ORIGINAL RESEACH COMMUNICATION

**GLUTATHIONE IS THE RESOLVING THIOL FOR∫ THIOREDOXIN PEROXIDASE ACTIVITY OF 1-CYS PEROXIREDOXIN WITHOUT BEING CONSUMED DURING THE CATALYTIC CYCLE**

**Pedrajas, J. R.****1a, McDonagh, B.2, Hernández-Torres1b, F.1, Miranda-Vizuete, A.3, González-Ojeda, R.4,5, Martínez-Galisteo E.4,5, Padilla, C.A.4,5, Bárcena, J.A.4,5\***

1)Biochemistry and Cellular Signaling Group (a) and Cardiovascular Research Group (b), Department of Experimental Biology, University of Jaén, 23071-Jaén, Spain.

2)MRC-Arthritis Research UK Centre for Integrated Research into Musculoskeletal Aging (CIMA), Skeletal Muscle Pathophysiology Group, Institute of Ageing and Chronic Disease, University of Liverpool, Liverpool L69 3GA, United Kingdom.

3)Instituto de Biomedicina de Sevilla (IBIS), Hospital Universitario Virgen del Rocío/CSIC/Universidad de Sevilla, 41013 Sevilla, Spain,

4)Department of Biochemistry and Molecular Biology, University of Córdoba, 14071-Córdoba, Spain.

5)Córdoba Maimónides Institute for Biomedical Research, IMIBIC, 14071-Córdoba, Spain.

**ABBREVIATED TITLE**: Protective and catalytic role of GSH on 1-Cys Prx.

**ADDRESS FOR CORRESPONDENCE**: Prof. J. A. Bárcena, Department of Biochemistry and Molecular Biology, Ed "Severo Ochoa”, Pl. 1, Campus de Rabanales, University of Córdoba, 14071-Córdoba, Spain. Tel.: +34 947 218590; Fax: +34 957 218592; e-mail: <ja.barcena@uco.es>

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**Abstract**

***Aims***: A three-step catalytic cycle is common to all Peroxiredoxins (Prxs) despite structural and kinetic differences. The second step in 1-Cys type Prxs is a matter of debate since they lack an additional cysteine to play the resolving role as happens with the 2-Cys Prxs. The aim of this study was to elucidate the role of glutathione (GSH) in the thioredoxin dependent peroxidase activity of *Saccharomyces cerevisiae* mitochondrial Prx1p, a 1-Cys type Prx. ***Results***: The peroxidatic Cys91 residue of two Prx1p peptides can be linked by a disulfide, which can be reduced by thioredoxin and by GSH (Km=6.1 µM). GSH forms a mixed disulfide with the peroxidatic cysteine spontaneously *in vitro* and *in vivo*. Mitochondrial Trx3p deglutathionylates Prx1p without the formation of GSSG, so that GSH is not consumed in the process. The structural unit of native Prx1p is a dimer whose subunits are not covalently linked, but a hexameric assembly of 3 disulfide-bound dimers can also be formed. ***Innovation***: GSH is presented as a protective cofactor of Prx1p, which is not consumed during the peroxidase reaction, but provides a robust mechanism as the “resolving cysteine” and efficiently preventing Prx1p overoxidation. GSH exerts these roles at concentrations well below those commonly considered necessary for its antioxidant and redox buffering functions. ***Conclusion***: A 1-Cys peroxide scavenging mechanism operates in yeast mitochondria involving an autonomous glutathione molecule and the thioredoxin system, that could be of universal validity. Prx1p is fairly well protected from overoxidation, questioning its role in a “floodgate” mechanism for H2O2 signaling.

**Introduction**

Peroxiredoxins (Prx) are ubiquitous enzymes catalyzing non-heme thiol dependent scavenging of peroxides. They constitute an abundant family of proteins in most organisms, constituted by 6 subfamilies based on structural characteristics ([37](#_ENREF_37)). Despite structural and kinetic differences, all Prx share the same catalytic cycle based on a conserved active-site cysteine residue, the peroxidatic Cys (Cp) whose function is to directly reduce peroxides. The catalytic cycle can be divided into three steps: 1) peroxidation, 2) resolution and 3) recycling ([18](#_ENREF_18)). In the first step, Cp-S- is oxidized to a sulfenic acid (Cp-SOH) by the substrate peroxide; in the second step a disulfide is formed between Cp and a second resolving Cys (CR) provided by either the same protein (2-Cys mechanism) or by another protein or small molecule (1-Cys mechanism). Finally, in step three, recycling occurs by reduction of the disulfide by another protein or small thiol molecule regenerating the free thiols CP-SH and CR-SH.

Regarding the second step of their catalytic cycle, Prxs are divided into two groups: 1-Cys and 2-Cys Prxs. In 2-Cys Prxs, the resolving thiol is provided by the very same protein forming an intermolecular or intramolecular disulfide bond. However, 1-Cys Prxs do not have a second cysteine to act as the resolving residue near the peroxidatic cysteine and for this reason different mechanisms involving other proteins or small molecules have been postulated. Therefore, the resolving mechanism in 1-Cys Prxs has been a matter of debate for a long time. We have shown that mitochondrial Trx3p, together with the thioredoxin reductase Trr2p, support the peroxidase activity of the 1-Cys Prx of *Saccharomyces cerevisiae*, Prx1p, *in vitro* ([14](#_ENREF_14); [29](#_ENREF_29); [30](#_ENREF_30)). This would be expected for a canonical peroxiredoxin mechanism, although evidence indicates that Trx3p is not implicated *in vivo* ([14](#_ENREF_14)). We have also shown that Grx2p can also support recycling of Prx1p with the compulsory involvement of GSH in the catalytic process ([30](#_ENREF_30)). Other proposed mechanisms imply the formation of mixed disulfides with diverse proteins like Trr2p ([14](#_ENREF_14)) or πGST ([24](#_ENREF_24)).

There are characteristics of methionine sulfoxide reductases (Msrs) that resemble those of Prxs described above in some ways. Msrs catalyze the reduction of Met-SO in a thiol dependent reaction. The catalytic cycle of group-B (MsrBs) can be divided into the same 3 steps of oxidation, resolution and recycling and the members are also classified into two groups: 2-Cys and 1-Cys MSRB. As for 1-Cys Prxs, resolution of 1-Cys MSRB is an open question. It has been proposed that thioredoxin directly reduces the sulfenate formed at CP ([22](#_ENREF_22); [39](#_ENREF_39)), thus providing both resolution and recycling. Resolution by glutathionylation of CP and recycling by glutaredoxin has also been postulated ([38](#_ENREF_38)). Thioredoxin was proposed to directly reduce the peroxidatic sulfenic acid in *E.* coli BCP ([21](#_ENREF_21)) and PDI that of human glutathione peroxidase Gpx7 ([5](#_ENREF_5)).

Sulfiredoxins (Srxs) catalyze the ATP-dependent reduction of overoxidized sulfinate to sulfenate on the CP of some Prxs. Recently, a model of 1-Cys sulfiredoxin obtained by elimination of the second cysteine of yeast Srx was reported to be efficiently resolved and recycled by GSH/Grx/glutathione reductase system ([6](#_ENREF_6)).

In all cases where GSH was involved, it was the ultimate electron donor and was consumed stoichiometrically per catalytic cycle. We now report evidence that support a thioredoxin-dependent mechanism for 1-Cys Prxs in which glutathione is involved but is not consumed, thus playing a catalytic role in the second step of the catalytic cycle.

The relationships of glutathione with Prxs have another aspect as a mechanism for regulation of the quaternary structure of Prxs. Glutathionylation of the peroxidatic Cys of human PRXD1 converts the decamer to its dimeric structure in parallel with loss of its activity as a chaperone ([8](#_ENREF_8); [27](#_ENREF_27)). It has also been described how glutathionylation of type-II Prx from poplar, homologous to human PRXD5 induced the dissociation of non-covalently bound dimeric Prx ([26](#_ENREF_26)).

Some Prxs are sensitive to inactivation by their peroxide substrate by overoxidation of Cp to sulfinic (Cp-SO2H) or sulfonic (Cp-SO3H) before resolution can take place, rendering inactive forms of the enzyme. Evidence suggests that structural features that favor overoxidation of some members of the Prx family have been selected for during evolution to provide a “peroxide floodgate” to allow for signal transduction at concentrations of hydrogen peroxide above resting levels ([42](#_ENREF_42)). Evidence also suggests that Prx overoxidation allows them to function as molecular chaperones ([20](#_ENREF_20)).

In a previous work, we demonstrated that the GSH/Grx2/Glr system could efficiently support recycling of Prx1p and that GSH affected the oligomeric state of a form of Prx1p ([30](#_ENREF_30)). In the present study we have employed a form of Prx1p with one single cysteine (Cys91) in its sequence, the peroxidatic cysteine, which more accurately represents the native mitochondrial peroxiredoxin according to its signal peptide ([41](#_ENREF_41)). We report new data on the properties of Prx1p related to its catalytic cycle and subunit interaction that present GSH as a protective cofactor which is not consumed during the peroxidase reaction, but provides a fast universal mechanism acting as the “resolving cysteine” and efficiently preventing Prx1p from overoxidation.

**Results**

1. *Prx1p shares characteristics with members of the Prx6 subfamily of peroxiredoxins*

Prx1p is a mitochondrial peroxiredoxin coded by the YBL064C ORF of *Saccharomyces cerevisiae*. The full sequence contains 3 cysteine residues at positions 6, 38 and 91 (Fig. 1A). To study the molecular and biochemical properties of Prx1p we have produced 3 recombinant forms (Fig. 1A). N21Prx1p was designed according to a predicted mitochondrial matrix peptidase cutting-site between Q21 and A22. This protein has 2 cysteines at positions Cys38 and Cys91 and has been used and described before ([14](#_ENREF_14); [29](#_ENREF_29); [30](#_ENREF_30)). We have also produced a site directed mutated form, N21(S)Prx1p, in which Cys38 had been substituted by serine, and a form lacking the first 38 residues, N39Prx1p. The latter is similar to that employed to study the role of ascorbate as a reductant of peroxiredoxins ([25](#_ENREF_25)). Actually, the N-terminus of the mature Prx1p present in the mitochondrial matrix starts at residue 39 as determined experimentally ([41](#_ENREF_41)). Hence the mature protein contains only one cysteine, Cys91, and belongs to the 1-Cys group of peroxiredoxins included in the Prx6 subfamily, according to PREX (<http://www.csb.wfu.edu/prex/>) ([37](#_ENREF_37)). The functional signature of this subfamily, represented by human PRXD6 is highly conserved in yeast Prx1 (Fig. 1A). A structural modeling of Prx1p using human PRXD6 (PDB, 1prx; ([10](#_ENREF_10))) as a template, which shares 48% identity, produced a very reliable model with two different servers, SwissModel ([4](#_ENREF_4)) (QMEAN4 raw score=0,584, Z-score=-2,81 ([3](#_ENREF_3))) and I-TASSER ([43](#_ENREF_43)) (C-score = 0.78 ([34](#_ENREF_34))), where even the side-chains of the consensus active site triad are equally oriented (Fig. 1B). The peroxidatic cysteines of each subunit, which are the only cysteines in Prx1p, are ≈30Å apart from each other in the dimeric structure.

2. *The peroxidatic cysteines of two Prx1p peptides can be linked by a disulfide bond.*

Electrophoretic analysis of the three recombinant proteins showed differential behavior depending on the number of cysteines present in the protein. When denatured under reducing conditions, N21Prx1p appears as a band corresponding to a peptide of about 28 kDa (Fig.1C) but under non-reducing conditions behaves as a set of bands of different sizes corresponding to one (28 kDa), two (56 kDa), four (≈100 kDa), six, etc. peptides. However, when Cys38 was substituted by serine in N21(S)Prx1p only the bands of 28 and 56 kDa were present under non-reducing conditions and the same held true for N39Prx1p (Fig. 2A). These results indicate that the peroxidatic Cys91 establishes a disulfide bond between two Prx1p peptides and that these covalent dimers can bind each other by extra disulfide bridges if Cys38 is present.

Most of the purified recombinant N39Prx1p protein is in the form of Prx1-Prx1 covalent dimers under non-reducing electrophoresis (Fig. 2A, 2B and 2C, lanes “0” and “-“, respectively), but is converted to the single peptide when denatured under reducing conditions in the presence of ß-mercaptoethanol (see Fig. 3A, lane 1), an indication that two N39Prx1p polypeptides are linked together by a disulfide bridge between their Cys91. Interestingly enough, the covalent dimers ~~is~~ are also converted to the single peptide by reduction with GSH at low concentrations (<1mM) (Fig 2B). The effect of GSH is concentration dependent reaching 50% reduction at around 10:1 GSH:Prx1p ratio under the assay conditions, making it physiologically feasible (Fig. 2A). The complete mitochondrial thioredoxin system composed of Trx3p, Trr2p and NADPH, is also an efficient reductant of the disulfide bond between the unique Cys91 residues of each subunit in N39Prx1p (Fig. 2B, lane 4). However, in the absence of Trx3p, Trr2p alone cannot reduce this Prx1-Prx1 disulfide linkage between the catalytic cysteines (Fig. 2B, lane 5). Other non-physiological reductants like TCEP (Fig.2B, lane 6) can reduce this disulfide bridge but not ascorbate (Fig.2B, lane 7), which had been postulated as a possible universal reductant of peroxidases ([25](#_ENREF_25)).

Formation of disulfide bond between two Prx1p peptides is induced by the peroxide substrate. Addition of low concentrations of H2O2 to a preparation of N39Prx1p previously reduced with TCEP, regenerated the 56 kDa covalent dimer (Fig. 2C), although the proportion of covalent dimer formed was lower at relatively high concentrations of peroxide. However, when the peroxiredoxin was previously reduced with GSH, the covalent dimer could not be regenerated under the same oxidative conditions (data not shown). This effect was confirmed by another experimental approach. When a preparation of N39Prx1p reduced by the thioredoxin system was mixed with a peroxide generating system formed by xanthine, XO and SOD, the 56 kDa covalent dimer was formed with time, as the reducing power of the thioredoxin system was exhausted (Fig. 2D). However, when GSH was also added, the formation of the covalent dimer was abrogated, even at long incubation periods (Fig. 2E) likely due to glutathionylation of Cys91. Formation of the disulfide-linked dimer was inhibited in the presence of the sufenic acid specific reagent dimendone (Fig. 2F). These results indicate that formation of a disulfide bond between two Prx1p peptides is dependent on mild oxidative conditions that oxidize Cys91 to sulfenic acid in a proportion of Prx1p peptides, while the rest are still reduced, so that both can react between them to form an intermolecular disulfide and protect themselves from overoxidation..

*3. Prx1p forms a mixed disulfide with GSH, which Trx3p reduces.*

Evidence that Prx1p is glutathionylated was obtained by Western blot using anti-GSH antibodies and confirmed by mass spectrometry. As shown in Fig. 3A, only the single peptides obtained after treatment of the covalent dimer with GSH are reactive towards anti-GSH antibodies, indicating that GSH breaks the disulfide bond between two peptides and keeps bound through a mixed disulfide bond to each peptide. Incubation of glutathionylated Prx1p with the mitochondrial thioredoxin system resulted in a decrease of Prx1p reactivity with anti-GSH antibodies (Fig. 3B) indicating that Trx3p has deglutahionylase activity. Electrophoretic bands of untreated samples of N39Prx1p or treated with ß-ME, GSH or TCEP were analyzed by LC-MS/MS upon proteolytic digestion with AspN protease. A glutahionylated form of the Cys91 containing peptide was prominently detected in the band of Prx1p treated with glutathione (Fig. 3C), while the protein reduced with ß-ME, formed a mixed disulfide between Cys91 and ßME, *e. g*., “destreak” posttranslational modification (Fig. 3D).

To test whether Prx1p was also glutathionylated *in vivo*, cell free extracts from WT and double mutant ∆grx2∆Trx3 lacking mitochondrial thioredoxin and glutaredoxin systems were prepared in the presence of NEM to block any thiol present. These extracts were then run on non-reducing SDS-PAGE and a portion of gel between 20 and 30 kDa was cut, digested with AspN protease and analyzed by LC-MS/MS. The peptide containing glutathionylated Cys91 was detected in the double mutant extract but not in the WT, whereas the reduced (NEM-modified) peptide was conspicuous in WT but almost undetectable in the double mutant (Suppl. Fig. 2).

These results clearly demonstrate that GSH reacts with oxidized Prx1p and forms a mixed disulfide with Cys91 spontaneously, as a means to protect and to resolve the catalytic CP and that Trx3p can reduce this disulfide.

*4. Native Prx1p shows different oligomeric states whether it is glutathionylated or not.*

In a previous study we observed that N21Prx1p, which has 2 cysteines per peptide, behaved as an oligomer of hexameric structure or higher when analyzed by size exclusion chromatography, but behaved as a dimer in the presence of GSH ([30](#_ENREF_30)). In the present study we have carried out a similar analysis with N39Prx1p, which has only 1 cysteine per peptide, after treatment or not with GSH, to confirm the role of cysteines in the oligomerization process. Under non-denaturing conditions the untreated N39Prx1p eluted as a molecule of approximately 150 kDa, which would be consistent with a hexamer (Fig. 4A, chromatogram a). An aliquot of this eluted fraction (F1) appeared as a 56 kDa band, corresponding to a covalent dimer, when analyzed by non-reducing SDS-PAGE (Fig. 4A, inset). When N39Prx1p was glutathionylated by preincubation with GSH eluted as a 50-60 kDa protein (Fig. 4A, chromatogram b) corresponding to a dimer, but this eluted fraction (F2) resulted in a single band of 28 kDa when analyzed by non-reducing SDS-PAGE, corresponding to the size of a single peptide (Fig. 4A, inset).

These results demonstrate that the structural unit of native N39Prx1p is a dimer whose subunits are not covalently linked. However, the subunit peroxidatic cysteine of one non-covalent dimer can form a disulfide bridge with the equivalent cysteine of another non-covalent dimer giving rise to a hexameric assembly of 3 disulfide bound dimers.

*5. Oligomerization state of Prx1p in vivo.*

To check whether the hexamers could be formed *in vivo* or not, cell free extracts of yeast were analyzed by size exclusion chromatography (Fig. 4B) and the eluates were analyzed by Western blot with anti-Prx1 antibodies (Fig. 4B, inset). In these samples, Prx1p was only detected in the fractions corresponding to the dimer and appeared as a single peptide by non-reducing SDS-PAGE, indicating that the peptides are non-covalently associated physiologically. No Prx1p could be detected in the fractions of the eluate corresponding to a 150 kDa molecule.

Next we checked whether Prx1p would form disulfide-bridged dimers *in vivo*. Cell free extracts from mutants of mitochondrial redoxins or gamma glutamylcysteine synthetase (∆GSH1) were analyzed by non-reducing SDS-PAGE followed by detection of Prx1p by Western blot (Fig. 5A). Covalent dimers could not be detected by SDS-PAGE in a ∆GSH1 mutant lacking γ-glutamylcysteine synthetase or in mutants lacking the mitochondrial redoxins Grx2p or Trx3p or both. The covalent dimer could not be detected even in total cell-free extracts or mitochondrial subcellular fractions under different oxidative conditions (data not shown). Covalent dimers could only be detected in the ∆GSH1 mutant when the cell free extract was dialyzed before loading onto the gel (Fig. 5B). This result indicates that a metabolite is necessary to prevent Prx1p from forming an inter-subunit disulfide bond in the absence of GSH. The covalent dimer was reduced back to its monomeric form when the dialyzed extract from ∆GSH1 was incubated with NADPH (Fig. 5C)

From the results presented in Fig. 2D we postulate that this metabolite should be NADPH acting through the Trx system.

*6. GSH stimulates the thioredoxin dependent peroxidase activity of Prx1p.*

Given two options to resolve the peroxidatic cysteine, either through a disulfide bond between two Cys91 of two subunits or by a mixed disulfide with glutathione, it was worth comparing their kinetic efficiency when thioredoxin acted as the final thiol reductant for the recycling step of the catalytic cycle. As shown in Fig. 6A, reduction of peroxide is faster if GSH is present in the reaction mixture than when it is absent and the effect is immediate after addition of GSH to the reaction mixture (Fig. 6B). The peroxidase activity slows down progressively when GSH is absent, likely due to overoxidation under the conditions of the assay, but the reaction rate ~~duplicates~~ doubles when GSH is added (Fig. 6B).

Since glutathione plays a catalytic role in this reaction, a very small stoichiometric concentration of GSH was sufficient to reach maximum activity with a very small Km value of 6.1 µM (Fig. 6C). However, when the final reductant was Grx2, higher concentrations of GSH were necessary for maximum activity (Fig. 6D). As expected, the activity with Grx2 was highly dependent on GSH concentration, since GSH is also a reactant, not just a catalyst. These results demonstrate that the cysteine of glutathione is endowed with the highest efficiency ever reported for a resolving CR of a peroxidase system.

*7. Structural considerations of Prx1p glutathionylation*

As shown in the first paragraph of this section, the structure of yeast Prx1p can be modeled onto the folding pattern of human PRXD6 with high confidence. We have performed docking simulations of GSH on a model of monomeric Prx1p and on dimeric PRXD6 with SwissDock server with good fitness. A set of poses of docked GSH on the modeled structure of Prx1p are in the vicinity of the active site close enough to react and form a disulfide bond with Cys91 (Fig. 7A). It is worth mentioning the presence of Arg175 at less ~~that~~ than 5 Å from Cys91, which could serve as a counterion to stabilize a thiolate during the catalytic cycle, thus explaining the high reactivity of Cys91 with GSH ([12](#_ENREF_12)).

Peroxidatic cysteine (Cys47) of human PRXD6 lies in the bed of a wide and long groove at the subunits interphase (Fig. 7B and C). Docked GSH fits fairly well in this groove of the dimer with access to the catalytic site (Fig. 7D). In this structure of PRXD6 (PDB: 1PRX) the peroxidatic Cys47 is oxidized in the form of -SOH and the active site is in the fully folded (FF) conformation ([18](#_ENREF_18)). GSH access to the active site could even improve in the locally unfolded (LU) state needed for the resolution step of the catalytic cycle ([18](#_ENREF_18)).

**Discussion**

The results shown in Figs. 1C and 2 demonstrate that the Cys91 of two Prx1p peptides can form a disulfide bond between them as a means to carry out the resolving phase of its catalytic cycle and to provide Trx3p with a substrate to act on for regeneration. Formation of this disulfide between peroxidatic cysteines could also serve as a protective mechanism to avoid overoxidation. This uncovers a novel dual role for this residue both as “peroxidatic cysteine” (CP) and “resolving cysteine” (CR). In 2-Cys type peroxiredoxins, these roles are played by two different cysteine residues of the same polypeptide ([17](#_ENREF_17)). In human PRXD6, which is homologous to yeast Prx1p, the catalytic cysteine is “resolved” by heterodimerization with glutathione transferase πGST ([24](#_ENREF_24); [32](#_ENREF_32)) but this mechanism is unlikely to be operative in yeast because an equivalent transferase is not present in yeast mitochondria ([36](#_ENREF_36)).

Contrary to previous reports (Greetham & Grant, 2009 Should this be ref number?; Fig 4 therein) Cys38 is not required to form disulfide-stabilized dimers of yeast Prx1p. A basic covalent dimer can be established by disulfide linkage between peroxidatic Cys91 from two peptides as demonstrated by the behavior of N21(S)Prx1p and N39Prx1p, which contain only one cysteine residue. Disulfide linked chains of an apparently unlimited number of these basic covalent dimers could be formed if Cys38 were present in the peptide (Fig. 1C). However, if the protein possesses only one cysteine, as is the case of mature yeast Prx1p ([41](#_ENREF_41)), it would not be possible to form higher order disulfide linked oligomers once both Cys91 of two peptides are already involved in a disulfide bond. On the other hand, the homologous human cytosolic PRXD6 has an additional cysteine, which may explain the observation of high molecular weight forms of this protein under low reducing conditions ([13](#_ENREF_13)).

A summary of the catalytic cycle of Prx1p in the light of the data presented here is shown in Scheme 1A. The ability of Prx1p to form a disulfide between the peroxidatic cysteines, which can be reduced by Trx3p ~~explains~~ explaining the peculiar thioredoxin peroxidase activity of Prx1p, which has not been detected in other 1-Cys-Prxs. Moreover, the disulfide-linked dimer can also be broken by reduced glutathione resulting in the glutathionylation of the peroxidatic cysteines, as confirmed by mass spectrometry (Fig. 3). We had already postulated the formation of a mixed disulfide between glutathione and peroxidatic cysteine to explain the glutaredoxin-dependent peroxidase activity of Prx1 ([30](#_ENREF_30)). Mixed disulfides between glutathione and Prx1p or other peroxiredoxins have been detected before by mass spectrometry, but in all cases glutathionylation was forced under non-physiological conditions by incubation with relatively high concentrations of GSSG. In this study, we demonstrate that glutathionylation of Prx1p at its peroxidatic cysteine takes place directly, in a non-enzymatic mode, at very low concentrations of reduced glutathione, compatible with the cellular physiological environment. Moreover, Prx1p was glutathionylated in cells lacking mitochondrial thioredoxin and glutaredoxin systems (Suppl Fig. 2) demonstrating that the protective action of GSH is fully operative *in vivo*.

We also confirm that mitochondrial Trx3p catalyzes the deglutathionylation of Prx1p with very high efficiency. The canonical activity of Trx’s is the reduction of inter- and intra-molecular protein disulfide bonds, while reduction of protein-glutathione mixed disulfides is essentially carried out by glutaredoxins ([7](#_ENREF_7); [19](#_ENREF_19)). However, denitrosylating activity of human Trx1 and Trx2 ([2](#_ENREF_2)) and deglutathionylating activity by h-type Trxs in *A. thaliana* ([1](#_ENREF_1)) have also been observed. Cytosolic Trx1 and Trx2 from *S. cerevisiae* were also shown to contribute to lowering the general levels of protein glutathionylation although no specific substrates were identified ([16](#_ENREF_16)). The conformation of Cys91 glutathionylated Prx1p, likely in the locally unfolded state (LU) ([18](#_ENREF_18)), may favor the interaction with Trx3 in the same way as with the disulfide linked Prx1p dimer.

GSH stimulates Trx3p dependent peroxidase activity of Prx1p (Fig. 6). This could be explained simply by the fact that in the presence of GSH each peroxidatic cysteine would be available for catalysis on its own, whereas in the absence of GSH both peroxidatic cysteines have to be involved at the same time in a single catalytic cycle. In other words, the presence of GSH would be equivalent to duplication of the number of active sites. Actually, addition of small amounts of GSH exactly duplicates the activity of Prx1p. The high affinity of GSH for oxidized Cys91 is remarkable, likely as a consequence of a very high reactivity of the Cys91-SOH formed after the first step of the catalytic cycle, since the Km of GSH is 6.1 µM and the concentration needed for maximum thioredoxin-dependent peroxidase activity is roughly equimolar with the concentration of Prx1p.

It has been reported that certain plant Trxs display atypical active site sequences with altered residues between the two conserved cysteines *e.g.* WCRKC and it was argued that this sequence context allows them to use one single cysteine for a “monothiol” catalytic mechanism with glutathione as a reductant, since replacement of this sequence by the canonical one, WCGPC was sufficient to suppress its capacity to employ GSH ([9](#_ENREF_9)). We demonstrate here that this is not the case for mitochondrial Trx3p, whose active site sequence is, indeed, WCGPC but displays full activity with GSH as a cofactor. We propose a typical “dithiol” mechanism for Trx3p in this glutathione dependent recycling step of Prx1p involving both cysteines of the catalytic site as depicted in Scheme 1B. This mechanism would be consistent with the low amounts of GSH required for maximum activity (see Fig. 6). If a monothiol mechanism would take place, a second GSH molecule would be required to regenerate reduced Trx3p(SH)2, GSH would be consumed as a reactant and would be required at higher concentration as with the Grx2p system (Fig. 6D).

Formation of a mixed disulfide between Prx1p and Trx3p has been inferred from the detection of a ≈65kDa band by non reducing Western blot with anti-Prx1 antibodies in a sample of yeast cells containing a *Trx3::C58S* mutation ([15](#_ENREF_15)). In the dithiolic mechanism we propose, Cys55 of Trx3p could form a transient mixed disulfide with Cys91 of Prx1p as detected by MALDI-TOF mass spectrometry of a reaction mixture (Suppl. Fig. 1).

Mitochondrial thioredoxin reductase Trr2p cannot support recycling of the Prx1p catalytic cycle without the participation of Trx3p as shown in Fig. 6A. However, recycling of Prx1p by a system containing Trr2p and GSH in the absence of Trx3p has been observed with presumed formation of a mixed disulfide between Trr2p and the N21Prx1p, which bears an additional cysteine residue at position 38 ([14](#_ENREF_14)). This discrepancy may be due to the lack of extra cysteine residues in the more physiological N39Prx1p that we have used in this study.

In the structure of the non-covalently bound dimer of human PRXD6, the peroxidatic cysteines of each subunit are ~~very~~ relatively far away from each other (> 30Å) and the same holds true for the modeled structure of yeast Prx1p. It seems likely that the Prx1p covalent dimer is formed by “peptide swapping” between two non-covalent dimers and would require a conspicuous conformational change. It appears that a hexameric assembly provides the best arrangement to stabilize the new folding pattern of Prx1p covalent dimer (Fig. 4A). This supramolecular structure is similar to the oligomeric structures of other Prxs endowed with chaperone activity ([27](#_ENREF_27)). Even if the hexameric structure of Prx1p can perform chaperone activity, its existence *in vivo* would require the unlikely depletion of glutathione and NADPH from the mitochondrial matrix.

It has been proposed that oxidation of Trx3p induces apoptosis through “caspase-like protein” activation, but harsh forced oxidative conditions under respiratory growth and the presence of Prx1p were needed ([15](#_ENREF_15)). Whether the hexameric assembly of Prx1p dimers could take part in this pro-apoptotic or other processes, as a signaling element would be an interesting matter for further research.

Our results demonstrate that multiple surveillance systems protect Prx1p in the mitochondria to avoid its inactivation by overoxidation. First, under conditions of high reducing power and low peroxide pressure, the Trx3/Trr2 system would keep the peroxidatic Cys91 reduced. Second, if Cys91 is sulfenated, as occurs in the first step of the catalytic cycle, GSH would react rapidly with it to halt further oxidation. Third, in the unlikely event of these two protective mechanisms failing, Prx1p would self-protect itself by formation of a disulfide bridge between the peroxidatic cysteines of two molecules. The later would constitute an emergency mechanism that would keep Prx1p in a latent but active state, until GSH or Trx3p(SH)2 could activate it ready for another catalytic cycle.

Mitochondrial Prx1 from *S. cerevisiae* ~~had~~ has developed robust mechanisms to resist against peroxide pressure and to avoid overoxidation and inactivation, consisting of a high reactivity of Cys91-SOH towards the omnipresent thiol GSH. The presence of just small amounts of GSH in the mitochondria would suffice to quickly react with Cys91-SOH and protect it from further oxidation. Resistance to hyperoxidation is a property shared with human mitochondrial PRDX3 ([11](#_ENREF_11)) and may be a general characteristic of mitochondrial Prxs, although the mechanisms of resistance may differ. As long as reducing power is available to complete the third step of its catalytic cycle, the mitochondrial build-up and spillover of H2O2, as the “floodgate hypothesis” postulates ([42](#_ENREF_42)), would not take place in the presence of a resistant Prx1p. NADPH would eventually provide the required reducing power through the glutathione/Grx2 or Trx3/Trr2 systems, which would then represent the real limiting factors for H2O2 signaling from mitochondria.

**Innovation**

1-Cys Prx1p from yeast has the peculiarity of using its unique cysteine residue to accomplish the functions of both peroxidatic and resolving cysteine, when GSH is not available. The action of GSH on the thioredoxin dependent peroxidase activity described here for yeast Prx1p reveals a role of this ubiquitous small molecule at concentrations well below those commonly considered necessary for its antioxidant and redox buffering functions, in the same way as is required for iron sulfur cluster assembly ([23](#_ENREF_23)). Moreover, GSH exerts this function by itself as a cofactor, which is not consumed in the process and is recovered after every catalytic cycle. This protective action is operative *in vivo* and given the widespread abundance of glutathione in all ~~life~~ kingdoms of life and aksi considering that mitochondrial dysfunction caused by oxidative insults is ~~on~~ the basis of many diseases from cancer to neurodegenerative disorders, it would be worth checking whether this phenomenon is a general characteristic applicable to other peroxiredoxins.

**Materials and Methods**

*Materials:* NADPH, GSH, GSSG, TCEP, hydrogen peroxide, tert-butyl hydroperoxide, glutathione reductase from baker yeast, xanthine, xanthine oxidase from bovine milk, superoxide dismutase from *E. coli*, lysozyme, DNase I, human plasma thrombin, protease inhibitor cocktail and PMSF were purchased from Sigma, St. Louis, MO. Yeast strains Y10000 (WT), Y11934 (∆TRR2), Y17097 (∆GSH1), Y17197 (∆TRX3), and Y13090 (∆PRX1) were obtained from Euroscarf collection. Yeast strain MML44 (∆GRX2) was a kind gift from Prof. E. Herrero, University of Lleida, Spain ([33](#_ENREF_33)). The strains JR018 and JR024, carrying the double mutations ∆GRX2/∆TRX3 and ∆GRX2/∆TRR2, were obtained mating MML44 Mata haploid cells with Y17197 Mat or with Y11934 Matα, respectively as described elsewhere ([30](#_ENREF_30)).

*Cloning, expression and purification of the recombinant proteins:* Cloning into the pET-15b expression vector (Novagen) of the TRR2, TRX3, PRX1 (for the expression of N21Prx1p) and GRX2 genes of *Saccharomyces cerevisiae* was performed as described elsewhere ([28](#_ENREF_28); [29](#_ENREF_29); [31](#_ENREF_31)). To clone PRX1 gene for the expression of N39Prx1p, two primers (forward: 5’-GGGCATATGAAACAATTCAAACAAAGTGATCAACC-3’; reverse: 5’-GGGGGATCCTTATTTCGACTTGGTGAATCTTAAATAGG-3’) were designed, containing the NdeI and BamHI sites (underlined), respectively. The primers were used to amplify the corresponding fragment from *S. cerevisiae* DNA by PCR with Expand Long Template System (Roche Molecular Biochemicals). The PCR product was cloned into the pGEM-Te Vector System (Promega) and sequenced. The amplified fragment was subcloned into the NdeI/BamHI sites of the pET-15b expression vector, fusing the cloned fragment to a sequence that codes for a polypeptide of 20 amino acids at the N terminus, containing six histidine residues (His-tag) and a thrombin-cleavage site. *E. coli* TOP10 cells (Invitrogen) were transformed with this construct and colonies of transformed cells were obtained plating on Luria-Bertani medium containing 1 µg/ml ampicillin.

The expression of the recombinant proteins was performed in cultures of transformed *E. coli* BL21-DE cells in LB medium containing 1 g/ml ampicillin, growing at 37 ºC until a OD600 of 0.5 was attained. Then, the recombinant proteins were induced by addition of 0.5 mM isopropyl-ß-D-thiogalactoside (IPTG) to the cultures and cells were allowed to continue growing at 25 ºC overnight. The collected cells were diluted in 20 mM TrisHCl, pH 8.0, 0.1 M NaCl, 5 mM ß-mercaptoethanol, 0.6 mg/ml lysozyme, 1.2 µg/ml DNaseI. Lysis was performed as described previously ([30](#_ENREF_30)). Histidine-tagged proteins were purified from the extract by chromatography on a TALON™ Metal Affinity Resin column (Clontech), and the His-tag was subsequently removed adding 1 Unit/ml of thrombin to the purified preparations, dialyzed and passed through the affinity column again. Purity of the recombinant proteins was checked by SDS-PAGE and concentration was determined according to their theoretical extinction coefficients.

*SDS-PAGE and Western blotting*: SDS-PAGE was performed with homogeneous 10% (w/v) acrylamide gels. Samples were submitted to protein denaturalization at 100 ºC with 2.5 % (w/v) sodium dodecyl sulphate and in the presence or not of 0.715 M ß-mercaptoethanol. After electrophoresis, proteins were stained with Coomasie Blue or transferred to nitrocellulose membrane with a semi-dry electrophoretic transfer system. The membranes were processed by the method of Towbin et al. ([40](#_ENREF_40)). For the detection of Prx1p in samples, polyclonal Prx1p-antibodies were used at 1:5000 dilution. The Prx1p antibodies were purified from blood serum of immunized rabbits using a column of CNBr-activated Sepharose 4B with N39Prx1p coupled following the procedure recommended by the manufacturer (GE Healthcare). For the detection of glutathionylated proteins, mouse monoclonal anti-GSH (Virogen, Watertown, MA) was used at 1:1500 dilution. Membranes were incubated with the primary antibodies overnight at 4º C, then washed and incubated with the secondary antibodies, goat anti-rabbit (Bio-Rad) for anti-Prx1p detection and goat anti-mouse (Sigma) for anti-GSH. Chemiluminescent signal was induced using Clarity Western ECL Substrate (Bio-Rad).

*Activity assays*: Thioredoxin peroxidase activity was determined spectrophotometrically by measuring the oxidation of NADPH in a reaction mixture containing 250 µM NADPH, 0.5 mM GSH, 0.5 µM Trr2p thioredoxin reductase, 5 µM Trx3p thioredoxin, 5 µM N39Prx1p peroxiredoxin and 1 mM H2O2, in 20 mM TrisHCl, pH 8, 100 mM NaCl. The glutaredoxin dependent peroxidase activity of the peroxiredoxin was also determined spectrophotometrically in a reaction mixture containing 250 µM NADPH, 0.5 mM GSH, 1 Unit/ml of glutathione reductase, 5 µM Grx2p glutaredoxin, 5 µM N39Prx1p peroxiredoxin and 1 mM *tert*-butyl hydroperoxide, in 20 mM TrisHCl, pH 8, 100 mM NaCl. One unit of peroxidase activity is defined as the oxidation of 1 µmol of NADPH/min.

*Preparation of cell-free extract from yeast and subcellular fractionation*: *S. cerevisiae* cells grown in YPD medium up to the stationary phase were harvested and resuspended in 50 mM TrisHCl, pH 8, 100 mM NaCl, 0.1% v/v Triton X-100, 1 mM PMSF, 0.5% (v/v) of protease inhibitor cocktail and 0.5% (w/v) zymolyase 20T (USBiological). Cells were lysed by sonication during one hour while immersed in ice cold water. Homogenates were clarified by centrifugation at 12,000 x *g* for 30 minutes. Dialysis of cell-free extracts was performed with D-Tube Dialyzer Mini (Novagen) at 4ºC overnight. Protein concentration was determined spectrophotometrically with Bio-Rad Protein Assay using BSA as a standard.

*Gel-filtration chromatography*: A Superose 6 HR 10/30 column coupled to an FPLC instrument (Biologic Duo Flow, Bio-Rad) was equilibrated with 20 mM Tris-HCl, pH 8.0, 100 mM NaCl. A 100 µl sample was loaded for chromatography, which was performed at a flow rate of 0.3 ml/min. Samples contained 50 µM N39Prx1p, incubated or not with 1 mM GSH for 30 min at room temperature before loading onto the column. Samples of cell-free extracts contained a protein concentration of 2.5 mg/ml. Several standard preparations were run to calibrate the column: Blue dextran (2,000 kDa), ferritin (450 kDa), alcohol dehydrogenase (150 kDa), BSA (66 kDa) and carbonic anhydrase (29 kDa). Absorbance at 280nm after elution was monitored.

*Mass spectrometry*: Coomassie stained bands of interest from the SDS-PAGE gels kept in LCMS water were cut into pieces, destained, dehydrated with acetonitrile and dried in a speed vacuum. The reduction-alkylation step was omitted. AspN protease (Promega) was reconstituted as indicated by the manufacturer and was added to the sliced gel bands in a calculated ratio of 1:100 enzyme to protein in a total volume of 100µl and incubated overnight at 37ºC. The resulting digest was processed following the manufacturer’s protocol, dried in a speed vacuum and frozen. When required, the digest was resuspended in 3% acetonitrile, 0.1% TFA and analyzed by LC-MS/MS by injection in an ABSciex 5600+ Triple TOF after a 60 min chromatographic gradient. A MS survey scan was acquired followed by MS/MS scan for the 35 more intense detected ions. Files were searched using Mascot search engine against a *S. cerevisiae* database (2015/02/14 6718 sequences and 3,023940 residues) including glutathione, destreak (corresponding to mercaptoethanol), cabamidomethyl and N-ethylmaleimide dynamic modifications. Targeted analysis of the 1-Cys Prx1p peptide containing Cys91, DFTPVCTTEVSAFAKLKPEF, was performed using open source software Skyline ([35](#_ENREF_35)). The peak areas of [M+1] precursor ions identified as the Cys91 peptide containing either glutathione (2535.1782), destreak (2306.1083) or carbamidomethyl (2287.1315), were used for relative quantification of the oxidation sate of Cys91 ([35](#_ENREF_35)).

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**AUTHORS DISCLOSURE STATEMENT**

No competing financial interests exist.

**ABBREVIATIONS**

Grx or GRX, glutaredoxin; Msr or MSR, methionine sulfoxide reductase; ßME, ß-mercaptoethanol; N21Prx1p and N39Prx1p recombinant Prx1p devoid of a 21 and 39 residues N-terminal signal peptide, respectively; N21(S)Prx1p same as N21Prx1p in which Cys38 has been substituted by Serine; Prx or PRX, peroxiredoxin; *t*-BuOOH, *tert*-buthyl hydroperoxide; Srx or SRX, Sulfiredoxin; SOD, Superoxide dismutase; TCEP, Tris(2-carboxyethyl)phosphine hydrochloride;Trr or TRR, thioredoxin reductase; Trx or TRX, thioredoxin; XO, xanthine oxidase.

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**Legends to schemes and figures**

**Scheme 1: A)** Catalytic cycle of Prx1p as deduced from the data presented in this study. Each Prx1p polypeptide is represented by a large “P” with their peroxidatic Cys91 highlighted in the different redox states: sulfhydryl –SH, sulfenic –SOH, disulfide –S-S- and glutathionylated –SSG. Each step of the catalytic cycle is indicated with a circled number from 1 to 3; numbers with grey shadow represent the most likely steps under normal physiological conditions with GSH playing a catalytic and protective role in the resolving step and the mitochondrial thioredoxin system contributing the reducing recycling step. For more details see the main text. **B)** GSH dependent recycling of Prx1p by Trx3p should take place through a dithiolic mechanism in which reduced Trx3p(SH)2 would form a transient mixed disulfide with Prx1p. The mixed disulfide formed at the proximal Cys should be resolved by the distal cysteine to recover GSH and producing oxidized Trx3p. Trr2p Should eventually transfer reducing equivalents from NADPH to regenerate reduced Trx3p(SH)2.

**Figure 1**: **A)** Alignment of Prx1 with human PRXD6; the sequences of the three sites that constitute the functional signature of Prx6 subfamily according to PREX are highlighted in three different colors. The two cleavage sites assumed to design the recombinant proteins N21Prx1p and N39Prx1p are marked. In recombinant N21(S)Prx1p Cys38 was substituted by Serine. **B)** Structural modeling of mature Prx1 (dark blue) using human PRXD6 (PDB: 1prx.A; green) as a template; the side chains of three consensus residues of the Prx6 subfamily signature underlined in B) are shown and the functional structural signatures are highlighted in the same colors as in B). **C)** SDS-PAGE of N21Prx1p (a) and N21(S)Prx1p (b) denatured under reducing (ß-mercaptoethanol) and non-reducing conditions as described in Materials and Methods. The first lane contains Mw markers as indicated. (\*) Samples were preincubated with 1mM GSH before denaturing. (To see this illustration in color the reader is referred to the web version of this article at [www.liebertonline.com/ars](http://www.liebertonline.com/ars)).

**Figure 2**: Non-reducing SDS-PAGE of N39Prx1p after various treatments. **A**) Samples of 20 µM N39Prx1p pretreated with increasing concentrations of GSH (0-1000 µM). **B**) Samples of 20 µM N39Prx1p untreated (-) or pretreated with 1mM GSH, 0.5mM GSSG, a complete Thioredoxin System (TS), Thioredoxin System without Trx3p, 1mM TCEP or 1mM ascorbate. The Thioredoxin System is composed of 0.25mM NADPH, 0.5µM Trr2p and 2µM Trx3p. **C)** 20µM N39Prx1p untreated (-); or pre-treated first with 0.1mM TCEP followed by 0, 10, 20, 40, 60, 80, 100, 250 and 500 µMH2O2. The graphic shows an average densitometry of the bands corresponding to 56 kDa (black bars) and 28 kDa (grey bars) from the images of three gels. D.A.U., densitometric arbitrary units. **D)** 20µM N39Prx1p pre-incubated with a Thioredoxin System (TS) followed by addition of a Peroxide Generating System (PGS). Aliquots were taken from the incubation mixture at the indicated times, were denatured without reductant and loaded onto SDS-PAGE; **E)** same as in D), but with 100µM GSH added to the TS. **F)** 20 µM N39Prx1p was preincubated with 0.1 mM TCEP for 30 min at room temprerature followed by addition of 1 mM xanthine (-) or complete PGS without (0) or with dimedone at the indicated concentrations (5, 10, 15, 20) and further incubated for 1 h. Aliquots were taken, processed and analyzed as in D) and E).

TS: 50µM NADPH, 0.5µM Trr2p and 5µM Trx3p. PGS: 1mM xanthine, 0.5 mU/ml xanthine oxidase and 120 U/ml SOD.

**Figure 3**: **A**) Western blot of N39Prx1 showing in the left panel the Coomassie blue staining of a SDS-PAGE; sample 1 was denatured under standard reducing conditions and samples 2-4 under non-reducing conditions. Samples 3 and 4 were pre-treated with 1mM GSH and 0.1 mM TCEP, respectively, before denaturing. The bands **a** to **d** were picked for the mass spectrometry analysis; the right panel shows the Western blot of those samples with specific anti-GSH antibody. **B)** Deglutathionylase activity of Trx3p. 20 µM N39Prx1p samples pre-incubated or not with 0.5 mM GSH for 30 min as indicated, and then either treated or not with a mitochondrial thioredoxin reducing system, were subjected to non-reducing SDS-PAGE and then analyzed by Western blot with anti-GSH antibodies. Ponceau red stained membrane and developing with specific anti-GSH antibodies are shown. Mw markers were loaded on the first lane. The Thioredoxin System consisted of 250 µM NADPH, 0.5 µM Trr2p and 5 µM Trx3p. **C)** the protein bands a-d shown in A) were picked, digested with AspN proteinase and analyzed by LC-MS/MS. The MS/MS spectrum of glutathionylated 86-105 peptide identified in band **c** is shown; the detected fragment ions of the *y* and *b* series are indicated; Ions *y*152+, *y*172+and *y*182+, marked with circles, which are exclusive of the glutathionylated peptide, are conspicuous. **D)** Abundance of several redox modifications of the Cys91 containing peptide was measured and is shown in arbitrary units; “Destreak” is the name of the PTM resulting when when ßME remains bound to cysteine; -IAM is the carbamidomethylated cysteine produced by reaction with iodoacetamide; -SSG is the glutathionylated form.

**Figure 4**: Oligomerization of Prx1p. **A)** Analysis of 20µM samples of N39Prx1p by size exclusion chromatography without treatment (**a**) or pre-treated with 1mM GSH (**b**); numbered arrows indicate the elution times of Mw markers: 1, Blue dextran (2.000 kDa); 2, ferritin (450 kDa); 3, alcohol dehydrogenase (150 kDa); 4, bovine serum albumin (66 kDa); 5, carbonic anhydrase (29 kDa). F1 and F2 indicate fractions collected during the marked time intervals from chromatographies a and b, respectively. Inset, non-reducing SDS-PAGE of those fractions. **B)** Size exclusion chromatographies of cell-free extracts from WT (dotted line) and ∆PRX1 (solid line) yeast. Grey lines are the elution profiles of recombinant N39Prx1p shown in A) here superimposed for comparison. Fractions from both samples, as indicated, were collected and analyzed by Western blot with anti-Prx1 antibody (right panel).

**Figure 5**: Western blot detection of Prx1p in cell-free extracts from different yeast strains. **A)** The extracts were denatured under non-reducing conditions and 12.5 µg protein aliquots were analyzed by Western blot. **B)** Same as in A) but the extracts were further dialyzed at 4ºC overnight. **C)** Lanes 1 and 2, cell free extracts from ∆GSH1 mutant cells treated as in A) or B), respectively; lane 3, the dialyzed extract was incubated with 0.5 mM NADPH for 1 h at room temperature; lane 4, cell free extract from ∆PRX1 mutant. The migration positions of Mw markers are shown.

**Figure 6**: Thioredoxin peroxidase activity of N39Prx1p. The standard thioredoxin peroxidase assay was performed as described in Materials and Methods. At the time indicated by an arrow was added 1mM H2O2. **A)** The decrease in absorbance at 340 nm due to NADPH oxidation was recorded along time in an assay without Trx3p (upper line); in a normal standard assay (lower line); without GSH as indicated; and with GSH added at the time indicated by an arrow. **B)** The activity was measured without GSH, but at the times indicated by the arrows 0.5mM GSH was added to the reaction mixture, the reaction progress was monitored and the initial rates are shown in the inserted table. **C)** Thioredoxin peroxidase activities at different GSH concentrations. The inset is a double reciprocal plot from the data at the lower concentrations, which indicated an apparent Km of 6.1 µM for GSH. **D)** Peroxidase activity of N39Prx1p dependent on Grx2p. The composition of the assay was as described in Materials and Methods.

**Figure 7**: **A)** Predicted positions of GSH (purple) docked on a model of Prx1p (green) around the active site Cys91. Side-chains of residues from the catalytic triad according to PREX are shown. Arg175 and its distance to Cys91 are also depicted. The distance between Cys91 and GSH is also shown. **B)** and **C)** structure of dimeric human PRXD6 (PDB, 1prx). The surface of each subunit is colored differently; Cys47 from the yellow subunit is shown in purple color in B). **C)** Space filling of a wide and long groove in the interphase around Cys47 is shown in green. **D)** A closer look with GSH docked on the groove next to the catalytic site, where residues of the catalytic triad are shown. In this structure, Cys47 is oxidized in the form of -SOH and the active site is in the fully folded (FF) conformation. (To see this illustration in color the reader is referred to the web version of this article at [www.liebertonline.com/ars](http://www.liebertonline.com/ars)).

**Legend to Supplementary Figures**

**Supl. Fig. 1.** An incubation mixture containing 20 µM Prx1p and Thioredoxin System: TS: 50µM NADPH, 0.5µM Trr2p and 5µM Trx3p was vacuum-dried, resuspended in 0.1% TFA and 1 µl was spotted onto a MALDI plate. When dried, 0.6 µl sinapinic acid (10mg/ml in 0.3% TFA, 30% AN) were added and let until crystallization. The molecular mass analysis was carried out in a ABSciex MALDI-TOF/TOF 5800 mass spectrometer. Data acquisition was done using a High Mass method in a mass range between 10 - 75 kDa and the focused mass was fixed at at 35 kDa. 7500 shots were directed to the sample with a fixed lasser intensity of 4800 and a speed of movement over the sample of 700 µm/s.

**Suppl. Fig. 2. A)** Non-reducing SDS-PAGE of recombinant N39Prx1p treated with GSH (lane 1) and TCEP (lane 2) and cell free extracts from WT (lane 3) and ∆Grx2∆Trx3 double mutant (lane 4). Cells were grown in YPD-glucose medium and cell-free extracts were prepared in 50 mM Tris/HCl, pH 7.4, 1% SDS and 25 mM NEM to block any thiol present. Portions of gels between 20 and 30 kDa (dotted squares) were cut, digested with AspN protease and analyzed by LC-MS/MS as described in Material& Methods. **B)** Fragmentation spectrum of Cys91 containing peptide from cell-free extract of ∆Grx2∆Trx3 double mutant (sample 4). Prx1p was identified with 61.69% will give you exact % sequence coverage) and the Cys91-SSG containing peptide was detected by Mascot with high confidence. Two fragment ions unique for the glutathionylated peptide (*y(15)++* and *y(17)++*) were clearly identified and are marked with circles in the figure. **C)** MS/MS spectrum of NEM-modified Cys91 containing peptide from WT cell free extract (sample 3) , showing high confidence identification and good fragmentation with detection of unique ions (*y(15), y(16), y(17), b(6)++* and *b(9)++*).