Peroxiredoxins in Parasites

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Abstract

Significance: Parasite survival and virulence relies on effective defenses against reactive oxygen and nitrogen species produced by the host immune system. Peroxiredoxins (Prxs) are ubiquitous enzymes now thought to be central to such defenses and, as such, have potential value as drug targets and vaccine antigens. Recent Advances: Plasmodial and kinetoplastid Prx systems are the most extensively studied, yet remain inadequately understood. For many other parasites our knowledge is even less well developed. Through parasite genome sequencing efforts, however, the key players are being discovered and characterized. Here we describe what is known about the biochemistry, regulation, and cell biology of Prxs in parasitic protozoa, helminths, and fungi. At least one Prx is found in each parasite with a sequenced genome, and a notable theme is the common patterns of expression, localization, and functionality among sequence-similar Prxs in related species. Critical Issues: The nomenclature of Prxs from parasites is in a state of disarray, causing confusion and making comparative inferences difficult. Here we introduce a systematic Prx naming convention that is consistent between organisms and informative about structural and evolutionary relationships. Future Directions: The new nomenclature should stimulate the crossfertilization of ideas among parasitologists and with the broader redox research community. The diverse parasite developmental stages and host environments present complex systems in which to explore the variety of roles played by Prxs, with a view toward parlaying what is learned into novel therapies and vaccines that are urgently needed. Antioxid. Redox Signal. 17, 608–633.

Introduction and Scope

Peroxiredoxins (Prxs; EC 1.11.1.15) are exquisitely efficient cysteine-dependent peroxidases functioning in antioxidant, regulatory, and signaling systems (90, 116, 206, 254). They are ubiquitous and frequently present at high abundance (268). Broad reviews of Prx structure, function, and physiology are available (90, 92, 250), but here we focus on Prxs of the single-celled protozoa, multicellular helminths, and intracellular fungi that parasitize humans and are major contributors to the global burden of disease (267). In these organisms, Prxs are important for defense against endogenous and host-derived reactive oxygen species (ROS) and peroxynitrite (ONOO⁻) and are possibly involved in cellular signaling. We begin with an overview of structure-function relationships among Prxs and, based on the six Prx subfamilies, propose a systematic and informative nomenclature for parasite Prxs. In the survey section of this review, we list the Prxs of relevant parasites and review studies of their biochemical properties and cellular roles. Finally, we briefly discuss Prxs as potential drug and vaccine targets.

General Features of Prxs

Redox stress is a fact of life and organisms are exposed to oxidizing species from within and without, experiencing harmful oxidation of DNA, proteins, and lipids. Prxs, not recognized as widespread enzymes until 1994 (42), appear to be the first and dominant line of defense against hazardous ROS (263) and also are able to efficiently detoxify peroxynitrite/peroxynitrous acid (ONOO $^-/ONOOH$; pKa=6.8), a powerful oxidant and nitration agent produced by the reaction of superoxide (O2 $^{\bullet-}$) with nitric oxide (NO $^{\bullet}$), as previously reviewed (240); NO $^{\bullet}$, ONOO $^-/ONOOH$, and related molecules are often collectively referred to as reactive nitrogen species (RNS). In many eukaryotes Prxs also play a role in regulating signaling networks that use hydrogen peroxide (H2O2) as a second messenger (79, 89, 126, 213, 265).

Distinct conserved sequence patterns allow Prxs to be categorized into six subfamilies—Prx1/AhpC, Prx6, Prx5, PrxQ/BCP, Tpx, and AhpE (183). For simplicity, we will use Prx1 to refer to the Prx1/AhpC subfamily, and PrxQ to refer to the PrxQ/BCP subfamily. Conveniently, a Web-based searchable database, the PeroxiRedoxin classification indEX (PREX), has

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FIG. 1. A structurally detailed view of the Prx peroxidation reaction mechanism. Adapted from Ref. (92).

been developed that organizes Prxs into these families (235) (http://csb.wfu.edu/prex).

Catalytic properties of Prxs

Prxs catalyze the direct reduction of a variety of hydroperoxides (ROOH) to the corresponding alcohols (ROH; or H_2O in the reduction of H_2O_2) and water (184, 268). This occurs via a direct displacement S_N2 reaction, with a Prx active-site cysteine sulfur atom carrying out nucleophilic attack on the distal oxygen atom of the peroxide (77). The reaction proceeds through a planar transition state, releasing the alcohol and leaving the Prx cysteine oxidized to sulfenic acid (S_POH) (Fig. 1). To regenerate the reduced active-site cysteine for the next round of catalysis, the Prx participates in a series of reactions involving disulfide intermediates.

The complete catalytic cycle is diagrammed in Figure 2. In the first step of the reaction, within a fully folded (FF) Prx active site, the Prx peroxidatic cysteine (C_P) in its thiolate form (Prx^{FF}-S_P⁻) reacts with ROOH to yield ROH and Prx^{FF}-S_POH (Step 1, peroxidation). The Prx-S_POH pK_a value has been measured as 6.1 (208) and 6.6 (103); however, its protonation state during catalysis is unclear. Next, the Prx undergoes a local unfolding (LU) of its active site (Prx^{LU}-S_POH) to expose the S_POH group for reaction with a resolving thiol (R'-S_RH) to form a disulfide bridge (Prx^{LU}-S_P-S_R-R') and release water (Step 2, resolution). Prx^{LU}-S_P-S_R-R' is then reduced by a cysteine thiol from a different protein (R"-SH), yielding the disulfide R"-S-S_R-R' and restoring the reduced Prx^{LU}-S_P⁻ (Step 3, recycling). Finally, to complete the cycle, R"-S-S_R-R' is reduced and the Prx active site refolds to the FF-conformation required for peroxide reduction. Certain Prxs present in eukaryotes (268) are sensitive to overoxidation of the peroxidatic S_POH, which occurs when a second molecule of peroxide reacts with S_POH to form cysteine-sulfinic acid (S_PO₂H; Step 4). The S_PO₂H of such sensitive Prxs can in many organisms be reactivated (Step 5) by ATP-dependent reduction by sulfiredoxin (Srx) or, perhaps, the less-characterized sestrin (46, 113). How these Prxs facilitate peroxide signaling is not yet well understood but may involve a floodgate-like mechanism (265) and/or the formation of larger aggregates with molecular chaperone or other activities (12, 89, 197). Prx1 activity has also been reported to be regulated by phosphorylation (264).

Prxs can contain one or two participating Cys residues. In addition to the C_P possessed by all Prxs, 2-Cys Prxs have the resolving thiol (R'- S_R H) present as a cysteine (C_R) either on the

same chain (atypical 2-Cys Prxs) or on the other chain of a homodimer (typical 2-Cys Prxs). Figure 3 illustrates the rearrangements involved in a typical 2-Cys Prx catalytic cycle that allow the disulfide to form. R" is often thioredoxin (Trx) (41), a Trx equivalent such as tryparedoxin (Txn) (187), plasmoredoxin (Plrx; 24), or a Trx-like domain of a Prx reductase as in the bacterial alkyl hydroperoxide reductase component F (AhpF) (205). For 1-Cys Prxs, molecules such as Trx (193) or GSH (157) provide the R' thiol although the reducing equivalents can also be provided by ascorbate (171). The Trx or Trx equivalent is reduced by an NADPH-dependent reductase such as Trx reductase (TrxR). The NADPH is produced by the pentose phosphate pathway, connecting Prx activity to primary metabolism.

Early work on Prxs reported typical catalytic efficiency $(k_{\text{cat}}/K_{\text{m}})$ values of $10^5 \, \text{M}^{-1} \text{s}^{-1}$. Such reported catalytic rates, much lower than those of catalase and glutathione peroxidase (Gpx), led to a perception that Prxs were only physiologically

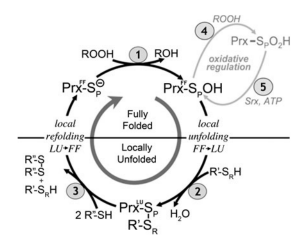


FIG. 2. The Prx catalytic cycle, showing both chemical steps and local unfolding/refolding steps. Shown are the three chemical steps (peroxidation [Step 1], resolution [Step 2], and recycling [Step 3]) of peroxide reduction, and the two chemical steps (overoxidation [Step 4] and ATP-dependent Srx-catalyzed recovery [Step 5]) seen for the oxidative regulation of sensitive, eukaryotic floodgate-type 2-Cys Prxs. The generic Prx is represented as a monomer with S_P designating the sulfur atom of C_P and S_R the resolving thiol. See the text for further descriptions. Adapted from Ref. (92).

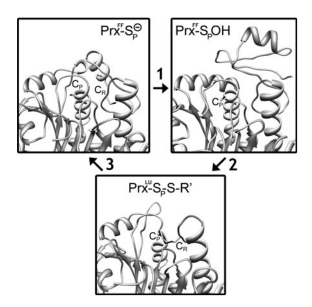


FIG. 3. Structures of three intermediates of the Prx catalytic cycle, using examples of dominant Prx1 and Prx6 family members. Steps in the cycle are numbered as in Fig. 2. Clockwise from upper left, the structures are *Pv*Prx1a (PDB Code 2I81), *Hs*Prx6 (1PRX), and *Pv*Prx1a (2H66).

relevant in organisms lacking the efficient heme- $(k_{\rm cat}/K_{\rm m} \sim 10^7~{\rm M}^{-1}{\rm s}^{-1})$ or seleno-peroxidases $(k_{\rm cat}/K_{\rm m} \sim 10^8~{\rm M}^{-1}{\rm s}^{-1})$ (77, 81). However, recent work has found that $k_{\rm cat}/K_{\rm m}$ values of $\sim 10^7~{\rm M}^{-1}{\rm s}^{-1}$ are typical for Prxs, with lower rates occurring when recycling of the Prx (Fig. 2, Step 3) by non-physiological redox partners is rate limiting (190). Given their high rate constants and high expression levels [as high as 0.5 mM (77)], Prxs are now considered the preferential reducing agent of ROOH, ONOO $^-$, and lipid peroxides in the cell (263), a view supported by a network model of cellular redox reactions (2). Prxs may also play a role in detoxifying peroxides of amino acids and proteins (196).

Remarkably, Prxs employ a simple cysteine sulfur atom as the nucleophile in the catalytic chemical reduction of ROOH yet achieve catalytic efficiencies nearly six orders of magnitude greater than expected for a thiolate (77). Recent structural analyses reveal that this catalytic power arises from an exquisitely structured active site that preferentially stabilizes the transition state; it does not simply activate the thiolate, but also orients and activates the peroxide substrate for attack (Fig. 1) (92).

Structural features of Prxs

The first Prx crystal structure (49) revealed the basic Prx fold (Fig. 4), which is related to that of Trx (192). Using a family-based naming of secondary structure elements (Fig. 4) (90), the C_P is always located in helix $\alpha 2$ on one face of the β -sheet, cradled between helices $\alpha 3$ and $\alpha 5$ (90). For 2-Cys Prxs, the C_R location tends to be characteristic of a given Prx subfamily, and as noted below can be near the C-terminus, or in $\alpha 2$, $\alpha 3$, or $\alpha 5$.

Prxs form two types of dimers. A-type dimers, thought to represent the ancestral mode of dimerization, interact in a tipto-tip manner at strands $\beta 1$ and $\beta 2$, and at the loops preceding helices $\alpha 2$, $\alpha 3$, and $\alpha 4$ (Fig. 5A) (222). B-type dimers, seen for Prx1 and Prx6 subfamily members, form by an edge-to-edge β -sheet interaction (the B-interface) to make an extended, 10-strand β -sheet (Fig. 5B) (90). The B-type dimers commonly associate via the A-interface into ring-shaped decamers (or dodecamers or octamers; Fig. 5C) to stabilize the FF active site and allow maximal peroxidatic activity (23, 191). The LU form of the enzyme favors dimers, suggesting that decamer–dimer transitions occur during the catalytic cycle (266); regulation by phosphorylation may occur via decamer disruption (268).

The following sections briefly describe structure–function characteristics of each subfamily.

Prx1. The Prx1 subfamily includes all "typical" 2-Cys Prxs. This subfamily is the largest and most widely distributed, with members found in archaea, bacteria, and all

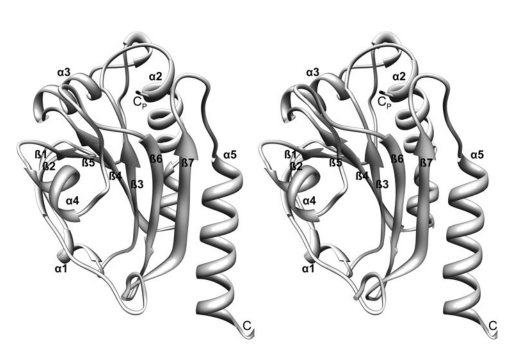
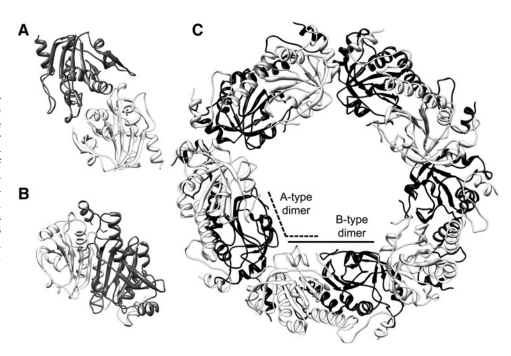


FIG. 4. The standard Prx fold. Shown is a stereo view of one chain of PDB entry 2I81 with labels giving the family numbering for secondary structural elements.

FIG. 5. Quaternary structure of Prxs. Shown are (A) an A-type dimer, (B) a B-type dimer, and (C) a decamer built using both A- and B-type interfaces; examples of A- and B-type dimers comprising the decamer are indicated by broken and solid lines, respectively. Structures shown are (A) *Pf*Prx5 (PDB entry 1XIY), (B) *Pf*Prx1m (2C0D), and (C) *Tcru*Prx1a (1UUL).



eukaryotic classes (183). Members include bacterial AhpC proteins; Txn peroxidases; yeast TSA1 and TSA2; and human Prxs I, II, III, and IV. These form doughnut-shaped decamers (Fig. 5C) and have C_R in a C-terminal extension of the partner chain of a B-type dimer (89). Thus, the disulfide form is a covalent dimer, and nonreducing SDS gels provide a simple method to test for Prxs in this form. Prx1s are typically the most abundant Prxs in an organism and, in eukaryotes, may be sensitive to overoxidation and involved in signaling. They generally prefer H_2O_2 over organic ROOH substrates (195).

Prx6. The Prx6 subfamily is named after human PrxVI (49). This subfamily is present in archaea, bacteria, and eukaryotes (183). Prx6 proteins resemble those of the Prx1 subfamily, containing a C-terminal extension, forming B-type dimers and, in some cases, oligomers (90). However, the vast majority of Prx6s are 1-Cys Prxs (61). The physiological reductant of most Prx6 members is unknown, and Trx notably fails to reduce human PrxVI (115).

Prx5. The Prx5 subfamily derives its name from human PrxV. Prx5s are present in bacteria, yeast, fungi, plants, and mammals (183), and have been localized to mitochondria, peroxisomes, and the cytoplasm (128). Members of the Prx5 subfamily form A-type dimers and include 1-Cys Prxs as well as 2-Cys Prxs that have C_R on helix α 5 of the same chain (60, 90, 222). A unique feature of the Prx5 active site is a π -helical bulge (52) in helix α 2 immediately following C_P (222). With the possible exception of the bacterial Prx5s fused to glutaredoxin (Grx) (124), the reductant for Prx5 subfamily members is Trx. Human PrxV prefers organic ROOH substrates and is an efficient reductant of ONOO $^-$ (250).

PrxQ. The PrxQ subfamily was found first in plant chloroplasts (132), and identified in *Escherichia coli* as the bacterioferritin comigratory protein (BCP), a name given before its function was known (109). This subfamily is predominantly bacterial, but also occurs in archaea and some

eukaryotes, but not in mammals (183). This subfamily includes primarily monomeric Prxs (183) but some form A-type dimers (*e.g.*, Protein Data Bank codes 2CX4, 2YWN). The PrxQ subfamily includes both 2-Cys and 1-Cys Prxs (257). For the 2-Cys members, the common location for C_R is in helix $\alpha 2$, five residues after C_P ; however, about 7% of the subfamily have C_R in helix $\alpha 3$ (147, 183).

Tpx. The Tpx subfamily is smaller and less phylogenetically diverse than the previous subfamilies. Known Tpx members are all bacterial (except for two trichomonad enzymes identified below) and are almost exclusively 2-Cys Prxs with C_R located in helix $\alpha 3$. Structural transitions involved in the local unfolding of the active site during catalysis have been examined in detail (93). Tpxs form A-type dimers that are unaffected by redox state and are essential for activity, as the substrate binding pocket includes residues from both subunits (93). Tpx proteins are commonly reduced by Trx. *E. coli* Tpx prefers hydrophobic, bulky cumene-OOH over H_2O_2 ($K_m \sim 9 \, \mu M \, vs. \, 1.7 \, mM$) (20), and, based on sequence conservation, a similar preference is expected for all Tpxs (91). This suggests a role in detoxifying lipid hydroperoxides (40).

AhpE. The AhpE subfamily, named after *Mycobacterium tuberculosis* AhpE, contains the fewest members, all in aerobic gram-positive bacteria of the order Actinomycetales (183). It appears to include both 1-Cys and 2-Cys Prxs. *M. tuberculosis* AhpE has been structurally characterized (146), but its natural reducing partner remains unclear (103). As no parasites have a Prx from this subfamily, we will not describe it further.

Occurrence and Studies of Prxs in Parasites Affecting Human Health

In this review, parasites are organized into groups and subgroups as follows: protozoa—the single-celled apicomplexans, kinetoplastids, and other protists; helminths—the multicellular nematode, cestode, and trematode worms; and

parasitic fungi. Detailed life cycle information for each parasite discussed here can be found on the U.S. Centers for Disease Control & Prevention Web site (http://dpd.cdc.gov/ DPDx). To identify all relevant Prxs, a primary search of our selected parasite genera (Appendix 1) used the PREX database (235); the EuPathDB bioinformatics databases (14–16, 85, 98) were consulted as a secondary source. One challenge in developing a list of Prxs is discerning whether highly similar Prx sequences from a single organism represent distinct gene products. Our simplistic solution to this problem was, for a given species, to list only the longest sequence within groups of Prxs with ≥95% amino acid sequence identity [as determined by Clustal Omega (231)]. For surveying the literature, the genera list (Appendix 1) was used to query National Center for Biotechnology Information (NCBI) PubMed (226) in combination with the terms "peroxiredoxin," "thiol peroxidase," "thioredoxin peroxidase," "tryparedoxin peroxidase," "thiol-specific antioxidant protein," and "peroxidoxin."

Of 97 unique parasite Prxs identified (Table 1), subfamily Prx1 is overwhelmingly the most common, with 80 members and at least one in every parasite considered. In some species, the multiple Prx1 isozymes may have similar functional properties but distinct localization or expression patterns such as is seen for human Prxs I-IV, all in the Prx1 subfamily (97, 105). Representatives of the Prx6, Prx5, PrxQ, Tpx, and AhpE subfamilies are much less frequently found, with 7, 4, 4, 2, and 0 occurrences, respectively. Plasmodia, the apicomplexan malaria parasites, are uniquely well endowed with a range of Prxs: two Prx1 proteins, a Prx6, a Prx5, and a PrxQ are present. Apart from *Plasmodia*, a PrxQ occurs only in the fungus Cryptococcus neoformans, a Prx5 is found only in the apicomplexan Toxoplasma gondii, and Prx6s were identified only among C. neoformans, T. gondii, and the nematode worms. The Tpx subfamily was represented only in the protozoan Trichomonas vaginalis.

As Table 1 was assembled, it became apparent that the nomenclature of Prxs from these parasites is highly inconsistent, confusing, and provides little or no information about the subfamily to which a given Prx belongs. For instance, 20 parasite Prx proteins named Tpx in the literature are not Tpx subfamily members, and names for members of the Prx1 subfamily include CPX, Prx3, Px2, Pxn2, Pxn3, Tpx-2, Trx-Px1, TRYP6, TSA, and TxnPx, among others (Table 1). To resolve this confusion, we here propose a systematic, informative, and forward-looking nomenclature in which Prxs are named according to the subfamily to which they belong (e.g., Prx6). Lettered suffixes (e.g., Prx1a and Prx1b) are used in the Prx1 and Tpx subfamilies since multiple Prxs of these families can be found in a single organism; even if just one Prx1 is currently known in an organism, a suffix is assigned based on its sequence similarity to other Prxs (e.g., Fig. 8). The suffix "m" (i.e., Prx1m) is given to Prxs assigned as mitochondrially

Table 1 lists these newly assigned names and their corresponding literature names, and notes experimental results that cannot be uniquely assigned to a specific Prx. To refer to a Prx from a certain organism, an italicized species-specific prefix is used (e.g., TcruPrx1a for the first cytosolic Prx1 subfamily enzyme of Trypanosoma cruzi). We assume unless proven otherwise that the equivalent Prxs in organisms of the same genus will behave similarly, and so studies of such en-

zymes are grouped together. We also assume, in the absence of contrary evidence, that the members of a Prx subfamily will share the features of that subfamily, so studies confirming such properties are only mentioned briefly.

As we begin the survey, a few general observations can be made. First, despite the presence in many Prx1 subfamily members of motifs very similar to the "GGLG" and "YF" motifs associated with the overoxidation shunt (265), there is no biochemical evidence that any protozoan parasite Prx1 is susceptibile to overoxidation. Also, no protozoan parasite appears to have a gene encoding sulfiredoxin. A second point is that for many Prxs, the recycling step can occur via multiple pathways and so sorting out the main path is nontrivial and perhaps even of little importance. Overall, the most common path seems to be through Trx via an NADPH-TrxR-Trx-Prx cascade. As noted above, the apicomplexans and trypanosomatids feature Trx equivalents (i.e., Plrx and Txn), which complement or replace the common system (Fig. 6). A third point is that among the Prxs listed in Table 1, there are only six for which crystal structures are available (Table 2). For the others, the amino acid sequence identity to crystallographically characterized Prxs is generally high (at least 40% for all Prxs except the PrxQs, which have greater than 34% identity to a known structure), so their three-dimensional structures can be inferred with reasonable accuracy.

Finally, we note that determining the physiological roles of Prxs can be difficult. Since all Prxs will turnover (although differentially) H₂O₂, ONOO⁻, and organic ROOH, it must be realized that when any given Prx is artificially overexpressed in a tissue, it will by its very nature protect that tissue from oxidative assault. While it is tempting to say that this result proves that the Prx in question is involved in protection from oxidative stress, it does not. Such a result gives no insight into the true physiological role of the Prx, because adding an excess of any peroxidase will provide such protection. In contrast, the true physiological role of a given Prx depends on where and to what extent the enzyme is expressed under natural conditions. Similarly, when a *Prx* gene is knocked out or knocked down (i.e., by RNA interference [RNAi]), the lack of a specific phenotype provides limited information about its physiological role, because with multiple antioxidant enzymes present, there is typically a high level of redundancy.

The following sections are grouped by parasite type and species, reviewing what is known about enzyme kinetic and substrate profile studies (available for ~ 50 Prxs), spatial and temporal protein localization experiments (immunoblot and immunolocalization, done for ~ 40 Prxs), proteomic and gene expression studies (reverse transcriptase–polymerase chain reaction [RT-PCR], performed for ~ 40 Prxs), and biological roles as dissected by gene overexpression, knockout (deletion), and knockdown (RNAi) studies (reported for ~ 30 Prxs). In many studies, enzymatic activity is evaluated indirectly by monitoring the prevention of peroxide-induced inactivation of glutathione synthase (GS) or the nicking of DNA (123, 148).

Prxs in apicomplexans

Apicomplexans are obligate intracellular parasites containing an apical complex that aids in entering host cells (28). Many also contain a chloroplast-derived apicoplast (216), which, for those that have it, is essential and is the site of

Table 1. Prxs Found in Parasites of Humans

PROTOZOA Apicomplexans 2, Trx-Px1 Kinetoplastids ExnPx2	Species ^a	Name	Literature name(s) Identifier (number	Identifier (number of loci) ^b	References
Prx1	yptosporidium <u>h</u> ominis yptosporidium <u>m</u> uris	Prx1a Prx1a Prx1m	PROTOZOA Apicomplexar		
lesi Prizia Michi, intri Prizia Prizia ICys-Prx Prizia ICys-Prx Prizia ICys-Prx Prizia ICys-Prx Prizia ICys-Prx Prizia Prizia MCP1 Prizia Iran Prizia Prizia Iran Prizia Prizia Iran Prizia	yptosporidium parvum asmodium falciparum	Prx1a Prx1a Prx1m Prx6 Prx5	TrxPx Prx-1, Px2, TPx-1, TPx-2, Trx-Px1 TPx-2 1-Cys-Prx, Prx-2, Tpx-1 AOP1 AOP1 AOP1	cgd4_740 cgd4_740 PF14_0368 PFL0725w PF08_0131 MAL7P1.159	(142) (4, 118, 121, 129, 130, 137, 186, 210, 271) (27, 121, 210, 271) (94, 117, 120, 121, 130, 137, 181, 185, 271) (121, 185, 222)
Price The The The The The Price Price Price Price Price The The The Price The The The Price The The The Price The Price Price Price Price Price Price Price Price Price Tre-Price Price Tre-Price Price Tre-Price Price Tre-Price Price Pr	asmodium <u>k</u> nowlesi	PrxQ Prx1a Prx6 Prx5	I-Cys-Prx AOP MCP1	FF1U_0208 PKH_126740 PKH_143220 PKH_011610 PKH_021360	(127, 214)
ii Prxla Prx, Trx-Px1 Prxla Prx, Trx-Px1 Prx6 Trx-Px2, Prx2, PrxII Prx5 AOP Ensis Prx1a Pxn1 Prx1a Pxn2 Prx1a CTxnPX Prx1a CTxnPX Prx1a TxnPX Prx1a TxnPX Prx1a TxnPX Prx1a Prx2, Pxx1, Pxx1 Prx1a Prx2, Pxx2 Prx1b Prx3, Pxx2 Prx1b Prx3, Pxx3 Prx1b Prx3, Pxx3 Prx1b Prx1x Prx1b Prx3, TxnPx Prx1b Prx1x Prx1b Prx1x Prx1b Prx3, TxnPx Prx1b Prx1b Prx3, TxnPx Prx1b Prx1b Prx1, TxnPx Prx1b Prx1b Prx1, TxnPx Prx1b Prx1b Prx1, TxnPx Prx1b Prx1b Prx1b, TxnPx Prx1b	asmodium <u>v</u> ivax	Prx1a Prx1m Prx6 Prx5	TPX-1 TPX-2 1-Cys-Prx	PVX_118545 PVX_123435 PVX_093630 PVX_081760	
ensis Prxla Pxnl Prxlb Pxn2 Prxla CTxnPX fiensis Prxla mTxnPX fiensis Prxla TxnPX Prxla TxnPX Prxlm Prxla TxnPX Prxla TxnPX, Prxl, Pxnl Prxlb Prx2, Pxn2 Prxlb Prx3, Pxn3 ng* Prxla TxnPx ^d Prxlm mPrx, TxnPx ^d Prxlm mPrx, TxnPx ^d Prxlm Pxxln mTxnPx, TRYP ^d	xoplasma gondii	Prx1a Prx1a Prx6 Prx5	x-Px1 2, Prx2, PrxII		(5, 141, 159) (141) (5, 61, 66, 95, 141)
tiensis Prx1a cTxnPX Prx1m mTxnPX Prx1a TxnPX Prx1a TxnPX Prx1a TxnPX, Prx1, Pxn1 Prx1b Prx2, Pxn2 Prx1b Prx3, Pxn3 nf* Prx1a TxnPx ^d Prx1m mPrx, TxnPx ^d Prx1m Pxx1 TxnPx ^d Prx1a Pxx1x TxnPx ^d Prx1b Pxx1a Pxx1x TxnPx ^d Prx1b Pxx1x TxnPx ^d Pxx1b Pxx1b Pxx1x TxnPx ^d Pxx1b	shmania <u>ae</u> thiopica	Prx1a Prx1b		, ,	(112) (112)
Prx1a TxnPx, Prx1, Pxn1 Prx1b Prx2, Pxn2 Prx1c Prx3, Pxn3 ni* Prx1a TxnPx ^d Prx1m mPrx, TxnPx ^d Prx1a PxN1, TxNPx ^d Prx1a PxN1, TXN2, TxnPx ^d Prx1b PxN2, TXN2, TxnPx ^d Prx1a CTxnPx, TRYP ^d Prx1a mTxnPx, TRYP ^d	shmania <u>am</u> azonensis shmania <u>b</u> raziliensis	Prx1a Prx1m Prx1a	cTxnPX mTxnPX TxnPX	AAX47428 AAX47429 LbrM.15.1080 (2)	(101, 149) (101, 149) (55)
Prx1a TxnPx ^d Prx1m mPrx, TxnPx ^d Prx1a PXN1, TXN1, TxnPx ^d Prx1b PXN2, TXN2,TxnPx ^d Prx1a cTxnPx, TRYP ^d Prx1a mTxnPx, TRYP ^d	shmania <u>c</u> hagasi	Prx1a Prx1b Prx1c	TxnPx, Prx1, Pxn1 Prx2, Pxn2 Prx3, Pxn3	LUTA:22:000 AAK69586 AAK69587	(21, 22)
Prx1a CTxnPx, TRYP ^d Prx1m mTxnPx, TRYP ^d	shmania <u>d</u> onovani ^c shmania <u>g</u> uyanensis	Prx1a Prx1a Prx1a Prx1a	TxnPx ^d mPrx, TxnPx ^d PXN1, TXnIy, TxnPx ^d PXN1, TxnPx ^d PXN1, TxnPx ^d	AAK00633 AAX73294 AAX18168 AAX21766	(80, 106, 111), (125, 194, 270) ^d (96), (125, 194, 270) ^d (1), (258) ^d (1), (258) ^d
THE STATE OF THE PARTY OF THE P	shmania <u>i</u> nfantum	Prx1a Prx1a Prx1m	CTxnPx, TRYP ^d mTxnPx, TRYP ^d m xnPx r x x x x x x x x x x x x x x x x x x	AAV51703 Linj.15.1100 (3) Linj.23.0050	(37, 38, 149, 220), (51) ^d (35–38, 54), (51) ^d (35–38, 54), (51) ^d
Letshmania <u>maj</u> or PrxIa IryPI, IKYP6, IXNP× LmjF15.1060 (7) PrxIm PrxIm Leishmania <u>mex</u> icana PrxIa LuxM.15.1040 (2) PrxIm Leishmania mexicana PrxIa	shmania <u>maj</u> or shmania <u>mex</u> icana	Prxla Prx1m Prx1a Prx1m	ItyP1, TRYP6, TXNPx	LmjF15.1060 (7) LmjF.23.0040 LmxM.15.1040 (2) LmxM.23.0040	(58, 72, 133, 144)

Table 1. (Continued)

		T TITLE T	(CONTINOED)	
Species ^a	Name	Literature name(s)	Identifier (number of loci) ^b	References
<u>L</u> eishmania <u>t</u> ropica	Prx1a Prx1h		AAZ23601 AAZ23600	
$\overline{\underline{\Gamma}}$ rypanosoma $\overline{\underline{b}}$ rucei	Prx1a Prx1m	c-TxnPx, CPX, TDPX, TryP, TRYP1 m-TcTxnPx, TRYP2		(32, 33, 39, 65, 135, 201, 247, 249, 260) (39, 135, 175, 201, 247)
<u>Trypanosoma cruzi</u>	Prx1a	c-TcTxnPx, CPx	Tc00.1047053504839.44 (2)	(39, 78, 87, 135, 163, 188, 199–203, 249, 261, 275), (9, 17, 64, 151, 230)
	Prx1m	m-TcTxnPx, MPx	Tc00.1047053509499.14	(39, 188, 198–201, 261), (9, 17, 64, 151, 230) ^d
\overline{B} lastocystis h ominis	Prx1a Prx1m	Othe	Other protozoa CBK22665 CBK25198	
<u>E</u> ntamoeba <u>d</u> ispar	Prx1a Prx1b Prx1c		EDI_034430 (1–17) EDI_182130 (1–7) EDI_053480 (1–2)	(50, 153, 242) ^d (50, 153, 242) ^d (50, 153, 242) ^d
Entamoeba luistolytica Entamoeba moshkovskii Giardia <u>lamb</u> lia	Prx1a Prx1a Prx1a Prx1b	Eh29, TSA	EH123510 (1–8) EH13210 (1–8) BAD66879 GL50803_14521 (2) GL50803_15383	(30, 102, 110), (13, 48, 50, 57, 104, 153, 212, 221) (47, 48) ^d
<u>Naegleria fowleri</u> Trichomonae vaginalie	Prx1a Prv1a	Trv-Drv	n/a TVAC 114310	(122) (53 143) (200 276) ^d
	Prx1b	117.117	TVAG_350540	(20), 130), (20), 2, 0) (20), 27(6) (20), 27(6) (20), 27(7)
	Prx1d		TVAG_095250	(209, 276) (209, 276) ^d
	Prxle Prx1f		I VAG_0/3420 TVAG_487570 TVAC_05530	(209, 276) ⁴ (209, 276) ⁴
	Tpx-a Tpx-b	Thiol peroxidase	TVAG_165690 (1–2)	(143) ^d
		HEI	HELMINTHS Nematodes	
<u>B</u> rugia <u>m</u> alayi	Prx1a Prx1b	TPx-2, TPX TPx-1	Q17172 P48822	(10, 86, 154, 155) (86)
<u>Loa loa</u>	Prx6 Prx1a Prx1h	TPX	AAF21098 EFO27148 FFO36191	
Onchocerca volvulus	Prx1a Prv6	PXN-2, Tpx-1, Tpx-2, TSA Tpx-2	AAC32810 AAC37392	(152, 227, 278) (44)
Wuchereria <u>b</u> ancrofti	Prx1b	TPx-1	, ,	(++)
Echninococcus granulosus Echninococcus multilocularis	Prx1a Prx1a	TPX TRX	Cestodes AAL84833 BAC11863	(100, 145, 158, 218, 234)
<u>T</u> aenia <u>crassiceps</u> <u>Taenia so</u> lium	Prx1a Prx1a	2-CysPrx 2-CysPrx	ACM89282 ACM89281	(252) (170, 252)

Table 1. (Continued)

Species ^a	Name	Literature name(s)	Identifier (number of loci) ^b	References
		Tr.	Trematodes	
Clonorchis sinensis	Prx1b	Prx2	AEK86199	(19, 114, 274)
1	Prx1m	Prx3	AEK86200	(19)
Fasciola gigantica	Prx1b	Prx-1, Prx2	ACU27401 (2)	(43, 211, 277)
\overline{F} asciola hepatica	Prx1a	FhePrx, Prx-1, Prx-2, Tpx	AAB71727	(67, 68, 108, 160, 161, 217, 229)
. 1	Prx1b		ACI04165	
Opisthorchis viverrini	Prx1b		ACB13822	(239)
Schistosoma japonicum	Prx1a	TPx-1	CAX78585	(138, 139)
	Prx1b	TPx-2	AAW25625	(138, 139)
	Prx1m	Prx-3, TPx-3	AAW25436	(138, 139)
Schistosoma mansoni	Prx1a	Prx1, TPx1	AAG15507	(140, 176, 177, 224, 225)
1	Prx1b	Prx2, TPx2	AAG15508	(68, 140, 176, 177, 224, 225, 256)
	Prx1c		XP_002577572	
	Prx1d		XP 002577887	
	Prx1m	Prx3, TPx3	AAG15506	(68, 140, 224, 225, 256)
			FUNGI	
Cryptococcus neoformans	Prx1	Tsa1 ^d	AAP68994	(165, 167), (166) ^d
I	Prx6	Tsa3 ^d	XP_567781	$(166)^{d}$
	PrxQ	${ m Tsa4^d}$	XP_569732	$(166)^{d}$
Encephalitozoon cuniculi	Prx1	TDX	ECU03_1190	
Encephalitozoon intestinalis	Prx1	TSA1	$Ein03_1090$	

"Species names shown in bold type are those with complete genome sequences available (see, e.g., http://eupathdb.org, http://www.sanger.ac.uk/resources/downloads/protozoa, and http://www.ncbi.nlm.nih.gov/genomes/leuks.cgj). Underlined portions of the name are used in the shorthand names of proteins from that organism.

"The EuPathDB gene ID, if available, is listed in italic type; otherwise, the NCBI accession number is provided. Where relevant, in parentheses is shown the number of loci (or range of possible loci) encoding gene products of ≥95% identity.

"The genome sequencing for this organism is in progress but is not publicly available.

"The genome sequencing for this organism is in progress but is not clearly correspond to a systematic name.

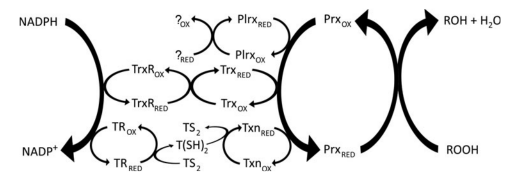


FIG. 6. Prx redox cascades showing special participants of *Plasmodia* (top pathway) and kinetoplastids (bottom pathway). TrxR, thioredoxin reductase; Plrx, plasmoredoxin; Trx, thioredoxin; TR, trypanothione reductase; TS₂, trypanothione; Txn, tryparedoxin.

isoprenoid, fatty acid, and heme biosynthesis (228). The apicomplexan parasites of humans are in the genera *Plasmodia*, *Toxoplasma*, and *Cryptosporidia*, and all have Prxs.

Plasmodia. Parasites of the genus *Plasmodia* are the cause of malaria. Their complex life cycle involves the blood-meal ingestion of male and female gametocytes by the *Anopheles* mosquito, wherein they multiply and form oocysts, which rupture to release infective sporozoites. In the human host, these sporozoites infect liver cells to form schizonts that rupture to release merozoites, which in turn infect red blood cells and develop into trophozoites. The trophozoites develop into schizonts that rupture to again release merozoites, which reinfect red blood cells to perpetuate the disease-causing blood stage or else differentiate into gametocytes.

While inside erythrocytes, *Plasmodia* are exposed to high levels of oxidative stress from host-defense ROS and the by-products of heme degradation. Plasmodial antioxidant enzymes are thought to be crucial for pathogenesis and are considered antimalarial therapeutic targets (62). Although *Plasmodia* possess a mitochondrial superoxide dismutase (SOD), they lack catalase and Gpx so their redox defense seems to rely heavily on Prxs (180). Their Prx recycling systems (Fig. 6) feature three Trxs (Trx1, Trx2, and Trx3) and also Plrx (24) and Grx (185), all found in characteristic cellular compartments (Fig. 7) (121). TrxR, the normal reductant for Trxs, has been shown to be nonessential (31); Plrx is known to reduce *Pf*Prx1a (186) and *Pf*Prx5 (185), but its reductant is unknown. Fascinatingly, *Plasmodia* are also reported to import host *Hs*PrxII (131) and other redox enzymes and equivalents (62).

Each plasmodial species considered here has a set of five Prxs: two from the Prx1 subfamily (Prx1a and Prx1m), one PrxQ, one Prx5, and one Prx6. The Prx1a and Prx1m enzymes cluster more closely with their cognates in other species than they do with each other (Fig. 8A), indicating that the Prx1a and Prx1m groups had already diverged in a common ancestor of the *Plasmodia*. *Plasmodia* spp. affecting nonhuman mammals have this same Prx complement, and are included in the discussion where relevant.

Localization studies: Studies using GFP fusion proteins (121) have localized *Plasmodium falciparum* Prx1a and Prx6 to the cytosol, Prx1m to the mitochondrion [see also Refs. (27, 62, 271)], and Prx5 to the apicoplast [see also Ref. (222)]. *Pf*PrxQ is exclusively localized to the nucleus (214), superceding previous work placing it in the cytosol and elsewhere (127). In the nucleus, it associates with chromatin (a novel observation for any Prx), and is enriched in coding regions (214).

Expression levels and other in vivo studies: PfPrx1a is constitutively expressed throughout the life cycle, with higher levels in the erythrocyte stage (118, 119, 271). A PfPrx1a knockout grows normally but is hypersensitive to exogenous ROS and RNS (129) and PfPrx1a mRNA and protein levels increase under oxidative stress (4). Also, the $\Delta prx1a$ mutant in the mouse parasite $Plasmodium\ berghei$ shows a 90% decrease in sporozoite formation, a 60% decrease in gametocyte yield, and attenuated infectivity in mice, although with no evidence of direct oxidative damage (272, 273).

PfPrx1m is present in the highly metabolically active trophozoite and schizont stages, and PfPrx6 is constitutively expressed, with transcript and protein levels elevated in the trophozoite and early schizont stages (271). Recombinant

TABLE 2. AVAILABLE PRX CRYSTAL STRUCTURES

PRX	PDB code	Resolution (Å)	Redox state	Conformation	Oligomeric state ^a	Reference
PvPrx1a	2H66	2.5	S ₅₀ -S _{170′}	LU	$(\alpha_2)_5$	(255) ^b
PvPrx1a	2I81	2.5	SH ₅₀ , SH ₁₇₀	FF	$(\alpha_2)_5$	n/a ^c
PyPrx1a	2H01	2.3	SH ₄₄ , SH ₁₆₄	LU	$(\alpha_2)_4$	n/a ^c
<i>Pf</i> Prx1m	2C0D	1.8	S ₆₇ -SH ₁₈₇	LU(alt) ^d	$\alpha_2(B)$	(27)
<i>Ťcru</i> Prx1a	1UUL	2.8	SH ₅₂ , SH _{173′}	FF `	$(\alpha_2)_5$	(202)
PfPrx5	1XIY	1.8	SO_3H_{59}	FF	$\alpha_2(A)$	(222)
РуРrx6	1XCC	2.3	SH_{47}	FF	$\alpha_2(B)$	$(255)^{b}$

^aAn A-type or B-type dimer is noted in parentheses. For octamers and decamers, both interfaces are used.

^bThe publication summarizes a number of structures solved by the Structural Genomics Consortium (SGC; http://sgc.utoronto.ca) and shows only structure snapshots without specific discussion of *PvPrx1a* or *PyPrx6*.

^cThe structure has been solved and deposited in the Protein Data Bank (http://pdb.org) by the SGC and is unpublished to date.

^dThe C_P loop has shifted, as is expected to occur after decamer dissociation.

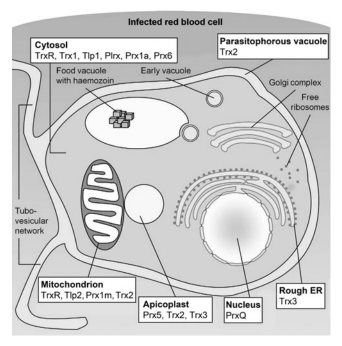


FIG. 7. Compartmentalization of Prxs and related redox cascade enzymes in *Plasmodia*. Adapted from Ref. (121). Plrx, plasmoredoxin; Tlp, thioredoxin-like protein.

*Pf*Prx6 coprecipitated with agarose-bound ferriprotoporphyrin IX (117) and so it was thought to protect against heme detoxification, although *Pf*Prx6 cellular distribution does not match that of heme (62). *Pf*Prx5 is expressed in all erythrocyte stages, with peak mRNA levels in trophozoites [microarray data PlasmoDB (16)], mirroring Grx expression (185).

Biochemical characterizations: All five *P. falciparum* Prxs have been biochemically characterized, along with certain homologs in other mammalian parasites, for example, *Plasmodium yoelii* Prx6 (119).

Prx1a. As expected for a Prx1, PfPrx1a is a doughnut-shaped (α_2)₅ decamer (4) and forms disulfide-linked dimers during its catalytic cycle (118). Mutagenesis confirms the locations of C_P and C_R (186), and proven substrates include H_2O_2 (4), $ONOO^-$ (186), t-butyl-OOH, and cumene-OOH (210). PfPrx1a can be reduced by E. coli Trx (118), plasmodial Trx (4), and Plrx (186). Stopped-flow kinetics studies, with Trx as the reductant, gave k_{cat}/K_m =6.7 × 10⁶ M^- 1s⁻¹ for H_2O_2 (4) and H_2 1 and H_2 2 for H_2 3 and H_2 3 for H_2 4. Structural genomics projects have produced crystal structures for H_2 4 for H_2 5 for H_2 6 for H_2 6. Structural genomics projects have produced crystal structures for H_2 6 for H_2 7 for H_2 7 for H_2 8. Structures for H_2 8 for H_2 9 for

Prx1m. Recombinant *Pf*Prx1m is a functional peroxidase that is reduced by the mitochondrial *Pf*Trx2 about 10-fold more effectively than the cytosolic *Pf*Trx1 (27); neither Grx nor lipoamide could directly support *Pf*Prx1m activity. The crystal structure of *Pf*Prx1m has been solved in its LU disulfide form (Table 2), with the protein in the crystal forming a B-type dimer (27). The mitochondrial targeting sequence (residues 1–19) is disordered and thus seems unimportant in the folding or activity of the mature protein.

Prx6. Despite showing activity in a GS-oxidation protection assay (117, 185), recombinant Prx6 from P. falciparum exhibits low peroxidase activity with both H_2O_2 and t-butyl-OOH. Its functional reducing partners include Grx and a Trx, but not GSH, even though one of its eight Cys residues can be glutathionylated $in\ vitro\ (185)$. The crystal structure of PyPrx6 has been determined by a structural genomics group (Table 2). Interestingly, PyPrx6 in the FF state with a reduced active site is present in the crystal as a B-type dimer, suggesting that not

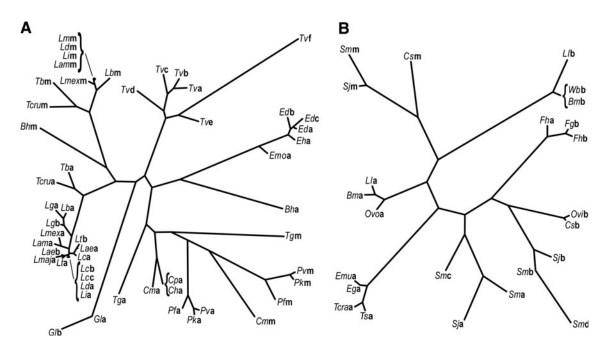


FIG. 8. Relatedness tree of Prx1 subfamily members. Shown are (A) a tree of the Prx1 subfamily members from protozoan parasites, and (B) a tree of the Prx1 subfamily members from helminthic parasites. For each Prx1 we show only its italicized species-specific prefix followed by its suffix in bold type. The tree was generated using Clustal Omega 1.0.2 multiple sequence alignment (231) followed by PHYML3.0 phylogeny calculation (88) and displayed using phylip 3.67: drawtree (76).

all Prx6 subfamily members form decamer-like ring complexes during their catalytic cycle.

Prx5. Recombinant *Pf*Prx5 functions as a 1-Cys Prx and has a preference for the organic ROOH substrates *t*-butyl-OOH and phosphatidylcholine-OOH, with H₂O₂ and cumene-OOH reduced less efficiently (185). Since the apicoplast is the site of fatty acid synthesis, this suggests that *Pf*Prx5 is important for the reduction of fatty acid peroxides. The recycling of *Pf*Prx5 is efficiently carried out by *Pf*Grx, and to a lesser extent *Pf*Trx1 and Plrx (185). The putative apicoplast proteins Trx2 and Trx3 (185) have not been tested. The existence of bacterial Grx-Prx5 fusion proteins (124) suggests that Grx is a plausible physiological redox partner of *Pf*Prx5. The *Pf*Prx5 crystal structure has been determined (Table 2) and is extensively described (222). This structure provided the first view of a C_P doubly overoxidized to C_P-SO₃⁻, thought to be an artifact of long-term storage in an oxidizing environment.

PrxQ. PfPrxQ has an N-terminal Prx domain and an uncharacterized, lysine-rich C-terminal domain (214). The recombinant Prx domain reduces H_2O_2 or cumene-OOH with $K_m \sim 10$ - $30~\mu M$ and a low specific activity of $6~\mu mol/min/mg$ protein. Its preferred reductant appears to be PfGrx1 over PfTrx1 (which does not show saturation). Its nuclear location and genome-wide chromatin association (214) suggests that it could play a sentinel role against oxidative DNA damage.

Toxoplasma. *T. gondii*, the causative agent of toxoplasmosis, infects about one-third of the world's population, causing serious illness in immune-challenged or pregnant individuals. The life cycle involves a bradyzoite stage with sexual reproduction in cats and an asexual, tachyzoite stage in cats, rodents, birds, or humans. The *T. gondii* genome contains four Prxs: Prx1a, Prx1m, Prx5, and Prx6. Except for PrxQ₁ these are a one-to-one match to those found in *Plasmodia*. Unlike *Plasmodia*, *T. gondii* has a functional cytosolic catalase (66).

Localization studies: Immunofluorescence microscopy has localized *Tg*Prx1a and *Tg*Prx6 to the cytosol and *Tg*Prx1m to the mitochondria (141). An independent study confirms this and identifies cytosolic Trx as a Prx reductant (5). Finally, a Prx (presumably *Tg*Prx1m) and an SOD isozyme undergo dual targeting to the apicoplast and mitochondria (204).

Expression levels and other in vivo studies: T. gondii was shown to express Prx1a and Prx1m constitutively and Prx6 in the tachyzoite and bradyzoite stages (141). A study both deleting and overexpressing TgPrx6 and catalase showed that both similarly protected against oxidative stress (141).

Biochemical characterizations: Biochemical characterizations exist for *TgPrx1a* and *TgPrx6*.

Prx1a. Recombinant TgPrx1a turns over H_2O_2 and t-butyl-OOH with a reported K_m in the nanomolar range and $k_{\rm cat}/K_m \sim 10^6-10^7~{\rm M}^{-1}{\rm s}^{-1}$ (5). As no nanomolar substrate concentrations were tested, these K_m values could be artificially low if the recycling reaction is rate limiting. Also, treatment of macrophages in vitro with recombinant TgPrx1a increases their replication rates and influences their developmental fate (159). TgPrx1a is also reported to interact with a histone lysine methyltransferase to influence chromatin structure and regulate the expression of antioxidant defenses (223).

Prx6. Unusual for this subfamily (183), TgPrx6 appears to be a 2-Cys Prx with Cys209 serving as C_R (61). The enzyme appears active by GS protection assay, but direct peroxide reduction using Trx, Grx, or GSH could not be shown. The

protein mostly forms dimers, but also associates in tetramers and hexamers (61). In a drug candidate screen, TgPrx6 was covalently modified at C_P by the drug conoidin, but the inhibition of TgPrx6 was not involved in conoidin bioactivity (95).

Cryptosporidia. Three Cryptosporidia species cause the water-borne disease cryptosporidiosis, characterized by gastroenteritis and diarrhea. The genomes of Cryptosporidium parvum and Cryptosporidium hominis encode only a Prx1a and that of Cryptosporidium muris a Prx1a and a Prx1m. Although the localization of CmPrx1m has not been investigated, its sequence contains a weak mitochondrial localization signal and clusters nearer to the plasmodial Prx1m enzymes than the other cryptosporidial Prx1s (Fig. 8A). The simpler Prx set in Cryptosporidia compared with Plasmodia and Toxoplasma makes sense in light of their further reductive evolution including loss of the apicoplast. Also, the absence of Prx1m in C. parvum and C. hominis is noteworthy in light of their simpler mitochondrion-like organelles (mitosomes) that lack DNA and most ATP synthase subunits and that do not carry out oxidative phosphorylation (169).

The only characterized cryptosporidial Prx is Prx1a from *C. parvum*. Proteomic analysis after high doses of gamma radiation revealed nearly 10-fold elevated CpPrx1a mRNA levels, peaking approximately 48 h after exposure (142). As radiation exposure can produce toxic levels of H_2O_2 (18), this induction may be an oxidative stress response.

Prxs in kinetoplastids

The kinetoplastid parasites (order Trypanosomatida) are named for the kinetoplast, a characteristic granular substructure associated with the basal body of the flagellum and containing the DNA of the lone, extended mitochondrion. They include *Leishmania* species responsible for disfiguring (cutaneous and mucocutaneous) and fatal (visceral) leishmaniases, as well as *Trypanosoma brucei* and *T. cruzi*, respectively the causes of African and American trypanosomiasis (sleeping sickness and Chagas disease).

As intracellular pathogens, the kinetoplastids must defend against ROS and ONOO⁻, with the latter demonstrated to be necessary and sufficient to control *Leishmania donovani* infections in a mouse model (59, 182). Recently, using *T. cruzi*, direct support for ONOO⁻ as an intraphagosomal cytotoxin in host defense has been obtained (9). This implies that enzymes like Prxs capable of reducing ONOO⁻ are crucial for virulence.

The kinetoplastid redox system is comprised of subfamily Prx1 peroxiredoxins (commonly called Txn peroxidases in the literature), their reducing partner Txn, the small-molecule dithiol trypanothione [TS₂, oxidized; T(SH)₂, reduced], and trypanothione reductase (TR), as depicted in Figure 6 (65, 135). Each species with a sequenced genome has one cytosolic Prx1 (*i.e.* Prx1a) and one Prx1m, and we suppose that the other species also have Prx1m enzymes. In these kinetoplastids, from two to seven highly similar, tandem cytosolic Prx1 genes are present. Although few of these are recognized as distinct by our 95% criteria (*e.g.*, only two of seven loci in *Leishmania major*), it is certain that at least some actually are distinct (*e.g.*, *Leishmania aethiopica* and *Leishmania chagasi* Prx1a-type genes are known to be differentially expressed).

Leishmania. The *Leishmania* life cycle involves the development of the infectious promastigote in the gut of the phlebotomine sandfly and development of the disease-causing amastigote in human macrophages.

Localization studies: The L. chagasi enzymes LcPrx1a and LcPrx1b (22), and L. amazonensis, LamPrx1a were localized to the cytosol (101, 149). LdPrx1m has been shown in promastigotes to be found in the kinetoplast region of the mitochondrion and throughout the entire mitochondrion in amastigotes (96). An interesting twist occurs for Leishmania braziliensis, for which a Prx was identified as secreted (55). The exact Prx was not specified, but as it was homologous to LgPrx1b, we denote it LbPrx1a. Because the LbPrx1a sequence has no apparent signal peptide, and also because about 98% of the L. donovani secretome lacks classical N-terminal secretion signals (232), it was suggested that Leishmania may employ nonclassical secretion pathways.

Expression levels and other in vivo studies: In *L. aethiopica*, Prx1a expression is restricted to amastigotes and stationary phase promastigotes, whereas Prx1b is expressed throughout the life cycle (112). Similarly, in *L. chagasi*, Prx1a is expressed in amastigotes, with Prx1b and Prx1c predominantly expressed in promastigotes, the latter at a much lower level (21). Prx1m expression levels in *L. donovani* were shown to increase strongly in promastigotes in the late logarithmic phase and in amastigotes (96). Also, *Lam*Prx1a and *Lam*Prx1m were upregulated by exposure to arsenite (101, 149). In *L. major*, Prx1a is expressed in promastigotes, and accounts for 1%–4% of total cellular protein (133).

Studies in *Leishmania guyanensis* show that oxidative challenges are present throughout normal growth, as *Lg*Prx1a and *Lg*Prx1b populate the disulfide form (*i.e.*, covalent dimers) during all developmental stages. Provocatively, it was observed that more of the disulfide form was seen in an aggressively metastatic *L. guyanensis* strain than in a nonmetastatic strain (1). In another isolate, an unspecified cytosolic *Lg*Prx1 also was identified as a virulence factor (258).

Prx overexpression, as expected, protects parasites from oxidative challenge. Cell culture studies include *Leishmania amazonensis* cells overexpressing *Lam*Prx1a or *Lam*Prx1m (101), and *L. chagasi* cells overexpressing *Lc*Prx1a (21), and *in vivo* results are available for *Leishmania infantum* overexpression of either *Li*Prx1a or *Li*Prx1m (38). In *L. donovani*, overexpression of *Ld*Prx1a in the promastigote results in protection against a combination of H₂O₂ and NO• and increased virulence (106) and overexpression of *Ld*Prx1m decreases H₂O₂-induced apoptosis (96). In *L. infantum*, *Li*Prx1a overexpression preferentially protects against H₂O₂ challenge and *Li*Prx1m overexpression protects only against *t*-butyl-OOH (38).

Biochemical characterizations: Prx1a and Prx1b. The cytoplasmic Prxs from L. chagasi and L. donovani have been biochemically characterized. Recombinant LcPrx1a has been shown to reduce H_2O_2 , alkyl ROOH, ONOO $^-$, and possibly even hydroxyl radicals or nitric oxide (21), whereas LcPrx1b was seen to only reduce H_2O_2 (22). Biochemical characterization of LdPrx1a, done with reducing partners Txn1 and Txn2 from model trypanosome Crithidia fasciculata, revealed its ping-pong reaction kinetics and very high activity with t-butyl-OOH ($k_{\rm cat}/K_{\rm m} = 2.4 \times 10^7 \ {\rm M}^{-1} {\rm s}^{-1}$); specific activity using t-butyl-OOH was similar with H_2O_2 and cumene-OOH, but was relatively low with linoleic acid-OOH and phos-

phatidylcholine-OOH (80). Interestingly, the specific activity of *Ld*Prx1a with *t*-butyl-OOH using the *Cf*Txn1 and *Cf*Txn2 is on the same order of magnitude as that of the native *C. fasciculata* Prx in this system.

Prx1m. Mitochondrial Prxs from *L. donovani* and *L. infantum* have been characterized. *Ld*Prx1m was shown to reduce peroxides and protect DNA from oxidative damage (96). Recombinant *Li*Prx1m had efficient peroxidase activity in a redox system comprised of NADPH, *T. cruzi* TR, T(SH)₂, and *C. fasciculata* Txn1 and Txn2 (35). *Cf*Txn2 bound *Li*Prx1m well ($K_{\rm m}$ =31.9 μ M), and the catalytic efficiency ($k_{\rm cat}/K_{\rm m}$) for *t*-butyl-OOH was 4.1×10^6 M⁻¹ s⁻¹. Curiously, *Li*Prx1m was reported to be inhibited in a time- and concentration-dependent manner by linoleic acid-OOH and phosphatidylcholine-OOH (35).

Trypanosoma. The trypanosomes have complex life cycles involving the tsetse fly (for *T. brucei*) or the triatomine bug (for *T. cruzi*) in which the epimastigote form develops and then differentiates into the infective metacyclic trypomastigote (MT). In the human host, *T. brucei* MTs develop into bloodstream trypomastigotes and disseminate to a variety of tissues (including blood, lymph, and spinal fluid). In contrast, *T. cruzi* MTs develop into amastigotes, and differentiate into trypomastigotes at the insect wound site, and then enter the bloodstream. Interestingly, *T. brucei* and *T. cruzi* lack Grx and Trx (135, 136), and the complete Prx cascade has been shown in *T. brucei* to be essential for parasite survival [reviewed in Ref. (8)] and is considered an important target for drug design. Both *T. brucei* and *T. cruzi* have two Prxs: Prx1a and Prx1m.

Localization studies: As expected, *T. cruzi* and *T. brucei* Prx1a and Prx1m are localized to the cytosol and mitochondrion, respectively (247, 261).

Expression levels and other in vivo studies: TbPrx1a and TbPrx1m are both expressed in the bloodstream and procyclic stages of the parasite (247). Similarly, TcruPrx1a was seen to be expressed throughout the life cycle (203). In support of the role of Prxs in maintaining infectivity, overexpression of TcruPrx1a both protect against exogenous ONOO⁻ and yield higher growth rates under ONOO⁻ challenge (199). These protective effects are lost if the C_P residue is mutated. T. cruzi parasites overexpressing both TcruPrx1a and TcruPrx1m exhibit increased virulence (199) and levels of TcruPrx1a and TcruPrx1m directly correlate with degree of parasitemia in mouse models (200).

Biochemical characterizations: Prx1a. Recombinant *Tb*Prx1a, in the presence of the recombinant native Prx redox cascade components (NADPH, T(SH)₂, and *T. brucei* Txn and TR), efficiently reduced H₂O₂ and could reduce ONOO⁻ with a $k_{\rm cat}/k_{\rm m}=9\times10^5~{\rm M}^{-1}{\rm s}^{-1}$ (249). Similarly, the reduction of ONOO⁻ by *Tcru*Prx1a was reported to be 7.2×10⁵ M⁻¹s⁻¹ (249) and independently around 1×10⁶ M⁻¹s⁻¹ (201). In the latter study, H₂O₂ reduction by *Tcru*Prx1a was very efficient ($k_{\rm cat}/k_{\rm m}=3\times10^7~{\rm M}^{-1}{\rm s}^{-1}$; 201); *t*-butyl-OOH reduction in a different study was much less so ($k_{\rm cat}/k_{\rm m}=3.2\times10^4~{\rm M}^{-1}{\rm s}^{-1}$) (87).

The crystal structure of TcruPrx1a has been determined in its FF thiol form, providing the first structure of an active Prx known to have efficient $ONOO^-$ reductase activity (202). It adopted the expected toroidal pentamer of dimers. Recently, molecular dynamics simulations have explored active-site pK_a values and oligomerization interactions (275).

Prx1m. A recent, comparative kinetics study showed that the remarkably efficient reduction of ONOO⁻ by TcruPrx1m (k_{cat}/K_{m} of 1.8×10^{7} M⁻¹s⁻¹) is about 10-fold greater than that by TcruPrx1a and more than double its own k_{cat}/K_{m} for the reduction of H₂O₂ (201).

Prxs in other selected protozoan parasites of humans

Here we cover localization, *in vivo* studies, and biochemical characterization of Prxs in metamonads (*T. vaginalis* and *Giardia lamblia*), *Naegleria fowleri*, and species of the genus *Entamoeba*.

T. vaginalis. *T. vaginalis* causes the sexually transmitted infection trichomoniasis, considered the most common parasitic infection in industrialized countries. It is a metamonad, which are flagellate anaerobic/microaerobic protozoa with a life cycle including swimming trophozoite and cystic stages. *T. vaginalis* prefers oxygen concentrations below $0.25~\mu M$ and is unable to tolerate oxygen concentrations above $60~\mu M$, or about $5\%~O_2$ (189). As an anaerobe, it lacks mitochondria and instead has a specialized organelle called the hydrogenosome. Hydrogenosomes, particularly well studied in *T. vaginalis*, produce ATP and hydrogen gas using an oxygen-sensitive pyruvate:ferredoxin oxidoreductase and a hydrogenase (179).

To maintain low intracellular oxygen concentrations, *T. vaginalis* employs cytosolic and hydrogenosomal oxidases and SOD (70). *T. vaginalis* lacks detectable Gpx or heme-dependent peroxidase activity (70), but has multiple Prxs: at least six Prx1 subfamily members and two Tpxs. There may be even more, as one of the Prx1 gene sequences filtered by our > 95% identity cutoff appears to represent a distinct gene. Of note, this is the only eukaryote with a Tpx subfamily Prx (183). Proteomics identified *TvT*px-a and a Trx as components of the *T. vaginalis* hydrogenosome (209), and later, two TrxR enzymes bearing novel internal hydrogenosome targeting sequences were also discovered (162). Thus, a complete Prx system exists in the hydrogenosome.

In expression studies of *Tv*Prx1a, a Western blot detected two protein bands of 22 and 20 kDa. The former is the expected size for *Tv*Prx1a; the latter may be a crossreacting Prx. High oxygen or the absence of ascorbate led to elevated Prx1a mRNA and increases in both Western bands. Adding cysteine slightly suppressed Prx1a mRNA levels, abolishing the 22 kDa protein band, but increasing the 20 kDa band (53).

In a proteomic study of antiprotozoal 5-nitroimidazole drugs, *Tv*Prx1a and *Tv*Tpx-b were two of only seven proteins identified as covalently modified by metronidazole and tinidazole (143). A twofold reduction in peroxidase activity was observed in extracts of metronidazole-treated cells, but a similar loss in TrxR activity was seen, so the Prxs are not necessarily implicated. Also, metronidazole-resistant cells had elevated levels of Prx1a, Prx1b, Prx1c, and a protein 96.4% identical to Prx1c (EuPathDB gene ID TVAG_455310), which could be a distinct gene product (143). An siRNA knockdown of an unspecified Prx and TrxR led to no clear detrimental effects although the cell cycle was somewhat extended [according to an abstract (276)].

Only one T. vaginalis Prx, TvPrx1a, has been studied in vitro. Recombinantly expressed TvPrx1a, TvTrxR, and TvTrx, together, were found to effectively reduce H_2O_2 , cumene-OOH, and t-butyl-OOH (53).

G. lamblia. *G. lamblia* [synonymous with *G. intestinalis* (174)], also a metamonad, is the cause of giardiasis, "beaver fever." By our criteria, it has two distinct Prx1 subfamily members. However, we found no associated published studies. The *Gl*Prx1b sequence is strongly predicted by SignalP-HMM (26, 71) to encode a secreted protein. As the *G. lamblia* Prxs are not closely related to those of *T. vaginalis* (Fig. 8A), they need not behave similarly.

N. fowleri. N. fowleri is a free-living amoeba that causes primary amoebic meningoencephalitis, a rare but acute and frequently fatal central nervous system disease. The genome has not yet been sequenced, but an 18-kDa Prx (matching a Prx1 from Aedes aegypti by its N-terminal sequence) was identified as an excretory–secretory pathogenic factor in N. fowleri infection (122). While it was not cloned and so may not be a Prx1, the sequenced genome of the nonpathogenic Naegleria gruberi (84) contains a Prx1 and a Prx5 (NCBI accession numbers XP_002678814 and XP_002680895, respectively). Though neither is characterized, the TargetP server (71) identifies a mitochondrial transit peptide in NgPrx5.

Entamoebae. *Entamoeba histolytica* is an anaerobic protozoan responsible for amoebic dysentery and amoebic liver abscess. Two morphologically indistinguishable, nonpathogenic relatives, *Entamoeba dispar* (63) and *Entamoeba moshkovskii* (7), commonly co-occur with *E. histolytica* and are also discussed here. *E. histolytica* does not possess mitochondria and lacks GSH as well as the enzymes central to GSH metabolism (73). The *E. histolytica* and *E. dispar* genomes are sequenced but not yet fully assembled and, using our 95% identity criteria, have one and three Prx1 subfamily members, respectively, all very similar in sequence (Fig. 8a). With 8 and 20 possible loci in the two species, we suspect that more of these will represent unique *Prx* genes, especially since seven were cloned from a single strain of *Entamoeba nuttalli*, a rhesus monkey parasite (243, 244).

EhPrx1a was initially discovered as a 29-kDa antigen Eh29. It is by far the primary source of free thiol on the protein surface (82), and formed the basis of a vaccine conferring protection in an animal model (236). The protein was shown to be a Prx (30) dependent on Trx and TrxR (207) and could also be reducible by the *E. histolytica* trypanothione–dependent TR system (245). EhPrx1a is 50 times more abundant than the EdPrx1 and is mostly present on the outer surface of cells (poised to counteract host redox defenses), and this may in part explain the invasiveness of E. histolytica versus E. dispar and E. moshkovskii (50). The only studied EdPrx1 seems restricted to the cytoplasm (50) and EmoPrx1a to the cytoplasm and nucleus (48); EhPrx1a may also be present in the cytoplasm (30, 242). EhPrx1a was found to be upregulated about twofold in trophozoites under oxygen stress conditions (3) or exposure to 50 mM Trichostatin A, a histone deacetylase inhibitor (104). Increased expression of EhPrx1a (along with SOD) is also associated with metronidazole resistance (246, 259).

The recombinant EdPrx1 (N-terminally fused to $E.\ coli\ Trx$) yielded a k_{cat}/K_m of $\sim 10^5\ M^{-1}s^{-1}$ for H_2O_2 . Its turnover of various organic ROOHs was 10%–40% less efficient (50). At saturating H_2O_2 , the rate of a corresponding EhPrx1a was about half that of the EdPrx1 (50). Independently, EhPrx1a was found by DNA nicking assay to be slightly less active than EmoPrx1a (48). The Prxs from $E.\ histolytica$ and $E.\ dispar$

have a distinctive \sim 40 residue cysteine-, lysine-, and glutamaterich sequence at the N-terminus, reportedly essential to enzyme activity, although some apparently truncated loci lack this sequence (30).

Prxs in helminths

The helminths are multicellular, parasitic worms, and those examined here are roundworms (Phylum Nematoda) and two classes of flatworms (Phylum Platyhelminthes), namely, tapeworms (Class Cestoda) and flukes (Class Trematoda). In total, 25 Prxs—all in subfamily Prx1 or Prx6—were identified in 17 species. No crystal structures of helminth Prxs have been solved.

One important question is whether, as multicellular eukaryotes, helminths have floodgate-type Prxs (89) that participate in cell signaling *via* an overoxidation shunt. While overoxidation-sensitive Prxs have been documented in *Schistosoma mansoni* (225), there are no recognizable sestrin genes in helminths, including *Caenorhabditis elegans* (248). Also, although *C. elegans* possesses a sestrin homolog and a Prx sensitive to overoxidation, its subsequent reduction is very slow. Furthermore, under ordinary conditions, accumulation of the overoxidized Prx could be detected neither in wild-type *C. elegans* nor in a sestrin deletion strain (248).

Nematoda. Nematodes cause widespread, disabling diseases known as filiariases. Of the five species surveyed, the filarial parasites with genomes sequenced have two or three Prxs each. Of these, Brugia malayi Prx1b and Onchocerca volvulus Prx1a have been expressed in vitro, and found to have peroxidase activity by DNA protection assay (86, 152). These two Prxs are transcribed and translated in larvae and adult worms. In the adult, BmPrx1b was localized to the hypodermis and lateral chord and was not secreted by or at the surface of the larval or adult worms (86). In contrast, OvoPrx1a is found in the larval and adult hypodermis and cuticle and appears to be secreted (152, 278). The closely related homolog BmPrx1a (86) (Fig. 8B) could be likewise surface-localized or secreted, as it is antigenic in mice (10) and in humans (154, 155). The 1-Cys BmPrx6 has been cloned (44) and patented as a vaccine antigen (U.S. Patent 6,352,836); its ortholog from the dog heartworm Dirofilaria immitis is secreted and has been found in developmental stages and localization patterns similar to the filiarial Prx1a enzymes, above (45).

Although no Prx has been identified in *Ascaris lumbricoides* (which lacks a sequenced genome), a closely related Prx1 (accession number Q9NL98) from the swine parasite *Ascaris suum* has been shown to be active *in vitro* by DNA nicking assay and to be expressed in adult female worms (251). Two uncharacterized Prxs (accession numbers YP_198397.1 and YP_197960.1) are found in the genus *Wolbachia*, the *B. malayi* bacterial endosymbiont (83).

Cestoda. Cestodes cause cysticercosis when larvae penetrate the intestinal wall and disseminate into muscle and central nervous system tissues. Although no genome of the four cestodes surveyed has been sequenced, each has one identified Prx. These are all Prx1 subfamily members and cluster by sequence (Fig. 8B). *Echinococcus granulosus* Prx1a is expressed in all tissues of the protoscolex and brood capsules and has DNA protection assay activity (100, 145). Its expression in the juve-

nile stage can be inferred from its isolation from a protoscolex cDNA library (NCBI accession number AAL84833). In cystic echinococcosis patients, *Eg*Prx1a elicits a strong humoral immune response, but since anti-*Eg*Prx1a antibodies are not protective against infection, their role, if any, is unclear (234).

Echninococcus multilocularis Prx1a is expressed at least in the larval stage, having been cloned from a metacestode cDNA library (NCBI accession number BAC11863). Taenia solium and Taenia crassiceps have high H₂O₂ tolerance, and constitutively express Prx1a (252). Antibodies against TsPrx1a were used to show its presence throughout development, as well as the presence of homologs in T. crassiceps cysticerci and Taenia saginata adults (170). Recombinant TsPrx1a was found to have Trx-dependent peroxidase activity with H₂O₂ and cumene-OOH (170).

Trematoda. Trematodes include the blood and liver flukes, which cause schistosomiasis and fascioliasis, respectively. The six species included in our survey are two blood flukes from the genus *Schistosoma* and four liver flukes, two from the genus *Fasciola* and two from the family Opisthorchiidae. All of these have Prxs only from subfamily Prx1. Trematodes do not possess catalase and their Gpx enzymes are generally restricted to the vitelline glands and intrauterine eggs (168) and exhibit little activity against H₂O₂ (156).

Schistosoma: The schistosomal species S. mansoni and Schistosoma japonicum are among the causative agents of schistosomiasis (bilharzia), a persistent infection occurring primarily in poor communities in the tropics, with over 200 million people infected and 700 million at risk. Three schistosomal Prx1 enzymes have been studied, referred to here as Prx1a, Prx1b, and Prx1m (the latter with a mitochondrial targeting sequence). The orthologous enzyme pairs from the two species cluster together in the relatedness tree (Fig. 8B), suggesting their similar function. The PREX database lists two additional Prx1 enzymes in S. mansoni, with SmPrx1c resembling SmPrx1a, and SmPrx1d resembling SmPrx1b (Fig. 8B).

Transcription of the three studied *S. mansoni* Prxs increases during development (224), peaking in the redox-stress tolerant adult forms (224). *Sm*Prx1a was expressed at a higher level than the others. In *S. japonicum*, all Prxs are constitutively expressed throughout development (138). *Sj*Prx1a has been localized to the surface of the miracidium and in the space between the miracidium and eggshell. In the adult fluke, it is found in the tegument. *Sj*Prx1a is secreted in culture and in host tissues surrounding eggs (138). Likewise, *Sm*Prx1a is present in male and female adult worms (140), and is also found in and secreted by eggs (262). Eggs also secrete a reducing partner of *Sm*Prx1a, *Sm*Trx1 (6). *Sj*Prx1b is found in eggs only inside the miracidia; in the adult, it is found in subtegumental tissues, the parenchyma, the vitelline gland, and the gut epithelium (138).

Knockdown of *Sm*Prx1a expression in culture results in sevenfold decreased survival of parasites and increased protein and lipid oxidation (224). Also, whereas knocking down *Sm*Prx1a and *Sm*Prx1b individually in *S. mansoni* sporocysts exhibited no phenotype, a combined *Sm*Prx1a/*Sm*Prx1b knockdown resulted in significantly smaller larvae (176) and reduced survival upon exposure to H₂O₂ or when cultured together with susceptible snail hemocytes (177). Similarly, with *Sj*Prx1a knocked down, larval growth is retarded by organic ROOH treatment, whereas a knockdown of *Sj*Prx1b

had less pronounced effects (139). Study of recombinant SmPrx1a, SmPrx1b, and SmPrx1c showed that each exhibits a catalytic efficiency (k_{cat}/K_m) of $\sim 10^4 - 10^5$ M $^{-1}s^{-1}$ (225). SmPrx1a is robust to oxidative inactivation by H_2O_2 , whereas SmPrx1b and SmPrx1m are sensitive to H_2O_2 and have a preference for organic ROOH substrates (225), so the latter enzymes may play a role in redox signaling, if this occurs in Schistosoma. Also, SmPrx1b and SmPrx1m could be reduced by both Trx and, unusually, GSH (225).

Fasciola: The Fasciola species F. gigantica and F. hepatica lack catalase and have little or no Gpx (43, 160). Three Prx1 genes from the genus have been identified (43). One F. hepatica Prx is secreted and so we denote it Prx1a (229). Although it lacks a secretion signal peptide (71), FhPrx1a is strongly predicted by the Secretome Server (25) to follow a nonclassical eukaryotic secretion pathway and FhPrx1b is not. Although not biochemically characterized, FhPrx1b (which is 99% identical in sequence to FgPrx1b) has a weakly predicted mitochondrial transit peptide of 24 residues at its N-terminus, and FhPrx1a does not (71). FhPrx1a has demonstrated efficacy as a vaccine antigen in an animal model (161), whereas an FgPrx1b-based vaccine has failed in a separate animal trial (211).

FhPrx1a protein is abundant in the parasite and has been cloned, recombinantly expressed, and characterized: it was shown to be active by DNA protection assay; to reduce H_2O_2 , cumene-OOH, and t-butyl-OOH with catalytic efficiencies $(k_{\rm cat}/K_{\rm m}) \sim 5 \times 10^5 \ {\rm M}^{-1} {\rm s}^{-1}$; and to be inactivated via overoxidation at $0.5 \ {\rm mM} \ H_2O_2$ (217, 229). FgPrx1b gene expression is constitutive throughout development, and localized to the gut epithelium, tegument, and reproductive system (43). In very early stage juveniles, the gut epithelium exhibits increased transcript levels (43).

Opisthorchiidae: Despite the lack of sequenced genomes, several Prxs have been identified in the liver flukes Clonorchis sinensis and Opisthorchis viverrini. In these organisms, thus far only a Prx1b and a Prx1m have been found. Both Prx genes are expressed during the entire life cycle, with CsPrx1b transcript levels observed (by semiquantitative RT-PCR) to increase steadily from the juvenile to the adult stages and the CsPrx1m gene expressed at a constant, much lower level (19). Although both proteins can be detected throughout the worm tissues, CsPrx1b is much more abundant and is also secreted (19), as confirmed by a secretion proteomics study (114). The *Ovi*Prx1b gene is expressed in all parasite tissues and the protein may also be secreted, as it was found in host secondary bile ducts too small for infection (239). Thus, OviPrx1b and CsPrx1b appear to combine the properties of the Prx1a and Prx1b of schistosomes. Recombinantly expressed CsPrx1b and CsPrx1m showed activity in a DNA protection assay (239) and yielded $k_{\rm cat}/K_{\rm m}$ values on the order of 10^{-3} – 10^{-4} M⁻¹s⁻¹ for H₂O₂, cumene-OOH, and 13-hydroperoxy-octadecadienoic acid (19). E. coli and yeast reducing partners were used, however, so these probably underestimate their true physiological rates.

Prxs in fungi

The intracellular pathogenic fungi with identified Prxs are *C. neoformans* and the *Encephalitozoon* species, *E. cuniculi* and *E. intestinalis*. The sequenced genomes show that *C. neoformans* has three Prxs (from subfamilies Prx1, Prx6, and PrxQ), and each *Encephalitozoon* species has a single Prx from subfamily Prx1. In *C. neoformans*, *Cn*Prx1a was shown to be in-

ducible by H_2O_2 (164, 165) and the gene encoding each Prx has been individually deleted: the deletion of the Prx1a gene exhibits reduced growth rate, increased sensitivity to ROS and RNS, and decreased virulence in a mouse model; deleting the genes encoding Prx6 and PrxQ yields no related phenotype (164). *E. cuniculi* lacks catalase and has no detectable peroxisome, and is expected to rely on Prx1a along with a GSH-dependent peroxidase and SOD for coping with ROS (75). One proteomics study found EcPrx1a to be expressed in the late sporogonial stages (29).

Prxs as Potential Drug, Vaccine, or Diagnostic Targets

Certain Prx systems seem to be essential for life, but current drugs tend to target enzymes upstream of the Prxs (8, 107). For example, arsenic-based drugs for African trypanosomiasis and antimony-based therapeutics used against leishmaniasis are thought to target T(SH)₂ and TR (56, 74, 178, 269). Similarly, in *L. infantum* a cytosolic Txn has been shown to be essential (215).

Although few selective Prx inhibitors are known, a structure-activity relationship study with derivatives of conoidin A, a covalent inhibitor of T. gondii Prx6, led to analogs with IC₅₀ values ranging from about 10 to $100\,\mu M$ (150). Although conoidin is toxic to T. gondii cells, TgPrx6 is not its molecular target (95). The inhibition of mammalian PrxI and PrxII by conoidin makes it less attractive as an antiparasitic lead compound, yet since few Prx inhibitors are known, Conoidin A and its derivatives may be of value as probes in Prx chemical genetics studies (238).

One validated Prx drug target is TbPrx1a, shown to be essential by RNAi (260). Inspired by the potential of some heteroaromatic quinols as cancer drugs, a series of substituted quinols were tested against T. brucei bloodstream trypomastigotes (134). Three of the derivatives exhibited half-maximal effective concentration (EC₅₀) values below 100 nM, although all exhibited poor specificity for T. brucei cells (only two-to threefold compared with MRC5 human fibroblasts). The most promising compound (PMX464) was shown to be trypanocidal at five times the EC₅₀ value. Other molecular targets of PMX464 were a Gpx-like protein (with an inhibition profile nearly identical to TbPrx1a) and T(SH)₂ (with 14-fold greater affinity than GSH and cysteine).

A likely barrier to the development of clinical Prx inhibitors is the strong conservation of the FF active site among all Prxs (90), which makes specificity an issue. Some authors suggest exploiting species-specific features such as the angle of approach of the C_R in the resolving step of the reaction [human erythrocyte TPx-B compared with TcruPrx1a (202)] or specific characteristics of the dimerization interface(s).

Prxs have been implicated in several instances of drug resistance. Antimony resistance resulted from overexpression of *LdPrx1a* in *L. donovani* promastigotes and amastigotes (106). *TcruPrx1a* and *TcruPrx1m* expression was found to be increased in benznidazole-resistant *T. cruzi* strains (11, 188). Likewise, metronidazole-resistant *E. histolytica* exhibited an upregulated Prx1 (259).

Prxs have shown promise as vaccine antigens. *Lmaj*Prx1a is part of a heterologous prime-boost vaccine that promotes long-term protection against *L. major* infection in mice (237), a vaccine protective against infection in mouse and monkey models (34), and a trivalent vaccine successful in phase I

clinical trials (51). *Ld*Prx1a has been patented as a vaccine antigen candidate (U.S. Patent 7795406).

Prx-based helminth vaccine development has also been pursued. The *Bm*Prx1a gene has been used in a DNA cocktail vaccine efficacious in mice (10) and a peptide derived from a *Bm*Prx1a epitope is strongly immunogenic in mouse and human cell lines, and strongly and selectively immunoreacts to sera from immune and infected individuals (154, 155). An *Fh*Prx1a-derived vaccine protected goats from liver damage after *Fasciola* infection (161), but the *Fg*Prx1b-derived vaccine failed to demonstrate efficacy in buffaloes (211). Also, a multiple antigen peptide combining *S. mansoni* aldolase with *Sm*Prx1a elicited a specific immune response in mice (69). On a related note, Prxs may have diagnostic value in testing for *Leishmania* species (219, 233), *E. granulosus* (158), *F. gigantica* (43, 277), and *Taenia* species (253).

Treatment, prevention, and diagnostic tools are very limited for many parasitic diseases, especially those that disproportionately affect people in resource-poor communities (99). Despite the urgent need for improved interventions, financial support for relevant product development is generally weak (172, 173), limiting the progression of candidate drugs, vaccines, and diagnostics through clinical testing. Fundamental research in parasite biology and biochemistry is therefore essential to ensure that the therapeutic, vaccination, and diagnostic approaches pursued in the clinic are the ones most likely to succeed. Thus, it is encouraging to survey the extensive work on parasite Prxs and to note numerous efforts underway to understand and exploit their properties for the benefit of hundreds of millions of patients and people at high risk of parasitic infection.

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References

- 1. Acestor N, Masina S, Ives A, Walker J, Saravia N, and Fasel N. Resistance to oxidative stress is associated with metastasis in mucocutaneous leishmaniasis. *J Infect Dis* 194: 1160–1167, 2006.
- Adimora N, Jones D, and Kemp M. A model of redox kinetics implicates the thiol proteome in cellular hydrogen peroxide responses. *Antioxid Redox Signal* 13: 731–743, 2010.
- Akbar MA, Chatterjee NS, Sen P, Debnath A, Pal A, Bera T, and Das P. Genes induced by a high-oxygen environment in *Entamoeba histolytica*. Mol Biochem Parasitol 133: 187–196, 2004.
- Akerman S and Muller S. 2-Cys peroxiredoxin PfTrx-Px1 is involved in the antioxidant defence of *Plasmodium falcipar*um. Mol Biochem Parasitol 130: 75–81, 2003.
- Akerman S and Muller S. Peroxiredoxin-linked detoxification of hydroperoxides in *Toxoplasma gondii*. J Biol Chem 280: 564–570, 2005.
- Alger H, Sayed A, Stadecker M, and Williams D. Molecular and enzymatic characterisation of *Schistosoma mansoni* thioredoxin. *Int J Parasitol* 32: 1285–1292, 2002.
- Ali IKM, Hossain MB, Roy S, Ayeh-Kumi PF, Petri WA, Haque R, and Clark CG. Entamoeba moshkovskii infections

- in children, Bangladesh. Emerging Infect Dis 9: 580–584, 2003.
- Alphey MS, König J, and Fairlamb AH. Structural and mechanistic insights into type II trypanosomatid tryparedoxin-dependent peroxidases. *Biochem J* 414: 375, 2008.
- Alvarez M, Peluffo G, Piacenza L, and Radi R. Intraphagosomal peroxynitrite as a macrophage-derived cytotoxin against internalized *Trypanosoma cruzi*: consequences for oxidative killing and role of microbial peroxiredoxins in infectivity. *J Biol Chem* 286: 6627–6640, 2011.
- Anand S, Murugan V, Prabhu P, Anandharaman V, Reddy M, and Kaliraj P. Comparison of immunogenicity, protective efficacy of single and cocktail DNA vaccine of *Brugia* malayi abundant larval transcript (ALT-2) and thioredoxin peroxidase (TPX) in mice. Acta Trop 107: 106–112, 2008.
- Andrade H, Murta S, Chapeaurouge A, Perales J, Nirde P, and Romanha A. Proteomic analysis of *Trypanosoma cruzi* resistance to Benznidazole. *J Proteome Res* 7: 2357–2367, 2008.
- Aran M, Ferrero D, Wolosiuk A, Mora-Garcia S, and Wolosiuk RA. ATP and Mg²⁺ promote the reversible oligomerization and aggregation of chloroplast 2-Cys peroxiredoxin. *J Biol Chem* 286: 23441–23451, 2011.
- 13. Arias D, Carranza P, Lujan H, Iglesias A, and Guerrero S. Immunolocalization and enzymatic functional characterization of the thioredoxin system in *Entamoeba histolytica*. *Free Radic Biol Med* 45: 32–39, 2008.
- 14. Aslett M, Aurrecoechea C, Berriman M, Brestelli J, Brunk BP, Carrington M, Depledge DP, Fischer S, Gajria B, Gao X, Gardner MJ, Gingle A, Grant G, Harb OS, Heiges M, Hertz-Fowler C, Houston R, Innamorato F, Iodice J, Kissinger JC, Kraemer E, Li W, Logan FJ, Miller JA, Mitra S, Myler PJ, Nayak V, Pennington C, Phan I, Pinney DF, Ramasamy G, Rogers MB, Roos DS, Ross C, Sivam D, Smith DF, Srinivasamoorthy G, Stoeckert CJ, Subramanian S, Thibodeau R, Tivey A, Treatman C, Velarde G, and Wang H. TriTrypDB: a functional genomic resource for the Trypanosomatidae. Nucleic Acids Res 38: D457–D462, 2010.
- 15. Aurrecoechea C, Brestelli J, Brunk BP, Carlton JM, Dommer J, Fischer S, Gajria B, Gao X, Gingle A, Grant G, Harb OS, Heiges M, Innamorato F, Iodice J, Kissinger JC, Kraemer E, Li W, Miller JA, Morrison HG, Nayak V, Pennington C, Pinney DF, Roos DS, Ross C, Stoeckert CJ, Sullivan S, Treatman C, and Wang H. GiardiaDB and TrichDB: integrated genomic resources for the eukaryotic protist pathogens Giardia lamblia and Trichomonas vaginalis. Nucleic Acids Res 37: D526–D530, 2009.
- Aurrecoechea C, Brestelli J, Brunk BP, Dommer J, Fischer S, Gajria B, Gao X, Gingle A, Grant G, Harb OS, Heiges M, Innamorato F, Iodice J, Kissinger JC, Kraemer E, Li W, Miller JA, Nayak V, Pennington C, Pinney DF, Roos DS, Ross C, Stoeckert CJ, Treatman C, and Wang H. PlasmoDB: a functional genomic database for malaria parasites. *Nucleic Acids Res* 37: D539–D543, 2009.
- Avila A, Yamada-Ogatta S, da Silva Monteiro V, Krieger M, Nakamura C, de Souza W, and Goldenberg S. Cloning and characterization of the metacyclogenin gene, which is specifically expressed during *Trypanosoma cruzi* metacyclogenesis. *Mol Biochem Parasitol* 117: 169–177, 2001.
- 18. Azzam EI, de Toledo SM, and Little JB. Stress signaling from irradiated to non-irradiated cells. *Curr Cancer Drug Targets* 4: 53–64, 2004.
- 19. Bae Y, Kim S, Lee E, Sohn W, and Kong Y. Identification and biochemical characterization of two novel peroxir-

edoxins in a liver fluke, *Clonorchis sinensis*. *Parasitology* 138: 1143–1153, 2011.

- Baker LM and Poole LB. Catalytic mechanism of thiol peroxidase from *Escherichia coli*. Sulfenic acid formation and overoxidation of essential CYS61. *J Biol Chem* 278: 9203–9211, 2003.
- Barr S and Gedamu L. Cloning and characterization of three differentially expressed peroxidoxin genes from Leishmania chagasi. Evidence for an enzymatic detoxification of hydroxyl radicals. J Biol Chem 276: 34279–34287, 2001
- Barr SD and Gedamu L. Role of peroxidoxins in *Leishmania chagasi* survival. Evidence of an enzymatic defense against nitrosative stress. *J Biochem* 278: 10816–10823, 2003.
- 23. Barranco-Medina S, Lázaro J-J, and Dietz K-J. The oligomeric conformation of peroxiredoxins links redox state to function. *FEBS Lett* 583: 1809–1816, 2009.
- 24. Becker K, Kanzok S, Iozef R, Fischer M, Schirmer R, and Rahlfs S. Plasmoredoxin, a novel redox-active protein unique for malarial parasites. *Eur J Biochem* 270: 1057–1064, 2003.
- Bendtsen JD, Jensen LJ, Blom N, von Heijne G, and Brunak S. Feature-based prediction of non-classical and leaderless protein secretion. *Protein Eng Des Sel* 17: 349–356, 2004.
- Bendtsen JD, Nielsen H, von Heijne G, and Brunak S. Improved prediction of signal peptides: SignalP 3.0. J Mol Biol 340: 783–795, 2004.
- Boucher IW, Mcmillan PJ, Gabrielsen M, Akerman SE, Brannigan JA, Schnick C, Brzozowski AM, Wilkinson AJ, and Müller S. Structural and biochemical characterization of a mitochondrial peroxiredoxin from *Plasmodium falci*parum. Mol Microbiol 61: 948–959, 2006.
- 28. Bradley P and Sibley L. Rhoptries: an arsenal of secreted virulence factors. *Curr Opin Microbiol* 10: 582–587, 2007.
- Brosson D, Kuhn L, Delbac F, Garin J, Vivares CP, and Texier C. Proteomic analysis of the eukaryotic parasite *Encephalitozoon cuniculi* (microsporidia): a reference map for proteins expressed in late sporogonial stages. *Proteomics* 6: 3625–3635, 2006.
- 30. Bruchhaus I, Richter S, and Tannich E. Removal of hydrogen peroxide by the 29 kDa protein of *Entamoeba histolytica*. *Biochem J* 326: 785–789, 1997.
- Buchholz K, Putrianti ED, Rahlfs S, Schirmer RH, Becker K, and Matuschewski K. Molecular genetics evidence for the in vivo roles of the two major NADPH-dependent disulfide reductases in the malaria parasite. J Biol Chem 285: 37388– 37395, 2010.
- Budde H, Flohé L, Hecht H-J, Hofmann B, Stehr M, Wissing J, and Lünsdorf H. Kinetics and redox-sensitive oligomerisation reveal negative subunit cooperativity in tryparedoxin peroxidase of *Trypanosoma brucei brucei*. *Biol Chem* 384: 619–633, 2003.
- Budde H, Flohe L, Hofmann B, and Nimtz M. Verification of the interaction of a tryparedoxin peroxidase with tryparedoxin by ESI-MS/MS. *Biol Chem* 384: 1305–1309, 2003.
- 34. Campos-Neto A, Porrozzi R, Greeson K, Coler R, Webb J, Seiky Y, Reed S, and Grimaldi G. Protection against cutaneous leishmaniasis induced by recombinant antigens in murine and nonhuman primate models of the human disease. *Infect Immun* 69: 4103–4108, 2001.
- Castro H, Budde H, Flohé L, Hofmann B, Lünsdorf H, Wissing J, and Tomás A. Specificity and kinetics of a mitochondrial peroxiredoxin of *Leishmania infantum*. Free Radic Biol Med 33: 1563–1574, 2002.

36. Castro H, Romao S, Gadelha F, and Tomás A. *Leishmania infantum*: provision of reducing equivalents to the mitochondrial tryparedoxin/tryparedoxin peroxidase system. *Exp Parasitol* 120: 421–423, 2008.

- Castro H, Sousa C, Novais M, Santos M, Budde H, Cordeiro-da-Silva A, Flohé L, and Tomás A. Two linked genes of *Leishmania infantum* encode tryparedoxins localised to cytosol and mitochondrion. *Mol Biochem Parasitol* 136: 137–147, 2004.
- Castro H, Sousa C, Santos M, Cordeiro-da-Silva A, Flohé L, and Tomás A. Complementary antioxidant defense by cytoplasmic and mitochondrial peroxiredoxins in *Leishmania* infantum. Free Radic Biol Med 33: 1552–1562, 2002.
- 39. Castro H and Tomás A. Peroxidases of Trypanosomatids. *Antioxid Redox Signal* 10: 1593–1606, 2008.
- Cha M-K, Kim W-C, Lim C-J, Kim K, and Kim I-H. *Escherichia coli* periplasmic thiol peroxidase acts as lipid hydroperoxide peroxidase and the principal antioxidative function during anaerobic growth. *J Biol Chem* 279: 8769– 8778, 2004.
- 41. Chae H, Chung S, and Rhee S. Thioredoxin-dependent peroxide reductase from yeast. *J Biol Chem* 269: 27670–27678, 1994.
- 42. Chae H, Robison K, Poole L, Church G, Storz G, and Rhee S. Cloning and sequencing of thiol-specific antioxidant from mammalian brain: alkyl hydroperoxide reductase and thiol-specific antioxidant define a large family of antioxidant enzymes. *Proc Natl Acad Sci U S A* 91: 7017–7021, 1994.
- 43. Chaithirayanon K and Sobhon P. Molecular cloning and characterization of two genes encoding 2-Cys peroxiredoxins from *Fasciola gigantica*. *Exp Parasitol* 125: 106–113, 2010.
- 44. Chandrashekar R, Curtis K, Lu W, and Weil G. Molecular cloning of an enzymatically active thioredoxin peroxidase from *Onchocerca volvulus*. *Mol Biochem Parasitol* 93: 309–312, 1998.
- 45. Chandrashekar R, Tsuji N, Morales TH, Carmody AB, Ozols VO, Welton J, and Tang L. Removal of hydrogen peroxide by a 1-cysteine peroxiredoxin enzyme of the filarial parasite *Dirofilaria immitis*. *Parasitol Res* 86: 200–206, 2000.
- 46. Chang T-S, Jeong W, Woo HA, Lee SM, Park S, and Rhee SG. Characterization of mammalian sulfiredoxin and its reactivation of hyperoxidized peroxiredoxin through reduction of cysteine sulfinic acid in the active site to cysteine. *J Biol Chem* 279: 50994–51001, 2004.
- 47. Cheng X and Tachibana H. Molecular cloning and characterization of peroxiredoxin from *Entamoeba moshkovskii*. *Arch Med Res* 31: S65–S66, 2000.
- Cheng X-J, Yoshihara E, Takeuchi T, and Tachibana H. Molecular characterization of peroxiredoxin from Entamoeba moshkovskii and a comparison with Entamoeba histolytica. Mol Biochem Parasitol 138: 195–203, 2004.
- Choi HJ, Kang SW, Yang CH, Rhee SG, and Ryu SE. Crystal structure of a novel human peroxidase enzyme at 2.0 A resolution. *Nat Struct Biol* 5: 400–406, 1998.
- 50. Choi M-H, Sajed D, Poole L, Hirata K, Herdman S, Torian BE, and Reed SL. An unusual surface peroxiredoxin protects invasive *Entamoeba histolytica* from oxidant attack. *Mol Biochem Parasitol* 143: 80–89, 2005.
- 51. Coler RN, Goto Y, Bogatzki L, Raman V, and Reed SG. Leish-111f, a recombinant polyprotein vaccine that protects against visceral Leishmaniasis by elicitation of CD4+ T cells. *Infect Immun* 75: 4648–4654, 2007.

- Cooley RB, Arp DJ, and Karplus PA. Evolutionary origin of a secondary structure: pi-helices as cryptic but widespread insertional variations of alpha-helices that enhance protein functionality. J Mol Biol 404: 232–246, 2010.
- 53. Coombs GH, Westrop GD, Suchan P, Puzova G, Hirt RP, Embley TM, Mottram JC, and Müller S. The amitochondriate eukaryote *Trichomonas vaginalis* contains a divergent thioredoxin-linked peroxiredoxin antioxidant system. *J Biol Chem* 279: 5249–5256, 2004.
- 54. Cordeiro-da-Silva A, Cardoso L, Araujo N, Castro H, Tomas A, Rodrigues M, Cabral M, Vergnes B, Sereno D, and Ouaissi A. Identification of antibodies to *Leishmania* silent information regulatory 2 (SIR2) protein homologue during canine natural infections: pathological implications. *Immunol Lett* 86: 155–162, 2003.
- 55. Cuervo P, De Jesus J, Saboia-Vahia L, Mendonça-Lima L, Domont G, and Cupolillo E. Proteomic characterization of the released/secreted proteins of *Leishmania (Viannia) braziliensis* promastigotes. *J Proteomics* 73: 79–92, 2009.
- Cunningham ML, Zvelebil MJ, and Fairlamb AH. Mechanism of inhibition of trypanothione reductase and glutathione reductase by trivalent organic arsenicals. *Eur J Biochem* 221: 285–295, 1994.
- 57. Davis P, Zhang X, Guo J, Townsend R, and Stanley S. Comparative proteomic analysis of two *Entamoeba histolytica* strains with different virulence phenotypes identifies peroxiredoxin as an important component of amoebic virulence. *Mol Microbiol* 61: 1523–1532, 2006.
- 58. de Oliveira A, Ruiz J, Cruz A, Greene L, Rosa J, and Ward R. Subproteomic analysis of soluble proteins of the microsomal fraction from two *Leishmania* species. *Comp Biochem Physiol Part D Genomics Proteomics* 1: 300–308, 2006.
- Dea-Ayuela M, Ordonez-Gutierrez L, and Bolas-Fernandez F. Changes in the proteome and infectivity of *Leishmania* infantum induced by in vitro exposure to a nitric oxide donor. Int J Med Microbiol 299: 221–232, 2009.
- Declercq JP, Evrard C, Clippe A, Stricht DV, Bernard A, and Knoops B. Crystal structure of human peroxiredoxin 5, a novel type of mammalian peroxiredoxin at 1.5 Å resolution. *J Mol Biol* 311: 751–759, 2001.
- Deponte M and Becker K. Biochemical characterization of *Toxoplasma gondii* 1-Cys peroxiredoxin 2 with mechanistic similarities to typical 2-Cys Prx. *Mol Biochem Parasitol* 140: 87–96, 2005.
- Deponte M, Rahlfs S, and Becker K. Peroxiredoxin systems of protozoal parasites. In: *Peroxiredoxin Systems*, edited by Flohé L and Harris JR. New York: Springer, 2007, pp. 219–229.
- Diamond LS and Clark CG. A redescription of Entamoeba histolytica Schaudinn, 1903 (Emended Walker, 1911) separating it from Entamoeba dispar Brumpt, 1925. J Eukaryot Microbiol 40: 340–344, 1993.
- 64. Diaz M, Solari A, and Gonzalez C. Differential expression of *Trypanosoma cruzi* I associated with clinical forms of Chagas disease: overexpression of oxidative stress proteins in acute patient isolate. *J Proteomics* 74: 1673–1682, 2011.
- Diechtierow M and Krauth-Siegel R. A tryparedoxindependent peroxidase protects African trypanosomes from membrane damage. Free Radic Biol Med 51: 856–868, 2011.
- 66. Ding M, Kwok LY, Schluter D, Clayton C, and Soldati D. The antioxidant systems in *Toxoplasma gondii* and the role of cytosolic catalase in defence against oxidative injury. *Mol Microbiol* 51: 47–61, 2004.
- 67. Donnelly S, O'Neill S, Sekiya M, Mulcahy G, and Dalton J. Thioredoxin peroxidase secreted by *Fasciola hepatica* in-

- duces the alternative activation of macrophages. *Infect Immun* 73: 166–173, 2005.
- Donnelly S, Stack C, O'Neill S, Sayed A, Williams D, and Dalton J. Helminth 2-Cys peroxiredoxin drives Th2 responses through a mechanism involving alternatively activated macrophages. FASEB J 22: 4022–4032, 2008.
- El Ridi R and Tallima H. Schistosoma mansoni ex vivo lungstage larvae excretory-secretory antigens as vaccine candidates against schistosomiasis. Vaccine 27: 666–673, 2009.
- Ellis JE, Yarlett N, Cole D, Humphreys MJ, and Lloyd D. Antioxidant defences in the microaerophilic protozoan Trichomonas vaginalis: comparison of metronidazoleresistant and sensitive strains. Microbiology 140: 2489– 2494, 1994.
- 71. Emanuelsson O, Brunak S, von Heijne G, and Nielsen H. Locating proteins in the cell using TargetP, SignalP and related tools. *Nat Protoc* 2: 953–971, 2007.
- Eslami G, Frikha F, Salehi R, Khamesipour A, Hejazi H, and Nilforoushzadeh M. Cloning, expression and dynamic simulation of TRYP6 from *Leishmania major* (MRHO/IR/ 75/ER). *Mol Biol Rep* 38: 3765–3776, 2011.
- 73. Fahey R, Newton G, Arrick B, and Overdank-Bogart T. Entamoeba histolytica: a eukaryote without glutathione metabolism. Science 224: 70–72, 1984.
- 74. Fairlamb AH, Henderson GB, and Cerami A. Trypanothione is the primary target for arsenical drugs against African trypanosomes. *Proc Natl Acad Sci U S A* 86: 2607–2611, 1989.
- 75. Fast NM, Law JS, Williams BAP, and Keeling PJ. Bacterial catalase in the microsporidian *Nosema locustae*: implications for microsporidian metabolism and genome evolution. *Eukaryot cell* 2: 1069–1075, 2003.
- 76. Felsenstein J. PHYLIP—Phylogeny Inference Package (Version 3.2). *Cladistics* 5: 164–166, 1989.
- 77. Ferrer-Sueta G, Manta B, Botti H, Radi R, Trujillo M, and Denicola A. Factors affecting protein thiol reactivity and specificity in peroxide reduction. *Chem Res Toxicol* 24: 434–450, 2011.
- 78. Finzi J, Chiavegatto C, Corat K, Lopez J, Cabrera O, Mielniczki-Pereira A, Colli W, Alves M, and Gadelha F. *Trypanosoma cruzi* response to the oxidative stress generated by hydrogen peroxide. *Mol Biochem Parasitol* 133: 37–43, 2004.
- Flohé L. Changing paradigms in thiology from antioxidant defense toward redox regulation. *Methods Enzymol* 473: 1– 39, 2010.
- 80. Flohé L, Budde H, Bruns K, Castro H, Clos J, Hofmann B, Kansal-Kalavar S, Krumme D, Menge U, Plank-Schumacher K, Sztajer H, Wissing J, Wylegalla C, and Hecht H. Tryparedoxin peroxidase of *Leishmania donovani*: molecular cloning, heterologous expression, specificity, and catalytic mechanism. *Arch Biochem Biophys* 397: 324–335, 2002.
- 81. Flohé L, Budde H, and Hofmann B. Peroxiredoxins in antioxidant defense and redox regulation. *Biofactors* 19: 3–10, 2003.
- 82. Flores BM, Batzer MA, Stein MA, Petersen C, Diedrich DL, and Torian BE. Structural analysis and demonstration of the 29 kDa antigen of pathogenic *Entamoeba histolytica* as the major accessible free thiol-containing surface protein. *Mol Microbiol* 7: 755–763, 1993.
- 83. Foster J, Ganatra M, Kamal I, Ware J, Makarova K, Ivanova N, Bhattacharyya A, Kapatral V, Kumar S, Posfai J, Vincze T, Ingram J, Moran L, Lapidus A, Omelchenko M, Kyrpides N, Ghedin E, Wang S, Goltsman E, Joukov V, Ostrovskaya

O, Tsukerman K, Mazur M, Comb D, Koonin E, and Slatko B. The *Wolbachia* genome of *Brugia malayi*: endosymbiont evolution within a human pathogenic nematode. *PLoS Biol* 3: e121, 2005.

- 84. Fritz-Laylin LK, Prochnik SE, Ginger ML, Dacks JB, Carpenter ML, Field MC, Kuo A, Paredez A, Chapman J, Pham J, Shu S, Neupane R, Cipriano M, Mancuso J, Tu H, Salamov A, Lindquist E, Shapiro H, Lucas S, Grigoriev IV, Cande WZ, Fulton C, Rokhsar DS, and Dawson SC. The genome of *Naegleria gruberi* illuminates early eukaryotic versatility. *Cell* 140: 631–642, 2010.
- 85. Gajria B, Bahl A, Brestelli J, Dommer J, Fischer S, Gao X, Heiges M, Iodice J, Kissinger JC, Mackey AJ, Pinney DF, Roos DS, Stoeckert CJ, Wang H, and Brunk BP. ToxoDB: an integrated *Toxoplasma gondii* database resource. *Nucleic Acids Res* 36: D553–D556, 2008.
- 86. Ghosh I, Eisinger S, Raghavan N, and Scott A. Thioredoxin peroxidases from *Brugia malayi*. *Mol Biochem Parasitol* 91: 207–220, 1998.
- 87. Guerrero S, Lopez J, Steinert P, Montemartini M, Kalisz H, Colli W, Singh M, Alves M, and Flohe L. His-tagged try-paredoxin peroxidase of *Trypanosoma cruzi* as a tool for drug screening. *Appl Microbiol Biotechnol* 53: 410–414, 2000.
- 88. Guindon S, Dufayard JF, Lefort V, Anisimova M, Hordijk W, and Gascuel O. New algorithms and methods to estimate maximum-likelihood phylogenies: assessing the performance of PhyML 3.0. *Syst Biol* 59: 307–321, 2010.
- 89. Hall A, Karplus PA, and Poole LB. Typical 2-Cys peroxiredoxins—structures, mechanisms and functions. *FEBS J* 276: 2469–2477, 2009.
- Hall A, Nelson KJ, Poole LB, and Karplus PA. Structurebased insights into the catalytic power and conformational dexterity of peroxiredoxins. *Antioxid Redox Signal* 15: 795– 815, 2011.
- 91. Hall A, Parsonage D, Horita D, Karplus PA, Poole LB, and Barbar E. Redox-dependent dynamics of a dual thioredoxin fold protein: evolution of specialized folds. *Biochemistry* 48: 5984–5993, 2009.
- 92. Hall A, Parsonage D, Poole LB, and Karplus PA. Structural evidence that peroxiredoxin catalytic power is based on transition-state stabilization. *J Mol Biol* 402: 194–209, 2010.
- 93. Hall A, Sankaran B, Poole LB, and Karplus PA. structural changes common to catalysis in the Tpx peroxiredoxin subfamily. *J Mol Biol* 393: 867–881, 2009.
- 94. Hamza A. Homology modeling and docking mechanism of the mercaptosuccinate and methotrexate to *P. falciparum* 1-Cys peroxiredoxin: a preliminary molecular study. *J Biomol Struct Dyn* 20: 7–20, 2002.
- 95. Haraldsen J, Liu G, Botting C, Walton J, Storm J, Phalen T, Kwok L, Soldati-Favre D, Heintz N, Müller S, Westwood N, and Ward G. Identification of conoidin A as a covalent inhibitor of peroxiredoxin II. Org Biomol Chem 7: 3040, 2009.
- Harder S, Bente M, Isermann K, and Bruchhaus I. Expression of a mitochondrial peroxiredoxin prevents programmed cell death in *Leishmania donovani*. Eukaryot Cell 5: 861–870, 2006.
- 97. Hattori F and Oikawa S. Peroxiredoxins in the central nervous system. In: *Peroxiredoxin Systems*, edited by Flohé L and Harris JR. New York: Springer, 2007, pp. 357–374.
- 98. Heiges M, Wang H, Robinson E, Aurrecoechea C, Gao X, Kaluskar N, Rhodes P, Wang S, He C-Z, Su Y, Miller J, Kraemer E, and Kissinger JC. CryptoDB: a *Cryptosporidium* bioinformatics resource update. *Nucleic Acids Res* 34: D419–D422, 2006.

 Hotez P and Pecoul B. "Manifesto" for advancing the control and elimination of neglected tropical diseases. PLoS Negl Trop Dis 4: e718, 2010.

- 100. Hou Q, Wang H, Zhang Z, Cao W, Zhang F, and Zhang W. [Immunolocalization of the antioxidant protein TPx of Echinococcus granulosus] (abstract of Chinese-language article). Xi Bao Yu Fen Zi Mian Yi Xue Za Zhi 23: 998–1000, 2007.
- 101. Hsu J-Y, Lin Y-C, Chiang S-C, and Lee S. Divergence of trypanothione-dependent tryparedoxin cascade into cytosolic and mitochondrial pathways in arsenite-resistant variants of *Leishmania amazonensis*. Mol Biochem Parasitol 157: 193–204, 2008.
- 102. Hughes M, Lee C, Holm C, Ghosh S, Mills A, Lockhart L, Reed S, and Mann B. Identification of *Entamoeba histolytica* thiol-specific antioxidant as a GalNAc lectin-associated protein. *Mol Biochem Parasitol* 127: 113–120, 2003.
- 103. Hugo M, Turell L, Manta B, Botti H, Monteiro G, Netto L, Alvarez B, Radi R, and Trujillo M. Thiol and sulfenic acid oxidation of AhpE, the one-cysteine peroxiredoxin from Mycobacterium tuberculosis: kinetics, acidity constants, and conformational dynamics. Biochemistry 48: 9416–9426, 2009.
- 104. Isakov E, Siman-Tov R, Weber C, Guillen N, and Ankri S. Trichostatin A regulates peroxiredoxin expression and virulence of the parasite Entamoeba histolytica. Mol Biochem Parasitol 158: 82–94, 2008.
- 105. Ishii T and Yanagawa T. Stress-induced peroxiredoxins. In: Peroxiredoxin Systems, edited by Flohé L and Harris JR. New York: Springer, 2007, pp. 375–384.
- 106. Iyer J, Kaprakkaden A, Choudhary M, and Shaha C. Crucial role of cytosolic tryparedoxin peroxidase in *Leishmania donovani* survival, drug response and virulence. *Mol Microbiol* 68: 372–391, 2008.
- Jaeger T and Flohe L. The thiol-based redox networks of pathogens: unexploited targets in the search for new drugs. *Biofactors* 27: 109–120, 2006.
- 108. Jefferies J, Campbell A, van Rossum A, Barrett J, and Brophy P. Proteomic analysis of *Fasciola hepatica* excretory-secretory products. *Proteomics* 1: 1128–1132, 2001.
- 109. Jeong W, Cha MK, and Kim IH. Thioredoxin-dependent hydroperoxide peroxidase activity of bacterioferritin comigratory protein (BCP) as a new member of the thiolspecific antioxidant protein (TSA)/Alkyl hydroperoxide peroxidase C (AhpC) family. J Biol Chem 275: 2924–2930, 2000.
- 110. Jiménez-Delgadillo B, Chaudhuri PP, Baylón-Pacheco L, López-Monteon A, Talamás-Rohana P, and Rosales-Encina JL. Entamoeba histolytica: cDNAs cloned as 30 kDa collagenbinding proteins (CBP) belong to an antioxidant molecule family. Protection of hamsters from amoebic liver abscess by immunization with recombinant CBP. Exp Parasitol 108: 7–17, 2004.
- 111. Jing B, Deng S, Zhang R, and Zhang J. [Comparative proteomic analysis of the promastigotes and amastigotes of *Leishmania donovani*] (abstract of Chinese-language article). *Zhongguo Ji Sheng Chong Xue Yu Ji Sheng Chong Bing Za Zhi* 27: 102–106, 2009.
- 112. Jirata D, Kuru T, Genetu A, Barr S, Hailu A, Aseffa A, and Gedamu L. Identification, sequencing and expression of peroxidoxin genes from *Leishmania aethiopica*. *Acta Trop* 99: 88–96, 2006.
- 113. Jönsson T and Lowther W. The peroxiredoxin repair proteins. In: *Peroxiredoxin Systems*, edited by Flohé L and Harris JR. New York: Springer, 2007, pp. 115–141.

- 114. Ju J, Joo H, Lee M, Cho S, Cheun H, Kim J, Lee Y, Lee K, Sohn W, Kim D, Kim I, Park B, and Kim T. Identification of a serodiagnostic antigen, legumain, by immunoproteomic analysis of excretory-secretory products of *Clonorchis sinensis* adult worms. *Proteomics* 9: 3066–3078, 2009.
- 115. Kang S, Baines I, and Rhee S. Characterization of a mammalian peroxiredoxin that contains one conserved cysteine. *J Biol Chem* 273: 6303–6311, 1998.
- Kang S, Rhee S, Chang T-S, Jeong W, and Choi M. 2-Cys peroxiredoxin function in intracellular signal transduction: therapeutic implications. *Trends Mol Med* 11: 571–578, 2005.
- 117. Kawazu S, Ikenoue N, Takemae H, Komaki-Yasuda K, and Kano S. Roles of 1-Cys peroxiredoxin in haem detoxification in the human malaria parasite *Plasmodium falciparum*. *FEBS J* 272: 1784–1791, 2005.
- 118. Kawazu S, Komaki K, Tsuji N, Kawai S, Ikenoue N, Hatabu T, Ishikawa H, Matsumoto Y, Himeno K, and Kano S. Molecular characterization of a 2-Cys peroxiredoxin from the human malaria parasite *Plasmodium falciparum*. *Mol Biochem Parasitol* 116: 73–79, 2001.
- 119. Kawazu S, Nozaki T, Tsuboi T, Nakano Y, Komaki-Yasuda K, Ikenoue N, Torii M, and Kano S. Expression profiles of peroxiredoxin proteins of the rodent malaria parasite *Plasmodium yoelii. Int J Parasitol* 33: 1455–1461, 2003.
- 120. Kawazu S, Tsuji N, Hatabu T, Kawai S, Matsumoto Y, and Kano S. Molecular cloning and characterization of a peroxiredoxin from the human malaria parasite *Plasmodium falciparum*. *Mol Biochem Parasitol* 109: 165–169, 2000.
- 121. Kehr S, Sturm N, Rahlfs S, Przyborski JM, and Becker K. Compartmentation of redox metabolism in malaria parasites. *PLoS Pathog* 6: e1001242, 2010.
- 122. Kim J, Yang A, Sohn H, Kim D, Song K, and Shin H. Immunodominant antigens in *Naegleria fowleri* excretory—secretory proteins were potential pathogenic factors. *Parasitol Res* 105: 1675–1681, 2009.
- 123. Kim K, Kim IH, Lee KY, Rhee SG, and Stadtman ER. The isolation and purification of a specific "protector" protein which inhibits enzyme inactivation by a thiol/Fe(III)/O2 mixed-function oxidation system. *J Biochem* 263: 4704–4711, 1988.
- 124. Kim SJ, Woo JR, Hwang YS, Jeong DG, Shin DH, Kim K, and Ryu SE. The tetrameric structure of Haemophilus influenza hybrid Prx5 reveals interactions between electron donor and acceptor proteins. *J Biochem* 278: 10790–10798, 2003.
- 125. Kima P, Bonilla J, Cho E, Ndjamen B, Canton J, Leal N, and Handfield M. Identification of *Leishmania* proteins preferentially released in infected cells using change mediated antigen technology (CMAT). *PLoS Negl Trop Dis* 4: pii e842, 2010.
- Klomsiri C, Karplus PA, and Poole LB. Cysteine-based redox switches in enzymes. *Antioxid Redox Signal* 14: 1065– 1077, 2011.
- 127. Klotz FW, Hadley TJ, Aikawa M, Leech J, Howard RJ, and Miller LH. A 60-kDa *Plasmodium falciparum* protein at the moving junction formed between merozoite and erythrocyte during invasion. *Mol Biochem Parasitol* 36: 177–185, 1989.
- 128. Knoops B, Loumaye E, and Van Der Eecken V. Evolution of the peroxiredoxins. In: *Peroxiredoxin Systems*, edited by Flohé L and Harris JR. New York: Springer, 2007, pp. 27–40.
- 129. Komaki-Yasuda K, Kawazu S, and Kano S. Disruption of the *Plasmodium falciparum* 2-Cys peroxiredoxin gene ren-

- ders parasites hypersensitive to reactive oxygen and nitrogen species. FEBS Lett 547: 140–144, 2003.
- 130. Komaki-Yasuda K, Okuwaki M, Kano S, Nagata K, and Kawazu S. 5' sequence- and chromatin modification-dependent gene expression in *Plasmodium falciparum* erythrocytic stage. *Mol Biochem Parasitol* 162: 40–51, 2008.
- 131. Koncarevic S, Rohrbach P, Deponte M, Krohne G, Prieto JH, Yates J, Rahlfs S, and Becker K. The malarial parasite *Plasmodium falciparum* imports the human protein peroxiredoxin 2 for peroxide detoxification. *Proc Natl Acad Sci U S A* 106: 13323–13328, 2009.
- 132. Kong W, Shiota S, Shi Y, Nakayama H, and Nakayama K. A novel peroxiredoxin of the plant *Sedum lineare* is a homologue of *Escherichia coli* bacterioferritin co-migratory protein (Bcp). *Biochem J* 351: 107–114, 2000.
- 133. Konig J and Fairlamb A. A comparative study of type I and type II tryparedoxin peroxidases in *Leishmania major*. *FEBS J* 274: 5643–5658, 2007.
- 134. Konig J, Wyllie S, Wells G, Stevens M, Wyatt P, and Fairlamb A. Antitumor quinol PMX464 is a cytocidal antitrypanosomal inhibitor targeting trypanothione metabolism. *J Biol Chem* 286: 8523–8533, 2011.
- 135. Krauth-Siegel L, Comini M, and Schlecker T. The trypanothione system. In: *Peroxiredoxin Systems*, edited by Flohé L and Harris JR. New York: Springer, 2007, pp. 231–251.
- 136. Krauth-Siegel R, Bauer H, and Schirmer R. Dithiol proteins as guardians of the intracellular redox milieu in parasites: old and new drug targets in trypanosomes and malariacausing plasmodia. Angew Chem Int Ed Engl 44: 690–715, 2005.
- 137. Krnajski Z, Walter R, and Müller S. Isolation and functional analysis of two thioredoxin peroxidases (peroxiredoxins) from *Plasmodium falciparum*. *Mol Biochem Parasitol* 113: 303–308, 2001.
- 138. Kumagai T, Osada Y, and Kanazawa T. 2-Cys peroxiredoxins from *Schistosoma japonicum*: the expression profile and localization in the life cycle. *Mol Biochem Parasitol* 149: 135–143, 2006.
- 139. Kumagai T, Osada Y, Ohta N, and Kanazawa T. Peroxiredoxin-1 from *Schistosoma japonicum* functions as a scavenger against hydrogen peroxide but not nitric oxide. *Mol Biochem Parasitol* 164: 26–31, 2009.
- 140. Kwatia M, Botkin D, and Williams D. Molecular and enzymatic characterization of *Schistosoma mansoni* thioredoxin peroxidase. *J Parasitol* 86: 908–915, 2000.
- 141. Kwok L, Schlüter D, Clayton C, and Soldati D. The antioxidant systems in *Toxoplasma gondii* and the role of cytosolic catalase in defence against oxidative injury. *Mol Microbiol* 51: 47–61, 2003.
- 142. Lee SU, Joung M, Nam T, Park WY, Ji YH, and Yu JR. *Cryptosporidium parvum*: radiation-induced alteration of the oocyst proteome. *Exp Parasitol* 127: 25–30, 2011.
- 143. Leitsch D, Kolarich D, Binder M, Stadlmann J, Altmann F, and Duchene M. *Trichomonas vaginalis*: metronidazole and other nitroimidazole drugs are reduced by the flavin enzyme thioredoxin reductase and disrupt the cellular redox system. Implications for nitroimidazole toxicity and resistance. *Mol Microbiol* 72: 518–536, 2009.
- 144. Levick M, Tetaud E, Fairlamb A, and Blackwell J. Identification and characterisation of a functional peroxidoxin from *Leishmania major*. *Mol Biochem Parasitol* 96: 125–137, 1998.
- 145. Li J, Zhang W, Loukas A, Lin R, Ito A, Zhang L, Jones M, and McManus D. Functional expression and characteriza-

tion of *Echinococcus granulosus* thioredoxin peroxidase suggests a role in protection against oxidative damage. *Gene* 326: 157–165, 2004.

- 146. Li S, Peterson N, Kim M, Kim C, Hung L, Yu M, Lekin T, Segelke B, Lott J, and Baker E. Crystal Structure of AhpE from Mycobacterium tuberculosis, a 1-Cys peroxiredoxin. J Mol Biol 346: 1035–1046, 2005.
- 147. Liao S-J, Yang C-Y, Chin K-H, Wang AHJ, and Chou S-H. Insights into the alkyl peroxide reduction pathway of *Xanthomonas campestris* bacterioferritin comigratory protein from the trapped intermediate-ligand complex structures. *J Mol Biol* 390: 951–966, 2009.
- 148. Lim YS, Cha MK, Kim HK, Uhm TB, Park JW, Kim K, and Kim IH. Removals of hydrogen peroxide and hydroxyl radical by thiol-specific antioxidant protein as a possible role in vivo. Biochem Biophys Res Commun 192: 273–280, 1993.
- 149. Lin Y-C, Hsu J-Y, Chiang S-C, and Lee S. Distinct over-expression of cytosolic and mitochondrial tryparedoxin peroxidases results in preferential detoxification of different oxidants in arsenite-resistant *Leishmania amazonensis* with and without DNA amplification. *Mol Biochem Parasitol* 142: 66–75, 2005.
- 150. Liu G, Botting C, Evans K, Walton J, Xu G, Slawin A, and Westwood N. Optimisation of conoidin A, a peroxiredoxin inhibitor. *ChemMedChem* 5: 41–45, 2010.
- 151. Lopez J, Carvalho T, de Souza W, Flohe L, Guerrero S, Montemartini M, Kalisz H, Nogoceke E, Singh M, Alves M, and Colli W. Evidence for a trypanothione-dependent peroxidase system in *Trypanosoma cruzi. Free Radic Biol Med* 28: 767–772, 2000.
- 152. Lu W, Egerton G, Bianco A, and Williams S. Thioredoxin peroxidase from *Onchocerca volvulus*: a major hydrogen peroxide detoxifying enzyme in filarial parasites. *Mol Biochem Parasitol* 91: 221–235, 1998.
- 153. MacFarlane R and Singh U. Identification of differentially expressed genes in virulent and nonvirulent *Entamoeba* species: potential implications for amebic pathogenesis. *Infect Immun* 74: 340–351, 2006.
- 154. Madhumathi J, Anugraha G, Prince P, Pradiba D, and Kaliraj P. Proliferative responses of *Brugia malayi* TPX-1 and its epitopic peptide(29–43) in an endemic population of human lymphatic filariasis. *Microbes Infect* 13: 602–606, 2011.
- 155. Madhumathi J, Prince P, Gayatri S, Aparnaa R, and Kaliraj P. Identification of a highly immunoreactive epitope of *Brugia malayi* TPx recognized by the endemic sera. *J Parasitol* 96: 1228–1229, 2010.
- 156. Maiorino M, Roche C, Kiess M, Koenig K, Gawlik D, Matthes M, Naldini E, Pierce R, and Flohe L. A seleniumcontaining phospholipid-hydroperoxide glutathione peroxidase in *Schistosoma mansoni*. Eur J Biochem 238: 838–844, 1996.
- 157. Manevich Y and Fisher AB. Peroxiredoxin 6, a 1-Cys peroxiredoxin, functions in antioxidant defense and lung phospholipid metabolism. *Free Radic Biol Med* 38: 1422–1432, 2005.
- 158. Margutti P, Ortona E, Delunardo F, Tagliani A, Profumo E, Rigano R, Buttari B, Teggi A, and Siracusano A. Thioredoxin peroxidase from *Echinococcus granulosus*: a candidate to extend the antigenic panel for the immunodiagnosis of human cystic echinococcosis. *Diagn Microbiol Infect Dis* 60: 279–285, 2008.
- Marshall E, Elshekiha H, Hakimi M-A, and Flynn R. Toxoplasma gondii peroxiredoxin promotes altered macrophage

- function, caspase-1-dependent IL-1 β secretion enhances parasite replication. *Vet Res* 42: 80, 2011.
- 160. McGonigle S, Curley G, and Dalton J. Cloning of peroxiredoxin, a novel antioxidant enzyme, from the helminth parasite Fasciola hepatica. Parasitology 115: 101–104, 1997.
- 161. Mendes R, Perez-Ecija R, Zafra R, Buffoni L, Martinez-Moreno A, Dalton J, Mulcahy G, and Perez J. Evaluation of hepatic changes and local and systemic immune responses in goats immunized with recombinant Peroxiredoxin (Prx) and challenged with Fasciola hepatica. Vaccine 28: 2832–2840, 2010
- 162. Mentel M, Zimorski V, Haferkamp P, Martin W, and Henze K. Protein import into hydrogenosomes of *Trichomonas vaginalis* involves both N-terminal and internal targeting signals: a case study of thioredoxin reductases. *Eukaryot Cell* 7: 1750–1757, 2008.
- 163. Mielniczki-Pereira A, Chiavegatto C, Lopez J, Colli W, Alves M, and Gadelha F. *Trypanosoma cruzi* strains, Tulahuen 2 and Y, besides the difference in resistance to oxidative stress, display differential glucose-6-phosphate and 6-phosphogluconate dehydrogenases activities. *Acta Trop* 101: 54–60, 2007.
- 164. Missall T, Lodge J, and Mcewen J. Mechanisms of resistance to oxidative and nitrosative stress: implications for fungal survival in mammalian hosts. *Eukaryot Cell* 3: 835–846, 2004.
- 165. Missall T, Moran J, Corbett J, and Lodge J. Distinct stress responses of two functional laccases in *Cryptococcus neo*formans are revealed in the absence of the thiol-specific antioxidant Tsa1. Eukaryot Cell 4: 202–208, 2005.
- 166. Missall TA, Pusateri ME, Donlin MJ, Chambers KT, Corbett JA, and Lodge JK. Posttranslational, translational, and transcriptional responses to nitric oxide stress in *Cryptococcus neoformans*: implications for virulence. *Eukaryot Cell* 5: 518–529, 2006.
- 167. Missall TA, Pusateri ME, and Lodge JK. Thiol peroxidase is critical for virulence and resistance to nitric oxide and peroxide in the fungal pathogen, *Cryptococcus neoformans*. *Mol Microbiol* 51: 1447–1458, 2004.
- 168. Mkoji GM, Smith JM, and Prichard RK. Antioxidant systems in *Schistosoma mansoni*: correlation between susceptibility to oxidant killing and the levels of scavengers of hydrogen peroxide and oxygen free radicals. *Int J Parasitol* 18: 661–666, 1988.
- 169. Mogi T and Kita K. Diversity in mitochondrial metabolic pathways in parasitic protists *Plasmodium* and *Cryptospor-idium*. *Parasitol Int* 59: 305–312, 2010.
- 170. Molina-López J, Jiménez L, Ochoa-Sánchez A, and Landa A. Molecular cloning and characterization of a 2-Cys peroxiredoxin from *Taenia solium*. J Parasitol 92: 796–802, 2006.
- 171. Monteiro G, Horta B, Pimenta D, Augusto O, and Netto L. Reduction of 1-Cys peroxiredoxins by ascorbate changes the thiol-specific antioxidant paradigm, revealing another function of vitamin C. *Proc Natl Acad Sci U S A* 104: 4886– 4891, 2007.
- 172. Moran M. Funding for research and development and the financial crisis. *Lancet Infect Dis* 10: 214–215, 2010.
- 173. Moran M, Guzman J, Ropars A-L, McDonald A, Jameson N, Omune B, Ryan S, and Wu L. Neglected disease research and development: how much are we really spending? *PLoS Med* 6: e30, 2009.
- 174. Morrison HG, McArthur AG, Gillin FD, Aley SB, Adam RD, Olsen GJ, Best AA, Cande WZ, Chen F, Cipriano MJ, Davids BJ, Dawson SC, Elmendorf HG, Hehl AB, Holder

- ME, Huse SM, Kim UU, Lasek-Nesselquist E, Manning G, Nigam A, Nixon JE, Palm D, Passamaneck NE, Prabhu A, Reich CI, Reiner DS, Samuelson J, Svard SG, and Sogin ML. Genomic minimalism in the early diverging intestinal parasite *Giardia lamblia*. *Science* 317: 1921–1926, 2007.
- 175. Motyka S, Drew M, Yildirir G, and Englund P. Over-expression of a cytochrome b5 reductase-like protein causes kinetoplast DNA loss in *Trypanosoma brucei*. *J Biol Chem* 281: 18499–18506, 2006.
- 176. Mourao M, Dinguirard N, Franco G, and Yoshino T. Phenotypic screen of early-developing larvae of the blood fluke, *Schistosoma mansoni*, using RNA interference. *PLoS Negl Trop Dis* 3: e502, 2009.
- 177. Mourao Mde M, Dinguirard N, Franco G, and Yoshino T. Role of the endogenous antioxidant system in the protection of *Schistosoma mansoni* primary sporocysts against exogenous oxidative stress. *PLoS Negl Trop Dis* 3: e550, 2009.
- 178. Mukhopadhyay R, Dey S, Xu N, Gage D, Lightbody J, Ouellette M, and Rosen BP. Trypanothione overproduction and resistance to antimonials and arsenicals in *Leishmania*. *Proc Natl Acad Sci U S A* 93: 10383–10387, 1996.
- 179. Müller M. The hydrogenosome. *J Gen Microbiol* 139: 2879–2889, 1993.
- Müller S. Redox and antioxidant systems of the malaria parasite *Plasmodium falciparum*. Mol Microbiol 53: 1291–1305, 2004
- 181. Murphy S, Samuel B, Harrison T, Speicher K, Speicher D, Reid M, Prohaska R, Low P, Tanner M, Mohandas N, and Haldar K. Erythrocyte detergent-resistant membrane proteins: their characterization and selective uptake during malarial infection. *Blood* 103: 1920–1928, 2004.
- 182. Murray H and Nathan C. Macrophage microbicidal mechanisms *in vivo*: reactive nitrogen versus oxygen intermediates in the killing of intracellular visceral *Leishmania donovani*. *J Exp Med* 189: 741–746, 1999.
- 183. Nelson KJ, Knutson ST, Soito L, Klomsiri C, Poole LB, and Fetrow JS. Analysis of the peroxiredoxin family: using active-site structure and sequence information for global classification and residue analysis. *Proteins* 79: 947–964, 2010.
- 184. Netto L, Chae H, Kang S, Rhee S, and Stadtman E. Removal of hydrogen peroxide by thiol-specific antioxidant enzyme (TSA) is involved with its antioxidant properties. TSA possesses thiol peroxidase activity. *J Biol Chem* 271: 15315–15321, 1996.
- Nickel C, Rahlfs S, Deponte M, Koncarevic S, and Becker K. Thioredoxin networks in the malarial parasite *Plasmodium falciparum*. *Antioxid Redox Signal* 8: 1227–1239, 2006.
- 186. Nickel C, Trujillo M, Rahlfs S, Deponte M, Radi R, and Becker K. *Plasmodium falciparum* 2-Cys peroxiredoxin reacts with plasmoredoxin and peroxynitrite. *Biol Chem* 386: 1129–1136, 2005.
- 187. Nogoceke E, Gommel D, Kiess M, Kalisz H, and Flohe L. A unique cascade of oxidoreductases catalyses trypanothione-mediated peroxide metabolism in *Crithidia fasci*culata. Biol Chem 378: 827–836, 1997.
- 188. Nogueira F, Ruiz J, Robello C, Romanha A, and Murta S. Molecular characterization of cytosolic and mitochondrial tryparedoxin peroxidase in *Trypanosoma cruzi* populations susceptible and resistant to benznidazole. *Parasitol Res* 104: 835–844, 2009
- Paget TA and Lloyd D. Trichomonas vaginalis requires traces of oxygen and high concentrations of carbon dioxide for optimal growth. Mol Biochem Parasitol 41: 65–72, 1990.

- 190. Parsonage D, Karplus PA, and Poole LB. Substrate specificity and redox potential of AhpC, a bacterial peroxiredoxin. *Proc Natl Acad Sci U S A* 105: 8209–8214, 2008.
- 191. Parsonage D, Youngblood DS, Sarma GN, Wood ZA, Karplus PA, and Poole LB. Analysis of the link between enzymatic activity and oligomeric state in AhpC, a bacterial peroxiredoxin. *Biochemistry* 44: 10583–10592, 2005.
- 192. Pedone E, Limauro D, D'Ambrosio K, De Simone G, and Bartolucci S. Multiple catalytically active thioredoxin folds: a winning strategy for many functions. *Cell Mol Life Sci* 67: 3797–3814, 2010.
- 193. Pedrajas J, Miranda-Vizuete A, Javanmardy N, Gustafsson J, and Spyrou G. Mitochondria of *Saccharomyces cerevisiae* contain one-conserved cysteine type peroxiredoxin with thioredoxin peroxidase activity. *J Biol Chem* 275: 16296–16301, 2000.
- 194. Pescher P, Blisnick T, Bastin P, and Spath G. Quantitative proteome profiling informs on phenotypic traits that adapt *Leishmania donovani* for axenic and intracellular proliferation. *Cell Microbiol* 13: 978–991, 2011.
- 195. Peskin A, Low F, Paton L, and Maghzal G. The high reactivity of peroxiredoxin 2 with H_2O_2 is not reflected in its reaction with other oxidants and thiol reagents. *J Biol Chem* 282: 11885–11892, 2007.
- 196. Peskin AV, Cox AG, Nagy P, Morgan PE, Hampton MB, Davies MJ, and Winterbourn CC. Removal of amino acid, peptide and protein hydroperoxides by reaction with peroxiredoxins 2 and 3. *Biochem J* 432: 313–321, 2010.
- 197. Phalen T, Weirather K, Deming P, Anathy V, Howe A, Van Der Vliet A, Jonsson T, Poole L, and Heintz N. Oxidation state governs structural transitions in peroxiredoxin II that correlate with cell cycle arrest and recovery. *J Cell Biol* 175: 779–789, 2006.
- 198. Piacenza L, Alvarez M, Peluffo G, and Radi R. Fighting the oxidative assault: the *Trypanosoma cruzi* journey to infection. *Curr Opin Microbiol* 12: 415–421, 2009.
- 199. Piacenza L, Peluffo G, Alvarez M, Kelly J, Wilkinson S, and Radi R. Peroxiredoxins play a major role in protecting *Trypanosoma cruzi* against macrophage- and endogenously-derived peroxynitrite. *Biochem J* 410: 359–368, 2008.
- 200. Piacenza L, Zago M, Peluffo G, Alvarez M, Basombrio M, and Radi R. Enzymes of the antioxidant network as novel determiners of *Trypanosoma cruzi* virulence. *Int J Parasitol* 39: 1455–1464, 2009.
- 201. Piñeyro M, Arcari T, Robello C, Radi R, and Trujillo M. Tryparedoxin peroxidases from *Trypanosoma cruzi*: High efficiency in the catalytic elimination of hydrogen peroxide and peroxynitrite. *Arch Biochem Biophys* 507: 287–295, 2011.
- 202. Piñeyro M, Pizarro J, Lema F, Pritsch O, Cayota A, Bentley G, and Robello C. Crystal structure of the tryparedoxin peroxidase from the human parasite *Trypanosoma cruzi. J Struct Biol* 150: 11–22, 2005.
- 203. Piñeyro MD, Parodi-Talice A, Portela M, Arias DG, Guerrero SA, and Robello C. Molecular characterization and interactome analysis of *Trypanosoma cruzi* Tryparedoxin 1. *J Proteomics* 74: 1683–1692, 2011.
- 204. Pino P, Foth BJ, Kwok LY, Sheiner, L, Schepers R, Soldati T, and Soldati-Favre D. Dual targeting of antioxidant and metabolic enzymes to the mitochondrion and the apicoplast of *Toxoplasma gondii*. PLoS Pathog 3: e115, 2007.
- 205. Poole LB. Bacterial defenses against oxidants: mechanistic features of cysteine-based peroxidases and their flavoprotein reductases. Arch Biochem Biophys 433: 240–254, 2005.

 Poole L and Nelson K. Discovering mechanisms of signalingmediated cysteine oxidation. Curr Opin Chem Biol 12: 18, 2008.

- 207. Poole LB, Chae HZ, Flores BM, Reed SL, Rhee SG, and Torian BE. Peroxidase activity of a TSA-like antioxidant protein from a pathogenic amoeba. *Free Radic Biol Med* 23: 955–959, 1997.
- 208. Poole LB and Ellis HR. Identification of cysteine sulfenic acid in AhpC of alkyl hydroperoxide reductase. In: *Methods in Enzymology*, edited by Sies H and Packer L. Maryland Heights, MO: Academic Press, 2002, pp. 122–136.
- 209. Putz S, Gelius-Dietrich G, Piotrowski M, and Henze K. Rubrerythrin and peroxiredoxin: two novel putative peroxidases in the hydrogenosomes of the microaerophilic protozoon *Trichomonas vaginalis*. *Mol Biochem Parasitol* 142: 212–223, 2005.
- 210. Rahlfs S and Becker K. Thioredoxin peroxidases of the malarial parasite *Plasmodium falciparum*. *Eur J Biochem* 268: 1404–1409, 2001.
- 211. Raina O, Nagar G, Varghese A, Prajitha G, Alex A, Maharana B, and Joshi P. Lack of protective efficacy in buffaloes vaccinated with *Fasciola gigantica* leucine aminopeptidase and peroxiredoxin recombinant proteins. *Acta Trop* 118: 217–222, 2011.
- 212. Ramos-Martinez E, Olivos-Garcia A, Saavedra E, Nequiz M, Sanchez E, Tello E, El-Hafidi M, Saralegui A, Pineda E, Delgado J, Montfort I, and Perez-Tamayo R. *Entamoeba histolytica*: oxygen resistance and virulence. *Int J Parasitol* 39: 693–702, 2009.
- 213. Rhee SG and Woo HA. Multiple functions of peroxiredoxins: peroxidases, sensors and regulators of the intracellular messenger H₂O₂, and protein chaperones. *Antioxid Redox Signal* 15: 781–794, 2011.
- 214. Richard D, Bartfai R, Volz J, Ralph SA, Müller S, Stunnenberg HG, and Cowman AF. A genome-wide chromatin-associated nuclear peroxiredoxin from the malaria parasite *Plasmodium falciparum*. *J Biol Chem* 286: 11746–11755, 2011.
- 215. Romao S, Castro H, Sousa C, Carvalho S, and Tomas AM. The cytosolic tryparedoxin of *Leishmania infantum* is essential for parasite survival. *Int J Parasitol* 39: 703–711, 2009.
- 216. Roos D, Crawford MJ, Donald RG, Fraunholz M, Harb OS, He CY, Kissinger JC, Shaw MK, and Striepen B. Mining the Plasmodium genome database to define organellar function: what does the apicoplast do? Philos Trans R Soc Lond B Biol Sci 357: 35–46, 2002.
- 217. Salazar-Calderon M, Martin-Alonso J, Ruiz de Eguino A, Casais R, Marin M, and Parra F. *Fasciola hepatica*: heterologous expression and functional characterization of a thioredoxin peroxidase. *Exp Parasitol* 95: 63–70, 2000.
- 218. Salinas G, Fernandez V, Fernandez C, and Selkirk M. *Echinococcus granulosus*: cloning of a thioredoxin peroxidase. *Exp Parasitol* 90: 298–301, 1998.
- Santarem N, Silvestre R, Cardoso L, Schallig H, Reed S, and Cordeiro-da-Silva A. Application of an improved enzymelinked immunosorbent assay method for serological diagnosis of canine leishmaniasis. *J Clin Microbiol* 48: 1866–1874, 2010
- 220. Santarem N, Tomas A, Ouaissi A, Tavares J, Ferreira N, Manso A, Campino L, Correia J, and Cordeiro-da-Silva A. Antibodies against a *Leishmania infantum* peroxiredoxin as a possible marker for diagnosis of visceral leishmaniasis and for monitoring the efficacy of treatment. *Immunol Lett* 101: 18–23, 2005.

221. Santi-Rocca J, Weber C, Guigon G, Sismeiro O, Coppee J, and Guillen N. The lysine- and glutamic acid-rich protein KERP1 plays a role in *Entamoeba histolytica* liver abscess pathogenesis. *Cell Microbiol* 10: 202–217, 2008.

- 222. Sarma GN, Nickel C, Rahlfs S, Fischer M, Becker K, and Karplus PA. Crystal structure of a novel *Plasmodium falci-parum* 1-Cys peroxiredoxin. *J Mol Biol* 346: 1021–1034, 2005.
- 223. Sautel C, Ortet P, Saksouk N, Kieffer S, Garin J, Bastien O, and Hakimi M. The histone methylase KMTox interacts with the redox-sensor peroxiredoxin-1 and targets genes involved in *Toxoplasma gondii* antioxidant defences. *Mol Microbiol* 71: 212–226, 2009.
- 224. Sayed AA, Cook SK, and Williams DL. Redox balance mechanisms in *Schistosoma mansoni* rely on peroxiredoxins and albumin and implicate peroxiredoxins as novel drug targets. *J Biol Chem* 281: 17001–17010, 2006.
- 225. Sayed AA and Williams DL. Biochemical characterization of 2-Cys peroxiredoxins from *Schistosoma mansoni*. J Biol Chem 279: 26159–26166, 2004.
- 226. Sayers EW, Barrett T, Benson DA, Bolton E, Bryant SH, Canese K, Chetvernin V, Church DM, DiCuccio M, Federhen S, Feolo M, Fingerman IM, Geer LY, Helmberg W, Kapustin Y, Landsman D, Lipman DJ, Lu Z, Madden TL, Madej T, Maglott DR, Marchler-Bauer A, Miller V, Mizrachi I, Ostell J, Panchenko A, Phan L, Pruitt KD, Schuler GD, Sequeira E, Sherry ST, Shumway M, Sirotkin K, Slotta D, Souvorov A, Starchenko G, Tatusova TA, Wagner L, Wang Y, Wilbur WJ, Yaschenko E, and Ye J. Database resources of the National Center for Biotechnology Information. Nucleic Acids Res 39: D38–D51, 2011.
- 227. Schrum S, Bialonski A, Marti T, and Zipfel P. Identification of a peroxidoxin protein (OvPXN-2) of the human parasitic nematode *Onchocerca volvulus* by sequential protein fractionation. *Mol Biochem Parasitol* 94: 131–135, 1998.
- Seeber F and Soldati-Favre D. Metabolic pathways in the apicoplast of apicomplexa. *Int Rev Cell Mol Biol* 281: 161– 228, 2010.
- 229. Sekiya M, Mulcahy G, Irwin J, Stack C, Donnelly S, Xu W, Collins P, and Dalton J. Biochemical characterisation of the recombinant peroxiredoxin (FhePrx) of the liver fluke, Fasciola hepatica. FEBS Lett 580: 5016–5022, 2006.
- 230. Sela D, Yaffe N, and Shlomai J. Enzymatic mechanism controls redox-mediated protein-DNA interactions at the replication origin of kinetoplast DNA minicircles. *J Biol Chem* 283: 32034–32044, 2008.
- 231. Sievers F, Wilm A, Dineen D, Gibson TJ, Karplus K, Li W, Lopez R, McWilliam H, Remmert M, Söding J, Thompson JD, and Higgins DG. Fast, scalable generation of high-quality protein multiple sequence alignments using Clustal Omega. *Mol Syst Biol* 7: 539, 2011.
- 232. Silverman J, Chan S, Robinson D, Dwyer D, Nandan D, Foster L, and Reiner N. Proteomic analysis of the secretome of *Leishmania donovani*. *Genome Biol* 9: R35, 2008.
- 233. Silvestre R, Santarem N, Cunha J, Cardoso L, Nieto J, Carrillo E, Moreno J, and Cordeiro-da-Silva A. Serological evaluation of experimentally infected dogs by LicTXNPx-ELISA and amastigote-flow cytometry. *Vet Parasitol* 158: 23–30, 2008.
- 234. Siracusano A, Rigano R, Ortona E, Profumo E, Margutti P, Buttari B, Delunardo F, and Teggi A. Immunomodulatory mechanisms during *Echinococcus granulosus* infection. *Exp Parasitol* 119: 483–489, 2008.
- Soito L, Williamson C, Knutson ST, Fetrow JS, Poole LB, and Nelson KJ. PREX: PeroxiRedoxin classification indEX,

- a database of subfamily assignments across the diverse peroxiredoxin family. *Nucleic Acids Res* 39: D332–D337, 2011
- 236. Soong CJ, Torian BE, Abd-Alla MD, Jackson TF, Gatharim V, and Ravdin JI. Protection of gerbils from amebic liver abscess by immunization with recombinant *Entamoeba histolytica* 29-kilodalton antigen. *Infect Immun* 63: 472–477, 1995.
- 237. Stober C, Lange U, Roberts M, Alcami A, and Blackwell J. Heterologous priming-boosting with DNA and modified vaccinia virus Ankara expressing tryparedoxin peroxidase promotes long-term memory against *Leishmania major* in susceptible BALB/c Mice. *Infect Immun* 75: 852–860, 2007.
- 238. Stockwell BR. Chemical genetics: ligand-based discovery of gene function. *Nat Rev Genet* 1: 116–125, 2000.
- 239. Suttiprapa S, Loukas A, Laha T, Wongkham S, Kaewkes S, Gaze S, Brindley P, and Sripa B. Characterization of the antioxidant enzyme, thioredoxin peroxidase, from the carcinogenic human liver fluke, *Opisthorchis viverrini*. *Mol Biochem Parasitol* 160: 116–122, 2008.
- Szabó C, Ischiropoulos H, and Radi R. Peroxynitrite: biochemistry, pathophysiology and development of therapeutics. Nat Rev Drug Discov 6: 662–680, 2007.
- 241. This reference has been deleted.
- 242. Tachibana H and Cheng XJ. *Entamoeba dispar*: cloning and characterization of peroxiredoxin genes. *Exp Parasitol* 94: 51–55, 2000.
- 243. Tachibana H, Yanagi T, Akatsuka A, Kobayashi S, Kanbara H, and Tsutsumi V. Isolation and characterization of a potentially virulent species *Entamoeba nuttalli* from captive Japanese macaques. *Parasitology* 136: 1169–1177, 2009.
- 244. Tachibana H, Yanagi T, Pandey K, Cheng X, Kobayashi S, Sherchand J, and Kanbara H. An *Entamoeba* sp. strain isolated from rhesus monkey is virulent but genetically different from *Entamoeba histolytica*. *Mol Biochem Parasitol* 153: 107–114, 2007.
- 245. Tamayo EM, Iturbe A, Hernandez E, Hurtado G, de Lourdes Gutierrez XM, Rosales JL, Woolery M, and Ondarza RN. Trypanothione reductase from the human parasite Entamoeba histolytica: a new drug target. Biotechnol Appl Biochem 41: 105–115, 2005.
- 246. Tazreiter M, Leitsch D, Hatzenbichler E, Mair-Scorpio G, Steinborn R, Schreiber M, and Duchene M. Entamoeba histolytica: response of the parasite to metronidazole challenge on the levels of mRNA and protein expression. Exp Parasitol 120: 403–410, 2008.
- 247. Tetaud E, Giroud C, Prescott A, Parkin D, Baltz D, Biteau N, Baltz T, and Fairlamb A. Molecular characterisation of mitochondrial and cytosolic trypanothione-dependent tryparedoxin peroxidases in *Trypanosoma brucei*. Mol Biochem Parasitol 116: 171–183, 2001.
- 248. Thamsen M, Kumsta C, Li F, and Jakob U. Is overoxidation of peroxiredoxin physiologically significant? *Antioxid Redox Signal* 14: 725–730, 2011.
- 249. Trujillo M, Budde H, Piñeyro MD, Stehr M, Robello C, Flohé L, and Radi R. *Trypanosoma brucei* and *Trypanosoma cruzi* tryparedoxin peroxidases catalytically detoxify peroxynitrite via oxidation of fast reacting thiols. *J Biol Chem* 279: 34175–34182, 2004.
- 250. Trujillo M, Ferrer-Sueta G, Thomson L, Flohé L, and Radi R. Kinetics of peroxiredoxins and their role in the decomposition of peroxynitrite. In: *Peroxiredoxin Systems*, edited by Flohé L and Harris JR. New York: Springer, 2007, pp. 83–113.

- 251. Tsuji N, Kasuga-Aoki H, Isobe T, and Yoshihara S. Cloning and characterisation of a peroxiredoxin from the swine roundworm *Ascaris suum*. *Int J Parasitol* 30: 125–128, 2000.
- 252. Vaca-Paniagua F, Parra-Unda R, and Landa A. Characterization of one typical 2-Cys peroxiredoxin gene of *Taenia solium* and *Taenia crassiceps. Parasitol Res* 105: 781–787, 2009.
- 253. Vaca-Paniagua F, Torres-Rivera A, Parra-Unda R, and Landa A. *Taenia solium*: antioxidant metabolism enzymes as targets for cestocidal drugs and vaccines. *Curr Top Med Chem* 8: 393–399, 2008.
- 254. Veal E, Day A, and Morgan B. Hydrogen peroxide sensing and signaling. *Mol Cell* 26: 1–14, 2007.
- 255. Vedadi M, Lew J, Artz J, Amani M, Zhao Y, Dong A, Wasney G, Gao M, Hills T, Brokx S, Qiu W, Sharma S, Diassiti A, Alam Z, Melone M, Mulichak A, Wernimont A, Bray J, Loppnau P, Plotnikova O, Newberry K, Sundararajan E, Houston S, Walker J, Tempel W, Bochkarev A, Kozieradzki I, Edwards A, Arrowsmith C, Roos D, Kain K, and Hui R. Genome-scale protein expression and structural biology of *Plasmodium falciparum* and related apicomplexan organisms. *Mol Biochem Parasitol* 151: 100–110, 2007.
- 256. Vermeire J and Yoshino T. Antioxidant gene expression and function in *in vitro*-developing *Schistosoma mansoni* mother sporocysts: possible role in self-protection. *Parasitology* 134: 1369–1378, 2007.
- 257. Wakita M, Masuda S, Motohashi K, Hisabori T, Ohta H, and Takamiya K. The significance of type II and PrxQ peroxiredoxins for antioxidative stress response in the purple bacterium *Rhodobacter sphaeroides*. *J Biol Chem* 282: 27792–27801, 2007.
- 258. Walker J, Acestor N, Gongora R, Quadroni M, Segura I, Fasel N, and Saravia N. Comparative protein profiling identifies elongation factor-1beta and tryparedoxin peroxidase as factors associated with metastasis in *Leishmania guyanensis*. Mol Biochem Parasitol 145: 254–264, 2006.
- 259. Wassmann C, Hellberg A, Tannich E, and Bruchhaus I. Metronidazole resistance in the protozoan parasite *Entamoeba histolytica* is associated with increased expression of iron-containing superoxide dismutase and peroxiredoxin and decreased expression of ferredoxin 1 and flavin reductase. *J Biol Chem* 274: 26051–26056, 1999.
- 260. Wilkinson S, Horn D, Prathalingam S, and Kelly J. RNA interference identifies two hydroperoxide metabolizing enzymes that are essential to the bloodstream form of the African trypanosome. *J Biol Chem* 278: 31640–31646, 2003.
- 261. Wilkinson S, Temperton N, Mondragon A, and Kelly J. Distinct mitochondrial and cytosolic enzymes mediate trypanothione-dependent peroxide metabolism in *Trypa-nosoma cruzi*. *J Biol Chem* 275: 8220–8225, 2000.
- 262. Williams D, Asahi H, Botkin D, and Stadecker M. Schistosome infection stimulates host CD4(+) T helper cell and B-cell responses against a novel egg antigen, thioredoxin peroxidase. *Infect Immun* 69: 1134–1141, 2001.
- 263. Winterbourn C and Hampton M. Thiol chemistry and specificity in redox signaling. *Free Radic Biol Med* 45: 549–561, 2008.
- 264. Woo HA, Yim SH, Shin DH, Kang D, Yu DY, and Rhee SG. Inactivation of peroxiredoxin I by phosphorylation allows localized $\rm H_2O_2$ accumulation for cell signaling. *Cell* 140: 517–528, 2010.

265. Wood ZA, Poole LB, and Karplus PA. Peroxiredoxin evolution and the regulation of hydrogen peroxide signaling. *Science* 300: 650–653, 2003.

- 266. Wood ZA, Poole LB, Hantgan RR, and Karplus PA. Dimers to doughnuts: redox-sensitive oligomerization of 2-cysteine peroxiredoxins. *Biochemistry* 41: 5493–5504, 2002.
- 267. World Health Organization Department of Control of Neglected Tropical Diseases. Working to Overcome the Global Impact of Neglected Tropical Diseases: 1st WHO Report on NTDs. Geneva, Switzerland: WHO Press, 2010, p. 171.
- Wood ZA, Schröder E, Harris JR, and Poole LB. Structure, mechanism and regulation of peroxiredoxins. *Trends Bio-chem Sci* 28: 32–40, 2003.
- 269. Wyllie S, Cunningham ML, and Fairlamb AH. Dual action of antimonial drugs on thiol redox metabolism in the human pathogen *Leishmania donovani*. J Biol Chem 279: 39925– 39932, 2004.
- 270. Wyllie S, Mandal G, Singh N, Sundar S, Fairlamb A, and Chatterjee M. Elevated levels of tryparedoxin peroxidase in antimony unresponsive *Leishmania donovani* field isolates. *Mol Biochem Parasitol* 173: 162–164, 2010.
- 271. Yano K, Komaki-Yasuda K, Kobayashi T, Takemae H, Kita K, Kano S, and Kawazu S-I. Expression of mRNAs and proteins for peroxiredoxins in *Plasmodium falciparum* erythrocytic stage. *Parasitol Int* 54: 35–41, 2005.
- 272. Yano K, Komaki-Yasuda K, Tsuboi T, Torii M, Kano S, and Kawazu S. 2-Cys Peroxiredoxin TPx-1 is involved in gametocyte development in *Plasmodium berghei*. *Mol Biochem Parasitol* 148: 44–51, 2006.
- 273. Yano K, Otsuki H, Arai M, Komaki-Yasuda K, Tsuboi T, Torii M, Kano S, and Kawazu S. Disruption of the *Plasmodium berghei* 2-Cys peroxiredoxin TPx-1 gene hinders the sporozoite development in the vector mosquito. *Mol Biochem Parasitol* 159: 142–145, 2008.
- 274. Yoo W, Kim D-W, Ju J-W, Cho P, Kim T, Cho S-H, Choi S-H, Park H-S, Kim T-S, and Hong S-J. Developmental transcriptomic features of the carcinogenic liver fluke, *Clonorchis sinensis. PLoS Negl Trop Dis* 5: e1208, 2011.
- 275. Yuan Y, Knaggs MH, Poole LB, Fetrow JS, and Salsbury FA. Conformational and oligomeric effects on the cysteine pK_a of tryparedoxin peroxidase. *J Biomol Struct Dyn* 28: 51–70, 2010.
- 276. Zhang J, Fu Y, Xu X, Wu T, and Cao F. [RNA interference to the expression of peroxiredoxin-related genes in *Trichomonas vaginalis*]. (Abstract of Chinese-language article). *Zhongguo Ji Sheng Chong Xue Yu Ji Sheng Chong Bing Za Zhi* 23: 437–440, 2005.
- 277. Zhang W, Rogniaux H, Huang W, Chauvin A, and Moreau E. Analysis of thioredoxin peroxidase as a promising antigen for diagnosis of *Fasciola gigantica* infection: a preliminary study. *Parasitol Int* 60: 206–208, 2011.
- 278. Zipfel P, Schrum S, Bialonski A, and Buttner D. The peroxidoxin 2 protein of the human parasite *Onchocerca volvulus*: recombinant expression, immunolocalization, and demonstration of homologous molecules in other species. *Parasitol Res* 84: 623–631, 1998.

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Abbreviations Used

AhpF = alkyl hydroperoxide reductase component F

BCP = bacterioferritin comigratory protein

 C_P = peroxidatic cysteine

 C_R = resolving cysteine

 $EC_{50} = half$ -maximal effective concentration

FF = fully folded

Gpx = glutathione peroxidase

Grx = glutaredoxin

GS = glutathione synthase

LU = locally unfolded

MT = metacyclic trypomastigote

NCBI = National Center for Biotechnology Information

PDB = Protein Data Bank

Plrx = plasmoredoxin

PREX = the PeroxiRedoxin classification indEX

Prx = peroxiredoxin

Prx1 = a Prx subfamily or subfamily member

Prx5 = a Prx subfamily or subfamily member

Prx6 = a Prx subfamily or subfamily member

PrxQ = a Prx subfamily or subfamily member

RNAi = RNA interference

RNS = reactive nitrogen species

ROS = reactive oxygen species

RT-PCR = reverse transcriptase-polymerase chain reaction

SOD = superoxide dismutase

 $S_P = \text{sulfur atom of the peroxidatic Cys}$

 $S_POH =$ sulfenic acid form of the peroxidatic Cys

 S_PO_2H = sulfinic acid form of the peroxidatic Cys

 S_R = sulfur atom of the resolving thiol

Srx = sulfiredoxin

Tpx = a Prx subfamily named thiol peroxidase, or member of the same

Trx = thioredoxin

TrxR = thioredoxin reductase

TR = trypanothione reductase

 $TS_2 = trypanothione (oxidized)$

 $T(SH)_2 = trypanothione (reduced)$

Txn = tryparedoxin

Appendix 1. List of Parasite Genera and Important Pathogenic Species Included in Prx Searches

Genus Important species^a Acanthamoeba astronyxis, castellanii, divionensis, hatchetti, lenticulata, lugdunensis, polyphaga, rhysodes, ulbertsoni Ancylostoma duodenale Angiostrongylus cantonensis, costaricensis Anisakis simplex lumbricoides Ascaris Austrobilharzia variglandis Balamuthia mandrillaris Balantidium coli **Baylisascaris** procyonis Blastocystis hominis Brugia malayi, timori Capillaria hepatica, philippinensis Clonorchis sinensis, viverrini neoformans Cryptococcus Cryptosporidium parvum, hominis, muris Cyclospora cayetanensis Cystoisospora belli (formerly Isospora belli) Dicrocoelium dendriticum Dientamoeba fragilis Dioctophume renale Diphyllobothrium latum Dipylidium caninum Dracunculus medinensis Encephalitozoon cuniculi, intestinalis **Echinococcus** granulosus, multilocularis, oligarthrus, vogeli **Echinostoma** echinatum Entamoeba dispar, histolytica, moshkovskii Enterobius vermicularis, gregorii Fasciola hepatica, gigantica **Fasciolopsis** buski Giardia lamblia Gnathostoma hispidum, spinigerum Heterophyes heterophyes Hymenolepis diminuta, nana Leishmania aethiopica, amazonensis, braziliensis, chagasi, donovani, major, mexicana, guyanensis, infantum, panamensis, peruviana, tropica, venezuelensis Loa loa Mansonella streptocerca Metagonimus yokogawai fowleri Naegleria Necator americanus Onchocerca volvulus **Opisthorchis** viverrini, felineus Paragonimus africanus, caliensis, kellicotti, skrjabini, uterobilateralis, westermani Plasmodium falciparum, knowlesi, malariae, ovale, vivax Pneumocystis carnii (also known as P. jirovecii) decipiens Pseudoterranova Rhinosporidium seeberi Schistosoma haematobium, japonicum, mansoni, mekongi Spirometra erinacei, mansoni, mansonoides, ranarum Strongyloides stercoralis Taenia crassiceps, solium, saginata Toxocara canis, cati **Toxoplasma** gondii Trichinella britovi, nativa, nelsoni, spiralis regenti Trichobilharzia vaginalis **Trichomonas** Trichuris trichiura, vulpis

brucei, cruzi

bancrofti

Trypanosoma

Wuchereria

^aSpecies indicated in bold type are those with at least one Prx protein or gene that is mentioned in the review. For all others, we found no Prx gene or protein mentioned in the literature or present in sequence databases.