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Human Quiescin-sulfhydryl Oxidase, QSOX1: Probing Internal Redox Steps by Mutagenesis†

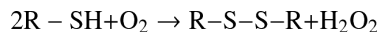
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Abstract

The flavoprotein Quiescin-sulfhydryl oxidase (QSOX) rapidly inserts disulfide bonds into unfolded, reduced proteins with the concomitant reduction of oxygen to hydrogen peroxide. This study reports the first heterologous expression and enzymological characterization of a human QSOX1 isoform. Like QSOX isolated from avian egg white, recombinant HsQSOX1 is highly active towards reduced ribonuclease A (RNase) and dithiothreitol but shows a >100-fold lower k_{cat}/K_m for reduced glutathione. Previous studies on avian QSOX led to a model in which reducing equivalents were proposed to relay through the enzyme from the first thioredoxin domain (C70–C73) to a distal disulfide (C509–C512), then across the dimer interface to the FAD-proximal disulfide (C449–C452), and finally to the FAD. The present work shows that, unlike the native avian enzyme, HsQSOX1 is monomeric. The recombinant expression system enabled construction of the first cysteine mutants for mechanistic dissection of this enzyme family. Activity assays with mutant HsQSOX1 indicated that the conserved distal C509–C512 disulfide is dispensable for the oxidation of reduced RNase or dithiothreitol. The four other cysteine residues chosen for mutagenesis, C70, C73, C449, and C452, are all crucial for efficient oxidation of reduced RNase. C452, of the proximal disulfide, is shown to be the charge-transfer donor to the flavin ring of QSOX, and its partner, C449, is expected to be the interchange thiol, forming a mixed disulfide with C70 in the thioredoxin domain. These data demonstrate that all the internal redox steps occur within the same polypeptide chain of mammalian QSOX and commence with a direct interaction between the reduced thioredoxin domain and the proximal disulfide of the Erv/ALR domain.

All eukaryotic sulfhydryl oxidases yet described utilize either metal ions or a flavin cofactor to couple the oxidation of thiols with the reduction of oxygen to hydrogen peroxide:



The copper- and iron-dependent enzymes are relatively poorly understood; there are no sequences, structures, or detailed analyses of the catalytic role of the metal (1–4). In contrast, a growing literature describes the structures, mechanisms, and roles in oxidative

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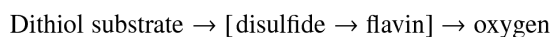
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SUPPORTING INFORMATION AVAILABLE Supplemental Figures S1–4 show sequence alignments of selected QSOXs; the full sequence of the human QSOX1b construct used for most of this work; the sequence of the construct used for the analytical ultracentrifugation; and analysis of ultracentrifugation data. This material is available free of charge via the Internet at <http://pubs.acs.org>.

protein folding of the flavin-dependent sulfhydryl oxidases, *e.g.*, Ero1 and members of the Erv/ALR family (4–11). The latter includes single-domain proteins, such as yeast Erv1p and Erv2p (5–8), mammalian augments of liver regeneration (ALR) (9–11), plant (12, 13) and viral (14, 15) sulfhydryl oxidases, as well as multi-domain Quiescin-sulfhydryl oxidases (QSOX) (3, 4, 16, 17), which are the subject of this contribution.

Rat seminal vesicle sulfhydryl oxidase, now known to be a member of the QSOX branch of the Erv/ALR family, was the first flavin-dependent sulfhydryl oxidase to be described (18). However most of the mechanistic work on these enzymes has utilized avian egg white QSOX (2, 17, 19–21). The initial anaerobic titrations of avian QSOX provided strong evidence for the presence of two redox centers: a flavin prosthetic group and a juxtaposed protein disulfide (19). This led to a proposal for the flow of reducing equivalents during QSOX catalysis (21):



Although this minimal model (19) showed close parallels to that established for lipoamide dehydrogenase, glutathione reductase, and thioredoxin reductase (22, 23), QSOX enzymes were subsequently found to be evolutionarily unrelated to the pyridine nucleotide-disulfide oxidoreductase family (24, 25). In addition, the amino acid sequence and domain organization of avian QSOX (Figure 1) suggested the presence of additional potential redox-active disulfides that would have been spectroscopically silent in the initial series of anaerobic titrations (17, 24). The domain organization of metazoan QSOX from the amino terminus is as follows: two thioredoxin-like domains, a region predicted to be helix-rich (HRR) but about which little else is known, an Erv/ALR FAD-binding module (7, 9, 13), a region of high sequence variability predicted to be partially unstructured, and a membrane spanning region that can be eliminated by differential splicing to yield soluble isoforms (3, 4, 25, 26). The CxxC motif, common in proteins that undergo dithiol/disulfide redox reactions, is found in three places in the enzyme (16, 17): in the first thioredoxin-like domain (Trx1), in the Erv/ALR domain, and directly downstream of the Erv/ALR domain (Figure 2).

Two additional lines of evidence suggested that the initial mechanistic proposal was an oversimplification (17). Partial proteolysis of the avian oxidase showed that protein substrates of the enzyme interacted primarily with the CxxC motif of the Trx1 domain and that reducing equivalents were subsequently transferred between the Trx1 and Erv/ALR domains (17). Further insights were provided by the crystal structure of yeast Erv2p (7). Fass and coworkers proposed a model for the flow of reducing equivalents in this oxidase depicted in Figure 2 (7). The initial reduction of Erv2p occurs *via* dithiol/disulfide exchange with the distal (C176–C178) disulfide bond in one subunit of the homodimer (7, 13, 27). A second disulfide exchange across the dimer interface leads to the reduction of the proximal disulfide (C121–C124) in the opposite subunit followed by reduction of the flavin (22, 28) and transfer of reducing equivalents to molecular oxygen (29, 30). Single cysteine to alanine mutations of each of these four cysteine residues (C121, 124, 176, and 178) abolished the ability of Erv2p to rescue the growth defects of a yeast strain harboring a temperature-sensitive Ero1p mutant, consistent with an important role for shuttling reducing equivalents between distal and proximal disulfide/dithiol centers in dimeric Erv2p *in vivo* (7).

Several observations suggested (17) that avian QSOX had incorporated key aspects of the mechanism proposed for Erv2p. First, a distal CxxC motif is conserved in all available QSOX sequences (16, 17) and is located at a comparable position in the amino acid sequence to the CxC disulfide in yeast Erv2p (7, 17). If this distal disulfide were to occupy a

position and play a role comparable to that depicted for Erv2p in Figure 2, then avian QSOX would be expected to be a dimer. Indeed both the full length egg white enzyme (19) and a proteolytic fragment containing the Erv/ALR domain (17) were found to be dimers by gel filtration. Finally, reductive titrations under forcing conditions showed that avian QSOX could accept a total of eight electrons (17), suggesting that the flavin and all three of the QSOX disulfides shown in Figure 3 ($C_{xx}C_{Trx}$, $C_{xx}C_{prox}$ and $C_{xx}C_{dist}$) can undergo reduction. These data led Raje et al. to a more detailed model for avian QSOX catalysis (17) which utilized all three $C_{xx}C$ disulfides and an intersubunit transfer step as depicted in Figure 3.

Further progress in understanding the mechanism of QSOX enzymes required the application of site directed mutagenesis to study the individual roles of these six cysteine residues. We report here the first heterologous expression and enzymatic characterization of human QSOX1 and compare its catalytic specificity to the avian egg white and bovine milk proteins. Surprisingly, site-directed mutagenesis of the human enzyme demonstrates that it does not follow the catalytic mechanism shown in Figure 3. Our results, taken with complementary observations from smaller members of the Erv/ALR family of oxidases to be detailed later, show that monomeric versions of these enzymes can utilize a pathway for flow of reducing equivalents that bypasses the distal disulfide altogether. These studies provide important new insight into these intriguing catalysts of oxidative protein folding.

MATERIALS AND METHODS

Materials

Primers for the HsQSOX1b construct were from the Great American Gene Company and primers for mutagenesis were from Integrated DNA Technologies. Ampicillin, tetracycline, chloramphenicol, riboflavin, protease inhibitor for His-tagged proteins, imidazole, RNase, catalase and glutathione were from Sigma. IPTG was from Promega; DTT was from Acros Organics; yeast, tryptone, NaCl, FAD, EDTA, and kanamycin sulfate were from Fisher.

Human QSOX1 construct and mutagenesis

Quiescin-sulfhydryl oxidase 1 (HsQSOX1) isoform b cDNA clone (ID 4447666) from human kidney mRNA was obtained from Invitrogen in the vector pCMV-SPORT6. The desired construct was amplified by PCR with an N-terminal forward primer omitting the signal sequence: 5'-CTGGGATCCCGGTCGGCGCTC-3' (BamHI site underlined) and a C-terminal reverse primer incorporating a stop codon: 5'-CTGAAGCTT TCAAATAAGCTC-3' (stop codon italicized, HindIII site underlined). The construct was blunt-end ligated into a TOPO TA cloning vector pCR2.1 (Invitrogen) and digested with restriction enzymes. The insert was then ligated into the expression vector pTRC HisA (Invitrogen) digested with BamHI and HindIII. This construct, including an N-terminal hexahistidine tag, was found to contain four sequence errors presumably as a result of low polymerase fidelity. These errors were corrected by site-directed mutagenesis (QuikChange, Stratagene) and the construct was re-sequenced to ensure no additional changes had been introduced. Site-directed cysteine mutations described in the text used the following primers: C70S; forward 5'-GCCTCCTGGAGCGGCCACTGCATCGCCTTCGCC-3', reverse 5'-GGCGAAGGCGATGCAGTGGCCGCTCCAGGAGGC-3'; C73S; forward 5'-GCCTCCTGGTGC GGCCACAGCATCGCCTTCGCCCGACGTGG-3', reverse 5'-CCACGTCGGGGCGAAGGCGATGCTGTGGCCGCACCAGGAGGC-3'; C449A; forward 5'-GGCTACGTGCACTACTTCTTCGGCGCCCGAGACTGCGCTAGCCAC-3', reverse 5'-GTGGCTAGCGCAGTCTCGGGCGCCGAAGAAGTAGTGCACGTAGCC-3'; C449S; forward 5'-GGCTACGTGCACTACTTCTTCGGCAGCCGAGACTGCGCTAGCCAC-3', reverse 5'-

GTGGCTAGCGCAGTCTCGGCTGCCGAAGAAGTAGTGCACGTAGCC-3'; C452S; forward 5'-GGCTGCCGAGACAGCGCTAGCCACTTCGAGCAG-3', reverse 5'-CTGCTCGAAGTGGCTAGCGCTGTCTCGGCAGCC-3'; C509A; forward 5'-CCCCGTGAACCTTGCTTCTGCCTGCCACAATGAACGC-3', reverse 5'-GCGTTCATTGTGGCAGGCAGAAGCAAGTTCACGGGG-3'; C512A; forward 5'-CGTGAACCTTGTCTGCCGCCACAATGAACGC-3', reverse 5'-GCGTTCATTGTGGGCGGCAGAACAAGTTCACG-3'; C509SC512S; forward 5'-CCCCGTGAACCTTAGTTCTGCCAGCCACAATGAACGCCGT-3', reverse 5'-ACGGCGTTCATTGTGGCTGGCAGAAGTTCACGGGG-3'. All constructs were fully sequenced to verify mutations and to ensure no additional changes in the sequence had been introduced.

Expression of QSOX1 and mutants

Expression of the QSOX1 construct in *E. coli* was first attempted in BL21 DE3 cells (Invitrogen) at 37 °C. Cells were grown to an OD₆₀₀ of 0.6 and then induced with 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG) for 12 h. This resulted in the expression of mostly insoluble material. Auto-induction using BL21 (Star) DE3 cells was also assessed as a means of expressing HsQSOX1 but provided a low yield of soluble protein. Attempts to refold insoluble protein from inclusion bodies using 8 M guanidine hydrochloride in 100 mM DTT followed by dilution into a matrix of solutions from a Fold-It kit (Hampton Research) did not yield active protein.

The construct was then transformed into the Rosetta-gami DE3 strain (Novagen) that contains genes for rare tRNAs and mutations in the *trxB* (thioredoxin reductase) and *gor* (glutathione reductase) genes to enhance disulfide bond formation in the cytoplasm. A large number of expression trials with this construct (varying the times of growth and induction, the IPTG concentration, the temperature, and the use of rich and minimal growth media) were aimed at maximizing solubility and yield of protein. The following expression conditions gave workable, if low (~0.3 mg/L), purified yields of active HsQSOX1. Cultures were grown overnight at 37 °C in 5 mL of LB media supplemented with 50 µg/mL ampicillin, 34 µg/mL chloramphenicol, 15 µg/mL kanamycin, and 12.5 µg/mL tetracycline and used to inoculate four 2 L flasks each containing 500 mL of LB media supplemented with the same antibiotics and 5 µM riboflavin. Inoculated cultures were grown at 37 °C to OD₆₀₀ 1.0 – 2.0. The cultures from the four flasks were then centrifuged and combined in fresh LB media supplemented as above and grown at 15 °C for 24 h in the presence of 0.2 mM IPTG. Cells were harvested at 5000 g at 4 °C and stored at – 20 °C before use.

Purification of QSOX1 and mutants

Harvested cells were resuspended at 0.25 g cell paste/mL of 50 mM potassium phosphate, pH 7.5, supplemented with 50 µM FAD, 0.1 mg/mL lysozyme, and a protease inhibitor cocktail for His-tagged proteins (Sigma: according to the manufacturer's instructions). The suspension was disrupted with two passes through a French press (10,000 psi) and clarified by centrifugation at 4 °C for 20 min at 8000 g. The supernatant (20 mL) was added to 2 mL of Pro Bond Ni-NTA resin (Invitrogen) previously equilibrated in 50 mM potassium phosphate pH 7.5 containing 50 mM imidazole and 250 mM NaCl. The mixture was rocked overnight at 4 °C and packed into an empty column with elution of unbound materials. The resin was washed with three column volumes of 50 mM potassium phosphate pH 6.0 containing 1 M NaCl and 50 mM imidazole, followed by three column volumes of 50 mM potassium phosphate pH 7.5 containing 50 mM imidazole without NaCl. The column was developed with 5 mL of 100, 300, and 500 mM imidazole in 50 mM potassium phosphate pH 7.5. For wild-type QSOX1, the eluted protein fractions (at 300 mM imidazole) were purified using a second Ni-NTA column, incubating the protein solution with resin for one

hour and using the same wash and elution steps. For QSOX1 mutants, the Ni-NTA column protein fractions were further purified on a Source 30 HR 5/5 cation exchange column using an AKTA FPLC (Amersham Pharmacia) with a 0 – 1 M NaCl gradient in 20 mM potassium phosphate pH 6.5, 1 mM EDTA. Following this second column purification, both wild-type and mutants were dialyzed against 50 mM potassium phosphate pH 7.5 containing 1 mM EDTA. Centrifuge ultrafiltration of recombinant human QSOX1, for protein concentration or buffer exchange, proved unsatisfactory because of aggregation at the membrane surface.

Size exclusion chromatography of the recombinant wild type enzyme was performed in 50 mM potassium phosphate pH 7.5 with 1 mM EDTA on a Superdex 200 HR 10/30 gel filtration column on the same FPLC system. The apparent native molecular weight of wild type recombinant enzyme was determined by comparison to a standard curve formed using: thyroglobulin, ferritin, aldolase, bovine serum albumin, hemoglobin, chicken egg ovalbumin, and cytochrome *c*. Apparent molecular weights of the recombinant wild type and mutant enzymes were determined by 10% SDS-PAGE gels (Biorad) stained with Coomassie Blue.

UV/VIS Spectroscopy

Absorption spectra were obtained with an HP8453 instrument blanked with assay buffer 50 mM potassium phosphate pH 7.5 with 1 mM EDTA. Wild type enzyme concentrations were determined using an extinction coefficient for bound flavin of $12.5 \text{ mM}^{-1} \text{ cm}^{-1}$ at 456 nm (see below). The same extinction coefficient was used for all the mutants except the C449A and C449S mutants, which had extinction coefficients of $11.6 \pm 0.4 \text{ mM}^{-1} \text{ cm}^{-1}$ and $11.2 \pm 0.1 \text{ mM}^{-1} \text{ cm}^{-1}$ respectively at 456 nm. Scatter corrections were performed using the HP8453 Agilent ChemStation software.

Reduced RNase Assays

Reduced RNase was prepared by a modification of a prior procedure (20). Here, reduction was performed overnight at 4 °C and the protein was desalted on a PD10 column equilibrated with deoxygenated 0.1% v/v acetic acid. Reduced RNase was stored anaerobically and standardized for thiol content prior to use (average of 8.0 DTNB-reactive thiols per RNase molecule). Sulfhydryl oxidase-mediated oxidation of RNase used the following concentrations: 35 μM to 1000 μM RNase thiols, 10 nM catalase, and 50 nM enzyme. Controls were run in the absence of enzyme. Samples were diluted 1:10 into 0.5 mM DTNB dissolved in 50 mM potassium phosphate pH 7.5, 1 mM EDTA and the thiol titer recorded at 412 nm.

Oxygen Electrode Assays

Mutant and wild type oxidase activities were assayed in a Clark-type oxygen electrode as before (19). DTT was prepared in sample buffer and standardized with DTNB. Before standardization ~ 100 mM stock solutions of GSH in buffer were returned to pH 7.5 by the addition of concentrated KOH.

Extinction Coefficient Determination and Thiol Titer of HsQSOX1

The UV/VIS spectrum of purified recombinant HsQSOX1 was recorded after dialysis into 100 mM Tris HCl pH 7.5, 1 mM EDTA and then re-measured 1 min after the addition of 0.1% SDS (from a 20% stock solution in water). Control experiments showed that the extinction coefficient of FAD ($11.3 \text{ mM}^{-1} \text{ cm}^{-1}$) was essentially unchanged in 0.1% SDS. The thiol titer of wild-type enzyme was determined after the addition of 0.5 mM DTNB to SDS-denatured protein.

Sequence Analysis and Modeling

DNA and amino acid sequence comparisons were done with LALIGN, ClustalW (31) and MultAlin (32) using default values. ClustalW and LALIGN were used to align the two thioredoxin domains of human QSOX1 against the sequence of the **a** and **b** domains of yeast Pdi1p ((33); PDB ID 2B5E) and of the oxidized and reduced forms of human thioredoxin 2 (PDB IDs 1W4V and 1W89, respectively). These alignments were submitted to SWISS-MODEL (34) and the resulting predictions viewed with Accelrys DS ViewerPro v5.0 and Swiss-Pdb viewer v3.7. MyHits Motif Scan (35) was used to help in domain boundary determination.

RESULTS AND DISCUSSION

Human QSOX1 Expression Construct: Domain Boundaries and Cysteine Content

Two transcripts of QSOX1 have been described, QSOX1a and 1b (3, 4, 25, 26). QSOX1a has a single transmembrane region, which is lacking in isoform 1b. In this study, the signal peptide of QSOX1b was replaced with a hexa-histidine tag followed by a 26 amino acid linker (Figure S2; Materials and Methods).

The twelve cysteines in the HsQSOX1 construct are depicted in Figure 1 and will be described in turn. The conserved CGHC motif in the Trx1 domain contains C70 and C73. A second cysteine pair (C101 and C110; C_xC, Figure 1) is conserved within the Trx1 domain in all of the metazoan and plant QSOX sequences so far examined but has a C_xC spacing in the *Trypanosoma* and *Leishmanii* QSOX proteins (supplemental Figure S1). The first cysteine in this C_xC motif corresponds to that of the C_xC disulfide bond found in the **a** domain of yeast Pdi1p (33). This particular disulfide (33, 36–38) has been suggested by Gilbert and coworkers (38) to play a regulatory role in Pdi1p by destabilizing the catalytic **a** domain disulfide. This presumed regulatory disulfide is encountered in Pdi1p homologues Eug1p (39) and Eps1p (40) and, with different spacings, in Mdp1p (C_xC; (41)) and Mpd2p (C_xC; (42)). The six-residue loop closed by the disulfide in yeast Pdi1p (33) might reasonably be expanded to eight residues in metazoan and plant QSOXs. The cysteines within the Trx2 domain, C165 and C237 (Figure 1), are not conserved (supplemental Figure S1) and are unlikely to play a catalytic role. They are distant in sequence and appear 12–14 Å apart in homology models based on the yeast Pdi1p (PDB ID 2B5E) and human thioredoxin 2 (PDB ID 1W4V) structures. Close to the N-terminus of the Erv/ALR domain are the conserved C393 and C405 residues (Figure 1 and supplemental Figure S1). Within the Erv/ALR domain are C449 and C452, which form the proximal C_{xx}C disulfide, named for its predicted location next to the bound flavin. By analogy to the pyridine nucleotide disulfide oxidoreductase family (22, 23) and homology to Erv2p and ALR, C452 interacts primarily with the flavin. Its partner, C449, would then serve as the interchange cysteine, forming mixed disulfides with other redox centers. The final conserved cysteine pair, C509 and C512, is in a comparable position in the HsQSOX1 primary structure as the C_xC motif of Erv2p and is expected to be in a flexible stretch of polypeptide chain prior to the beginning of the variable C-terminus.

Expression of Human QSOX1 in *Escherichia coli*

We used the *gor/trx* mutant *Escherichia coli* expression strain to produce HsQSOX1 because QSOX enzymes contain multiple disulfides. Induction at 37 °C led to expression of the protein as insoluble aggregates that could not be workably refolded using standard protocols. An increasing proportion of soluble enzyme was obtained as the induction temperature was lowered from 30 °C to 23 °C. However, the highest yields of active protein resulted from growth into log phase at 37 °C followed by re-suspension of the cells in fresh media containing 0.2 mM IPTG and growth at 15 °C for 24 h. HsQSOX1 was purified from crude

lysate by Ni-NTA chromatography, followed by application to a second Ni-NTA column for wild-type enzyme or to a Source 30 cation exchange column for mutant enzymes (see Experimental Procedures). Typically, 0.3 mg of HsQSOX1 was obtained with >90% purity on SDS-PAGE from 1 L of the original 37 °C culture. Although yields are low, the enzyme is stable for several months when stored at 4 °C in 50 mM phosphate buffer pH 7.5, containing 1 mM EDTA. The modest levels of protein forced us to curtail some obvious approaches and to conduct spectrophotometric experiments in microcells to conserve material.

Recombinant HsQSOX1 was found to share many properties with the previously characterized QSOX enzymes. The UV-visible spectrum of recombinant HsQSOX1 shows a typical unresolved flavin envelope (Figure 4) that was seen for both the rat seminal vesicle (RnQSOX1), the avian egg white (GgQSOX1) and the bovine milk (BtQSOX1) enzymes (1, 18, 19). The experimentally-determined extinction coefficient is $12.5 \pm 0.6 \text{ mM}^{-1} \text{ cm}^{-1}$ at 456 nm (see Methods), comparable to that obtained for GgQSOX1. HsQSOX1 contained 2.1 ± 0.1 DTNB-reactive thiols in denaturing conditions (see Methods), consistent with five disulfides and two free thiols. The apparent molecular weight of the enzyme on SDS-PAGE was 70 kDa, in good agreement with the expected value of 67.7 kDa from the sequence (supplemental Figure S2). Gel filtration, using a calibrated size-exclusion column under non-denaturing conditions, gave a corresponding value of 76 kDa. The striking result that recombinant HsQSOX1 is monomeric in solution was verified by equilibrium analytical ultracentrifugation (see Supplementary Materials).

Catalytic Activity of Recombinant HsQSOX1

Table 2 compares the catalytic activity of recombinant HsQSOX1 with native GgQSOX1 using three substrates: GSH, DTT, and RNase. As observed previously for GgQSOX1, reduced glutathione is a rather poor substrate of HsQSOX1 in terms of K_m (19–21). In contrast, reduced RNase is an excellent substrate of HsQSOX1, with catalytic efficiencies comparable to that observed with GgQSOX1 and BtQSOX1, consistent with the idea that reduced unfolded proteins may be the physiological substrates of this enzyme family (4, 19–21). We note that the K_m values for HsQSOX1, GgQSOX1, and BtQSOX1 towards GSH and DTT are comparable to those for the rat seminal vesicle enzyme described in 1980 (18). However the seminal vesicle enzyme is reported to have about 60- to 100-fold higher turnover numbers for GSH and DTT respectively (80,000 and 136,000 thiols oxidized per minute (18)). Our confidence in k_{cat} values in the 1000–4000/min range for vertebrate QSOXs (Table 2) is increased by our recent finding of similar values for native bovine QSOX1 (1). In summary, the recombinant human enzyme shows comparable activity to non-recombinant avian and bovine QSOXs (Table 2).

Role of three CxxC motifs in HsQSOX1 catalysis

The three CxxC motifs were previously implicated in catalysis of disulfide bond formation by GgQSOX1 (2, 17). To address the role of each CxxC motif in HsQSOX1, we made single Cys-to-Ala or -Ser mutants for all six cysteine residues; for one pair (509/512) we also generated an SxxS double-mutant (see Methods). Activities of mutant proteins were initially measured using 5 mM DTT or 38 μM reduced RNase (0.3 mM thiols) as substrates (Table 3).

When cysteines in the CxxC motif of the HsQSOX Trx1 domain were mutated to serine (C70S and C73S), residual activities of about 7.5% of wild-type with DTT and 1.5% with reduced RNase were observed (Table 3). These activities likely represent the basal oxidase activity of the Erv/ALR domain. An Erv/ALR domain fragment of avian QSOX1 and several smaller sulphydryl oxidases lacking Trx domains show turnover numbers from 20–

150 thiols oxidized per min at 5 mM DTT and, correspondingly, 0 – 8/min towards reduced RNase (6–8, 10, 17, 27, 43).

We next mutated the two cysteine residues that comprise the proximal disulfide C449–452 in HsQSOX1 (corresponding to C121–124 in Erv2p; Figure 1). In Erv2p, C121, the residue furthest from the isoalloxazine ring, would correspond to the interchange thiol (following the nomenclature developed for the pyridine nucleotide disulfide oxidoreductases (22, 23)) forming mixed disulfides with attacking thiolate nucleophiles. Scission of the proximal disulfide would then allow C124 to interact with the electron deficient isoalloxazine ring. Thiolate to flavin charge-transfer complexes have already been observed in static reductive titrations and during steady state turnover of avian QSOX1 (19, 21). Figure 5 confirms the identity of this charge-transfer donor as C452 as predicted from the sequence alignments and the structures of Erv/ALR family members (7, 9, 44). Thus, the spectrum of the C449A mutation shows a significant long wavelength feature centered at about 540 nm with the general signature of a charge-transfer interaction (45, 46). The C449S mutation also exhibits a long wavelength absorbance band but with a slightly different shape. While a serine residue may participate in hydrogen bonding interactions not found with alanine, a molecular explanation for this difference is not known at present. As expected the corresponding C452S mutation abolishes this charge-transfer interaction and leads to a native-like, but significantly blue shifted flavin envelope (Figure 5). Such blue shifts (here from 456 nm for the wild type to 449 nm) have been observed previously upon reduction of a disulfide adjacent to the flavin (2, 17, 19, 47, 48) and have been attributed to increased solvent access to the flavin that accompanies removal of the disulfide bond.

The C452S mutant shows very low activity with DTT and reduced RNase (0 and ~1% respectively; Table 3), as expected if this charge-transfer donor forms a covalent adduct with the flavin as a critical intermediate in the generation of dihydroflavin ((22, 28); Figure 6). While this adduct would normally be resolved by the interchange thiol (C449) as mentioned above, it might be attacked less effectively by one or more substrate thiols assuming the role of C449 in Figure 6. Thus the C449S mutant shows 4% and 2% of the wild type activity with DTT and reduced RNase respectively (Table 3).

Surprisingly, mutations of the distal disulfide (C509–512, previously implicated in catalysis, see above) had comparatively minor effects on the turnover numbers observed with either 5 mM DTT or 300 μ M reduced RNase thiols. Under these conditions the C509A mutant showed between 55 and 98% of the wild type activity and the corresponding C512A mutant between 26 and 59%. A double serine mutant retained 29–30% activity. These mutants were also evaluated by determining k_{cat} and K_m values for reduced RNase (Table 4). Again the impact of these mutations was minor, with k_{cat}/K_m values reduced by at most 4-fold (see later).

As mentioned earlier, the models for QSOX catalysis developed by analogy to Erv2p feature an intersubunit transfer of reducing equivalents from the distal (shuttle) to the proximal disulfides as depicted in Figures 1 and 3. However, we found that recombinant HsQSOX1 was monomeric. For example, gel filtration using a calibrated column gave an apparent molecular weight of 67 kDa (data not shown, see Methods). We also estimated the molecular weight by sedimentation equilibrium (see supplemental Material) as 70.7 kDa. Because the construct used for this study contained both an N-terminal His tag and a linker region of 26 amino acids, we also determined the molecular weight of a second construct of HsQSOX1 in which this extension to the N-terminus was removed (supplemental Figure S4). This version also behaves as a monomer (supplemental Figure S3). Thus the N-terminal His tag does not promote monomerization of a QSOX dimer. Consistent with these data, but in contrast to avian QSOX1 (19), rat seminal vesicle QSOX1 (25) and bovine QSOX1 (1)

also elute as monomers on gel filtration (18). Finally the turnover number of HsQSOX1 with 5 mM DTT is unaffected over a 40-fold range of concentrations (from 5 – 200 nM QSOX; data not shown). Hence the active form of HsQSOX is a monomer under these conditions.

Conclusions

Figure 7 presents a revised minimal catalytic mechanism for human QSOX1. We have shown that the HsQSOX1 monomer is active and that the conserved distal disulfide (16) does not have an essential role in catalysis of disulfide bond formation in dithiothreitol or reduced RNase. Distal disulfide mutants of HsQSOX1 show only small (up to 4-fold) decreases in k_{cat}/K_m in the oxidation of reduced RNase. It was observed previously that the distal disulfides of small Erv enzymes are also dispensable for certain activities. For example, Erv2p and AtErv1 lacking the distal disulfides oxidize DTT efficiently (27). More significantly, a monomeric mutant of Erv2p can oxidize model dithiol substrates and reduced Pdi1p without the obligatory intervention of the shuttle disulfide (27). This latter observation provides an important precedent in support of the model in Figure 7 in which a thioredoxin domain interacts directly with the proximal disulfide of a monomeric QSOX.

However, distal disulfides are essential for other activities of single-domain Erv/ALR sulfhydryl oxidases. For example, yeast strains expressing Erv2p without the distal disulfide no longer rescued a growth defect caused by the absence of the Ero1p sulfhydryl oxidase (7, 8). In addition, the distal disulfide of an Erv/ALR sulfhydryl oxidase from *Arabidopsis thaliana* (AtErv1) was essential for the *in vitro* oxidation of reduced thioredoxin (13). From our experiments, we cannot exclude the possibility that the distal disulfide of QSOX is involved in an alternate pathway for the oxidation of protein substrates whose importance depends on the type of protein oxidized, or that the distal disulfide is involved in the regulation of enzyme activity. If the distal disulfide indeed participates in electron transfer from certain substrates to the FAD-proximal disulfide in monomeric HsQSOX1, then it must differ in position from the distal disulfides of small, dimeric Erv family enzymes. The striking divergence of the QSOX sequence from that of Erv2p after the end of the fourth helix in the bundle (Figure 1) prevents prediction of the position of the QSOX distal disulfide on the basis of the Erv2p structure and opens the possibility that it approaches within dithiol-disulfide exchange distance of the active-site disulfide on the same polypeptide chain.

Finally, we note interesting parallels between monomeric metazoan QSOXs and yeast Pdi1p. Both proteins have four principal domains with the outermost domains being the ones that carry redox-active CxxC motifs. QSOX requires intramolecular redox communication between the Trx1 domain and the Erv/ALR module and, while inter-domain transfer is not expected to be an essential feature of PDI catalysis, the two CxxC domains of PDI face each other across a flexible cavity formed from the four domains. This twisted U shaped cavity is lined with hydrophobic residues presumably to facilitate binding of protein substrates and to discourage their aggregation. It will be interesting to learn whether the hydrophobic surfaces of QSOXs (19) play a similar role in handling client proteins during oxidative protein folding.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Abbreviations

ALR	augmenter of liver regeneration
DTNB	5,5'-dithiobis(2-nitrobenzoate)
DTT	dithiothreitol
Erv/ALR	the family of enzymes that comprises homologs of the yeast protein 'Essential for respiration and viability 1' including ALR
GSH	reduced glutathione
HRR	helix-rich region
IPTG	isopropyl- β -D-thiogalactopyranoside
PDI	protein disulfide isomerase
QSOX	flavin-dependent sulfhydryl oxidase homologous to Quiescin Q6.

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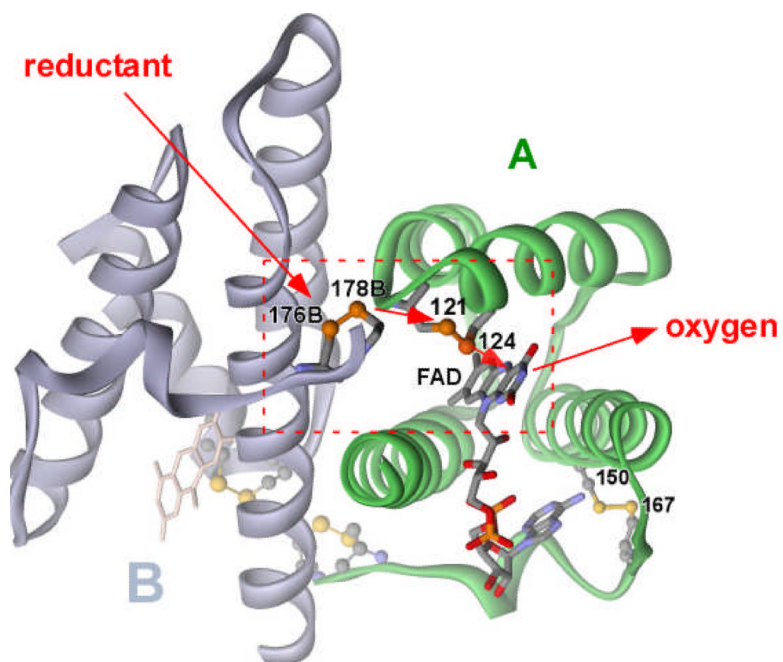


FIGURE 2. Crystal structure and the proposed flow of reducing equivalents in the yeast Erv2p homodimer. The proposed flow of reducing equivalents from the reductant (Pdi1p) to molecular oxygen is depicted from left to right using a series of arrows. An equivalent set of redox centers are shown unlabelled at the back of the homodimer. C150–167 is a structural disulfide in Erv2p that is absent in the multi-domain Quiescin-sulfhydryl oxidase (QSOX).

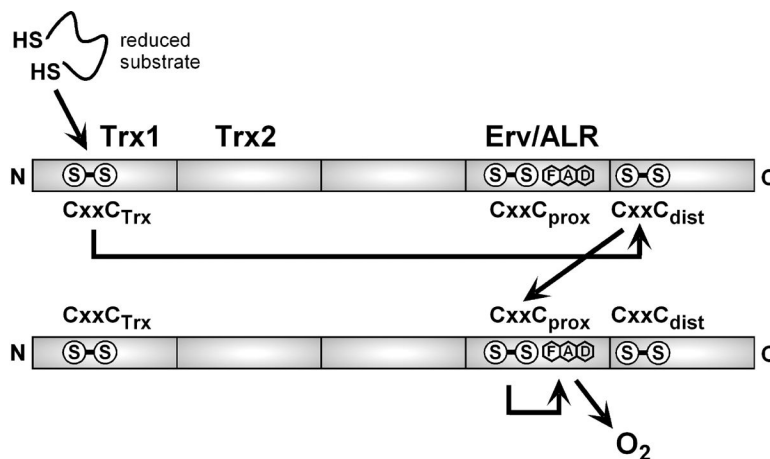


FIGURE 3. Suggested flow of reducing equivalents during oxidation of a reduced protein by avian QSOX1. The client reduced protein is oxidized by the Trx1 domain of QSOX followed by transfer of the pair of reducing equivalents to the Erv/ALR domain. The model incorporates an intersubunit disulfide exchange between distal and proximal CxxC motifs of a dimeric Erv/ALR domain as shown in Figure 2 (see Text).

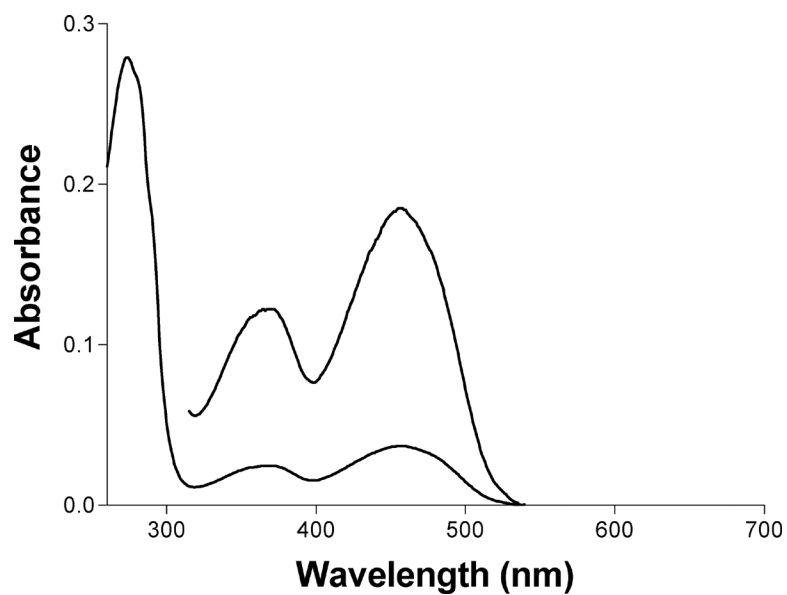


FIGURE 4.

UV-visible spectrum of recombinant human QSOX1. The spectrum of 2.9 μM enzyme was recorded in 50 mM phosphate pH 7.5, containing 1 mM EDTA and was corrected for small amounts of scattering material as described in the Experimental Procedures. The top spectrum is multiplied by 5 to show details of the flavin envelope.

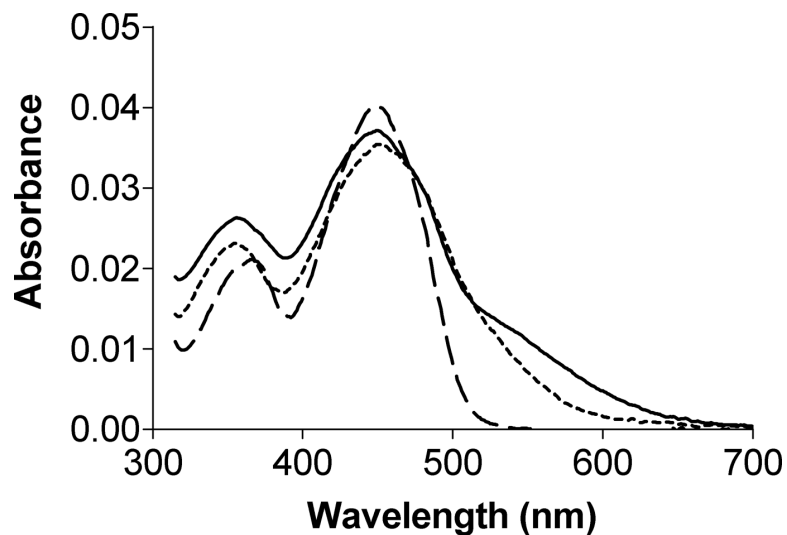


FIGURE 5. UV-Visible spectra of cysteine mutants of the proximal disulfide of HsQSOX1. Spectra of the following mutants of HsQSOX1 in phosphate buffer pH 7.5: C449A (solid line), C449S (short dashed line), and C452S (long dashed line). Spectra were normalized to correspond to a concentration of 3.2 μ M using the extinction coefficients reported in the Experimental Procedures.

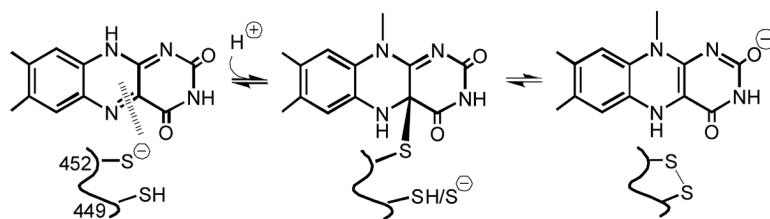
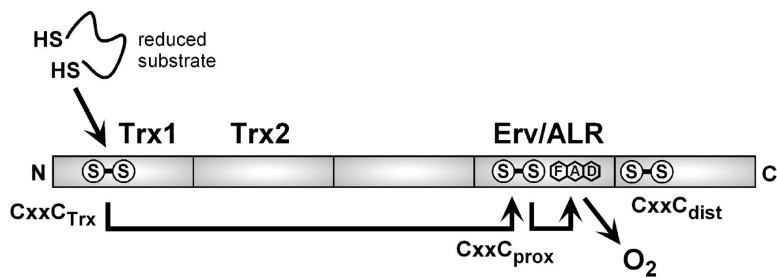


FIGURE 6. Proposed mechanism for the transfer of reducing equivalents between flavin and the reduced proximal disulfide in QSOX. The interchange and charge transfer thiols correspond to C449 and C452 respectively.

**FIGURE 7.**

Revised model for flow of reducing equivalents in QSOX. This model involves a transitory mixed disulfide between C70 of Trx1 and C452 of the proximal disulfide in the Erv/ALR domain (see the Text). The distal CxxC disulfide (C509–512) would be on the opposite face of the Erv/ALR domain more than 25 Å from C452.

Table 1

Proposed function and connectivity of cysteine residues in human QSOX1.

Cysteines	Proposed connectivity	Proposed function (domain location)	Corresponding residues in related proteins
C70	Disulfide, CxxC	Catalytic (Trx1)	C61/C64, yeast PDI1p
C73			
C101	Disulfide, Cx ₈ C	Structural/regulatory (Trx1)	C90/C97, yeast PDI1p Cx ₆ C C91/C99, yeast Mpd2p Cx ₇ C C87/C96, yeast Mpd1p Cx ₈ C
C110			
C165	Cysteines (not conserved)	(Trx2)	Not applicable
C237			
C393	Disulfide, Cx ₁₁ C	Unknown (Spacer-Erv/ALR interface)	Spacer domain unique to QSOXs
C405			
C449	Disulfide, CxxC	Catalytic (Erv/ALR)	C121/C124, Erv2p
C452			
C509	Disulfide, CxxC	Uncertain, see text (Erv/ALR-C terminus interface)	C176/C178, Erv2p
C512			

Table 2

Comparison of substrate kinetics for human, avian and bovine QSOX1.^a

	Human QSOX			Avian QSOX ^{b,c}			Bovine QSOX ^{b,c}		
	TN _{max} min ⁻¹	Km mM	TN _{max} /Km M ⁻¹ s ⁻¹	TN _{max} min ⁻¹	Km mM	TN _{max} /Km M ⁻¹ s ⁻¹	TN _{max} min ⁻¹	Km mM	TN _{max} /Km M ⁻¹ s ⁻¹
DTT	1240	0.10	2.0 × 10 ⁵	2066	0.150	2.30 × 10 ⁵	1880	0.086	3.66 × 10 ⁵
GSH	1480	12.4	1.98 × 10 ³	2770	20	2.30 × 10 ³	1760	4.9	5.94 × 10 ³
RNase	4320	0.32	2.2 × 10 ⁵	1220	0.115	1.76 × 10 ⁵	1340	0.060	3.74 × 10 ⁵

^aTurnover numbers are quoted in terms of thiols oxidized per min (and not disulfides generated per min as listed previously).

^{b,c}Taken from references to the avian (19, 20) and the bovine QSOX1 (1) enzymes expressing the turnover numbers as above.

Table 3

Turnover numbers for wild-type and cysteine mutants of HsQSOX1 using DTT and reduced RNase substrates.^a

Protein	5 mM DTT TN min ⁻¹	Relative TN	300 μM RNase thiols TN min ⁻¹	Relative TN
WT	1280 ± 40	1.00	2170 ± 14	1.00
C70S	88 ± 14	0.07	46 ± 24	0.02
C73S	108 ± 12	0.08	42 ± 16	0.02
C449S	48 ± 18	0.04	42 ± 20	0.02
C452S	0	0	16 ± 8	0.007
C509A	1260 ± 240	0.98	1200 ± 140	0.55
C512A	760 ± 10	0.59	570 ± 120	0.26
C509S,C512S	380 ± 60	0.30	630 ± 70	0.29

^aTurnover numbers are reported as thiols oxidized (not disulfides generated) per minute.

Table 4

Kinetic parameters of serine and alanine mutants of the distal disulfide.

	RNase TN _{max} min ⁻¹	Km μM per thiol	TN _{max} / Km M ⁻¹ s ⁻¹
WT	4320	320 ± 35	2.2 × 10 ⁵
C509A	5240	700 ± 230	1.3 × 10 ⁵
C512A	1680	420 ± 130	6.6 × 10 ⁴
C509S,C512S	740	230 ± 20	5.4 × 10 ⁴