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Moonlighting by different stressors: Crystal structure of the chaperone species of a 2-Cys peroxiredoxin

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SUMMARY

2-Cys peroxiredoxins (Prxs) play two different roles depending on the physiological status of the cell. They are thioredoxin-dependent peroxidases under low oxidative stress, and ATP-independent chaperones upon exposure to high peroxides concentrations. These alternative functions have been associated with changes in the oligomerization state from low (LMW) to high (HMW) molecular weight species. Here we present the structures of Schistosoma mansoni PrxI in both states: the LMW decamer and the HMW 20-mer, formed by two stacked decamers. The latter is the first structure of a 2-Cys Prx chaperonic form. Comparison of the structures sheds light on the mechanism by which chemical stressors, such as high H₂O₂ concentration and acidic pH, are sensed and translated into a functional switch in this protein family. We also propose a model to account for the in vivo formation of long filaments of stacked Prx rings.

INTRODUCTION

Oxidative stress is a potential hazard for protein folding. Under redox stress conditions, ATP-dependent foldases are no longer sufficient to cope with protein misfolding and cells employ ATP-independent holdases to counteract the new redox milieu (Kumsta and Jacob, 2009). Holdases are a particular branch of molecular chaperones, which prevent protein...
aggregation by placing client proteins in quarantine. Within this group, redox-regulated chaperones have a fascinating mechanism of action: they sense the redox level of the environment and translate this redox information into quaternary structure changes and functional activation. As an example, upon cysteine residues oxidation, heat shock protein 33 (HSP33) forms dimers and exposes hydrophobic patches at the N-terminus to bind non-native proteins (Winter et al., 2005).

Among the redox-regulated chaperones, peroxiredoxins (Prxs) have recently become the object of increasing attention, due to their ubiquitous distribution and implication in several human pathological conditions (Hofmann et al., 2002; Wood et al., 2003; Flohé et al., 2003; Sayed et al., 2006). Additionally, Prxs are moonlighting proteins: at high \( \mathrm{H}_2\mathrm{O}_2 \) concentrations they act as holdases, while in the absence of oxidative stress they are thioredoxin (Trx)-dependent peroxidases (Jang et al., 2004).

Typical 2-Cys Prxs are the largest and most widely distributed subfamily of Prx proteins (Soito et al., 2011). Their catalytic properties as antioxidant scavengers rely on the peroxidatic Cys (Cys\( _p \)), a redox-active cysteine located in the first turn of helix \( \alpha_2 \) in the N-terminal region of the enzyme. Cys\( _p \) reduces \( \mathrm{H}_2\mathrm{O}_2 \) by releasing \( \mathrm{H}_2\mathrm{O} \) and forming a sulfinic acid derivative, Cys\( _p\)-SOH. Cys\( _p\)-SOH can be converted to a disulfide by the resolving cysteine (Cys\( _r \)) located at the C-terminus of the other subunit in the obligate homodimer (Wood et al., 2003; Winterbourn and Hampton, 2008; Neuman et al., 2009). Oxidation of Cys\( _p \) unwinds the first turn of the \( \alpha_2 \) helix, thus allowing Cys\( _r \) to attack Cys\( _p \). The resulting disulfide bridge is reduced by Trx and/or glutathione, thus completing the enzymatic cycle (Wood et al., 2003; Neumann et al., 2009). The peroxidase activity of 2-Cys Prxs has been shown to be mediated by low-molecular weight (LMW) species. These can be dimers or (do)-decamers, made up of either six or five dimers arranged in a ring-like structure with pseudo six- or five-fold symmetry, respectively. In particular, the (do)-decamers are most abundant in the presence of reduced cysteines, whereas disulfide-bond formation shifts the equilibrium towards the dimers (Barranco-Medina et al., 2009).

At high \( \mathrm{H}_2\mathrm{O}_2 \) concentrations, before resolution by Cys\( _r \), Cys\( _p\)-SOH may react with additional \( \mathrm{H}_2\mathrm{O}_2 \) forming sulfinic acid, Cys\( _p\)-SO\( _2 \)H, and/or sulfonic acid, Cys\( _p\)-SO\( _3 \)H. This overoxidation is associated with a change in function resulting in the loss of peroxidase activity and onset of chaperone activity. This change in biochemical function is coupled to a shift in the oligomerization equilibrium towards higher-order oligomers (HMW; Kunsta and Jacob, 2009; Jang et al., 2004; Barranco-Medina et al., 2009; Moon et al., 2005). In particular, the chaperone activity of yeast cytosolic PrxI has been attributed to different HMW species whose structures, determined by electron microscopy, consist either of rings with 5-fold symmetry or spherically-shaped particles (Jang et al., 2004). Moreover, overoxidation of yeast PrxI to Cys\( _p\)-SO\( _3 \)H shifts the equilibrium towards a HMW form of about 530 kDa, which is proposed to be a double (do)-decamer that is responsible for the chaperone function (Lim et al., 2008). HMW species of several other 2-Cys Prxs have been shown by electron microscopy to be constituted of two or more stacked (do)-decameric rings (Wood et al., 2004; Gourlay et al., 2003). Further, in cultured epithelial cells under high oxidative stress, long filaments of stacked Prx rings have been detected and shown to alert cells of perturbations in peroxide homeostasis, thus determining cell cycle arrest (Gourlay et al., 2003; Phalen et al., 2006).

In vivo, formation of HMW forms is reversible, due to the presence of sulfiredoxin (Srx). Srx specifically reduces Cys\( _p\)-SO\( _2 \)H, restoring the peroxidase activity and favouring the dissociation of HMW complexes (Lowther and Haynes, 2011). Recently, the complex between human PrxI and Srx has been solved by X-ray crystallography (Jönsson et al., 2008). This structure shows that Cys\( _p \) of the Prx dimer is exposed on the protein outer...
surface and the C-terminus is completely unwound and lodged in a hydrophobic groove of Srx. Even though this 3D-complex has provided clues to understand the mechanism of recognition between 2-Cys Prxs and Srx, the principal substrate of Srx under physiological conditions is likely to be the HMW species (Lowther and Haynes, 2011).

At variance with the well characterized peroxidase activity of Prx, neither the atomic structures of HMW forms or the mechanism by which the quaternary assembly is modified and the holdase ability acquired have been elucidated yet. Here we present two distinct 3D crystal structures of cytosolic PrxI from the human parasite *Schistosoma mansoni* (GenBank: Thioredoxin peroxidase I, AF301002): the LMW decamer (herein called SmPrxI-LMW) and the HMW double decamer assembly (called SmPrxI-HMW). The latter represents the first crystallographic structure of a HMW assembly of a typical 2-Cys Prx.

*S. mansoni* is a causative agent of schistosomiasis, a debilitating parasitic disease affecting approximately 200 million people in tropical areas. The worm lacks catalase, and its glutathione peroxidases are not efficient scavengers of hydroperoxides; thus the oxidative stress generated by the host’s immune response is essentially counteracted by Prxs (Sayed et al., 2006; Sayed and Williams, 2004). As previously reported for other typical 2-Cys Prxs (Kumsta and Jacob, 2009; Jang et al., 2004), our experiments demonstrate that SmPrxI-LMW acts as a peroxidase, while SmPrxI-HMW is endowed with holdase activity. Given the high sequence and structural conservation of Prx proteins (Wood et al., 2003), the structure of SmPrxI-HMW may represent a general organization of 2-Cys Prx HMW species. Based on structural comparisons between SmPrxI-LMW and -HMW, we propose a mechanism explaining how chemical modifications of Cys₃₅ trigger the formation of HMW assemblies. Moreover, we propose a model of how the long filaments of stacked rings (Phalen et al., 2006) may be built.

**RESULTS**

**Quaternary Assembly of SmPrxI as a Function of pH and Oxidation State**

Size Exclusion Chromatography (SEC) experiments were carried out to assess the quaternary states of SmPrxI under different redox conditions and at different pH values. We analysed three samples: i) reduced SmPrxI at pH 7.4; ii) reduced SmPrxI at pH 4.2; iii) overoxidized SmPrxI-SO₂H/SO₃H at pH 7.4. The overoxidized SmPrxI derivatives were obtained by Trx-dependent reduction of the Cys₃₅-Cys₅₅ disulfide bonded derivative in the presence of 50 mM H₂O₂, following the previously reported procedure (Sayed et al.; 2006; Sayed and Williams, 2004).

Reduced SmPrxI at pH 7.4 eluted as one homogeneous peak centred around 250 kDa, compatible with a decameric assembly, and at pH 4.2 as a peak at 500 kDa (Figure 1A), indicating the presence of a 20-mer HMW species. The sample containing SmPrxI-SO₂H/ SO₃H at pH 7.4 eluted with almost indistinguishable profiles as a peak with intermediate MW, suggesting the presence of HMW and LMW species in rapid equilibrium relative to the time of the experiment. Each fraction was collected and dialysed in order to assay the peroxidase and chaperone enzymatic activities.

**Peroxidase and Chaperone Activities**

Reduced SmPrxI at pH 7.4 displays peroxidase activity with apparent steady state parameters $K_M(\text{H}_2\text{O}_2)=12\ \mu\text{M}$ and $k_{cat}=1.1\ \text{s}^{-1}$, at fixed concentrations of *E. coli* Trx (1.6 μM) and Trx-reductase (200 nM). The holdase activity of this form is negligible at pH 7.0 (Figure 1B), even in the presence of 10-fold excess of reduced SmPrxI. It was not possible to increase the molar ratio over 10:1 SmPrxI-monomer:citrate synthase (CS) due to the precipitation of reduced SmPrxI in the assay conditions (*i.e.*, absence of salts and 43°C).
These conditions are required to allow measurements of chaperone activity, since CS precipitation was prevented by even very small amounts of salts.

At pH 4.2, under reducing conditions, no peroxidase activity of SmPrxI could be detected, but a holdase activity, weaker than that observed for overoxidized Prx at pH 7.0, was present (Figure 1C). The apparent stoichiometry under these conditions is higher than for the overoxidized SmPrxI, approaching 20:1 SmPrxI-monomer:CS, possibly due to a reduction in affinity (Figure 1C).

Overoxidized SmPrxI-SO$_2$H/SO$_3$H at pH 7.0 was shown to be active as a holdase chaperone at stoichiometric concentrations of the SmPrxI-monomer with CS (Figure 1B). In the same conditions, peroxidase activity was not detected, as previously reported (Sayed et al., 2006).

At pH 4.2 SmPrxI-SO$_2$H/SO$_3$H was shown to be active as a holdase, even though to a lesser extent than at pH 7.0 (Figure 1C). The molar ratio SmPrxI-monomer:CS could not be increased over 10:1 due to the low solubility of SmPrxI-SO$_2$H/SO$_3$H in the assay conditions (see above).

Even though the holdase assay is semi-quantitative, it is possible to roughly estimate the relative activities of SmPrxI under the different conditions tested. Taking the ratio SmPrxI-monomer:CS 10:1 as reference condition, the protection activity of the different SmPrxI forms against CS precipitation is the following: 100% for overoxidized SmPrxI at pH 7; 50% for reduced SmPrxI at pH 4.2; 20% for overoxidized SmPrxI at pH 4.2; close to 0% for reduced SmPrxI at pH 7. This underestimates the actual holdase activity of overoxidized SmPrxI at pH 7, which reaches its asymptote at a lower SmPrxI-monomer:CS ratio than 10:1. Moreover, overoxidized SmPrxI increases the lag time of CS precipitation at pH 7, whereas at pH 4.2 the lag time varies scarcely, if at all.

**Crystal Structures of SmPrxI**

Following the results of SEC experiments, we set up crystallization trials in the experimental conditions required to stabilize either the LMW or the HMW species. We obtained two structures by molecular replacement: a decameric assembly (SmPrxI-LMW), under conditions similar to those yielding the SEC fraction showing peroxidase activity (i.e., reducing environment and pH 7.6); a double decamer (SmPrxI-HMW), under the same conditions yielding the single SEC peak with chaperone activity (i.e., reducing environment and pH 4.2).

**SmPrxI-LMW: the Single Decamer**

The 3.0 Å resolution electron density map of reduced SmPrxI-LMW allowed us to fit almost all 185 amino acids for each of the 10 polypeptide chains, which were identified with letters from A to J (see Tables 1 and S1). The overall structure of SmPrxI-LMW (Figure 2A–B) consists of a decameric ring with 52 point group symmetry and is superimposable with that of homologous proteins crystallised in the reduced form (Hall et al., 2009). The ring is 4.5 nm thick and has outer and inner diameters of 12 and 6 nm, respectively.

Each subunit shows the same secondary structure arrangement occurring in all typical 2-Cys Prxs solved to date (Wood et al., 2003). The overall topology (Figure S1) consists of a modified Trx fold with a seven-stranded β-sheet (β1–β7) sandwiched by: two α-helices (α1 and α5) and one short β-hairpin on the internal face of the decameric ring; four α-helices (α2, α3, α4 and α6) on the external face (Figure 2A–B). The two protomers of each obligate homodimer are correlated by a pseudo 2-fold axis perpendicular to the plane containing the extended β-sheet (Figure 2C) that, in turn, forms an angle of 27° with respect to the pseudo 5-fold symmetry axis. Both the monomer-monomer interface within each dimer and the
dimer-dimer interface (called B- and A-interfaces, respectively; Aran et al., 2009) resemble those observed in the structures of other typical 2-Cys Prx (Aran et al., 2009; Barranco-Medina et al., 2008; Nelson et al., 2011).

The intra-dimeric B-interface (Figures 2B-C and S2) is symmetrical and is contributed by: i) the antiparallel $\beta^7$-$\beta^7'$ strand interaction from each of the two subunits, through which the continuous $\beta$-sheet of the dimer is formed (the apex indicates structural elements from the interacting monomer); ii) on the internal face of the ring, the N-terminal region of one subunit interacts with the $\beta$-hairpin and part of the $\alpha^5'$ helix of the other subunit; iii) on the external face of the ring, the $\beta^7$-$\alpha^6$ loop interacts with the $\alpha^6'$ helix, and the interacting $\alpha^2$ helix and C-terminal region of the other subunit comprise the active site.

The active-site (Figure 2C) is in the fully-folded (FF) conformation (Wood et al., 2003): residues 47–50, including the reduced Cys$_p$ (Cys48), form the first turn of the $\alpha^2$ helix, and the C-terminal tail (residues 165–185), containing Cys$_r$ (Cys169), is folded in a hairpin-like structure.

The sulphur atom of Cys$_p$ is engaged in an electrostatic interaction with the NH$_2$ atom of Arg124 (3.6 Å), but it is too distant from the sulphur atom of Cys$_r$ to form a disulfide bridge with it (~13 Å in the A/B dimer). This implies that, during the catalytic cycle, a structural rearrangement must occur to allow the formation of a disulfide bridge between Cys$_r$ and Cys$_p$ (Wood et al., 2003).

The inter-dimeric A-interface is also symmetrical, and involves contacts between: i) the $\alpha^4$ helix of one subunit and $\alpha^4'$ helix and $\beta^3'$-$\alpha^2'$ loop of the adjacent subunit, on the external side of the decameric ring; and ii) the $\beta^5$-$\alpha^5$ loop of one subunit and the short $\beta$-hairpin of the other subunit on the ring internal side (Figures 2B–C and S3).

**The Structure of SmPrxI-HMW: Two Stacked Decamers**

Reduced SmPrxI-HMW crystallized at pH 4.2 is a 20-mer arranged in two stacked decamers (Figure 3). Since both decamers forming the SmPrxI-HMW have a 52 point group symmetry, they can stack one on top of the other without a preferential orientation. The 3 Å resolution electron density map was clearly visible for residues 3–46 and 50–165 for all 20 subunits, while density for residues 47–49 is missing in some subunits (Table S1). Only a few residues of the C-terminal tail could be fitted in the electron density map and they did not assume regular secondary structures.

The interface between the two decameric rings stacked on top of each other in SmPrxI-HMW is R (ring). The R-interface (Figures 3C–D and S4) comprises two regions: (i) the $\alpha^2$ and $\alpha^6$ helices of subunit B (upper ring) are in contact with the equivalent helices of subunit K (lower ring); and (ii) strand $\beta^2$ of subunit B interacts with the equivalent strand $\beta^2$ of subunit M (lower ring). If we compare each single decamer to a cogwheel, the secondary structure elements present in these regions may resemble paws through which the two rings are interlocked (Figure 3D–E). The observed joints are made possible by a rotation of 18° around the five-fold symmetry axes of one decamer with respect to the other (Figure 3B). The four helices $\alpha^2$ and $\alpha^6$ of the adjacent B and K monomers are oriented in such a way that all of their carbonyl groups, and hence the helix dipoles, point towards the R-interface. This creates a negatively charged area centred on residues Lys164 and His165, located at the end of $\alpha^5$ helix of both B and K subunits (Figure 3D). The dipoles of the peptide bonds are expected to raise the $pK_a$ of His165 at the end of the $\alpha^6$-helix. Thus, the positively charged side-chains of Lys164 and His165 are expected to stabilize the negative charge of the helix dipoles (Sali et al., 1988), thus contributing to the stabilization of the stacked decamers. On
the same pawl, Glu21 and Lys23, belonging to the β2 strands, stabilize the interface between monomers B and M with two polar contacts (Figure 3D).

Comparison Between SmPrxI-LMW and SmPrxI-HMW

The monomers constituting SmPrxI-HMW have the same overall fold as those in SmPrxI-LMW. Optimal superposition of the Cα atoms of residues in the common regions of LMW and HMW monomers (i.e., 3–46, 50–165, see Table S1) provided root mean square deviation (RMSD) values ranging between 0.8 and 1.0 Å (Table S2). The highest differences occur in the active site. The FF conformation observed in SmPrxI-LMW becomes locally unfolded (LU) in SmPrxI-HMW (Figure 4A), and electron density for a large number of C-terminal residues is missing in the electron density map (Table S1), similarly to what previously reported for other Prx structures in the dimeric form (Wood et al., 2003; Lowther and Haynes, 2011; Jönsson et al., 2008). Furthermore, unwinding of the first turn of the α2 helix in SmPrxI-HMW leads to exposure of Cys_p (Figure 4A and Movies S1 and S2).

The homodimer represents both the minimal structural unit of Prxs and the smallest unit that is populated during the catalytic cycle. Superposition of the SmPrxI-LMW and SmPrxI-HMW dimers in the conserved regions provided RMSD values ranging between 0.8 and 1.0 Å, very similar to those obtained for the monomers (Table S2). The main change associated with the structural transition from LMW to HMW form occurs at the B-interface. Indeed, the pattern of contacts between the α2 helix of one monomer and the C-terminus of the adjacent monomer at the B-interface is almost completely different between the two forms (Figure S2 and Movies S1 and S2). The two structural features distinctive of HMW dimers (i.e., unfolded C-terminus and unwound α2 helix) are strongly correlated, since a folded C-terminus would clash with the unwound α2′ helix at the B interface.

The HMW and LMW decamers differ because of both tertiary and quaternary structural changes. The RMSD values calculated after optimal superposition of the LMW and HMW forms in the common decamer regions are, on average, 1.1 Å, a value that is slightly but significantly higher than the average of those calculated after optimal superpositions of both dimers and monomers (Table S2). The local unfolding of the β3-α2 loop (Figure 4A) results in a pressure on the α3 helix of the adjacent dimer, which, in turn, determines a slight increase of the A-interface in the region near Cys_p. However, the A-interface is perturbed to a lesser extent than the B-interface (Figure S3).

The quaternary structural changes affecting the relative position of the dimers within LMW and HMW decamers can be visualized using as a marker the distance between residues belonging to the α6 helix, whose conformation does not change in the two forms. Figure 4C shows the α6 helices of monomers A (within the A/B dimer) and C (within the adjacent C/D dimer) after optimal superposition of the A/B dimers belonging to SmPrxI-LMW and SmPrxI-HMW, respectively (highlighted by an arrow in Figure 4B). In SmPrxI-HMW, the distance between the Cα atoms of the Phe161 residue within the α6 helix of any monomer and the equivalent residue in the adjacent dimer is on average shorter by 1.5 Å than in SmPrxI-LMW (Figure 4C). In view of the strong geometrical constraints imposed by the closed-ring arrangement of both SmPrxI-LMW and SmPrxI-HMW decamers, and the observed structural conservation of the A-interfaces, we believe that the structural changes highlighted in Figures 4B–C are significant.

Superposition of SmPrxI-LMW to each decamer of SmPrxI-HMW revealed that unfolding of the C-terminal region downstream of the α6 helix of each subunit is required to allow the stacking of the decamers, as it removes a protrusion on the ring surface that otherwise would clash with the other ring at the R-interface (Figure S4 and Movie S1).
In summary, the quaternary changes at the R-interface appear to be related to: i) tertiary changes (unfolding of the C-terminus and unwinding of the first turn of \( \alpha_2 \) helix), allowing the direct ‘vertical’ interaction between monomers B and K (Figure 3C–E); ii) propagation of tertiary changes ‘horizontally’ through the A-interfaces within the single ring. As an example, the \( \beta_3-\alpha_2 \) loop (residues 45–50, Figure 4A) of monomer B protrudes towards the \( \alpha_3 \) helix of monomer C; this in turn moves as a rigid body (see RMSD values in Table S2) making the \( \alpha_6 \) helix act as a pawl for interlocking the decamer cogwheels (Figure 4B–C).

**DISCUSSION**

Prxs sense the redox environment by changing the oxidation state of the catalytic cysteine thiol to a sulfinic/sulfonic acid (-SO\(_2\)H/-SO\(_3\)H). This chemical modification is at the basis of the moonlighting behaviour of Prxs, in that it induces the formation of HMW complexes, which are competent for the molecular chaperone activity and are implicated in cell signalling (Jang et al., 2004; Phalen et al., 2006; Hall et al., 2009). Neither the 3D structure of these HMW species nor their mechanism of assembly have been reported to date. Comparison of the structures of SmPrxI-LMW (the decameric ring) and SmPrxI-HMW (the double decamer) here presented reveal the quaternary transitions by which Prxs sense and respond to chemical stressors (high oxidant concentration and acidic pH), switching to HMW complexes endowed with chaperone activity. Moreover, they suggest where unfolded proteins may be bound by the HMW forms, and how these may pile up to form long filaments.

The following events are involved in the activation of SmPrxI and of typical 2-Cys Prxs: (i) overoxidation of the sulphur atom of Cys\(_p\), which causes destabilization of the \( \alpha_2 \) helix; (ii) concerted tertiary variations, i.e. unwinding of the first turn of the \( \alpha_2 \) helix; unfolding of the C-terminal tail; protrusion of the \( \beta_3-\alpha_2 \) loop towards the nearby dimer; and (iii) quaternary structural changes leading to the formation of the HMW species (Movies S1 and S2).

The first important questions to address are why both oxidative and acidic stimuli should cause the same tertiary and quaternary structural changes in SmPrxI, and whether SmPrxI-HMW crystallized at acidic pH is representative of the structure acquired by the same protein or other 2-Cys Prxs when exposed to oxidative stress. It is known that under oxidative stress SmPrxI, as well as other eukaryotic 2-Cys Prx, switches its structure and its function (from LMW/peroxidase to HMW/chaperone), both *in vitro* and *in vivo* (Kumsta and Jacob, 2009; Jang et al., 2004). SEC experiments here presented indicate that the HMW species is populated both at pH 4.2 and after H\(_2\)O\(_2\) treatment at physiological pH. We propose that overoxidation and protonation of Cys\(_p\), which is at the beginning of helix \( \alpha_2 \), trigger unwinding of this helix through two different mechanisms. In the case of overoxidation, the increased volume and hydrophilicity of Cys\(_p\)-SO\(_2\)H/SO\(_3\)H are likely to favour its removal from the compact hydrophobic pocket formed by Tyr40, Pro41, Ala42, Thr45 and Pro49 that, together with Arg124, surround the reduced Cys\(_p\) in the LMW species. Once Cys\(_p\) is removed from its pocket, the first turn of the \( \alpha_2 \) helix can unwind and the \( \beta_3-\alpha_2 \) loop assume a LU conformation (Figure 4A). This is favoured by the presence of a proline residue adjacent to Cys\(_p\) (Pro49), highly conserved in typical 2-Cys Prxs (Hofmann et al., 2002), which reduces the propensity of the first turn of the \( \alpha_2 \) helix to retain its secondary structure. In the case of acidic pH, a very similar structural change would be caused by the loss of the ionic bond between Cys\(_p\) and Arg124. Indeed in all typical 2-Cys Prxs, Cys\(_p\) is in contact with a conserved arginine residue, homologous to Arg124 of SmPrxI, which lowers the pKa of the catalytic residue from 8.5 to 5–6 (Winterbourn and Hampton, 2008; Neumann et al., 2009). Accordingly, in SmPrxI, one Cys residue, likely to be Cys\(_p\), has a pK\(_a\)=6.0 (Figure S5). At pH 4.2 this residue is fully protonated, and the ionic bond with Arg124 cannot be formed. We propose that the versatile chemical nature of the
pocket containing Cys\textsubscript{p} evolved to trigger the same local unfolding of the active site in response to stimuli as different as overoxidation and acidification.

Unwinding of the first turn of helix α2 has been previously observed in the structure of the dimeric form of human PrxI (HsPrxI) bound to human sulfiredoxin (HsSrx), the enzyme that restores the active form of HsPrxI by reducing the overoxidized Cys\textsubscript{p} (Jönsson et al., 2008). As a result, Cys\textsubscript{p} is exposed towards the Srx active site, similarly to what observed for the Cys\textsubscript{p} to Asp mutant in a different structure (Jönsson et al., 2008; Jönsson et al., 2009). The similarity between the A/B dimer of SmPrxI-HMW and the isolated dimer of HsPrxI bound to Srx, in particular in the β3-α2 loop and α2 helix region, is highlighted in Figure 5A. These observations suggest that unwinding of the first turn of the α2 helix and the increased accessibility of Cys\textsubscript{p} to solvent and/or interacting protein may be a general structural mechanism by which 2-Cys Prxs respond to chemical stressors.

The next point to address concerns the relationships between the tertiary structural change just described and the other observed changes in tertiary structure (i.e., the unfolding of the C-terminal tail and the concomitant protrusion of the β3-α2 loop towards the nearby dimer), and between these tertiary events and the change in the quaternary assembly from LMW to HMW. Superposition of SmPrxI-LMW and SmPrxI–HMW showed that the C-terminal region cannot assume the FF hairpin-like conformation observed in the LMW form when the first turn of α2 helix is unwound. As a consequence, unwinding of this helix necessarily triggers a conformational change of the C-terminus, and vice versa. Accordingly, in the structure of HsPrxI in complex with HsSrx, where helix α2 is unwound, the C-terminal arm of HsPrxI is unfolded and embraces HsSrx (Figure 5A; Jönsson et al., 2008; Jönsson et al., 2009).

Unwinding of the C terminus, in turn, results in: (i) changes in the surface of the ring decamer at the R-interface, allowing its interaction with another Prx decamer (Figures 3C–D and S4A); (ii) recognition of proteins with sulfiredoxin activity that restore the peroxidase activity through reduction of Cys\textsubscript{p}-SO\textsubscript{2}H and disassembly of the HMW species (Jönsson et al., 2008; Jönsson et al., 2009); (iii) recognition and binding of unfolded proteins.

We sequentially ordered the events above as local-tertiary-quaternary, but they might occur in a concerted way: on the one hand, oxidation/protonation of Cys\textsubscript{p} is driven by a large free energy change; on the other hand, the aggregation of FF dimers to the LMW or of LU dimers to HMW occurs at relatively low concentrations (see Experimental procedures), therefore it must be associated with a negative free energy. Consequently, both local (oxidation/protonation) and quaternary (aggregation) events are likely to contribute to the stabilization of the tertiary structural changes.

Tertiary changes analogous to those observed in SmPrxI-HMW, are known to take place when Cys\textsubscript{p} is disulfide bridged to Cys\textsubscript{r} (located in the C-terminal region) during the enzymatic cycle of Prxs. However, when this disulfide bridge is formed, the dimeric form of the protein is stabilized, while HMW states are not populated. Superposition of two copies of the eukaryotic Rattus norvegicus PrxI (RnPrx; Hirotsu et al., 1999), another 2-Cys Prx, to two adjacent dimers of SmPrxI-HMW (Figure 5B) shows that the presence of the disulfide bridge in RnPrx: (i) constrains the loop β3-α2 in a different orientation from that observed in SmPrxI-HMW, resulting in a perturbation of the contacts between the α3 helices that is likely to hamper formation of the A-interface; and (ii) prevents the complete unfolding of the C-terminus. These features are likely to block the propagation of the local unfolding of α2 helix and C-terminus, both horizontally through the A-interface and vertically at the R-interface, in the disulfide forms of Prx dimers.
The structure of the SmPrxI-HMW chaperone here presented resembles that of chaperonins such as HSP60 and its bacterial homologue GroEL. These are large complexes of two heptamer rings stacked on each other, enclosing a cavity where the correct folding of misfolded proteins takes place. The observed similarity is in agreement with the results of a recent study proposing that the most likely origin of the chaperone fold is the Prx-like fold (Dekker et al., 2011).

It has been hypothesized that the HMW species of Prxs bind unfolded substrates via unfolded hydrophobic regions, in analogy with other stress-regulated chaperones (Kumsta and Jacob, 2009). Although the binding site for unfolded polypeptides in the structure of SmPrxI-HMW cannot be identified with certainty, we propose a role for the C-terminus. This is unstructured and disordered in SmPrxI-HMW, but not in SmPrxI-LMW, which has no holdase activity. Additionally, the C-terminus of HsPrxI has been shown to interact with hydrophobic patches on the surface of partner proteins (Jönsson et al., 2008). Moreover, the presence of 20 unfolded C-termini in the 20-mer HMW, five at each rim and ten radiating from the equatorial portion of the structure, is consistent with the stoichiometry of the SmPrxI-monomer:CS ratio observed in our experiments.

Multi-ring stacking of 2-Cys Prxs has been observed both in vivo and in vitro by electron microscopy (Gourlay et al., 2003; Phalen et al., 2006; Wood et al., 2002; Harris et al., 2001), which has been correlated with cellular signalling. All Prx decamers intrinsically possess the structural information to lock with one another. Interlocking of SmPrxI decamers is mediated by two series of five pawls represented by the conserved α2 and α6 helices and β2 strands (Wood et al., 2003). Due to the pseudo 2-fold symmetric structure of the dimers, the pawls lean out above and below the plane of the ring. Like SmPrxI-LMW, SmPrxI-HMW (Figure 3C–E) has ten pawls available for interlocking with other decamers, leading to the formation of multi-ring structures. In these high-order assemblies the α2 and α6 helices longitudinally span through the rings and a discontinuous β-sheet, comprising the whole central β-sheet of the dimer, wraps the supramolecular structure in a left-handed helicoidal embrace (Figure 6). The external and internal diameters of this filament are in good agreement with those calculated by electron microscopy (Wood et al., 2002; Harris et al., 2001).

**Pathophysiological role**

The symptoms of schistosomiasis are due to egg secretion by the female worm residing in the blood stream. To leave the human body and close the parasite life cycle, the eggs both induce inflammation to penetrate the host tissues and, at the same time, prevent the formation of large granuloma that might hamper their escape (Green and Colley, 1981; Stadecker, 1992). We suggest that SmPrxI, one of the few proteins secreted by the eggs (Cass et al., 2007; Williams et al., 2001), is required to maintain this delicate equilibrium. The pH at a site of inflammation is below 6.0 (Poon et al., 2002); in addition, the granuloma contains immune cells that generate oxidative stress (neutrophils, eosinophils). Under these conditions SmPrxI might be in the stress-induced chaperone form here reported and may help controlling excessive precipitation of proteins associated with granuloma formation.

**EXPERIMENTAL PROCEDURES**

**Protein Preparation**

Recombinant 6-histidine-tagged SmPrxI, cloned from adult *S. mansoni* cDNA, was expressed in *Escherichia coli* and purified as previously described (Sayed et al., 2006; Sayed and Williams, 2004).

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To obtain the overoxidized derivative, recombinant 1 μM SmPrxI was incubated in 50 mM potassium phosphate, 2 mM EDTA pH 7.0 with 1 mM NADPH, 200 nM *E. coli* Trx-reductase, 4 μM *E. coli* Trx, and 50 mM H$_2$O$_2$ at 25°C for 3 hours in aerobic conditions. The mixture was then dialyzed by ultrafiltration using concentrator tubes (Amicon) with a 50 kDa cut-off, concentrated to 1 mg/mL, and used for further experiments.

**Size Exclusion Chromatography**

Size exclusion chromatography (SEC) was performed on HPLC (LabService Inc.) at 25°C with a flow rate of 0.5 mL/min using a Tosoh TSK G3000PWxl column equilibrated with phosphate buffered saline (PBS) and 100 mM sodium-phosphate/sodium-citrate either at pH 7.4 or at pH 4.2. Molecular weight standards were DNA-binding protein (DPS) from *Listeria innocua* (230 kDa), ferritin from *Pyrococcus furiosus* (500 kDa) and cytochrome C from horse heart (12.4 kDa). Three different protein samples were run on the column: 1.5 mg/mL of reduced SmPrxI in PBS/Na-citrate pH7.4; 1.3 mg/mL of reduced SmPrxI in PBS/Na-citrate pH4.2; 1.0 mg/mL of SmPrxI-SO$_2$/SO$_3$H in PBS/Na-citrate pH7.4. Each SEC run was repeated three times.

**Assays of Chaperone and Peroxidase Activities**

The elution fractions of reduced SmPrxI and SmPrxI-SO$_2$/SO$_3$H were dialyzed against 50 mM Hepes pH 7.0 and then concentrated and assayed for the peroxidase and holdase activities. Trx-dependent peroxidase activity of Prxs linked to NADPH oxidation was monitored by the decrease in absorbance at 340 nm as previously described (Sayed et al., 2006). The holdase activity was measured by using 0.8 – 1 μM porcine heart CS (Jang et al., 2004; Cremers et al., 2010). Turbidity, induced at 43°C, in the presence or absence of SmPrxI samples was monitored at 360 nm in a spectrophotometer (Hewlett Packard, USA) equipped with a thermostatic cell holder both at pH 7.4 and 4.2. Each experiment was repeated three times.

**Crystallization of SmPrxI-HMW and SmPrxI-LMW**

The reduced form of SmPrxI was obtained by treating a solution of the enzyme (7 mg/mL) in PBS with 5 mM DTT. Crystallization trials were set up using the vapour diffusion method by both the hanging and sitting drop procedures. 1 μL of the reservoir solution (100 mM potassium citrate, 20% PEG 3350 and 20 mM DTT) was mixed with an equal volume of the protein solution. In the case of SmPrxI-HMW, reducing agents were not used. SmPrxI-HMW (7 mg/mL) was crystallized in 100 mM PBS/citrate buffer pH4.2, 200 mM NaCl and 20% PEG 8000. Both forms of the crystals were cryoprotected by adding 30% PEG 200 to the mother liquor and flash cooling in liquid nitrogen.

**X-ray Data Collection, Processing and Refinement**

Diffraction patterns of both SmPrxI-LMW and SmPrxI-HMW were collected at 100K, λ=0.918Å at the beamline ID14.1 of BESSYII (Berlin, Germany). For SmPrxI-LMW, data processing and reduction were performed using MOSFLM and SCALA in the CCP4 suite (Battye et al., 2011; Evans, 2011; Winn et al., 2011). The crystal belonged to space group P1. The unit cell contained 10 monomers per asymmetric unit with an estimated solvent content of 52%. Molecular replacement was performed with MolRep (Vagin and Teplyakov, 2010), using human PrxIV (PDB code 2pn8) as search model. Model building and refinement were performed using Coot (Emsley et al., 2010) and REFMAC5 (Murshudov et al., 2011) respectively. Data processing and reduction of SmPrxI-HMW crystal were performed using HKL2000 (Otwinowski and Minor, 1997). PHENIX/Xtriage (Adams et al., 2010) revealed that the diffraction pattern belongs to space group P2$_1$ with pseudo-merohedral twinning (Yeates, 1997). The unit cell contained 20 monomers per asymmetric unit.
units with an estimated solvent content of 59.3%. Molecular replacement was performed using SmPrxI-LMW as search model. Model building and refinement were performed using Coot (Emsley et al., 2010) and PHENIX/Refine (Adams et al., 2010), respectively. The latest step of refinement revealed a twinning fraction of 0.5. NCS averaging on both structures was performed as a control and no differences were observed in the statistics and maps. Prior to deposition in the PDB, both structures were validated using MolProbity (Davis et al., 2004). In SmPrxI-LMW: 96.8% of residues are in the favoured regions, 3.2% in the allowed regions. In SmPrxI-HMW: 92.4% in favoured regions, 7.6% in allowed regions. A summary of data collection and refinement statistics are presented in Table 1. Figures were produced with PyMOL (Schrödinger LLC) and CCP4MG (Potterton et al., 2004).

**Structure Analyses**

Secondary structure assignment has been performed with DSSP (Kabsch and Sander, 1983) and solvent accessible surface area calculations with Naccess (http://www.bioinf.manchester.ac.uk/naccess/). Structure visualization and analyses were performed using the programs InsightII (Accelrys Inc.), PyMOL (Schrödinger LLC) and in house-built programs. The distance thresholds used for inter-atomic contacts were 4.0 and 3.6 Å for polar and hydrophobic atoms, respectively.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

**Acknowledgments**

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**ABBREVIATIONS**

- **Prx** Peroxiredoxin
- **Trx** Thioredoxin
- **Trx-reductase** thioredoxin reductase
- **CS** citrate synthase
- **LMW** low molecular weight
- **HMW** high molecular weight
- **PBS** phosphate buffer saline
- **DTT** dithiothreitol
- **SEC** size exclusion chromatography
- **EDTA** Ethylenediaminetetraacetic acid
PEG  polyethylene glycol
FF   fully folded
LU   locally unfolded

References


Structure. Author manuscript; available in PMC 2013 March 7.
Figure 1. Quaternary Assembly and Chaperone Activities of SmPrxI

(A) SEC of SmPrxI was performed using a TSK G3000PW XL column equilibrated with a mix of PBS and Na-citrate. Three different protein samples were fractionated on the column: (▲) reduced SmPrxI in PBS/Na-citrate pH 7.4; (☐) reduced SmPrxI in PBS/Na-citrate pH 4.2; (●) SmPrxI-SO\(_2\)H/SO\(_3\)H in PBS/Na-citrate pH 7.4. Horse heart cytochrome c was used as an internal reference (indicated by the arrow at 12.4 kDa).

(B) The chaperone activity of reduced and overoxidized SmPrxI was measured by the temperature-induced aggregation of citrate synthase (CS) from porcine heart (Sigma-Aldrich) at 43 °C in 50 mM Hepes pH 7.0: CS 1 µM (control) (●); CS 1 µM + reduced SmPrxI-monomer 10 µM (○); CS 1 µM + SmPrxI-SO\(_2\)H/SO\(_3\)H-monomer 0.5 µM (▲), 1 µM (☐), 2 µM (●).

(C) The chaperone activity of reduced SmPrxI in 50mM Na-citrate pH 4.2 was measured using the same procedure as in Panel B: CS 0.8 µM (control) (▲); CS 0.8 µM + SmPrxI-SO\(_2\)H/SO\(_3\)H-monomer 10 µM (+); CS 0.8 µM + reduced SmPrxI-monomer 10 µM (×), 20 µM (☐); 50 µM (●).
Figure 2. Crystal Structure of SmPrxI-LMW

(A) Ribbon representation of SmPrxI-LMW structure (top view). The pentameric arrangement of the dimers is highlighted. The internal and external diameters of the decamer are shown, together with the B-interface (A/B dimer) and A-interface (B/C monomers). Dimers are indicated by five different colours. Within each dimer, the two monomers have different shades. See also Figure S1.

(B) Ribbon representation of SmPrxI-LMW structure (front view). The A/B dimer and the thickness of the decamer are indicated. See also Figure S2.

(C) Overall fold of one SmPrxI-LMW dimer. The catalytically active Cysₚ (Cys48), within the first turn of the α2 helix, and Cysᵣ (Cys169), in the C-terminal tail, together with Arg124 (responsible for lowering the pKₐ of Cysₚ) are displayed as sticks. The active site is in the...
fully-folded (FF) conformation (see text). Structure elements that undergo conformational changes between SmPrxI-LMW and SmPrxI-HMW (i.e., α2 and α6 helices, β2 strand, C-terminus), together with the A-interface (helix α4 and β3–α2 loop) and B-interface (β7 strands), are indicated. See also Figure S3.
Figure 3. Crystal Structure of SmPrxI-HMW

(A) Overall surface of SmPrx-HMW. The HMW species is constituted by two decamers (grey and orange, respectively) locked one above the other along the five-fold symmetry axis.

(B) The upper and lower decamer of SmPrxI-HMW (view from top) appear to be rotated by 18° around the five-fold symmetry axis. To highlight this feature, the Cα atoms of residue Phe161 of adjacent monomers within each of the two decamers are joined by grey and orange lines, respectively.

(C) Secondary structure elements forming the pawls allowing the locking of the cogwheel rings at the R-interface, namely the α2 and α6 helices and the β2 strand from the B monomer (upper ring) and K and M monomers (lower ring), are shown in dark grey and orange, respectively. The thickness of the double-decamer is also indicated. See also Figure S4.

(D) Magnification of the contacts between the secondary structure elements of the two stacked decamers at the R-interface. The α2 and α6 helices create a negatively charged region due to the orientation of their carbonyl groups. The positively charged residues, Lys164 and His165 at the end of α6, stabilize this architecture. The β2 strands are joined by polar contacts between Glu21 and Lys23 of chain B and Glu21 of chain M. Structural elements of the B monomer are indicated with apexes.

(E) The cogwheel model. Schematic representation of the exploded linearized rings of SmPrxI-HMW. The secondary structure elements (α2, α6 and β2) implicated in the self-assembly of the HMW species are shown.
Figure 4. Tertiary and Quaternary Structural Changes Between SmPrxI-LMW and SmPrxI-HMW

(A) Superposition of the A/B dimers of SmPrxI-LMW (blue) and SmPrxI-HMW (orange) highlighting the changes in the active sites. The first turn of the α2 helix and the β3-α2 loop in the fully folded (FF) conformation of SmPrxI-LMW and in the locally unfolded (LU) conformation of SmPrxI-HMW are displayed as cartoons. The side-chains of Cysp of both structures are displayed as sticks. The C-terminal extension is visible in the electron density map only in SmPrxI-LMW.

(B) Changes in the quaternary structure between adjacent dimers of SmPrxI-LMW and SmPrxI-HMW decamers (view from the top). Distances between the Ca atoms of Phe161 in monomers A, C, E, G and I are shown as dashed lines after optimal superposition of the A/B dimers (highlighted by an arrow) of SmPrxI-LMW (cyan) and SmPrxI-HMW (orange). The displacement of the α6 helices increases with distance from the best-matched dimers.

(C) Changes in the quaternary structure between two adjacent dimers (view from the side of the ring). Only the A/B and C/D dimers of both SmPrxI-LMW (blue) and SmPrxI-HMW (orange) are displayed. The distances between the Ca atom of Phe161 within the α6 helix of monomer A and the equivalent atom of monomer C in SmPrxI-LMW and SmPrxI-HMW are shown as dashed lines, as a marker of the quaternary changes between the two structures.
Figure 5. Comparison With Other 2-Cys Prxs

(A) Comparison between the tertiary structures of the A/B dimer of SmPrxI-HMW (orange) and human PrxI (HsPrxI, magenta) in complex with human HsSrx (green; PDB ID: 2RII; Jönsson et al., 2008). SmPrxI-LMW (blue) and the hydrophobic surface of the HsSrx region involved in HsPrxI recognition are also shown (Jönsson et al., 2008 and 2009). The β3-α2 loops of SmPrxI-HMW and HsPrxI have a very similar conformation, which is different from that adopted by the same loop in SmPrxI-LMW, as highlighted by the relative position of the homologous residues Phe46 in SmPrxI and Phe50 in HsPrxI.

(B) Superposition between two adjacent dimers of SmPrxI-HMW (orange cartoons) and two copies of the oxidized dimer of Rattus norvegicus HBP23 (RnPrx, violet cartoons; PDB ID: 1QQ2). Equivalent aromatic residues contributing to the A-interface of SmPrxI-HMW and to the modelled interface in RnPrx are displayed as sticks.
Figure 6. Theoretical Model of the Long Filament Structure

Surface representation of the modelled filamentous-like structure of SmPrxI. This model has been built by joining two copies of the SmPrxI-HMW structure after optimal superposition of the lower decamer (chains K–T) of the first copy to the upper decamer (chains A–J) of a third copy of the molecule and of the upper decamer of the second copy (chains A–J) to the lower decamer (chains K–T) of the third copy. The α-helices (cyan) belonging to different decamers, which vertically span the filament, and a discontinuous β-sheet (red), which wraps the structure in a helicoidal embrace, are highlighted.
Table 1

Data Collection and Refinement Statistics

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**Refinement statistics**

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Values in the high resolution shell are shown in parentheses.