Methionine sulfoxide reductases preferentially reduce unfolded oxidized proteins and protect cells from oxidative protein unfolding*

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Running title: MSRs preferentially reduce unfolded oxidized proteins

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Keywords: Methionine sulfoxide; Methionine sulfoxide reductase; Oxidative stress; Protein unfolding; Yeast

Background: Methionine sulfoxide reductases have previously been studied mostly using low molecular weight substrates.

Results: Methionine sulfoxide reductases preferentially reduce unfolded oxidized proteins.

Conclusion: These enzymes serve a critical function in protein folding by repairing oxidized nascent polypeptides and unfolded proteins.

Significance: Understanding precise functions of methionine sulfoxide reductases will help define mechanisms of protein repair and identify their physiological substrates.

SUMMARY

Reduction of methionine sulfoxide (MetO) residues in proteins is catalyzed by methionine sulfoxide reductases A (MSRA) and B (MSRB), which act in a stereospecific manner. Catalytic properties of these enzymes were previously established mostly using low molecular weight MetO-containing compounds, whereas little is known about the catalysis of MetO reduction in proteins, the physiological substrates of MSRA and MSRB. In this work, we exploited an NADPH-dependent thioredoxin system and determined the kinetics parameters of yeast MSRA and MSRB using three different MetO-containing proteins. Both enzymes showed Michaelis-Menten kinetics with the $K_M$ lower for protein than for small MetO-containing substrates. MSRA reduced both oxidized proteins and low molecular weight MetO-containing compounds with similar catalytic efficiencies, whereas MSRB was specialized for the reduction of MetO in proteins. Using oxidized glutathione-S-transferase as a model substrate, we showed that both MSR types were more efficient in reducing MetO in unfolded than in folded proteins, and that their activities increased with the unfolding state. Biochemical quantification and identification of MetO reduced in the substrates by mass spectrometry revealed that the increased activity was due to a better access to oxidized MetO in unfolded proteins; it also showed that MSRA was intrinsically more active with unfolded proteins regardless of MetO availability. Moreover, MSRs most efficiently protected cells from oxidative stress that was accompanied by protein unfolding. Overall, this study indicates that MSRs serve a critical function in the folding process by repairing oxidatively damaged nascent polypeptides and unfolded proteins.
INTRODUCTION

Proteins may undergo various post-translational modifications altering their structure and function. Their sulfur-containing residue, methionine (Met), can be oxidized to \( R \)- and \( S \)-diastereoisomers of Met sulfoxide (MetO). This modification is reversible, as MetO can be reduced back to Met by methionine sulfoxide reductases (MSRs) A (MSRA) and B (MSRB), which are specific for the \( S \)- and \( R \)-forms of MetO, respectively. These enzymes are present in almost all living organisms and catalyze the reduction of their substrates at the expense of NADPH using thioredoxin (Trx) or glutaredoxin systems (1, 2).

Whereas the catalytic mechanisms of MSRs are well characterized (2, 3), their physiological functions remain elusive, mainly due to insufficient information on the identity of their cellular targets. The absence of clearly established substrates also limits studies on specificity of Met oxidation and MetO reduction in proteins. Met oxidation has been reported for proteins, which could be classified into three groups (4): (i) enzymes activated by Met oxidation, such as the calcium/calmodulin-dependent protein kinase II (5); (ii) proteins not impaired by Met oxidation, which could fulfill, together with MSRs, a protective (antioxidant) function though cyclic oxidation and reduction of Met (6); and (iii) proteins damaged by Met oxidation, such as those involved in neurodegenerative diseases (7, 8). The consequence of Met oxidation is determined by the effects of this modification on structure and function of these proteins; however, these effects have been characterized only for a handful of proteins. For example, in human prion, oxidation of two Met residues converts a cellular \( \alpha \)-helix rich form to the infectious \( \beta \)-sheet rich form by perturbing the network of stabilizing interactions (9). In addition, oxidation of two solvent-accessible Met in a human growth hormone increases its susceptibility to thermal denaturation (10), and oxidation of two Met in calmodulin prevents protein-protein interaction due to incompatibility of MetO for stable \( \alpha \)-helixes (11).

MSRs, which repair oxidative modifications, play important roles in the protection of proteins from oxidative stress in various eukaryotes (12–14). Similar protective effects were also observed in prokaryotes, particularly in the case of the stress induced by hypochlorite, a strong antimicrobial agent found in household bleach that is also produced by mammalian neutrophils to kill invading microorganisms (15). For instance, a Helicobacter pylori strain deficient in the expression of MSRs could not survive in a neutrophil cell culture (16). Hypochlorite is known to have a dual effect in provoking oxidation and unfolding of proteins, and trigger chaperone activation (17). These studies imply that MSR might directly participate in the protection from oxidative and unfolding stress through the reduction of MetO in proteins.

The MSR activity was quantified for several protein substrates, such as bacterial Ffh and several targets of plant MSRBs (13, 18, 19). However, little is known about enzyme kinetics of MSRs with oxidized protein substrates, their physiological targets. The yeast Saccharomyces cerevisiae possesses single MSRA and MSRB genes, as well as an \( f^{R} \)MSR that is specific for the reduction of the \( R \)-diastereoisomer of free MetO (20, 21). In this work, we took advantage of the NADPH-coupled Trx system and used different forms of MetO-containing proteins as substrates to examine functions of yeast MSRA and MSRB.

EXPERIMENTAL PROCEDURES

Cloning and site-directed mutagenesis - Sequences coding for NADPH-dependent thioredoxin reductase 1 (TRR1), Trx1, MSRB (from codon 30 to the stop codon), were amplified by PCR from the S. cerevisiae genomic DNA using Platinum® Pfx DNA Polymerase (Invitrogen, Carlsbad, CA, USA) and specific pairs of primers shown in Supplemental Table S1. Similarly, sequences coding for Met-rich protein 4 (MRP4) (from codon 23 to stop codon) and Met-rich protein 5 (MRP5) (from codon 21 to stop codon) were amplified from genomic DNA of Idiomarina loihiensis L2TR and Pseudomonas putida W619,
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respectively, using specific primers. Amplicons were purified and digested with NdeI and BamHI for cloning of TRR1 and Trx1 in pET15b or with BamHI and XhoI, or NheI and NotI, for cloning MSRB and MRP4 in pET21b (EMD Biosciences, Billerica, MA, USA), respectively. Met-rich protein 1 (MRP1) (22) and MRP5 were cloned similarly in p425 yeast expression vector under GPD promoter using BamHI and SalI restriction sites.

Site-directed mutagenesis of MSRA and MSRB was made by whole plasmid amplification with Phusion® High-Fidelity DNA Polymerase (Thermo Scientific, Billerica, MA, USA) using primers containing mutated bases (Supplemental Table S1). After amplification, the methylated template vector was digested by incubation with DpnI for 1 h at 37 °C. µl of the digested PCR product was used to transform NEB 5-alpha Competent Escherichia coli (High Efficiency) cells (New England Biolabs, Ipswich, MA, USA) and clones were selected on Luria-Bertani plates containing ampicillin (50 µg.ml⁻¹). All constructs were validated by DNA sequencing.

Expression and purification of recombinant proteins - SoluBL21™ E. coli (Gelantis, San Diego, CA, USA) cells were transformed with the expression vector and grown in Luria-Bertani containing ampicillin (50 µg.ml⁻¹) at 37 °C. When the OD₆₀₀ reached ~0.6, production of the recombinant protein was induced by addition of 100 µM isopropyl β-D-1-thiogalactopyranoside. After overnight incubation at 25 °C, cells were harvested by centrifugation. For TRR1, Trx1, MSRA, MSRB and MRP4 containing His₆-tag, pellets were resuspended in PBS containing 25 mM imidazole in the presence of Complete, EDTA-free, Protease inhibitor cocktail (Roche, Indianapolis, IN, USA), and for GST, the pellet was resuspended in PBS. Cells were disrupted by sonication, His₆-tagged proteins were purified on nickel-containing His•Bind® Resin (Novagen, Billerica, MA, USA), and GST was purified on Glutathione Sepharose 4 Fast Flow (GE Healthcare). Protein solutions were concentrated using 15 ml Amicon® Ultra concentrators with 30 kDa or 10 kDa cutoffs (Millipore, Billerica, MA, USA) and desalted in 30 mM Tris-HCl, pH 8, using 5 ml HiTrap™ Desalting columns (GE Healthcare). Protein concentrations were determined spectrophotometrically using the Pierce® BCA Protein assay kit (Thermo Scientific) and specific molar extinction coefficients at 280 nm. Protein purity was verified using SDS-PAGE gels stained with Imperial™ Protein Stain (Thermo Scientific).

Protein and N-acetyl-MetO preparation - MRP4, GST, bovine β-casein (Sigma-Aldrich, St. Louis, MO, USA), and Bacillus sp. α-amylase (Sigma-Aldrich) (all 1 mg.ml⁻¹) were oxidized by incubation with 100 mM H₂O₂ in PBS overnight at room temperature, concentrated and then desalted using HiTrap™ Desalting or Illustra NAP5™ columns (GE Healthcare) in 30 mM Tris-HCl, pH 8. For unfolding assays, oxidized proteins were incubated with urea at a final concentration ranging from 0.25 M to 7 M for 1 h at room temperature prior to the assays. N-acetyl-MetO was prepared as described (23).

MSR absolute stoichiometry and activity with dabsyl-MetO - Activities of recombinant MSRs (1 µM) were determined by monitoring the reduction of 0.5 mM dabsyl-MetO in the presence of 20 mM DTT. The absolute stoichiometry was determined similarly, 100 µM of reduced and desalted MSRs was incubated with 1 mM dabsyl-MetO for 1 h at room temperature without the reducing agent. Dabsyl-Met and dabsyl-MetO were separated by HPLC using a C₁₈ reverse phase column, SunFire™ 3.5 µm, 3.0 x 50 mm (Waters, Milford, MA, USA) as described (2).

MSR activity assays - MSR activity was measured following NADPH oxidation at 340 nm in the presence of the Trx system (200-400 µM NADPH, 2 µM TRR1, 25 µM Trx1) using 1-10 µM MSRA or MSRB in the presence of free MetO (0.25-20 mM), N-acetyl-MetO (62.5
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µM to 5 mM), oxidized MRP4 (0.8-200 µM), oxidized β-casein (6.25-200 µM), oxidized GST (9.4-150 µM) or urea-treated oxidized GST (4.8-150 µM) as substrates. Reactions were carried out at 25 °C in a 500 µl reaction volume. MSR activities were calculated from the slope after subtracting the background (absence of the enzyme), considering that 1 mol of oxidized NADPH corresponds to 1 mol of MetO reduced. The apparent stoichiometry was determined similarly using sub-saturating concentrations of substrates: 0.8-6.5 µM oxidized MRP4, 7-29 µM oxidized β-casein, 11-44 µM oxidized GST and 5.6-22 µM 4 M urea-treated oxidized GST. The amount of oxidized NADPH was determined after the rate of oxidation reached the basal level. To test the activity using the urea-treated oxidized protein, 10-25 µM substrate was used, and urea was added in control assays at the same concentration (less than 50 mM). Kinetics and catalytic parameters were calculated from non-linear regressions using GraphPad Prism 4.0 (GraphPad Software, Inc, La Jolla, CA, USA).

Fluorescence analyses - Emission spectra of intrinsic fluorescence were recorded in 200 µl of 30 mM Tris-HCl, pH 8.0, containing 10 or 25 µM non-oxidized, oxidized or urea-treated oxidized proteins with excitation at 280 nm. As a control, amounts of urea used in the urea-treated proteins were added to the oxidized protein samples (less than 50 mM). The same samples were used to determine 8-anilinonaphthalene-1-sulfonate (ANS) fluorescence. Emission at 466 nm (excitation at 377 nm) was recorded 15 min after addition of 50 µM ANS. Fluorescence was recorded in 96-well microplates using a SpectraMax M5 fluorescence microplate reader (Molecular Devices, Sunnyvale, CA, USA).

Mass spectrometry analysis - Oxidized MRP4, oxidized GST and urea-treated oxidized GST (100 µM) were incubated with or without 5 µM MSRA or MSRB in the presence of 10 mM DTT in 30 mM Tris-HCl, pH 8, for 1 h at 25 °C. Solution containing 50 µg substrate was incubated with 12.5 mM iodoacetamide for 30 min at 25 °C in 0.1 M ammonium sulfate, then with 1 µg Sequencing Grade Modified Trypsin (Promega, Madison, WI, USA) and 1 mM CaCl₂ overnight at 37 °C. An additional 1 µg of trypsin was added and the solution incubated for 2 h at 37 °C. The trypsin digests were frozen and stored at -80 °C until mass spectrometry analysis. LC-MS/MS analysis was performed on an LTQ-Orbitrap Discovery mass spectrometer (Thermo Fisher) coupled to an Agilent 1200 series HPLC system. Tryptic digest (30 µl) was pressure loaded onto a 250 µm fused silica desalting column packed with 4 cm of Aqua C₁₈ reverse phase resin (Phenomenex, Torrance, CA, USA). The peptides were then eluted onto a C₁₈ column (100 µm fused silica with a 5 µm tip, packed with 10 cm C₁₈) using a gradient 5–100% buffer B in buffer A (buffer A: 95% water, 5% acetonitrile, 0.1% formic acid; buffer B: 20% water, 80% acetonitrile, 0.1% formic acid) and into the mass spectrometer. The flow rate through the column was set to ~0.25 µl.min⁻¹ and the spray voltage was set to 2.75 kV. One full MS scan (FTMS) (400–1,800 MW) was followed by 7 data dependent scans (ITMS) of the nᵗʰ most intense ions with dynamic exclusion enabled.

Peptide identification - The tandem MS data were searched using the SEQUEST algorithm (24) using a concatenated target/decoy variant of the human and mouse International Protein Index databases modified to include the sequences for the proteins used in this study. A static modification of + 57.02146 on cysteine was specified to account for iodoacetamide alkylation and a differential modification of + 16 was specified on methionine to account for oxidation. SEQUEST output files were filtered using DTASelect 2.0 (25). Reported peptides were required to be fully tryptic and discriminant analyses were performed to achieve a peptide false-positive rate below 5%. The percentage of oxidation per Met was calculated using the redundancy of the peptide containing the specific Met, oxidized or not. The percentage
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of oxidation values corresponds to the number of time a MetO was found divided by the total number of times the peptide was found (coverage), multiplied by 100. This method allows quantifying the oxidation of each Met found in peptide containing several Met, which is not possible using area integration.

Yeast spotting assays - MSRA expressed under the GPD promoter from the high copy number yeast expression vector p425 and MSR null strains were described previously (21). WT yeast cells were transformed with MRP1, MRP5 and/or yeast MSRA constructs. After colony formation, single colonies were picked up from the plates and grown overnight in the media lacking histidine or leucine. The following day, these cells were re-transformed with MSRA and/or empty vectors and incubated on media lacking histidine and leucine. This procedure was also applied for empty vector transformation which was used as a control. In the spotting assay, the indicated strains were grown overnight in appropriate media and diluted to an OD_{600} of 0.3. Cells were washed with the pre-warmed PBS buffer and incubated in 1 ml PBS and indicated concentrations of sodium hypochlorite (NaOCl). Every 5 min, 5 µl of each culture was spotted on the selective media. Yeast MSR mutant strains lacking MSRA, MSRB or both genes, were also tested as indicated above for NaOCl sensitivity. All plates were incubated for 3 days at 30 °C and photographed. Halo assays were performed to assess viability of MSRA, MRP1 and MRP5 overexpressing yeast cells under conditions of H_2O_2 stress. Cells were prepared as indicated above and the OD_{600} was adjusted to 0.5. Cells were washed with the pre-warmed PBS and 1 ml of each culture spread onto agar plates missing appropriate amino acids for selection. Plates were dried for 1 h at room temperature, and filter paper discs were placed in the middle of each plate. 4 µl of 30% hydrogen peroxide were applied onto each paper disc, and the plates were incubated for 3 days at 30 °C and photographed. The diameter of cleared zones in each plate was measured with a ruler. This experiment was repeated three times.

Modeling, structural analysis of substrate proteins and determination of MSR hydrophobicity - The percentage of amino acids included in disordered regions was calculated using SPINE-D (26) (http://sparks.informatics.iupui.edu/SPINE-D/) and Multilayered Fusion-based Disorder predictor (MFDp) (27) (http://biomine-ws.ece.ualberta.ca/MFDp.html) web-servers. The structural coordinates of MSRA and GST were obtained from the PDB repository (pdb codes, 3PIL and 1DUG, respectively). To model MSRB, we used a fold-recognition algorithm, FAS03 (http://ffas.ljcrf.edu/), to generate alternative profile-profile alignments. MSRB were modeled with Modeller (http://salilab.org/modeller) using the alignments generated with FAS03 and 2K8D as template. Detailed atomic exposure calculations were calculated with Surface Racer© (28), http://apps.phar.umich.edu/tsodikovlab/index_files/Page756.htm). Structural profiles were generated as previously described (29). Briefly, all residues lying with one or more of their atoms found within 8 Å from the sulfur atom of the catalytic Cys residues of yeast MSRA and MSRB were considered and then separately analyzed for their hydrophobic content (implementing the standard Kyte-Doolittle scale, where each amino acid is described by numeric value ranging from negative, i.e., hydrophilic, to positive, i.e., hydrophobic, (Supplemental Fig S4)) (30), and compositional features (e.g., content of basic, acidic, or aromatic residues) through in-house Python (v2.6) scripts.

RESULTS

Yeast MSRA efficiently reduces oxidized proteins and free MetO whereas MSRB is specialized for the reduction of oxidized proteins - To characterize the kinetics of yeast MSRA and MSRB, we utilized the NADPH-coupled Trx system, which is physiologically relevant and allows comparative analyses of MetO-containing substrates. Catalytic parameters of MSRs are usually determined using low-molecular MetO-containing compounds, whereas little is known on the catalysis of MetO reduction in oxidized proteins. We examined
MSRAs and MSRBs preferentially reduce unfolded oxidized proteins and used free MetO and N-acetyl-MetO for comparison. These proteins were chosen based on their high Met content and structural features: both Met-rich protein 4 (MRP4) and β-casein are predicted to be completely disordered and possess 31 (22%) and 7 (3.1%) Met residues, respectively. A third protein, glutathione-S-transferase (GST), has the Met content similar to that of β-casein (9 Met, 3.8%), but is a highly structured protein (Supplemental Table S2). For all substrates, MSRA and MSRB catalysis followed the Michaelis-Menten kinetics (Table 1). MSRA displayed the \( k_{\text{cat}} \) values from \(~1 \text{s}^{-1}\) to \(~13 \text{s}^{-1}\). The \( K_M \) values were \(~0.5 \text{mM} \) for free MetO, N-acetyl-MetO and GST, and \(~10 \) fold lower for MRP4 and β-casein. The catalytic efficiency \( (k_{\text{cat}}/K_M) \) was \(~2.1 \text{mM}^{-1}\cdot\text{s}^{-1} \) for GST, and this value increased \(~5, ~13, ~15 \) and \(~200 \) fold in the case of free MetO, β-casein, N-acetyl-MetO and MRP4, respectively. For MSRB, the \( k_{\text{cat}} \) values were \(~1 \text{ s}^{-1}\) for all tested substrates, but a striking difference was observed in \( K_M \) values, which were \(~6.5 \text{ mM} \) for free MetO and \(~10 \) times lower for N-acetyl-MetO. The differences were more pronounced for proteins, \( i.e. \), the \( K_M \) values were 45-50 fold lower for oxidized proteins than for free MetO (Table 1). Analysis of these data reinforces the idea that MSRB is far more efficient in the reduction of MetO in oxidized proteins \( e.g. \), its catalytic efficiency was \(~1,200 \) fold higher for MRP4 than for free MetO), whereas MSRA reduces both proteins and MetO-containing compounds with similar efficiency.

Quantification of MetO reduction in protein substrates - The absolute stoichiometry displayed by each MSR was 1 mol of MetO reduced per mol of enzyme (Supplemental Fig. S1A). Two redox-active Cys were used by each enzyme, and mutation analyses verified the roles of Cys25 and Cys176 in MSRA, and Cys157 and Cys97 in MSRB as catalytic and resolving Cys, respectively (Supplemental Fig. S1B). We further estimated the number of MetO reduced by MSRA and MSRB in oxidized proteins as the apparent stoichiometry \( i.e., \), mol of NADPH oxidized per mol of substrate using sub-saturating concentrations of substrates. Figure 1 shows the data for MRP4. After addition of the substrate, NADPH consumption was followed until the rate reached the background level, representing the state when all MetO residues reducible by MSR were reduced (Fig. 1A). The amount of oxidized NADPH was then plotted as a function of substrate concentration (Fig. 1B). The slope of the calculated linear regression corresponded to the apparent stoichiometry. Using MSRA and increasing concentrations of MRP4, the apparent stoichiometry of \(~4 \text{ mol NADPH ox. mol sub}^{-1}\) was found (Table 1; Fig. 1B). With MSRB, this value was \(~7 \text{ mol NADPH ox. mol sub}^{-1}\) (Table 1; Fig. 1B). As changes in oxidized NADPH were directly proportional to those in MetO (18, 31), the data indicate that MSRA and MSRB reduced 4 and 7 MetO equivalents, respectively, in the oxidized MRP4. Since this protein has 31 Met, 13% and 23% Met equivalents were reduced, respectively. To compare catalytic parameters for different substrates, the data were normalized by multiplying the \( K_M \) values by the apparent stoichiometry, yielding values per MetO reduced, and thus allowing to remove variation due to the different numbers of MetO reduced in each substrate (Table 1). For MSRA, this normalization gave a \( k_{\text{cat}}/K_M \) ratio similar to that for N-acetyl-MetO and free MetO. For MSRB, the corrected catalytic efficiency was 15 and 150 fold higher than that for N-acetyl-MetO and free MetO, respectively (Table 1). Similar analyses with β-casein showed that 2.5 and 3.2 MetO equivalents, which correspond to 36% and 46% of the 7 Met in the protein, were reduced by MSRA and MSRB, respectively. The corrected catalytic efficiency was similar to that obtained for MSRA with free MetO and 50 fold higher in the case of MSRB (Table 1).

The analysis of apparent stoichiometries further indicated that 1.2 and 0.8 MetO equivalents were reduced per molecule of oxidized GST, corresponding to 13% and 9% of all Met in the case of MSRA and MSRB, respectively (Table 1). In the MSRA-catalyzed reaction, the corrected catalytic efficiency observed with GST as the substrate was 5-50 fold lower than with other tested substrates. In the MSRB-dependent reduction of oxidized GST, this correction gave a catalytic efficiency 45 fold higher than that
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obtained with free MetO, but similar to those determined for the two other proteins. It is noteworthy that the oxidized GST was a better substrate for MSRB: its catalytic efficiency was 2 fold higher than the MSRA value. Altogether, these results indicated that MSRA efficiently reduced both free and protein-bound MetO, whereas MSRB showed a dramatic preference for oxidized proteins compared to low molecular weight MetO-containing molecules, such as free MetO and N-acetyl-MetO.

Differential reduction of MetO residues by MSRA and MSRB - Quantification of MetO reduced by MSRs in the tested proteins suggested that the various oxidized Met were not equivalent substrates. We subjected MRP4 and GST to tryptic digestion and LC-MS/MS analyses to determine the oxidation state of each Met before and after reduction by MSRs (Table 2; Supplemental Table S3). The coverage of oxidized MRP4 was 64%, allowing us to determine precisely the oxidation state of 11 Met among the 31 Met present in the protein (Supplemental Table S3). All detected Met were highly oxidized (more than 90% of Met were in the form of MetO), with the exception of Met66 (48% oxidized). After the reduction by MSRA or MSRB, protein coverage was 89% in both cases, allowing us to estimate the oxidation status of 27 Met, which corresponded to 55% and 64% for the substrate reduced by MSRA and MSRB, respectively. Considering only the 11 Met, for which the oxidation state was determined in the oxidized MRP4 prior to MSR reduction, MSRA and MSRB reduced 28% and 21% of MetO, respectively. However, not all MetO were reduced with the same efficiency and their reduction also depended on the MSR used. For instance, oxidized Met31 was reduced only by MSRA, and oxidized Met66 only by MSRB. In addition, the last Met, Met125, was completely oxidized, but reduced by neither MSR (Supplemental Table S3).

Met residues were on average 70% oxidized in the oxidized GST, with an oxidation status varying from 16% (Met154) to 99% (Met168). Surface accessibility of each Met, calculated using protein structure, was found to correlate with the oxidation status, with the 4 buried Met being less than 80% oxidized (Table 2). The average percentage of reduction by MSRA and MSRB was 18% and 24%, respectively, but the data varied significantly, indicating that, as in the case of MRP4, not all MetO served as substrates for MSRs. For instance, oxidized Met69 and Met81 were efficiently reduced by MSRA and MSRB, respectively, whereas oxidized Met94, Met165 and Met168 were reduced by neither enzyme. These results indicate that MetO residues in the tested proteins were not equivalent MSR substrates and that sequence and structure properties influenced the capacity for their reduction. The data also suggest that the complete reduction of MetO in the protein substrate would require opening of the structure to give access to MSRs, as low reduction capacities were recorded for buried MetO.

MSRs preferentially reduce unfolded proteins - To examine the reduction of MetO present in the hydrophobic core of protein substrates, we assayed MSRA and MSRB activities with the three oxidized Met-rich proteins described above and Bacillus subtilis α-amylase that contains 12 Met (2.3%) and is highly structured (Supplementary Table S2), following treatment with 4 M urea, a chaotropic agent that leads to protein unfolding (Fig. 2). These proteins were treated with urea in a small volume, followed by dilution of the chaotropic agent when the treated proteins were transferred to the reaction mixture. As a control, the MSR activities were determined with untreated oxidized proteins added to the reaction mixture containing the same final amount of urea. No significant differences in MSRA or MSRB activities were observed in the case of urea-treated or untreated oxidized MRP4 and β-casein, both of which are unstructured proteins. In contrast, when oxidized α-amylase was used as a substrate, the MSRA and MSRB activities were significantly increased, by 1.5 and 2.3 fold, respectively, with the urea-treated substrate compared to the non-treated protein. These activity increases were even more dramatic in the case of urea-treated oxidized GST, which showed 2.8 and 3.6 fold increase in MSRA and MSRB activity, respectively, compared to the non-treated protein (Fig. 2A). Although the proteins...
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unfolded by 4 M urea were rapidly assayed following the transfer to the MSR activity assay mixture whereby diluting urea, we could not exclude a possibility of spontaneous partial refolding of proteins during the enzymatic reaction. To monitor the substrate folding state during activity measurements, we characterized their intrinsic fluorescence and used an 8-anilinonaphthalene-1-sulfonate (ANS) probe, which emits fluorescence upon binding to hydrophobic areas of proteins (Fig. 2B; Supplemental Fig. S2). We assayed non-oxidized, oxidized and urea-treated oxidized proteins. For all tested proteins, oxidation induced changes in intrinsic and ANS fluorescence (with the exception of MRP4, for which no intrinsic fluorescence could be detected), indicating modification of protein structure. However, when oxidized proteins were compared with the urea-treated oxidized proteins, changes were observed in intrinsic and ANS fluorescence (with the exception of MRP4 and β-casein, indicating that urea treatment did not induce changes in the folding state as these proteins are not structured (Supplementary Table S2). In contrast, following urea treatment, α-amylase and GST remained significantly changed under conditions used in the MSR activity assays (Fig. 2B; Supplemental Fig. S2). Altogether, these results indicate that for the tested proteins, MSRA and MSRB activities were higher with unfolded oxidized than with folded oxidized proteins.

GST was further used to investigate MSRA and MSRB activities with unfolded proteins in detail. The kinetics determined using increasing concentrations of urea-treated oxidized GST could be described by a regression curve similar to an allosteric sigmoidal (Equation 1), with the h values equal to 1.4 ± 0.3 and 1.5 ± 0.1 for MSRA and MSRB, respectively (Fig. 3). However, these values are close to 1 and the known MSR mechanism with Trx as a reductant (32) suggest that modification of this kinetics could be due to an experimental factor, especially for the low values, and does not reflect an allosteric behavior. In the case of MSRA (Table 1; Fig. 3A), the catalytic efficiency (kcat/KM) was 6 fold higher with the urea-treated oxidized GST than the oxidized GST (Table 1; Fig. 3A). Similarly, for MSRB, the catalytic efficiency revealed a 3.2 fold increase in activity with the urea-treated oxidized GST compared to the oxidized protein (Table 1; Fig. 3B). The apparent stoichiometries determined for MSRA and MSRB revealed that 3.2 and 4.7 MetO equivalents, corresponding to 36% and 52% of the 9 Met, were reduced, respectively. Normalization of catalytic parameters to the stoichiometry revealed that both MSRA and MSRB were more efficient in reducing the urea-treated oxidized GST considering the whole protein as a substrate. Moreover, MSRA was more efficient even when the activity values were corrected to the number of MetO reduced in the protein substrate.

The apparent stoichiometries indicated that a higher proportion of MetO was reduced in the urea-treated oxidized GST than in the oxidized protein. To determine the identity of MetO used as substrates following urea treatment, the unfolded protein was analyzed by mass spectrometry after incubation with MSRA or MSRB (Table 2). In both cases, the average percentage of reduction was almost doubled for the urea-treated oxidized GST compared to the native protein. In particular, buried Met154 and Met165 were more efficiently reduced by both MSRA and MSRB, indicating that urea treatment allowed better access to buried MetO (Table 2).

We further incubated oxidized GST with increasing concentrations of urea to examine MSRA and MSRB activities as a function of the unfolded state of protein substrate. To characterize the folding state, ANS and intrinsic fluorescence were measured for each urea concentration (Fig. 4). Following oxidation, ANS fluorescence increased with the increase in urea concentration, up to 4 M, presumably due to increased exposure of hydrophobic regions, and then decreased at higher concentrations, either due to a decrease in exposed hydrophobic parts or the fact that ANS did not bind the protein under conditions of high urea (Fig. 4A). The intrinsic Trp fluorescence fitted a sigmoidal curve with a half-maximum of 2.2 ± 0.1 M of urea (Fig. 4B). Interestingly, MSRA and MSRB
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activities using the same urea-treated GST samples followed similar sigmoidal curves, with the half-maxima of 2.3 ± 0.2 and 2.9 ± 0.3 M of urea, respectively (Fig. 4C,D). These results suggest that MSRA and MSRB activities were proportional to the urea concentration and increased with the unfolding state of oxidized GST.

Overall, these results showed that both MSRs were more efficient in the reduction of unfolded proteins due to a better access to MetO in the hydrophobic core of the substrate. Moreover, regardless of MetO exposure, MSRA was intrinsically more efficient in the reduction of MetO in the unfolded oxidized GST.

MSRs protect against oxidative unfolding stress induced by hypochlorite - To test if the observed preferential reduction of unfolded proteins by MSRs is physiologically relevant, we utilized sodium hypochlorite (NaOCl), which leads to concomitant oxidation and unfolding of proteins (17). First, we examined if cells deficient in MSRs were more susceptible to NaOCl stress than WT cells. Growth of MSRA-null yeast cells was dramatically inhibited by treatment with 20 µM NaOCl (Fig. 5A). Although MSRB deficient cells grew similarly to WT cells when treated with NaOCl, deletion of this gene in the context of MSRA deficiency made yeast cells more susceptible to NaOCl stress compared to the MSRA knockout cells. Thus, both MSRA and MSRB contributed to protection against NaOCl stress by reducing MetO formed in cellular proteins upon treatment with NaOCl. However, MSRA was more efficient in protecting yeast cells.

We further took advantage of previously identified Met-rich proteins (22) to determine if these proteins could protect cells through cyclic oxidation of Met and reduction of MetO by MSRs. We overexpressed a structured MRP1 possessing 38 Met and a completely disordered MRP5, which has 31 Met (Supplemental Table S2). Overexpression of MSRA, MRP1 and MRP5 independently protected yeast cells from oxidation. However, the most dramatic effect was observed when MSRA and an MRP were coexpressed (Fig. 5B). We conclude that MRPs alone or in combination with MSRs protect cellular proteins through cyclic oxidation of their Met residues during the oxidative and unfolding stress provoked by hypochlorite.

Finally, we tested the role of MSRA and MRP in the protection against H₂O₂, which induces oxidative stress but does not lead to obvious protein unfolding (17). Although the effects were less pronounced, overexpression of MSRA protected yeast cells from this oxidant; however, expression of MRPs did not offer additional protection (Supplemental Fig. S3). Thus, whereas the Met oxidation/reduction cycle protects yeast during an oxidative and unfolding stress, the effect is less clear in the case of oxidative stress alone. Altogether, these results showed that both MSR types protected yeast against oxidative and unfolding stress and that the protection offered by MSRA was the most efficient.

DISCUSSION

This study describes the first comparative kinetics analyses of MSRA and MSRB using MetO-containing proteins as substrates. MSRA reduced efficiently both free MetO and oxidized proteins, whereas MSRB was specialized for the reduction of oxidized proteins. Further experiments revealed that both MSR types were more efficient in the reduction of unfolded than folded oxidized proteins, and this effect was due to a better access to MetO present in the hydrophobic cores of substrate proteins. However, MSRA was intrinsically more efficient with unfolded proteins irrespective of MetO accessibility. The relevance of this finding was verified in vivo as yeast cells deficient in MSRs were highly sensitive to oxidative unfolding stress provoked by hypochlorite, whereas overexpression of MSRA and Met-rich proteins was highly protective. Our study suggests a new functional category of natural targets for MSRs, the unfolded proteins such as nascent polypeptides, proteins in route for subcellular compartments and proteins unfolded because of oxidative stress (Fig. 6).
Characterization of catalytic mechanisms of MSRA and MSRB was previously carried out using free MetO or its variants, such as N-acetyl-MetO or dabsyl-MetO (23, 32, 33). The only catalytic parameters with a protein substrate were known for bovine MSRA acting on oxidized calmodulin (35). We used three oxidized proteins as model substrates: an unstructured MRP4 that has an exceptionally high Met content (21%), another unstructured protein β-casein (3.1% Met) (36), and GST with 3.8% Met and a well-defined structure (37) (Supplemental Table S2). We showed that both MSRs displayed higher affinity for these proteins than for free MetO and N-acetyl-MetO (Table 1). Whereas MSRA was characterized by catalytic efficiencies similar for free MetO, N-acetyl-MetO and oxidized proteins, MSRB displayed a striking preference for oxidized proteins (Table 1). The catalytic parameters for the reduction of free MetO and N-acetyl-MetO by MSRA and MSRB were in the range of those determined for prokaryotic and plant MSRs (23, 31, 33, 34), consistent with the idea that the preference for MetO reduction in proteins is a common feature of MSRBs, whereas MSRAs act on any MetO-containing substrates. This finding also agrees with in vivo analysis showing that, despite the presence of 3 MSRB isozymes, human cells could not reduce the R-diastereoisomer of free MetO (38).

The use of NADPH-coupled Trx system allowed us to determine the fraction of MetO actually used as a substrate in oxidized proteins. As the overall stoichiometry was 1 mol of NADPH oxidized per mol of substrate reduced, NADPH consumption directly reflected MetO reduction using low substrate concentrations (18, 31). This assay is independent of enzyme concentration, and the use of increasing substrate concentrations allows carrying out linear regression whose slope represents the apparent stoichiometry and, thus, the number of MetO reduced in the protein (Fig. 1). MRP4 possesses 31 Met and the apparent stoichiometries were 4 and 7 mol NADPH ox. mol sub⁻¹, indicating that MSRA and MSRB reduced 13% and 23% of MetO, respectively (Table 1). Further analysis of the Met oxidation state in oxidized MRP4 by mass spectrometry showed that MSRA and MSRB reduced on average 33% and 24%, of MetO, respectively (Supplemental Table S3). In excellent agreement with MSRB, the value found for MSRA was 2.5 fold higher than that calculated by apparent stoichiometry. This could be due to the lower protein coverage observed in the oxidized sample, with 11 Met not covered, whereas only 4 were not covered in the sample repaired by MSRA or MSRB, indicating that Met oxidation affects efficiency of tryptic digestion as suggested previously (39). Similarly, the mass spectrometry analysis showed that MSRA and MSRB reduced 18% and 24% of MetO in GST. These values were higher than those determined by the calculation of apparent stoichiometry, i.e., 1.2 (13%) and 0.8 (9%) MetO reduced by MSRA and MSRB, respectively. This difference was likely due to enrichment of Met-containing peptides, as in almost all cases, the mass spectrometry signal for each peptide was higher for the MSR-reduced samples than for the oxidized proteins (Table 2). The amino acid immediately upstream of Met may influence both the susceptibility of Met to oxidation and its reduction by MSRs, particularly when a Pro flanks the Met (39). Our mass spectrometry analysis corroborated this observation as we found that the Met residues preceded by Pro residues in MRP4 and GST were completely oxidized by hydrogen peroxide but were not substrate for MSRs (Table 2; Supplemental Table S3).

Treatment of the oxidized GST with urea dramatically affected the MSR catalytic parameters, increasing catalytic efficiency (Table 1; Fig. 4). Higher MSRA and MSRB activities were also recorded with the urea-treated oxidized α-amylase than with the corresponding untreated oxidized protein (Fig. 3). Further characterization revealed that protein unfolding increased with the increase in urea concentration, following a sigmoidal curve with the half-maximum at ~2 M urea (Fig. 4); the maximum was reached when the oxidized protein was fully unfolded. In addition, a higher proportion of MetO was reduced in the 4 M urea-treated oxidized protein than in the untreated oxidized protein (Table 2). Moreover,
MSRs preferentially reduce unfolded oxidized proteins

unfolding allowed an almost complete reduction of MetO residues.

Correction of the $K_M$ for the apparent stoichiometry allowed estimation of the catalytic efficiency per each reduced MetO. MSRA was fold more efficient in the reduction of MetO in the unfolded GST than in the folded oxidized protein. Apparently, MSRA is intrinsically more efficient in the reduction of MetO in unfolded proteins regardless of the accessibility to MetO. In the case of MSRB, the catalytic efficiency was higher with the unfolded proteins than with the folded one, but this could be explained by improved accessibility of MetO. These observations are consistent with the finding that MSRA preferentially reduced MetO in a disordered region of oxidized calmodulin and that tryptic digestion was required to open up the protein structure for access to all MetO (35).

An analysis of yeast MSR structures showed that hydrophobicity was particularly pronounced in the active sites, which were also enriched in aromatic residues (Supplemental Fig. 4). These properties could make these enzymes better equipped to interact with unfolded proteins, which expose hydrophobic residues usually confined to the core regions of folded proteins (40, 41). Therefore, the high content of aromatic and aliphatic residues in MSRs could be an important factor promoting the ability of these enzymes to preferentially use unfolded protein substrates through hydrophobic or π-stacking interactions. This property should allow them to exhibit a higher affinity for hydrophobic regions of target proteins, such as protein cores and regions involved in protein-protein interactions through hydrophobic interactions, where Met is particularly enriched (40, 41). Contrary to the oxidation of surface-exposed Met in folded proteins, which may have little effect on protein function (42), Met oxidation in buried regions should dramatically affect folding and function of cellular proteins. Likewise, oxidation of Met in the regions involved in protein interactions is expected to affect protein structure and function as shown for calmodulin (35). MSRs may play a major role in the protein folding process by protecting Met from oxidation in nascent peptides.

Recently, the use of hypochlorite was found to induce Met oxidation in catalase, concomitant with inactivation and unfolding of this protein. This study showed that the reduction of MetO within the hydrophobic core of catalase was required for enzyme refolding by the chaperone GroEL (16). Hypochlorite treatment leads to both oxidative and unfolding stress as demonstrated by HSP33 activation (17). Indeed, whereas the concomitant treatment with $H_2O_2$ and thermal denaturation were required to activate HSP33, sodium hypochlorite alone led to its full activation. This could be due to these two oxidants oxidizing Met residues in different ways. For example, Met oxidation by NaOCl could result in dehydromethionine intermediates that are then converted to MetO (43).

An analysis of yeast cells grown in the presence of NaOCl showed an involvement of MSRs in protection against oxidative unfolding stress (Fig. 5). Indeed, the MSRA-null mutant, and especially MSRA/MSRB-null cells, were more sensitive than WT cells to hypochlorite treatment, and overexpression of MSRA protected cells from this stressor. Similar effects were observed in prokaryotes (15, 16). Interestingly, overexpression of two Met-rich proteins also conferred protection, very likely due to reversible Met oxidation/reduction. This observation shows that the higher MSR activity with unfolded proteins was relevant in vivo during the NaOCl stress that triggered unfolding of proteins. This property may be particularly important during the oxidative battle between neutrophils and pathogens, wherein both opponents induce protein oxidation and use MSRs as a sword and shield strategy (44, 45). Overall, our findings suggest that a major protective function of MSRs in the cell is to rescue and repair oxidized nascent polypeptides and unfolded proteins.
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Acknowledgments - We thank Dr. Pascal Rey (CEA-Cadarache) for the kind gift of dabsyl-MetO.

FOOTNOTES
*This work was supported by NIH grant AG021518. 3The abbreviations used are: ANS, 8-anilinonaphthalene-1-sulfonate; DTT, dithiothreitol; GST, glutathione-S-reductase; HPLC, high performance liquid chromatography; MRP, Methionine-rich protein; MSR, methionine sulfoxide reductase; MSRA, methionine sulfoxide reductase; MSRB, methionine sulfoxide reductase B; NaOCl, sodium hypochlorite; Trx, thioredoxin, TRR1, NADPH-dependent thioredoxin reductase 1; MetO, methionine sulfoxide.4Liang et al., submitted.
FIGURE LEGENDS

FIGURE 1. Apparent stoichiometry of MetO reduction in oxidized MRP4. (A) NADPH consumption was followed with 0.8-6.4 μM oxidized MRP4 as the substrate. The initial reaction mixture contained 250 μM NADPH, 2 μM TRR1, 25 μM Trx1 and 10 μM MSRA. After equilibration, the substrate was added (arrow) and NADPH consumption was followed until it reached the diaphorase level. The final OD was subtracted taking into account the diaphorase activity for each concentration of the substrate (two sided arrows a, b, c and d). (B) Oxidized NADPH plotted as a function of substrate concentration for MSRA (●) and MSRB (○).

FIGURE 2. MSR activities using oxidized and urea-treated oxidized proteins as substrates and characterization of their folding state by fluorimetry. (A) MSRA and MSRB activities were measured using the NAPDH-coupled Trx system with 25 μM oxidized protein (Ox.) and the oxidized protein treated with 4 M urea (Ox + Urea). As a control, for oxidized proteins, the same amount of urea present in the urea-treated oxidized proteins was added to the reaction mixture (less than 50 mM final concentration). *,**, and ***, significantly different with p < 0.05, p < 0.01 and p < 0.0001, respectively (t-test). (B) Intrinsic and ANS fluorescence of 25 μM oxidized or urea-treated oxidized protein was determined under conditions similar to those used for MSR activity assays.

FIGURE 3. Saturation curves of MSRA (A) and MSRB (B) activities using oxidized and urea-treated oxidized GST. MSRA and MSRB activities were measured using the NAPDH-coupled Trx system with 9.4-150 μM oxidized GST or 5-150 μM oxidized GST treated with 4 M urea. As a control, the amount of urea present in the sample of the urea-treated oxidized GST was added in the reaction mixture (less than 50 mM final concentration). The saturation curves obtained with the urea-treated oxidized GST best fit sigmoidal regression as described in Experimental Procedures with the $h = 1.4 \pm 0.3$ and $h = 1.5 \pm 0.1$ for MSRA and MSRB, respectively.

FIGURE 4. Analysis of the folding state of the urea-treated oxidized GST and MSR activities. 10 μM non-oxidized or oxidized GST treated with 0-7 M urea were incubated in 30 mM Tris-HCl pH 8.0 in the presence (A) or absence (B) of 50 μM ANS. Intrinsic and ANS fluorescence was then recorded. Relative MSRA (C) and MSRB (D) activities were recorded using 25 μM oxidized GST treated with 0-7 M urea.

FIGURE 5. Roles of MSRs and MRP in the protection of yeast cells from hypochlorite. (A) Viability of indicated MSR mutants and WT cells under conditions of hypochlorite stress. Yeast strains (5 µl each) were spotted onto agar plates incubated at 30°C. (B) Viability of cells overexpressing MRPs and/or MSRA. Cells overexpressing an indicated MRP and/or MSRA were treated with different concentrations of NaOCl. Cells were plated, incubated at 30 °C and photographed. Cells expressing empty vectors were used as controls.

FIGURE 6. Overview of MSR targets. Based on the effects of MetO oxidation on the structure and activity of proteins, three categories of MSR targets were proposed (4): 1) enzymes in which Met oxidation leads to an increased activity, such as calcium/calmodulin-dependent protein kinase II; 2) proteins that support cyclic Met oxidation and MetO reduction thereby providing antioxidant defense; and 3) native proteins damaged by formation of MetO, such as those involved in neurodegenerative diseases. Based on our data, we propose a fourth type of MSR targets: 4) unfolded proteins and nascent polypeptides whose protein core Met are susceptible to oxidation thereby affecting their folding, structure and function.
**TABLE 1. Kinetic parameters of MSRA and MSRB in the reduction of oxidized proteins.**
Assays were carried out under steady-state conditions following NADPH oxidation at 340 nm. Apparent stoichiometry (mol of NADPH oxidized per mol of substrate) was determined after full reduction of the substrate using sub-stoichiometric substrate concentrations as described in *Experimental procedures*. Data are represented as the means ± S.D.

|                | MSRA                                      | MSRB                                      |
|----------------|-------------------------------------------|-------------------------------------------|
|                | Apparent stoichiometry | $k_{cat}$ | $K_M$ | $k_{cat}/K_M$ | Apparent stoichiometry | $k_{cat}$ | $K_M$ | $k_{cat}/K_M$ |
| **mol NADPH**  | **ox. mol sub$^{-1}$** s$^{-1}$ | **μM** | **M$^{-1}$.s$^{-1}$** (×10$^3$) | **mol NADPH** | **ox. mol sub$^{-1}$** s$^{-1}$ | **μM** | **M$^{-1}$.s$^{-1}$** (×10$^3$) |
| Free MetO      | Measured                                  | 7.71 ± 0.03                               | 1.120 ± 16    | 6.9          | 0.59 ± 0.02                               | 13,014 ± 752   | 0.05                                |
|                | Corrected$^a$                            | 560 ± 8                                   | 13.8          | 15.0         | 0.80 ± 0.06                               | 1,348 ± 224    | 0.6                                    |
| N-acetyl-MetO  | Measured                                  | 13.23 ± 1.83                              | 896 ± 282    | 15.0          | 0.80 ± 0.06                               | 1,348 ± 224    | 0.6                                    |
|                | Corrected$^a$                            | 448 ± 141                                 | 30.0          | 674 ± 112    | 1.2                                    |
| MRP4           | Measured                                  | 4.16 ± 0.16                               | 33 ± 18       | 386.3        | 7.18 ± 0.66                               | 10 ± 3        | 106.7                                  |
|                | Corrected$^b$                            | 137 ± 76                                  | 95.3          | 70 ± 20      | 14.9                                    |
| β-Casein       | Measured                                  | 2.49 ± 0.15                               | 45 ± 13       | 25.1         | 3.19 ± 0.34                               | 54 ± 9        | 14.4                                    |
|                | Corrected$^b$                            | 113 ± 32                                  | 10.1          | 172 ± 28     | 4.5                                    |
| GST            | Measured                                  | 1.21 ± 0.02                               | 356 ± 105    | 2.1          | 0.78 ± 0.02                               | 142 ± 48      | 2.9                                    |
|                | Corrected$^b$                            | 428 ± 127                                 | 1.7           | 111 ± 37     | 3.7                                    |
| Urea-treated GST$^c$ | Measured                              | 3.25 ± 0.05                               | 78 ± 24       | 12.4         | 4.66 ± 0.24                               | 55 ± 11       | 9.5                                    |
|                | Corrected$^b$                            | 254 ± 78                                  | 3.8           | 256 ± 51     | 2.0                                    |

$^a$Considering that only the $S$- and $R$-diastereoisomers serve as substrates for MSRA and MSRB, respectively, and assuming that the other diastereoisomer does not act as inhibitor, the $K_M$ values were divided by 2.

$^b$For comparison, the $K_M$ values were multiplied by the apparent stoichiometry, allowing to remove variation due to the different numbers of MetO reduced in each substrate.

$^c$The regression curves obtained for MSRA and MSRB with the urea-treated oxidized GST fit a sigmoidal described by the equation 1 (*Experimental Procedures*) with the $h$ values equal to 1.4 ± 0.3 and 1.5 ± 0.1 for MSRA and MSRB, respectively.
**TABLE 2. Surface accessibility of Met residues in GST and their differential oxidation as revealed by mass spectrometry analyses.**

Surface accessibility of each Met to the solvent in Å² was determined using protein structure, and the percentage of oxidation of individual Met in the substrate was determined by mass spectrometry as described in *Experimental procedures*. Met *position*, position of the Met in the primary sequence, *Oxidized*, the GST was oxidized with 100 mM H₂O₂ prior to tryptic digestion, *Oxidized + MSRA*, the oxidized GST was reduced by MSRA prior to tryptic digestion, *Oxidized + MSRB*, the oxidized GST was reduced by MSRB prior to tryptic digestion.

| Met position | Secondary structure | Surface accessibility | Oxidized | Oxidized + MSRA | Oxidized + MSRB |
|--------------|---------------------|-----------------------|----------|----------------|----------------|
|              |                     | Å²                    | % oxidation (coverage^a) | % oxidation (coverage^a) | % reduction | % oxidation (coverage^b) | % reduction^b |
| Oxidized GST |                     |                       |          |                |                |                         |               |
| 1            | coil                | ND                    | 93 (25)  | 57 (65)        | 39            | 79 (29)                | 15            |
| 69           | α-helix             | 8.025                 | 67 (47)  | 57 (110)       | 14            | 66 (58)                | 2             |
| 81           | coil                | 2.071                 | 97 (12)  | 100 (22)       | 0^c           | 31 (29)                | 68            |
| 94           | α-helix             | 0                     | 53 (348) | 52 (383)       | 1             | 50 (326)               | 5             |
| 129          | α-helix             | 0                     | 80 (4)   | 0 (11)         | 100           | 50 (4)                 | 38            |
| 132          | α-helix             | 9.977                 | 100 (1)  | ND (0)         | ND            | ND (0)                 | ND            |
| 154          | coil                | 0                     | 16 (50)  | 19 (79)        | 0^c           | 6 (52)                 | 64            |
| 165          | coil                | 0                     | 26 (65)  | 23 (96)        | 10            | 26 (70)                | 0             |
| 168          | coil                | 37.720                | 99 (67)  | 99 (100)       | 0             | 99 (70)                | 1             |
| Average      |                     |                       | 7 ± 12   | 70 ± 33 (69)   | 51 ± 36 (96) | 18 ± 37                | 51 ± 30       |

| 4 M Urea-treated oxidized GST |                     |                       |          |                |                |                         |               |
| 1            | coil                | ND                    | 100 (2)  | 53 (15)        | 47            | 43 (7)                 | 57            |
| 69           | α-helix             | 8.025                 | 60 (30)  | 68 (31)        | 0^c           | 59 (37)                | 1             |
| 81           | coil                | 2.071                 | 100 (5)  | 50 (4)         | 50            | 0 (6)                  | 100           |
| 94           | α-helix             | 0                     | 57 (367) | 54 (393)       | 5             | 53 (374)               | 7             |
| 129          | α-helix             | 0                     | 100 (3)  | 67 (3)         | 33            | 50 (6)                 | 50            |
| 132          | α-helix             | 9.977                 | 100 (1)  | 100 (1)        | 0             | ND (0)                 | ND            |
| 154          | coil                | 0                     | 38 (24)  | 8 (64)         | 79            | 9 (22)                 | 76            |
| 165          | coil                | 0                     | 63 (27)  | 21 (87)        | 67            | 15 (26)                | 76            |
| 168          | coil                | 37.720                | 100 (28) | 99 (94)        | 1             | 88 (26)                | 12            |
| Average      |                     |                       | 7 ± 12   | 80 ± 25 (54)   | 58 ± 31 (77) | 30 ± 33                | 40 ± 30 (56)  |

^aCoverage represents the number of times the peptide in which Met, oxidized or not, was found.

^bThe percentage of reduction was calculated using the formula described in Supplemental Table S3.

^cDue to experimental approximation, calculation gave a slightly negative percentage of reduction when no reduction activity was observed.
MSRs preferentially reduce unfolded oxidized proteins

FIGURE 1

A

B

Time (min)

Oxidized MRP4 (μM)

A340 nm

0.8 mM

1.6 mM

3.2 mM

6.4 mM

No substrate

NaDH oxidized (μM)

MSRA

MSRB

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MSRs preferentially reduce unfolded oxidized proteins

FIGURE 2

A

B

Relative MSR activity

Relative fluorescence

MRP4  β-Casein  α-Amylase  GST

Ox.  Ox + Urea  Ox.  Ox + Urea  Ox.  Ox + Urea  Ox.  Ox + Urea

Intrinsic  ANS

Ox.  Ox + Urea  Ox.  Ox + Urea  Ox.  Ox + Urea  Ox.  Ox + Urea
FIGURE 3

MSRs preferentially reduce unfolded oxidized proteins
MSRs preferentially reduce unfolded oxidized proteins

FIGURE 4

A

B

C

D
MSRs preferentially reduce unfolded oxidized proteins

FIGURE 5

A

Time (min)

0 10 20 30 60

WT

msrAΔ

msrBΔ

msrABΔ

20 μM NaOCl

B

MRP1  MRP1+MSRA MRP5+MSRA WT MRP1  MRP1+MSRA MRP5+MSRA WT

NaOCl (μM)

0 40 50

5 min 10 min

0 40 50

15 min 20 min
FIGURE 6

MSRs preferentially reduce unfolded oxidized proteins
Methionine sulfoxide reductases preferentially reduce unfolded oxidized proteins and protect cells from oxidative protein unfolding
Lionel Tarrago, Alaattin Kaya, Eranthie Weerapana, Stefano M. Marino and Vadim N. Gladyshev

*J. Biol. Chem.* published online May 24, 2012

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