Overview on Peroxiredoxin

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Peroxiredoxins (Prxs) are a very large and highly conserved family of peroxidases that reduce peroxides, with a conserved cysteine residue, designated the “peroxidatic” Cys (Cₚ) serving as the site of oxidation by peroxides (Hall et al., 2011; Rhee et al., 2012). Peroxides oxidize the Cₚ–SH to cysteine sulfenic acid (Cₚ–SOH), which then reacts with another cysteine residue, named the “resolving” Cys (Cᵣ) to form a disulfide that is subsequently reduced by an appropriate electron donor to complete a catalytic cycle. This overview summarizes the status of studies on Prxs and relates the following 10 minireviews.

NOMENCLATURE

As in all biology, acronyms are overwhelming in Prx literature. A few of the more frequently used acronyms are TSA, AhpC, and Tpx (Rhee et al., 2005). Prx was initially identified in yeast in 1987 and named thiol-specific antioxidant (TSA), because it was thought to remove reactive sulfur species (like RS·, RSSR·, or RSOOH) rather than reactive oxygen species (like O₂·, H₂O₂, or ROOH) (Kim et al., 1988; 1989). Unlike enzymes known at the time to remove reactive oxygen species, purified TSA did not contain any redox cofactor such as a metal ion, heme, or flavin. Important clues to the actual enzymatic function of TSA were provided by database searches, which revealed a high sequence homology of TSA to AhpC (alkyl hydroperoxide reductase) identified in S. typhimurium in 1990 (Tartaglia et al., 1990), suggesting that TSA is also a peroxidase like AhpC. Subsequently, TSA was shown to reduce peroxides with thioredoxin (Trx) as the immediate hydrogen donor. Therefore the name of TSA was changed to thioredoxin peroxidase (TPx) in a manner analogous to glutathione peroxidase (GPx) (Chae et al., 1994a). It was then renamed Prx after realizing that certain members, for example 1-Cys Prxs, do not rely on Trx as the electron donor (Chae et al., 1994b). Proxiredoxin is now the name recommended by the Nomenclature Committee of the International Union of Biochemistry and Molecular Biology and is abbreviated as Prx or Prdx. As an example of typical Prx nomenclature, S. cerevisiae expresses five Prxs, which are two 2-Cys Prxs Tsa1 and Tsa2, two atypical 2-Cys Prxs Ahp1 and AhpE, and one 1-Cys mTpx (where m stands for mitochondria) [see the review by by Toldano and Huang (2016)].

CLASSIFICATION

The Cᵣ residue is conserved in all Prx enzymes. On the basis of the location or absence of the Cᵣ, Prxs are classified into 2-Cys, atypical 2-Cys, and 1-Cys Prx subfamilies (Chae et al., 1994b; Rhee et al., 2001; Wood et al., 2003b). 2-Cys Prx enzymes are homodimeric and contain two conserved (Cᵣ and Cₚ) cysteine residues per subunit. The Cᵣ–SOH reacts with the Cᵣ–SH of the other subunit to form an intersubunit disulfide. In atypical 2-Cys PrxV, the Cᵣ–SOH reacts with the Cᵣ–SH of the same subunit to form an intramolecular disulfide. Atypical 2-Cys is therefore also a 2-Cys Prx but is distinct from the 2-Cys Prx members that form an intermolecular disulfide on oxidation. Given the absence of a Cᵣ, Cᵣ–SOH of 1-Cys Prx cannot be resolved within the Prx molecules and instead forms a disulfide with Cᵣ–SH provided by other proteins or small thiol molecules (Fisher, 2011). Mammalian cells express six Prx isoforms (Prxl to PrxVI), which can be classified into 2-Cys (Prxl to PrxIV), atypical 2-Cys (PrxV), and 1-Cys (PrxVI) subfamilies (Rhee et al., 2001).

The subgrouping based on the position of the Cᵣ is mechanistically informative as mentioned earlier. As more Prx homologs/orthologs became known later, it was clear that Prx enzymes are ubiquitously expressed, with multiple isoforms present in most organisms (three in Escherichia coli, five in Saccharomyces cerevisiae, six in Homo sapiens, and nine in Arabidopsis thaliana). A different classification system has been suggested based on bioinformatic analysis of 29 crystal structures and >3500 sequences of Prx enzymes that had been determined by 2010 (Nelson et al., 2011). This “global evolutionary classification” divided Prx proteins into Prx1, Prx5, Prx6, Tpx, PrxQ, and AhpE subfamilies. The biochemical and structural features and phylogenetic distributions of each of these Prx subgroups are summarized in the review by Poole and Nelson (2016). Members of the Prx1 subgroup are those referred to as “typical” 2-Cys Prxs that include mammalian Prxl to PrxIV, bacterial AhpC, and yeast TSA. Mammalian PrxV and PrxVI belong to the Prx5 and Prx6 subfamilies, respectively. Mammalian cells do not express members of the Tpx and PrxQ subfamilies.

EXCEPTIONAL REACTIVITY OF PEROXIDATIC CYSTEINE

The Cᵣ of Prx is oxidized by peroxides like H₂O₂, lipid peroxide,
or peroxynitrite very rapidly with the second order rate constants for the formation of Cρ–SOH being in the range of $1 \times 10^6$ to $10^8 \text{M}^{-1}\text{s}^{-1}$, which are 5 to 7 orders of magnitude higher than those for small thiols (Winterbourn, 2013). The rate constants for next catalytic steps, the disulfide formation between Cρ–SOH and CR–SH and the reduction of CP–S–S–CP, were also measured for various Prx enzymes. Kinetic analysis also revealed that peroxides bind to Prx with submicromolar affinity. The methods used to obtain those kinetic constants for individual steps include the competitive kinetics with horseradish peroxidase or catalase or stopped flow monitoring of the change in intrinsic tryptophan fluorescence (Manta et al., 2009; Parsonage et al., 2015; Trujillo et al., 2007), which are described in the review by Winterbourn and Peskin (2016). Analyses of Prx structures with bound peroxide or peroxide-mimicking molecules revealed a model for the transition state of the peroxidase reaction (Hall et al., 2011; Nakamura et al., 2010; Perkins et al., 2015). In the model, all Prxs have an active site structure with a nearly universal sequence, PXXXTXXCρ (where X is any amino acid), as well as a conserved Arg that is distant in sequence but located nearby in the three-dimensional fold. The transition state involves an extensive hydrogen bond network, which comprises the Cρ thiolate anion, peroxide (ROOH), Thr and Pro residues located within a universal PXXXTXXCρ active site motif as well as the conserved Arg residue. This network of hydrogen bonds thus provides a binding site for peroxide that properly aligns the substrate for attack by the CP thiolate sulfur and lowers the activation energy of the bond-breaking and bond-forming processes by stabilizing the transition state intermediate.

**ANTIOXIDANT ROLE OF PRXS**

Undoubtedly the peroxidase activity of Prx enzymes towards H2O2, organic hydroperoxides and peroxynitrite is critical to protect cellular components from oxidative damage (Fisher, 2011; Knoops et al., 2011). However, the existence of multiple peroxide-removing enzymes such as catalase, GPx, Prx, and ascorbate peroxidase (APx) within a single organism, cell, or subcellular compartment, indicates that these peroxidases are not simply for oxidant defense. For example, chloroplasts contain three Prx, two GPx, and two APx isoforms (Dietz, 2016). They use different electron donors and are linked to distinct redox networks. The significance of those multiple peroxidase enzymes in the photosynthesizing chloroplast is discussed in context of various aspects including suborganellar localization, substrate preference, and metabolic coupling in the review by Dietz (2016).

**PRXS AS A REGULATOR OF LOCAL H2O2 CONCENTRATION**

Many mammalian cell types produce H2O2 for the purpose of intracellular signaling in response to stimulation through various cell surface receptors (Rhee, 2006). H2O2 propagates receptor signals by oxidizing proteinaceous thiols such as those in protein tyrosine phosphatases (PTPs) (Lee et al., 1998), the tumor suppressor PTEN (phosphatase and tensin homolog) (Kwon et al., 2004), and ASK1 (an upstream kinase of p38 and JNK mitogen activated kinases) (Saitoh et al., 1998; Nadeau et al., 2007). The targeted thiol groups of these H2O2 effector proteins react several orders of magnitude more slowly with H2O2 compared with the Cρ residue of Prxs (Winterbourn, 2013). Given that these effector proteins are at such a competitive disadvantage, it seems likely that neighboring Prx molecules must be transiently inactivated to allow them to react with H2O2. One example is the inactivation of lipid raft–associated mammalian Prx through phosphorylation at Tyr194 in cells stimulated with growth factors (Woo et al., 2010) or inactivation of centrosome-associated Prx through phosphorylation at Thr120 during early mitosis (Lin et al., 2015). Another example is the inactivation of 2-Cys Prx members through reversible hyperoxidation of Cρ–SH (see below). By switching “off” the highly active Prxs around the sites of H2O2 production, the oxidants produced could reach higher levels within these foci that could then permit the oxidation of protein targets involved in signaling, processes which would normally be much too slow to compete with the oxidation of Prx Cρ–SH.

**PRXS AS SENSORS AND TRANSDUCERS OF SIGNALING BY H2O2**

In an alternative scenario, redox-regulated proteins are not directly oxidized by H2O2, but rather their oxidation is mediated by Prx. In this case, the Cρ of Prx is rapidly and selectively oxidized by H2O2. The oxidized Prx, in either the sulfenic (Cρ–SOH) or disulfide state (Cρ–S–S–Cρ), forms an intermolecular disulfide-linked intermediate with a bound effector protein. Resolution of the disulfide by reaction with another Cys–SH of the effector results in regeneration of reduced Prx and oxidation of the effector protein, the latter process leading to a change in effector function. The sensor-transducer function of Prxs is extensively covered in three reviews by Netto and Antunes (2016), by Latimer and Veal (2016), and by Toledano and Huang (2016), taking yeast transcription factor Pap (Vivancos et al., 2005), mammalian transcription factor STAT3 (Sobotta et al., 2015), and ASK1 (Nadeau et al., 2007) as example of H2O2 target protein. Taking advantage of the extreme H2O2 sensitivity of the Cρ–SH of Prx, genetically encoded H2O2 fluorescent probes have been devised by fusing a Prx to green fluorescent protein (GFP) or yellow fluorescent protein (YFP). In the probes, the oxidation state of Prx is transferred to cysteine residues in GFP or YFP proteins. In the review by Van Laer and Dick (2016), sensitivity and design strategies for Prx-based probes are discussed.

**HYPEROXIDATION OF 2-CYS PRX**

During catalysis, the peroxidatic Cys–SOH of 2-Cys Prxs is occasionally further oxidized to Cys–SO2H before disulfide formation, resulting in inactivation of peroxidase activity (Yang et al., 2002). The hyperoxidation is reversed by the ATP-dependent enzyme sulfiredoxin, restoring peroxidase activity (Biteau et al., 2003; Woo et al., 2003). The reversible inactivation of 2-Cys Prx through hyperoxidation has been suggested to be an evolutionary adapted mechanism for eukaryotic cells and phototrophic bacteria: 1) to allow H2O2 to accumulate to substantial levels under certain circumstances for signaling purposes (Wood et al., 2003a); 2) to couple circadian rhythm to Prx function (O’Neill et al., 2011); or 3) to confer chaperone function to the 2-Cys Prxs in cells under severe oxidative stress (Jang et al., 2004).

**ROLE OF 2-CYS PRX HYPEROXIDATION IN H2O2 SIGNALING**

Hyperoxidized form of Prx III, a mitochondria-specific enzyme, was found to be abundant especially in the cortex of adrenal gland of mouse maintained under normal conditions. The adrenal cortex is where cholesterol is oxidized to corticosterone by cytochrome P450s in response to stimulation with adrenocorticotropic hormone. The conversion of cholesterol to corti-
cortesterone is accompanied by H₂O₂ generation as the result of leaky cytchrome c oxidase, which results in the hyperoxidation and inactivation of PrxIII (Kil et al., 2012). Inactivation of PrxIII triggers a sequence of events including accumulation of H₂O₂, activation of p38 mitogen-activated protein kinase (MAPK), suppression of steroidogenic acute regulatory protein synthesis, and inhibition of steroidogenesis (Kil et al., 2012). The coupling of CYP11B1 activity to PrxIII hyperoxidation thus provides a feedback regulatory mechanism for steroidogenesis. H₂O₂ has been increasingly recognized as an important component of cell signaling, including proliferation, differentiation, stereoidogenesis, metabolism, apoptosis, and senescence. Hampton and O’Connor (2016) suggest in their review that PrxIII hyperoxidation is a critical element of apoptotic process and that PrxIII–SO₂H can be a biomarker of redox changes during the initiation and progression of cell death as indicated by the fact that the timing of PrxIII–SO₂H accumulation during Fas-mediated apoptosis in Jurkat T-lymphoma cells corresponds to cytchrome c release from the mitochondria and subsequent caspase activation (Cox et al., 2008). Although numerous signaling pathways are known to be affected by H₂O₂, the underlying mechanisms are not well established. One of the better known H₂O₂–regulated pathways is the MAPK cascade. Latimer and Veal (2016) reviewed the evidence that Prx acts on multiple steps of MAPK signaling pathways in various organisms.

**2-CYS PRX HYPEROXIDATION AND CIRCADIAN RHYTHM**

As summarized in the review by Pulk and O’Neill (2016), it has been known for a while that the circadian clock regulates cellular redox state such as the ratio of NAD(P)H/NAD(P)⁺. The remarkable story of the circadian oscillation of 2-Cys Prx–SO₂H started with the discovery of the hyperoxidation of 2-Cys Prx as a transcription-independent circadian biomarker in a green alga (O’Neill et al., 2011) and human red blood cells (O'Neill and Reddy, 2011). The rhythmic variation in 2-Cys Prx–SO₂H abundance was subsequently detected in various organisms including fungus, worm, fly, and mouse, suggesting that Prx hyperoxidation cycles represent a conserved, ancestral circadian mechanism (Edgar et al., 2012; Olmedo et al., 2012). Interestingly, the amounts of PrxIII–SO₂H and Srx in the mitochondria of adrenal gland also showed daily variation (Kil et al., 2012). These results suggest that full circadian oscillation of corticosterone controlled by both the master and adrenal peripheral clocks requires the participation of the signaling pathway involving the reversible PrxIII hyperoxidation. The reversible PrxIII hyperoxidation appeared to play a critical role in linking metabolism also to other peripheral clocks in tissues like heart, lung, and brown adipose tissue (Kil, 2015). The oscillation in 2-Cys Prx–SO₂H abundance appears to be a cellular clock output driven by an underlying rhythm in oxidative metabolism (Causton et al., 2015).

**CHAPERONE FUNCTION OF HYPEROXIDIZED 2-CYS PRX**

In addition to the roles in H₂O₂ signaling and circadian output, hyperoxidation confers a new function on 2-Cys Prxs—namely, that of a protein chaperone. The first evidence for this gain of chaperone function came from the observation that yeast cells deficient in both TSAs (cytosolic 2-Cys Prxs) are highly sensitive to heat shock treatment, a phenotype often associated with organisms that lack crucial chaperones (Jang et al., 2004). The chaperone function was found to be independent of peroxidase activity. Chaperone activity was also subsequently demonstrated for hyperoxidized Prx in various species (Moon et al., 2005). As suggested initially for yeast TSAs, the chaperone activity observed in most species was attributed to high molecular weight species that appeared to correspond to double- or multiple-layered decameric rings, with the high molecular weight assembly being triggered by the C₇ hyperoxidation. As pointed out by Toledano and Huang in their review (Toledano and Huang, 2016), however, neither the hyperoxidation nor the higher order oligomerization was necessary for the chaperone function of 2-Cys Prxs in certain species. Toledano and Huang (2016) also point out that the in vivo relevance of chaperone function is yet to be established.

**PRXS AS PAMPS AND DAMPS**

During inflammation high levels of peroxides (H₂O₂, lipid hydroperoxide, and peroxynitrite) are produced by phagocytes to kill microorganisms. It has been well established that Prxs play cytoprotective antioxidant role in inflammation (Grete et al., 2012; Sun et al., 2010). Peroxides serve also in the complex regulation of inflammatory signaling pathways and Prxs are known to be critical modulator of the signaling peroxides (Diet et al., 2007). Recently, it has been proposed that Prxs may play key roles in innate immunity and inflammation. A 2-Cys malarial Prx from Plasmodium berghei was reported to act as a PAMP (pathogen-associated molecular pattern) by binding to a toll-like receptor on macrophages and triggering a pro-inflammatory response (Furuta et al., 2008). Moreover, exosomally released Prxs by different cells upon exposure to LPS and TNF-α has also been shown to act as a DAMP (host-derived damage-associated molecular patterns) to trigger inflammatory cytokines by macrophages (Mullen et al., 2015; Riddell et al., 2010; Salzano et al., 2014; Shichita et al., 2012). Those multiple, complex roles played by Prxs in inflammation are described in the review by Knoops et al. (2016).

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