conserved between the PyTrx-Px1 octamer and PvTrx-Px1 decamer (Figure 5D). On the other hand, the orientation (termed angle between monomers) and distance between the monomers of the dimer (termed core distance) is dramatically different between the octamer and decamer. The observation that the different oligomerization is attributable to the interface within the dimer suggests that the oligomerization is not an artifact of crystallization and that the dimer itself is unique at least in terms of its B-type interface.

Analysis of the structures of PyTrx-Px1 and PvTrx-Px1 identifies several key points of difference between the structures that may account for difference in oligomerization. First of all, the presence of a hydrogen bond between the dimers of PyTrx-Px1 (and not PvTrx-Px1) was already discussed (Figure 4C, D). Secondly, the C-terminal tails have different orientations, such that they are binding at different positions on the surface of their respective dimeric partners (Figure 6). The tails are bound by an intermolecular disulfide between Cp and Cr, as well as a series of hydrophobic interactions. The different binding orientations are linked to the orientation of the side chain of a conserved arginine (Arg142 and Arg148 for PyTrx-Px1 and PvTrx-Px1, respectively) (Figure 6A, B). In the case of PvTrx-Px1, the arginine is buried and the C-terminal tail adopts the typical binding pattern on the surface of its respective dimeric partner (Figure 6D). As opposed to PyTrx-Px1 where the equivalent arginine side chain is at the surface of the protein obstructing the typical pathway, such that the C-terminal tail adopts a different position at the surface of its partner (Figure 6C). Using the NCBI Molecular Modeling Database <http://www.ncbi.nlm.nih.gov/Structure/MMDB/mmdb.shtml>, a search was performed to identify 3D structures to similar PyTrx-Px1. A manual inspection of each the 2-Cys peroxiredoxins of the 143



low redundancy hits (from the 1532 total hits) showed no similarity in orientation to the *PyTrx-Px1* C-terminal tail, as expected as no other toroid octameric peroxiredoxins are known to date. A second point of differentiation between the two structures is a loop with the conserved sequence ¹⁴²NNLA(I/L)GRS¹⁴⁹ (numbering from *PvTrx-Px1* (PDB: 2H66)) that connects α 5 and β 7 (β -sheet involved in the peroxiredoxin B-type interface)

and contains the afore mentioned buried/surface Arg. The loops from each structure adopt different conformations that put Arg142 at the surface for *PyTrx-Px1* and the corresponding Arg148 buried for *PvTrx-Px1*. Although the *PvTrx-Px1* structure is from a full-length construct, only a structure with an N-terminal truncation of 6 residues crystallized sufficiently well for data collection in the case *PyTrx-Px1*. Although both

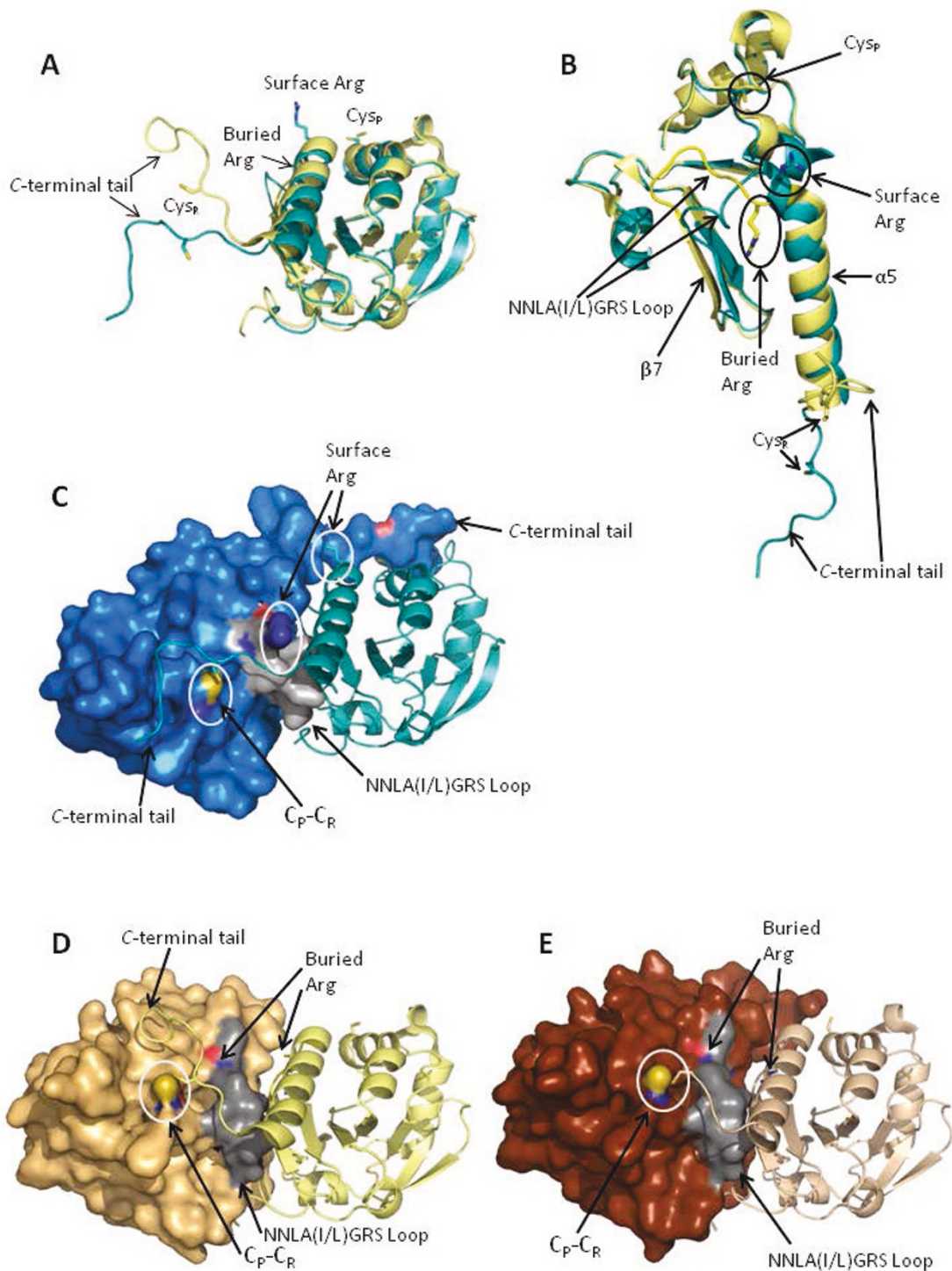


Figure 6 Role of C-terminal tail and NNLA(I/L)GRS-loop in the determination the oligomerization of *PyPrx-Tr1*. (A) and (B) Two views of the oxidized monomers of *PyTrx-Px1* (blue) and *PvTrx-Px1* (yellow) showing their structural differences: C-terminal tail adopts a different orientation in each structure, NNLA(I/L)GRS-loop also adopts a different orientation in each structure (see panel B), and the Arg adopts a surface position in *PyTrx-Px1* and a buried position in *PvTrx-Px1*. Note that chain I from PDB: 2H66 is used because it is most complete at the C-terminus. As well, portions of the structures in (B) are hidden to facilitate the view of the loop (including up to Tyr42 from the N-terminus and from Ala56 to Asp 77). (C) In order to demonstrate how the C-terminal tail positions itself in the dimer, a hybrid surface-cartoon rendering of *PyTrx-Px1* shows the orientation of the C-terminal tail and the NNLA(I/L)GRS loop in grey. (D) and (E) For comparison, the hybrid surface-cartoon renderings of *PvTrx-Px1* show the orientation of the C-terminal tail and the NNLA(I/L)GRS loop in grey. Note that (D) is from chains I and J which only shows a complete C-terminal chain for I. As well, for (E) chains C and D are used which both truncate around C_R for the C-terminal tail, but show that the C-terminus is the same up to this point.

constructs bear a *N*-terminal His₆-tag, the tag could only be cleaved from the *Pv*Trx-Px1 construct. Even after several days at room temperature (where typical conditions require an overnight reaction at 4°C), the His₆-tag remained intact on *Py*Trx-Px1. Although the tag is partially visible in *Py*Trx_Px1 case (³AFQG¹ from PDB: 2H01), it is not close enough to the conserved loop in both cases to affect its orientation. Further studies will determine what roles the structural features identified herein (*i.e.* the hydrogen bond between the dimers, the *N*-terminal tail, the surface/buried arginine, the NNLA(I/L)GRS-loop, and the *C*-terminal tail) play in the stoichiometry of oligomerization of these Trx-Px1 enzymes.

Crystal structure of *Py*Prx6 with product bound

The *Py*Prx6-SO₂H crystal structure has been solved at 2.3 Å (PDB ID: 3TB2). It is 47% identical to its closest human orthologue and shares 76% sequence identity with its *P. falciparum* orthologue. The closest available structure by sequence (at 48% sequence identity) is *Are-nicola marina* peroxiredoxin 6 which was further identified as a 2-Cys Prx [34], but it also shares 47% sequence identity with a solved structure for a human 1-Cys Prx [35]. Overall, the human (PDB ID: 1PRX) and *P. yoelii* structures are very similar with a core thioredoxin-fold and a *C*-terminal domain connected by both an extended helix α5 and a loop. The core rmsd is 1.38Å for 401 aligned residues in both structures. The *C*-terminal domain of *Py*Prx6 that comprised of 3 β-strands and a α-helix is larger than the *C*-terminal domain from the Prx1 subfamily (Figure 1D). This domain from each monomer extends over the other forming a domain swapped dimer.

The active site C_P (Cys47) is located at the bottom of a narrow pocket (~4 Å by ~7 Å) at the end of helix α2. There are 2 additional densities in each of the 4 active site pockets of the asymmetric unit (Figure 7) which are best filled by a glycerol molecule (from the purification buffer used which contained 5% glycerol) and fitting the C_P residues to their sulfinic acid derivatives (C_P-SO₂H). Previously, the C_P has been structurally characterized in the whole range of oxidation states (including C_P-SH, C_P-SOH, S_P-SO₂H, C_P-SO₃H, and C_P-SS-C_R) [36]; and our data agrees with those sulfinic acid structures previously studied. As well, nearly 20 peroxiredoxin structures have either substrate or what has been termed substrate analogue bound in their active site pockets, including H₂O₂, benzoate, acetate, dithiothreitol (oxidized), ethylene glycol, glycerol, sulfate, citrate, and formate [36].

In order to understand the reaction mechanism, our focus is on the comparison of structures with substrate (H₂O₂) or glycerol bound. The binding of H₂O₂ to

Aeropyrum pernix Tpx (thiol peroxidase) has been structural characterized (PDB: 3A2W and 3A2V) [38]. Further work on these structures and comparisons to other ligand bound peroxiredoxin structures previously showed that oxygen atoms of the ligands overlap with the proximal (relative to C_P) oxygen atom (O_A) and/or the distal oxygen atom (O_B) of bound H₂O₂ [37]. Interestingly, the glycerol molecule found in these structures can adopt all three possibilities: (1) in one monomer of *Ap*Trx (PDB: 3A2W) it is found with a single oxygen atom overlapping O_A (Figure 7C, pink); (2) in another monomer of *Ap*Trx (PDB: 3A2W), it is found to overlap with both O_A and O_B (Figure 7C, pink); and (3) reported for the first time, in our structure of *Py*Prx6, the glycerol is observed to overlay with only O_B (Figure 7C, purple). Previously, only 2 anionic ligands (sulfate and citrate) were observed to occupy O_B alone (see PDB: 1TP9 and 3DRN) [37]. Despite variations in the backbone orientation of the glycerol in our structure (which is also seen in the *Ap*Trx glycerol bound chains and presumably due to its conformational flexibility), the binding of all 4 glycerol molecules shows that each binds with the terminal hydroxyl in a similar position overlapping with O_B site (Figure 7B). Oxygen atoms at position O_A are postulated to be a mimetic for the substrate bound in a Michaelis complex ready for attack by the nucleophilic C_P, while oxygen atoms at position O_B are indicative of the leaving group (H₂O or alcohol) [37]. As such, it can be suggested that the glycerol positioned with its terminal hydroxyl at O_B is in a product bound configuration. Although there is variability in the binding of the remainder of the glycerol molecule, the alkyl group is always directed away from the active site pocket and the oxygen of the leaving group is in close proximity to a conserved threonine (Thr44). This arrangement suggests that it may be the proton donor, although others have suggested that this threonine functions as a hydrogen bond acceptor as it deprotonates the incoming substrate for attack by the C_P thiolate and that bulk solvent is responsible for the protonation of the leaving group [37]. Therefore, the active site pocket is adapted to accommodate different substrates (and thus products), which is exemplified through the structural flexibility exhibited in the product bound glycerol shown herein. As well, the O_B position appears to be designated for the oxygen of the leaving group and is well positioned for protonation by a conserved threonine.

The conserved residues of the C_P loop comprised of ⁴⁰PxxxxTxxC_P⁴⁷ and conserved Arg127 (numbering refers to *Py*Prx6) are implicated in catalysis (Figure 8). For the conserved arginine in both *Py*Prx6 (Arg127) and *Ap*Trx•H₂O₂ structures, it adopts position I, typical of Prx6, which bears a conserved arginine (Arg152) and

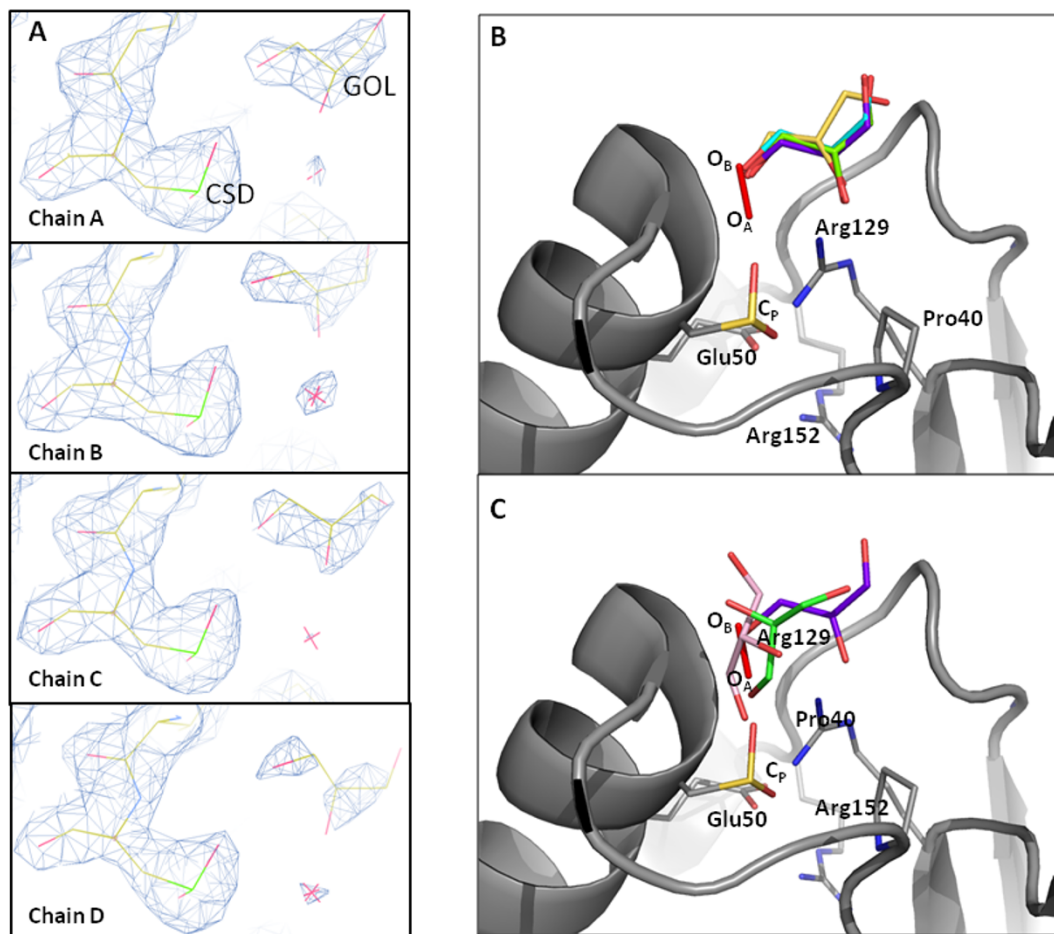
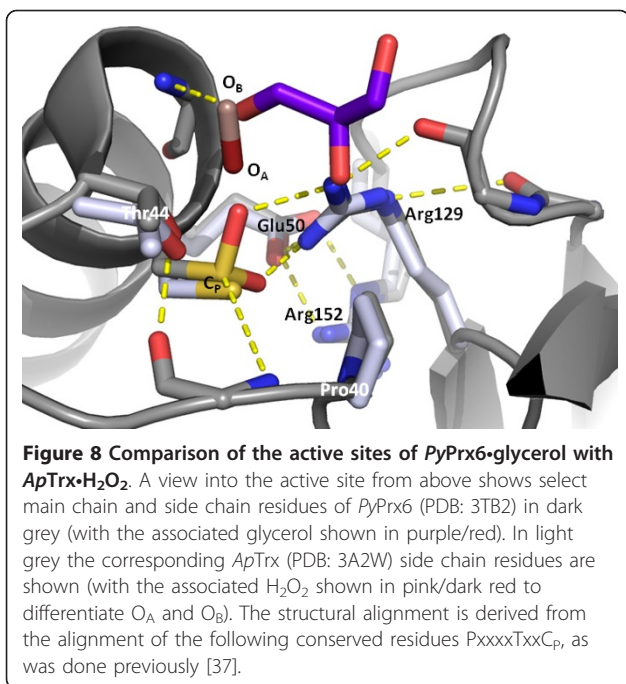


Figure 7 *PyPrx6* active site depicting the sulfinic acid and the binding orientations of glycerol. **(A)** Fc-Fo omit map (1.62 σ) of the active site from the 4 chains within the asymmetric unit of *PyPrx6* (PDB: 3TB2) showing the density assigned to both the sulfinic acid of C_P and the glycerol bound in the active site. Glycerol and sulfinic acid are labelled as GOL and CSD, respectively. Note that the carbons are yellow, sulfur is green, oxygen is red, and nitrogen is blue. **(B)** A comparison of the binding orientations of each of the glycerol molecules of *PyPrx6* relative to H_2O_2 derived from an alignment with the *ApTrx*- H_2O_2 structure (PDB: 3A2V) with our *PyPrx6* structure. The H_2O_2 is shown as the red oxygen atoms with positions O_A and O_B labelled, while the 4 glycerol molecules in the *PyPrx6* structure are shown with different coloured backbones and red oxygen atoms. Note that in each case, the terminal oxygen of the glycerol molecule aligns with position O_B . **(C)** For comparison, from a similar structural alignment the known binding orientations for glycerol (from PDB: 3A2W (*ApTrx*, chains A and C with the glycerol molecules depicted in pink and green, respectively) are compared to the same binding of H_2O_2 (PDB: 3A2V). Note that for *ApTrx* chain A, the glycerol terminal oxygen aligns with position O_A , while the glycerol molecule from *ApTrx* chain C has oxygen atoms aligning with positions O_A and O_B as was previously reported [37]. All structural alignments derived from the alignment of the following conserved residues PxxxxTxxC_P, as was done previously [37].

glutamate (Glu50) (or possibly a glutamine or histidine) and supports the positioning by a hydrogen bonding network, as previously described [37]. The active site geometry is fully folded (FF) and is virtually identical to that of the *ApTrx*· H_2O_2 . As was recently described, the nucleophilic C_P (as activated by a main chain amide N-H and the conserved arginine guanidinium) is expected to act as a thiolate and attack the substrate in an S_N2 fashion [37]. These hydrogen bonds, as well as those secondary ones from the backbone carbonyls and the glutamic acid/second conserved arginine, surrounding

the C_P are preserved between the *ApTrx* x· H_2O_2 and the *PyPrx6*·glycerol structures suggesting that the sulfinic acid form may be activated, and thus sufficiently nucleophilic (similar to its full reduced state) within the active site to undergo a further reaction with substrate. This would result in a subsequent oxidation of the active site to the sulfonic acid. Indeed, this form of the C_P has been structural characterized in other cases (PDB: 2CV4, 2NVL, and 1XIY). The conserved proline serves as a barrier to solvent, while the main chain of the C_P loop provides hydrogen bonds to the C_P and the



conserved threonine. Although at a reduced efficiency owing to the relative activity of an oxidized thiol relative to a thiolate, the preservation of the hydrogen bonding network about the active site cysteine, indicates that sequential H₂O₂ reductions are possible.

Discussion

Plasmodium lacks the two major antioxidant enzymes of eukaryotic cells, namely glutathione peroxidase and catalase [39]. As such, the parasite is likely to rely on the peroxiredoxin family to control peroxide production (as well as other reactive oxygen and nitrogen species) during critical stages of its lifecycle, for example during erythrocyte invasion when heme metabolism and immune response pathways ensue. During the trophozoite stage (feeding period), Prx enzymes accounts for 0.5% of the total expressed protein [40]. One recent study suggests that PfnPrx, the *P. falciparum* nuclear peroxiredoxin might be essential in the erythrocyte stage, as neither *P. falciparum* or *P. berghei* knock-out lines could be generated despite several attempts and success with generating tagged nPrx-GFP fusion cell lines [14]. Analysis of the growth of a *P. berghei* Trx-Px1 knockout also suggests that Trx-Px1 is not essential for growth, but *P. falciparum* and *P. berghei* show differences in their lifecycles [41]. There is a possibility that Trx-Px1 is essential during the asexual growth of *P. falciparum* [31]. Expression of Pfl-Cys Prx is elevated during the trophozoite and early schizont stages (when the parasites are maturing during the liver phase) suggesting that this subfamily detoxifies ROS, like those released

during heme metabolism [40]. Despite the important roles of peroxiredoxins, whether inhibitors targeting *Plasmodium* peroxiredoxins will lead to parasite death remains to be determined.

Structural characterization of the 2-Cys Prx enzymes has shown that the C-terminal tail (referred to as the C_R) is essential for stabilizing the octameric/decameric arrangement of peroxiredoxins. When Trx-Px1 (or another 2-Cys Prx) is oxidized to the disulfide form, the C_R loop is unfolded; and structural support of the octamer/decamer/dodecamer interfaces are weakened giving rise to dimer formation (as seen in part when our oxidized PflTrx-Px1 is run on a gel filtration column). When the C_R loop is folded as in the reduced form, the higher order oligomer is favoured because of increased stabilization for the B-type interface which supports oligomerization. The dramatic rearrangement of the C-terminal tail and ensuing changes in stability are clearly demonstrated in our structures from PyTrx-Px1 and PvTrx-Px1 (both oxidized and reduced forms). However, the high concentrations of protein used in crystallizing the enzyme (and also apparently during most of our gel filtration experiments) may account for the trapping of the disulfide forms of these peroxiredoxins in their predicted unfavoured octameric/decameric/dodecameric forms. Factors associated with the oligomeric forms primarily include reduction of the active site disulfide, but also, high or low ionic strength, low pH, high magnesium or calcium concentrations, and overoxidation of the peroxidic cysteine (Cys-SO₂H) [28]. At physiologically relevant concentrations, peroxiredoxins can be expected to exist as dimers awaiting reduction; and upon reduction, the catalytic cycle is complete and the reduced peroxiredoxin oligomerizes [23]. With data from the novel octameric configuration of PyTrx-Px1 studied herein, other specific structural features affecting oligomerization are considered. The molecular basis for octamer formation relies on the hydrogen bond between the dimers which is facilitated by the positioning of the C-terminal tail which in turn rests on interplay between the surface/buried arginine of the NNLA(I/L)GRS-loop and possibly the N-terminal tail. *In vivo* work directed at the further characterization of the different sizes and configurations of the oligomers will be necessary to fully understand the biological implications.

Aside from being antioxidant proteins, 2-Cys Prx (Prx1) has also been implicated in H₂O₂-mediated signal transduction. Eukaryotic 2-Cys Prx enzymes are sensitive to oxidative inactivation, while their bacterial orthologues are robust with respect to overoxidation [30]. The *Plasmodium* 2-Cys Prx enzymes accordingly have the 3 sequence motifs indicative of the H₂O₂-sensitive peroxiredoxins, and as shown herein are structurally identical with respect to sensitivity to H₂O₂ to known

H₂O₂-sensitive 2-Cys Prx enzymes, thus allowing low resting levels of H₂O₂, while permitting higher levels during signal transduction.

The *PyPrx6* structure presented herein has greatly enhanced our understanding of the chemistry of the peroxiredoxins. With a product bound configuration, the residues supporting the leaving group are further understood. There is flexibility for the alkyl chain, but the O_B position is indeed designated for the oxygen of the leaving group. As well, the retention of hydrogen bonds about the active site thiol (even in an oxidized state) indicates that it is poised for further reaction, albeit at a reduced efficiency owing to the reduced activity of an oxidized thiol relative to a thiolate.

Conclusions

Our structural data and mass spectroscopy confirms that *PvTrx-Px1* is H₂O₂-sensitive peroxiredoxin. The characterization of the oligomerization of *PyTrx-Px1* has identified structural features supporting its novel octameric oligomerization. Previously unreported abnormalities of the disulfide bond measurements in some of the Prx crystal structures are brought to the forefront. Finally, a crystal structure with an alcohol bound and the C_P oxidized gives a view to the product bound complex providing insight into leaving group and the susceptibility of some peroxiredoxins to overoxidation. These results enhance our understanding of the structural variations of the peroxiredoxin oligomers and the nature of the catalysis by these remarkable enzymes. Further work will lend insight into the biological implications of the oligomerization and how to exploit the active site features in drug discovery programs.

Methods

Cloning, expression, and purification

Full-length *P. falciparum* Trx-Px1 encoded by PlasmoDB ID: PF14_0368 <http://plasmodb.org/plasmo/>[42] was cloned from *P. falciparum* 3D7 genomic DNA with a His₆-tag with an integrated thrombin cleavage site (MGSSHHHHHHSSGLVPR*GS). Full-length *P. knowlesi* Trx-Px1 encoded by PlasmoDB ID: PKH_126740 was cloned from *P. knowlesi* H genomic DNA with a His₆-tag with an integrated TEV cleavage site (MGSSHHHHHHSSGRENLYFQ*G). Full-length *P. vivax* protein encoded by PlasmoDB ID: PVX_118545 with an *N*-terminal His₆-tag and TEV cleavage site (as above) was cloned from a *P. vivax* Salvador I cDNA library (generously provided by Prof. Liwang Cui of Penn State University). Full-length *P. yoelii* protein encoded by PlasmoDB ID: PY00414 (Trx-Px1) was cloned from *P. yoelii* 17XNL genomic DNA with an *N*-terminal His₆-tag and integrated TEV protease site (as above). Full-length *P. yoelii* protein encoded by PlasmoDB ID: PY04285 (Prx6) was

cloned from *P. yoelii* 17XNL genomic DNA with an *N*-terminal His₆ tag with different integrated TEV protease site (MGSSHHHHHHSSGRENLYFQ*GHM) and *C*-terminal addition (GS). All enzymes were expressed and purified according to methods described previously [26] except for *PyPrx6* which was expressed from Studier auto-induction media [27].

Characterization

All mass spectra were completed on an Agilent LC-MS-TOF (Model #G1969A) running in positive ion mode and integrated with an Agilent 1100 series HPLC using an Agilent Poroshell 300-SB-C3 column for fast binding/elution desalting. All gel filtration experiments were complete on a AKTA purifier chromatography system (GE Healthcare Life Sciences) equipped with a Superdex S200 gel filtration column that was calibrated with 4 samples (bovine thyroglobulin (670 kDa), bovine γ -globulin (158 kDa), chicken ovalbumin (44 kDa), and horse myoglobin (17 kDa) all from Bio-Rad) in 10 mM HEPES, pH 7.4 and 500 mM NaCl.

Crystallization and structure determination

PvTrx-Px1_ox with the His₆-tag intact was crystallized by mixing 1.5 μ L of protein (at a concentration of 8 mg/mL in a buffer of 10 mM HEPES, pH 7.5, 500 mM NaCl) with 1.5 μ L of reservoir solution containing 5% Peg 4 K, 50 mM NaAc, 100 mM NaAc, pH 4.6 in a hanging drop vapour diffusion setup with over 350 μ L of reservoir solution at 18°C in VDXm plates (Hampton Research). Crystals appeared overnight and were flashed-cooled in liquid nitrogen (N_{2(l)}) for data collection. Single wavelength data was collected at a synchrotron source (APS Beamline 17-ID) with a CCD detector (ADSC quantum 210). *PvTrx-Px1_red* with the His₆-tag intact was crystallized using the hanging drop vapour diffusion method in a VDXm plate with 350 μ L of mother liquor at 18°C. 1.5 μ L of the protein solution treated with 5 mM TCEP was mixed with 1.5 μ L of the reservoir solution containing 19% PEG 3350, 150 mM lithium citrate. Crystals appeared overnight. Data for crystals flash frozen in N_{2(l)} was collected at the synchrotron (APS Beamline 17-BM) with a CCD detector (MAR CCD 165 mm). *PyTrx-Px1* with the His₆-tag intact was crystallized at 4.3 mg/mL using the sitting drop vapour diffusion method in a Linbro plate with 300 μ L of mother liquor at 18°C. 1.5 μ L of the protein solution was mixed with 1.5 μ L of the reservoir solution containing 1.6 M ammonium sulfate, 100 mM HEPES, pH 6.8, 200 mM NaAc, 20 mM NaBr, 5% ethylene glycol. Crystals appeared in 3-5 days and were flash frozen in N_{2(l)} with data collected on a Rigaku FRE Superbright rotating anode with an RAXIS IV plate reader. *PyPrx6* with the His₆-tag intact at 15 mg/mL was crystallized by

Table 3 Data collection, phasing, and refinement statistics for the 2H66, 2I81, 2H01, and 3TB2

| Structure | PvTrx-Px1_ox | PvTrx-Px1_red | PyTrx-Px1_ox | PyPrx6-SO ₂ H |
|--|---------------------------|---------------------------|---------------------------|---------------------------|
| PDB Code | 2H66 | 2I81 | 2H01 | 3TB2 |
| Space Group | P21 | C2221 | P422 | C2221 |
| Cell Dimensions | | | | |
| a (Å) | 70.55 | 91.35 | 105.08 | 90.39 |
| b (Å) | 149.59 | 212.57 | 105.08 | 156.84 |
| c (Å) | 131.91 | 115.26 | 41.83 | 178.07 |
| α (°) | 90 | 90 | 90 | 90 |
| β (°) | 104.88 | 90 | 90 | 90 |
| γ (°) | 90 | 90 | 90 | 90 |
| Wavelength | 1.00 | 1.00 | 1.54178 | 0.97939 |
| Resolution (Å) | 50.00-2.48 (2.53-2.48) | 48.28-2.45 (2.55-2.45) | 50.00-2.30 (2.38-2.30) | 20.00-2.30 (2.38-2.30) |
| Unique reflections | 86441 | 41514 | 10337 | 53397 |
| R _{merge} | 0.115 (0.488) | 0.072 (0.460) | 0.153 (0.471) | 0.059 (0.397) |
| I/σ | 20.18 (1.3) | 15.51 (3.56) | 26.55 (4.29) | 24.04 (3.31) |
| Completeness (%) | 99.2 (99.8) | 99.9 (100.0) | 94.6 (99.3) | 99.7 (100.0) |
| Redundancy | 3.4 (3.3) | 6.4 (6.4) | 13.9 (12.8) | 4.3 (4.3) |
| Refinement | | | | |
| Resolution (Å) | 2.5 | 2.45 | 2.3 | 2.3 |
| Number of Reflections | 82137 | 39472 | 10337 | 53397 |
| Test Set Reflection numbers | 4302 | 2107 | 542 | 2858 |
| R _{work} /R _{free} | 0.194/0.232 | 0.217/0.265 | 0.208/0.232 | 0.186/0.207 |
| Number of Atoms (protein/ligand/water) | 13294/0/154 | 7190/0/80 | 1368/0/54 | 7055/89/364 |
| Mean B _{factor} | 45.74 | 38.6 | 37.8 | 40.0 |
| Ramachandran Favored (%) | 95.37 | 95.21 | 91.86 | 98.01 |
| Ramachandran Disallowed (%) | 0.48 | 0.33 | 0.58 | 0.00 |
| RMS deviations | | | | |
| Bond length (Å) | 0.0161 | 0.0083 | 0.0090 | 0.0086 |
| Bond angle (°) | 1.6765 | 1.2250 | 1.2343 | 1.1004 |

means of by hanging drop vapour diffusion in a VDXm plate. The plate was set with 1.5 μL protein plus 1.5 μL buffer in each drop and 350 μL reservoir volume per well. Crystals emerged in 23% Peg 3350, 0.1 M Bis-Tris pH 5.5, 200 mM (NH₄)₂SO₄ and 5% ethylene glycol at 20°C. MAD data from a crystal flash frozen in N₂(l) was collected at the synchrotron (APS Beamline 17-ID) with a CCD detector (ADSC Quantum 4).

Data were processed using the HKL2000 package [43]. Each structure was solved by molecular replacement using modified homology models created with the FFAS03 program [44]. The structures were refined by iterative rounds of manual building in Coot [45] and refinement using refmac5 from CCP4 package [46]. All structures were refined with good statistics and geome-

try, checked with MOLPROBITY [47]. Final statistics and data information for each structure can be found in Table 3. Figures for structural models were created using the Pymol visualization software <http://www.pymol.org>.

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Authors' contributions

WQ analyzed the structural data and solved the structures of PvTrx-Px1_{red} and PyTrx-Px1, as well, identified and assigned previously unknown densities within the PyPrx6 structure. AD collected diffraction data on each of the structures and assisted in the solving of the structures. JCP helped analyze all of the structural data and provided insight into the oligomerization and the characterization of the active site of PyPrx6. AB solved PyPrx6. JM helped solve PyTrx-Px1. AKW solved PvTrx-Px1_{ox}. TH investigated the PvTrx-Px1 and PyTrx-Px1 oligomerization. RH conceived the study, provided insight, and helped draft the manuscript. JDA analyzed the structures, drafted the manuscript and figures, as well as purified, characterized, and crystallized both PvTrx-Px1 structures and PyTrx-Px1. All authors read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

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