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Redox Sulfur Chemistry of the Copper Chaperone Atox1 is Regulated by the Enzyme Glutaredoxin 1, the Reduction Potential of the Glutathione Couple GSSG/2GSH and the Availability of Cu(I)

Jens Brose, Sharon La Fontaine, Anthony G Wedd and Zhiguang Xiao

Grx1 binds Cu(I) with femtomolar affinity and catalyses reversible redox sulfur chemistry of Atox1 with direction regulated by the cell potential and Cu(I) availability.
INTRODUCTION

Much of the copper in mammalian cells is imported by the high-affinity Cu(I) transporter Ctr1 and is then distributed by metallo-chaperones to intra-cellular destinations. A major pathway involves delivery of copper by Atox1 to the Type P1B ATPase transporters ATP7A/7B in the trans-Golgi network (TGN) for incorporation into enzymes, Atox1 and the six N-terminal metal binding domains (MBDs) of ATP7A/7B all feature the high affinity Cu(I)-binding motif Cys-xx-Cys incorporated into a ferredoxin-like βαββαβ fold. Both in vivo and in vitro experiments suggest that copper delivery by Atox1 involves direct protein-protein interactions mediated by Cu(I) binding and exchange between Cys-xx-Cys motifs of the protein partners. Cu(I)-mediated dimerisation of Atox1 is involved in its function as a transcription factor.

Cellular copper levels are maintained by regulation of Ctr1 levels at the plasma membrane and by reversible trafficking of ATP7A/7B from the TGN to vesicles and/or the plasma membrane for copper export. The latter process is controlled at the stage of copper transfer from Atox1 to ATP7A/7B via kinase phosphorylation at multiple sites on the ATPases.

Glutaredoxins (Grxs) are thiol,disulfide oxidoreductase enzymes proposed to catalyse (de)glutathionylation of protein thiols (P,SH) via cofactors glutathione GSH and its oxidised form GSSG (see recent reviews). However, such proposals have not been documented by in vitro experiments with physiologically relevant protein substrates. Model substrates including mercaptoethanol disulfide have been used as well as...
glutathionylated proteins generated by pure GSSG under non-physiological conditions.\textsuperscript{15}

Many Grxs such as human glutaredoxin 1 (hGrx1) adopt the thioredoxin fold and feature a solvent-exposed Cys-xx-Cys motif that is the enzyme active site (Cys\textsuperscript{3}-xx-Cys\textsuperscript{26} in Figure 1).\textsuperscript{16,17} The equivalent Cys-xx-Cys motifs in Atox1 and the ATP7A/B domains MBD1-6 have evolved to bind Cu\textsuperscript{I} with high affinity ($K_D \approx 10^{-17.5}$ M).\textsuperscript{18} However, it has been proposed that the hGrx1 active site (Cys-Pro-Tyr-Cys) is adapted to prevent binding of metal ions due to the presence of distorting Pro residues both within and near the active site.\textsuperscript{19}

Yeast two-hybrid and mammalian co-immunoprecipitation experiments provide evidence that both the metallo-chaperone Atox1 and the hGrx1 interact with the ATPases ATP7A/7B and that these interactions require copper and the MBDs of ATP7A/7B.\textsuperscript{20,21} Although other interpretations are possible, the observations have been rationalised in terms of direct regulation of the Cu-ATPases by hGrx1 with a consequent influence on copper homeostasis. Over-expression of hGrx1 in neuronal cells perturbs copper metabolism.\textsuperscript{22} However, complementary experiments on hGrx1-null cells are needed for definitive assessment.

GSH (0.1-10 mM in the cytosol)\textsuperscript{23} regulates the redox status of cells via inter-conversion with GSSG.\textsuperscript{12,13} It can also act as a buffer of Cu_{aq}$. The nature of Cu(I)-GSH interactions seems to vary with conditions and the specific complexes involved remain to be defined, together with the corresponding formation constants. However, at its typical cellular concentrations of > 1 mM, GSH appears to buffer free Cu_{aq} concentrations towards femtomolar levels.\textsuperscript{18} Although such affinities are too weak for GSH to compete effectively with copper chaperone proteins in the cytosol, recent evidence indicates that GSH may play a key role in copper uptake by Ctrl, possibly by mediating transfer of Cu(I) from Ctrl to the chaperones.\textsuperscript{24}

A recent report links copper homeostasis (via copper transporters/chaperones) with the GSH/GSSG couple in the cytosol.\textsuperscript{25} Redox enzymes of the thioredoxin family such as hGrx1 and glutathione reductase may participate by connecting the abundant thiol pool of GSH to the Cu(I)-binding thiol pool of proteins. It is also possible that the exposed Cys-xx-Cys motifs in Atox1 and MBDs may be protected from oxidation by glutathionylation (i.e., formation of Cys-S-SG bonds) mediated by hGrx1. Such processes are emerging as mechanisms of redox regulation and signalling, comparable to phosphorylation.\textsuperscript{26,27}

The present work has examined redox sulfur chemistry involving Atox1 and the couple GSSG/2GSH mediated by hGrx1 with the aim of defining a potential molecular role for hGrx1 in copper metabolism. It provides the first demonstration that hGrx1 actually binds Cu(I) with femtomolar affinity and catalyses the redox sulfur chemistry of Atox1 (a physiologically relevant substrate) in a way regulated by both the reduction potential of the couple GSSG/2GSH and the availability of Cu(I). The evidence suggests possible roles for hGrx1 in copper metabolism including redox regulation of copper trafficking and delivery of copper to the redox partners of copper transporters. The yeast homologue of Atox1 (Atox1) was identified originally as an antioxidant (Atx) in yeast cells lacking a functional superoxide dismutase 1 (Sod1).\textsuperscript{28} This work identifies a plausible molecular mechanism for such action that involves both hGrx1 and copper.

### RESULTS AND DISCUSSION

#### Protein generation and quantification

hGrx1 features five Cys residues (at positions 8, 23, 26, 79, 83). Two of them constitute the enzyme active site defined by Cys\textsuperscript{23}-xx-Cys\textsuperscript{26} (Figure 1a). To evaluate the contributions of these residues towards Cu(I)-binding and thiol-disulfide oxidoreductase activity, wild type hGrx1 and four variants (C23S, C26S, C23,26S and C8,79,83S) were generated and isolated. The identity of each protein was confirmed by DNA sequencing of the expression plasmids and electrospray ionization mass spectrometry (ESI-MS) analysis of the isolated proteins (Table S1). Each sample as isolated was converted to its fully reduced apo-form by incubation with excess...
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Consequently, two Cu(I) probes, bicinchoninic anion (Bca) and bathocuproine disulfonate anion (Bcs) (Figure 2), were employed to evaluate the protein domains such as the MBDs of ATP7A/7B. A solution with [Cu] = 12.7 was prepared by generating the anion [Cu(Bca)2]− (34 μM) in KPi buffer (50 mM, pH 7.0) (see caption of Figure 3). Increasing amounts of either wild type hGrx1 (hGrx1-wt) or the triple variant hGrx1-C87,98,99S (hGrx1-tm, in which the three Cys residues outside the active site Cys23-xx-Cys26 are replaced with Ser) were added to samples of this probe solution. This induced decreases in the concentration of [Cu(Bca)2]−, signalling transfer of Cu(I) from the probe complex to the proteins. The data in Figure 3a indicate that, when free Cu\text{aq}^{+} concentration was buffered at [Cu\text{aq}^{+}] = 12.7, hGrx1-wt bound more than one equivalent of Cu(I) while hGrx1-tm bound only one equivalent. When free Cu\text{aq}^{+} concentration was constrained further to [Cu\text{aq}^{+}] < 14, both hGrx1-wt and hGrx1-tm bound one equivalent of Cu(I) only (Figure 3b). Therefore, hGrx1-wt can bind at least two equivalents of Cu(I) with different affinities: \(K_D < 10^{-14}\) M for the highest affinity binding site and \(K_D = 10^{-15}-10^{-14}\) M for lower affinity binding of a second equivalent of Cu(I). On the other hand, the removal of the three Cys residues in hGrx1-tm outside the active site also removed the lower affinity binding.

Variants C23S and C26S (with one Cys residue in the active site Cys23-xx-Cys26 motif in hGrx1-wt replaced) displayed complete loss of the Cu(I) site of highest affinity but retained the weaker affinity (Table 1; Figure S1; supporting information). This weaker Cu(I) binding induced a tendency for the proteins to precipitate, most likely due to inter-protein linkage.

The above experiments demonstrate that both Cys23 and Cys26 are required for the highest affinity for Cu(I) (\(K_D < 10^{-14}\) M). Equivalent experiments with the free Cu\text{aq}^{+} concentration at the more stringent constraint of [Cu\text{aq}^{+}] = ~15 (increased total ligand concentration; see caption of Figure 4) led to an effective competition for Cu(I) between ligands Bca and P = hGrx1-wt or hGrx1-tm (Figure 4, data sets (ii)). This allowed an estimation of their Cu(I) affinities via eqns 1 and 2 (Figure 4):\[\begin{align*}
[CuL_2]^{3-} + P & \rightleftharpoons Cu^-P + 2 L^2- \\
\frac{[P]_{tot}}{[CuL_{tot}]} & = 1 - \frac{[CuL_2]}{[Cu]_{tot}} \\
+ K_{D}\beta_2 \left(\frac{[L]_{tot}}{[CuL_2]} - 2 \right)^2 \left(\frac{[CuL_2]}{1 - \frac{[CuL_2]}{[Cu]_{tot}}}\right)
\end{align*}\]
where the term [CuL₂] is the equilibrium concentration of probe complex [CuL₂]⁻ in eqn 1 and may be determined directly from the solution absorbance under the condition that this complex is the only absorbing species. The other terms in eqn 2 are the known total concentrations of the relevant species. As expected, a dilution of each solution by a factor of two resulted in partial transfer of Cu(I) from [Cu(Bca)]⁻ to the protein in order to maintain the relationship of eqn 2 under the new equilibrium conditions (Figure 4, data sets (iii)).

Curve fitting of each of the two sets of experimental data (ii and iii) for each protein to eqn 2 based on the known β₂ for [Cu(Bca)]⁻ (10⁻⁷.2 M⁻²)⁸ yielded the same Kᵯ = 10⁻¹⁵.⁵ M at pH 7.⁰ for both hGrx1-wt and hGrx1-tm. The individual estimates were indistinguishable within experimental error (Table 1). This affinity falls within the effective buffer range of probe Bcs (Figure 2), as well as that of Bca. Equivalents experiments with Bcs generated similar competition for Cu(I) and provided the same estimate (Kᵯ = 10⁻¹⁵.⁶ M) based on β₂ = 10⁻⁰.⁸ M⁻² for [Cu(Bcs)]⁻ (Figure S2; Table 1). On the other hand, variants C23S and C26S were unable to compete for Cu(I) under the same conditions (Figure 4, data sets (i)). These experiments demonstrated that the two Cys residues in the active site of hGrx1 contribute solely to its femtomolar affinity for Cu(I). The other three Cys residues may contribute to the weaker binding sites.

Interestingly, a conserved cis-Pro residue adjacent to the β₃ strand and another within the Cys-xx-Cys motif occur in the thioredoxin family of enzymes. They have been proposed to play a role in preventing assembly of Fe-S clusters in this site and possibly in preventing binding of other metal ions as well.¹⁹ Two such Pro residues are present in hGrx1 studied in this work (Pro71 and motif Cys-Pro-Tyr-Cys). However, the present work demonstrates that the enzyme is able to bind one equivalent of Cu(I) with femtomolar affinity. The thioredoxin from *E. coli* can do the same.³²

### Table 1. Apparent Cu(I) dissociation constants Kᵯ for the sites of highest affinity

<table>
<thead>
<tr>
<th>Protein</th>
<th>log Kᵯ</th>
<th>Competing L (µM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>hGrx1-wt</td>
<td>-15.5¹</td>
<td>500/250</td>
</tr>
<tr>
<td>hGrx1-tm</td>
<td>-15.5²</td>
<td>400/250</td>
</tr>
<tr>
<td>Atox1</td>
<td>-14.3¹</td>
<td>80</td>
</tr>
<tr>
<td>hGrx1-C23S</td>
<td>-15.6²</td>
<td>400/200</td>
</tr>
<tr>
<td>hGrx1-C26S</td>
<td>-15.6²</td>
<td>200</td>
</tr>
<tr>
<td>Atox1</td>
<td>-14.6³</td>
<td>200</td>
</tr>
</tbody>
</table>

¹ in KPi buffer; ² in Mes buffer; ³ with 7 M urea in the solution; ⁴ less reliable due to slow changes in spectra.
The reduction potential of Atox1 is more negative than that of hGrx1.

Preliminary experiments suggested that the extent of oxidation of the Cys-xx-Cys motifs of both Atox1 and hGrx1 could be controlled by the glutathione disulphide(glutathione redox couple GSSG/GSH that can buffer the effective reduction potential of the solution. This allowed estimation of the protein reduction potentials. The experiments were performed in a series of redox buffers containing varying molar ratios of GSSG/GSH at total concentrations in the range [GSH + 2GSSG] = 1.0-4.0 mM in KPi buffer (50 mM, pH 7.0). Each protein (10 µM) was incubated overnight in the redox buffers under anaerobic conditions to ensure attainment of equilibrium. The systems were quenched by alkylation with excess iodoacetamide, allowing estimation of the ratio of the fully reduced to fully oxidized protein forms (i.e., P-(SH)2P-(SS)) by ESI-MS analysis (see Figure 9 below and Figures S4, S5).15

At a fixed pH, the reduction potentials of the redox buffers and of the target proteins, respectively, may be calculated via the Nernst relationships of eqns 3 and 4:

\[
E = E_{\text{red}}^\circ - \frac{RT}{2F} \ln \left( \frac{[\text{GSH}]}{[\text{GSSG}]} \right) \quad (3)
\]

\[
E = E_{\text{red}}^\circ - \frac{RT}{2F} \ln \left( \frac{F_{\text{red}}}{1 - F_{\text{red}}} \right) \quad (4)
\]

\(E_{\text{red}}^\circ\) is the conditional standard reduction potential of the couple GSSG/2GSH which is pH-dependent and is equal to -240 mV at pH 7.030; \(E_{\text{red}}^\circ\) is the conditional standard reduction potential of the couple P-(SS)P-(SH)2 which is also pH-dependent; \(F_{\text{red}}\) is the fraction of the fully reduced protein to the sum of the fully reduced and fully oxidized protein forms (i.e., \(F_{\text{red}} = [P-(SH)2]/[P-(SH)2 + P-(SS)]\)). The minor form P-(SH)(SSG) (see Figure 9 below and Figure S4) was not included in eqn 4 since it was a redox intermediate in equilibrium with both fully reduced and fully oxidized forms and cancels out in the calculation.37 This redox intermediate reached a maximal fractional level of ~10% at ~50% oxidation-reduction (Figure 6, empty circles).

The fully reduced fraction \(F_{\text{red}}\) for both Atox1 and hGrx1-tm changed sensitively within different reduction potential windows (Figure 6). Curve-fitting of the experimental data to eqn 4 led to the estimates \(E_{\text{red}}^\circ = -188 \text{ mV and } -118 \text{ mV at pH 7.0 for Atox1 and hGrx1-tm, respectively. The reduction potential of Atox1 is 70 mV more negative than that of hGrx1-tm. This correlates with the lower \(pK_a\) of Cys23 in hGrx1 and its lower affinity for Cu(I). These differences are attributed mainly to the positively charged environment surrounding the catalytic residue Cys23 in hGrx1.12,16,34

The reduction potentials at pH 7.0 determined in this work for Atox1 (-188 mV) and hGrx1 (-118 mV) are, respectively, 40 mV and 100 mV more positive than recent literature values of -229 mV and -220 mV.25 A slightly more negative value for hGrx1 (-232 mV) was reported earlier.3 The reduction potential of a thiol group is pH-dependent.36 The two literature estimates and the present work were conducted in the same buffer using the same couple as redox buffer. The differences lie in the more complex approaches used previously to quench the reaction and quantify the protein oxidation states. The method of alkylation and ESI-MS analysis employed here is justified by the following experiments: (i) controls in which a solution of iodoacetamide with extra GSSG was added as quenching reagent provided the same results as those using iodoacetamide alone; this demonstrates that the rate of alkylation was rapid compared the rate of change of the redox equilibrium; (ii) both the alkylation/ESI-MS probe and the
solution probe $[\text{Cu}(\text{Bca})_2]^{1+}$ provided equivalent results in the kinetics experiments of Figure 8c given below (red crosses versus blue dots; see also Figure 9 below). The fact that the reduction potential of hGrx1 is 70 mV more positive than that of Atox1 (Figure 6) is consistent with all the experimental data presented below.

The active site variant hGrx1-C26S is more active in disulfide reduction than the native enzyme

A classic assay based on coupled enzyme reactions was employed (Figure 7a). The first reaction is reduction of mercaptoethanol disulfide (HOC$_4$H$_2$S)$_2$ to mercaptoethanol HOC$_4$H$_2$SH via oxidation of GSH to GSSG with hGrx1 as catalyst. The second reaction reduces product GSSG back to GSH with glutathione reductase (Grase) as catalyst. The oxidized product nicotinamide adenine dinucleotide phosphate (NADP$^+$) of reducing agent NADPH was detected directly by the absorbance change for NADPH at 340 nm ($\varepsilon = 6,319$ M$^{-1}$ cm$^{-1}$). The assay conditions were set to ensure that the overall oxidation rate of NADPH was proportional to hGrx1 concentration (Table 2). Under these conditions (Figure 7), the catalytic turnover rate for NADPH was ~ 280 min$^{-1}$ for hGrx1.

Conversion of the N-terminal Cys23 to Ser to form variant hGrx1-C23S led to ~ 90% loss of activity while substitution of both Cys23 and Cys26 to Ser in hGrx1-C23,26S led to complete loss (Figure 7b(i)). In contrast, substitution of the C-terminal Cys26 to Ser in hGrx1-C26S increased the activity by > 100% (Figure 7b(i, ii); Table 2). Addition of three equivalents of Cu(I) (relative to hGrx1) into the same assay mixture for hGrx1-wt decreased the activity by ~ 70% (Figure 7b(ii, iii); Table 2). The femtomolar affinity of the hGrx1 active site for Cu(I) (relative to hGrx1) into the same assay mixture for hGrx1-wt increased the activity by ~ 39,40.

Table 2 Comparison of catalytic rate of hGrx1 in KPi (pH 7.5)

<table>
<thead>
<tr>
<th>Protein</th>
<th>Turnover rate for NADPH (min$^{-1}$)</th>
<th>Turnover rate for (HOC$_4$H$_2$S)$_2$ (min$^{-1}$)</th>
<th>Relative to WT</th>
</tr>
</thead>
<tbody>
<tr>
<td>hGrx1</td>
<td>562</td>
<td>281</td>
<td>1.0</td>
</tr>
<tr>
<td>C23S</td>
<td>58</td>
<td>20</td>
<td>0.1</td>
</tr>
<tr>
<td>C26S</td>
<td>1280</td>
<td>640</td>
<td>2.3</td>
</tr>
<tr>
<td>C23,26S</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>hGrx1+3Cu(I)</td>
<td>176</td>
<td>88</td>
<td>0.3</td>
</tr>
</tbody>
</table>

It has been suggested that Cu(I) is not a direct substrate. Incubation with GSH is proposed to spontaneously generate the mixed disulfide HOC$_4$H$_2$S-GS that is the real substrate. A ping-pong mechanism has been proposed (Scheme I). The presence of Cys26 close to the Cys23 residue allows formation of an internal disulfide bond that inhibits the enzyme activity. Likewise, high affinity Cu(I) binding to the active site Cys$^{23,xx}$-Cys$^{26}$ motif may also suppress the
nucleophilic attack of the Cys\(^{33}\) thiolate on the mixed disulfide substrate and so inhibit catalytic activity.

The unique catalytic role of Cys\(^{23}\) is attributed primarily to its very low \(pK_a\) (~ 3.5)\(^{33}\) that ensures its existence at physiological pH as a thiolate ready for nucleophilic attack on disulfide bonds. Its solvent-exposed location over a surface cleft may also be important to allow it to act as a GSH binding site to facilitate formation of the mixed Cys\(^{33}\)S-SG (Figure 1b; Scheme I).

### hGrx1 catalyses oxidation of Atox1 by GSSG

One of the primary functions proposed for hGrx1 is to prevent irreversible oxidation of protein thiols by catalysing their reversible (de)glutathionylation employing cofactors GSSG/GSH.\(^{41}\) However, the reported activities of hGrx1 and similar enzymes have been dominated by assays based on model substrates such as (HOC\(_2\)H\(_2\)S)_\(_2\)

A new assay was developed here to evaluate how hGrx1 may mediate the sulfur redox chemistry of protein thiols involved in copper nutrition. It mimics cellular reactions and conditions based on catalytic oxidation of the fully reduced Cu(I)-binding protein Atox1 by GSSG with hGrx1 as catalyst. Its design is shown schematically in the inset of Figure 8. The catalytic oxidation reaction (framed by the green dotted line) was monitored and regulated by a binding/buffering reaction that provided competition for Cu(I) between reduced Atox1 and a chromophoric Cu(I) probe ligand L (L = Bca or Bcs; framed by the red dotted line). The availability of Cu(I) to Atox1 was controlled by ligand L and the oxidation rate was followed in real time by loss of high affinity for Cu(I) up on conversion of Atox1 to oxidised forms (internal disulfide or irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing 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oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidation of protein thiols by catalysing their irreversible oxidati

![Figure 8](Image336x351 to 537x742)

**Figure 8.** Comparison of the oxidation rate of Atox1 (35 µM) by GSSG (400 µM) catalysed by hGrx1-wt (blue trace), hGrx1-C26S (green trace) or no catalyst (red trace) in KPi buffer (20 mM; pH 7.0; NH\(_4\)OH; 1.0 mM). Concentration of hGrx1 = 2.0 µM. \(p[Cu^{2+}]\) was buffered at (a) ≤ 16 by buffer: \([Cu(I)]_{tot} = 35 \mu M, [Bca]_{tot} = 500 \mu M, (b) ≤ 17 M by buffer: \([Cu(I)]_{tot} = 35 \mu M, [Bcs]_{tot} = 140 \mu M, (c) no buffer. The rates in (a,b) were followed in real time by increase in concentration of the Cu(I) probe \([Cu(\text{Bca})]_3\) and the rate in (c) by quenching the oxidation at various reaction time points with the same probe \([Cu(\text{Bca})]_3\) but lower \([Bca]_{tot} = 100 \mu M \) (see Experimental section for details). The red crosses shown in (c) document an alternative quantification by quenching the oxidation via alkylation and ESI-MS analysis (see Figure 9 below). Inset: scheme for the assay that consists of the catalysis (framed in green dots) and the associated detection of Cu\(_{\text{aq}}^{2+}\)-buffering and Cu(I)-transfer (framed in red dots).
enhanced 20-fold to an initial turnover rate of ~2 min\(^{-1}\) (Figures 8b, S3b).

On the other hand, when substrate Atox1 was presented in the \textit{apo} form only (no added Cu(I)), both hGrx1-wt and hGrx1-C26S catalysed oxidation of Atox1 by GSSG much more efficiently (Figures 8c, S3c). hGrx1-C26S was again the most active enzyme (initial rate ~7.7 min\(^{-1}\) vs ~2.3 min\(^{-1}\)). The rates in these experiments were determined by transferring aliquots of reaction mixture at different reaction times into solutions containing Cu(I) probe \([\text{Cu}^{2+}]_{\text{aq}}\) to quench and quantify the oxidation. Alternatively, the oxidation rates could be determined by transferring aliquots of the same reaction mixture into solutions of excess iodoacetamide to quench and quantify by alkylation of the Cys thiols and ESI-MS analysis.35 The outcomes of the two approaches were the same, within the experimental error (blue dots versus red crosses in Figure 8c).

Equivalent experiments with hGrx1 variants as catalysts showed that the catalytic activity of hGrx1-tm was always lower than that of hGrx1-C26 but indistinguishable from that of hGrx1-wt. On the other hand, hGrx1-C23S and hGrx1-C23,26S exhibited no catalytic activity. It must be noted that, under the different conditions tested in Figure 8, free \([\text{Cu}^{2+}]_{\text{aq}}\) was buffered at such low concentrations \((p[\text{Cu}^{2+}] < 16)\) that all forms of hGrx1 could not compete for Cu(I). It can be concluded that the \([\text{Cu}^{2+}]_{\text{aq}}\) buffer itself had little impact on the enzyme activity and the observed catalytic activities may be attributed to the Cu-free forms of the hGrx1 and its variants.

Three conclusions may be drawn (the nature of the overall processes is depicted in Scheme II): (i) Cys23 in hGrx1 plays a central role in the catalytic oxidation of Atox1 by GSSG. The adjacent residue Cys26 may suppress the catalysis via formation of an internal disulfide bond with Cys23. The other non-active site Cys residues (Cys8,79,83) have little impact on the catalysis; (ii) \textit{apo}-Atox1 is thermodynamically vulnerable to oxidation but the process is slow unless catalysed by hGrx1 (even under oxidising conditions: initial concentration of GSSG, 400 \(\mu\)M); (iii) Binding of Cu(I) to Atox1 protects it from oxidation.

The internal disulfide forms of Atox1 and hGrx1 are thermodynamically favoured relative to glutathionylated forms

A number of different oxidation processes promoted by GSSG may lead to loss of high affinity binding of Cu(I) to the dihthiol Cys-xx-Cys motif in Atox1. They include formation of an internal disulfide bond leading to Atox1-(SS) and/or mixed disulfide forms such as Atox1-(SH)(SGG) and Atox1-(SSG).

To address this issue, the reaction product(s) formed during the course of catalytic oxidation of Atox1 by hGrx1-wt were characterised (Figure 8c; blue dots). The reactions were quenched by alkylation of protein thiols, as described above. The products were characterised directly by ESI-MS analysis (Figure 9).35 Two ions are expected for each of the reactant and products as the Atox1 samples were a mixture of molecules with and without an N-terminal methionine residue. Atox1 possesses three Cys residues (Cys12 and Cys15 in the Cys-xx-Cys motif and Cys41, Figure 1c) and the reduced substrate form was detected as the fully alkylated species Atox1-(SA)\(_3\) (Figure 9). The two Cys residues in the Cys-xx-Cys motif were oxidised to an internal disulfide bond: Atox1-(SA)(SS) (>96%) after 15 min with minimal production of glutathionylated forms: Atox1-(SGG)(SS) (< 2%); Atox1-(SA)\(_2\)(SSG) (< 2%).

To assess the oxidation products formed under non-catalytic conditions, Atox1 was incubated overnight in redox buffers containing different ratios of GSSG and GSH. The products were again alkylated and detected by ESI-MS. Under physiologically relevant conditions within the reduction potential window of normal cells \((E' \sim -210 \text{ to } -240 \text{ mV})\), Atox1 remained predominantly in its fully reduced form (Figure 10a). When the buffer was made more oxidising at \(E' \sim -190 \text{ mV}, \) about 50% of the protein appeared as a mixture of several oxidised forms including Atox1-(SA)(SS) (> 90%), Atox1-(SSG)(SS) (< 1%) and Atox1-(SA)\(_2\)(SSG) (< 10%) (Figure 10b). When the GSSG : GSH ratio was increased ~100 fold to the non-physiological value of \(E' > -100 \text{ mV}, \) essentially quantitative conversion to the disulfide form ensued (Figure 10c; ions detected: Atox1-(SA)(SS), > 98%; Atox1-(SSG)(SS), < 1%; Atox1-(SA)\(_2\)(SSG), < 1%). Even under the extreme conditions of a very high concentration of pure GSSG (25 mM), the oxidation forms of the Cys-xx-Cys motif in Atox1 were still dominated by the disulfide bond (Atox1-(SSG)(SS), ~ 75%; Atox1-(SSG)), ~ 23%) while the isolated single Cys41 was trapped largely in glutathionylated form (~ 98%) (Figure 10d).

Equivalent experiments with hGrx1-tm showed that the oxidised forms were dominated by hGrx1-(SS) (> 90%) under all physiologically relevant conditions. The content of the minor component hGrx1-(SH)(SGG) varied with conditions and reached up to ~10% of the total oxidation products after the protein was about 50% oxidised (Figure S4).
The above experiments demonstrated that: (i) only the protein thiols in the Cys-xx-Cys motifs are thermodynamically susceptible to oxidation by GSSG and isolated single Cys residues (such as Cys41 in Atox1) are susceptible only at very high GSSG concentrations (Figure 10). This is consistent with previous observations that single Cys residues in proteins are not easily glutathionylated under physiological conditions;\(^{25,42}\) (ii) the dominating oxidised form of the Cys-xx-Cys motif for P = Atox1 or hGrx1 (and likely for similar proteins) is P-(SS) under all conditions; (iii) under physiologically relevant conditions, the doubly glutathionylated form P-(SSG) was not observed and the singly glutathionylated form P-(SH)(SSG) was detected as a minor product. The content of the latter was minimal (< 2%) under catalytic conditions, but increased slightly under non-catalytic conditions. Apparently, it is the formation of the internal disulfide bond in Atox1 that provides the driving force for the oxidation of the Cys-xx-Cys motif by GSSG.

However, species P-(SH)(SSG) are proposed to appear as intermediates of both substrate Atox1 and catalyst hGrx1 in the overall oxidation (Scheme II). Their concentrations are kept minimal under catalytic conditions but may accumulate slightly under non-catalytic conditions. Consequently, previous conclusions of glutathionylation of ATP7A based on detection by immunoprecipitation may represent a small fraction only of the total ATP7A present.\(^{21}\) 

**hGrx1 can catalyse reduction of Atox1 by GSH but only in the presence of Cu(I)**

Oxidation of the Cys-xx-Cys motif in Atox1 under physiologically relevant conditions results in Atox1-(SS) (Figures 9, 10, S3; Scheme II). Consequently, this internal disulfide form is a potential substrate for reduction by GSH with hGrx1 as catalyst. It was prepared readily by oxidation of reduced Atox1 with 2-3 equivalents of K\(_3\)Fe(CN)\(_6\), followed by separation over a desalting column. The assay was similar to that employed in Figure 8 but operated in the opposite direction (inset, Figure 11). The assay was conducted in a metal buffer controlled by [Cu(I)(Bcs)]\(_3\)^2-/Bcs\(^2-\) (pCu\(^+\) = 15.5). Atox1-(SS) has little affinity for Cu(I) but reduction by GSH generates the reduced form that binds Cu(I) with high affinity (K\(_D\) = 10\(^{-17.4}\) M at pH 7.0).\(^{18}\) This form scavenges Cu(I) from the chromophoric probe [Cu(I)(Bcs)]\(_3\)^2- at a reaction rate much faster than the Atox1-(SS) reduction rate. Consequently, the reduction may be followed in real time via the decrease in probe concentration.

Reduction of Atox1-(SS) (50 µM) by GSH (800 µM) in metal buffer ([Cu(I)\(_{tot}\), 37 µM; Bcs\(_{tot}\), 100 µM] was slow and unaffected by addition of hGrx1-C23S, an inactive hGrx1 variant (Figure 11a). The rate of reaction increased upon addition of active catalysts hGrx1-tm or hGrx1-C26S (1.0 µM; Figure 11b, c). In contrast to the higher catalytic activity of
Reduction of Atox1-SS (50 µM) by GSH (800 µM) in KPi (50 mM, pH 7.0) containing Cu(I) probe [Cu(Bcs)]$^{3+}$ (composition: [Cu]$_{tot}$ = 37 µM; [Bcs]$_{tot}$ = 100 µM; [NH$_4$OH]$_{tot}$ = 100 µM) with catalyst (each 1.0 µM) to initiate the catalytic reduction: (a) no catalyst (no difference with addition of 1.0 µM hGrx1-C23S); (b) hGrx1-C26S; (c) hGrx1-tm; (d) the same as (c) but the catalyst hGrx1-tm was pre-mixed with Atox1$_{apo}$ and GSH at 105% concentrations in (c) for 20 min (dashed trace) and then the Cu(I) probe [Cu(Bcs)]$^{3+}$ was added at 20x concentration to the same final solution conditions in (c). The reduction was followed by monitoring Cu(I) transfer from the probe [Cu(Bcs)]$^{3+}$ to the reduced Atox1 generated in situ as shown in the reaction scheme of inset. The dashed black line shows the [Cu(Bcs)]$^{3+}$ concentration expected when 50 µM Atox1-SS is fully reduced (see Figure S6).

hGrx1-C26S relative to hGrx1-tm for the opposite reaction driven by excess GSSG (Figure 8; Scheme II), their activities are comparable under the reducing conditions controlled by excess GSH (800 µM). The initial Cu$_{eq}$ concentration of the assay solution was calculated to be p[Cu$^{+}$] = 15.5, but is decreased rapidly to a final p[Cu$^{+}$] ~ 16.7 with Atox1 reduction (Figure S6). Neither form of hGrx1 is able to bind Cu(I) effectively under the conditions (Figure S6) and the inhibited form hGrx1-(SS) appears to be activated efficiently by the high concentration of GSH.

In contrast, in a control reaction that contained Atox1-(SS), GSH and hGrx1-tm but no [Cu(Bcs)]$^{3+}$, reduction of Atox1-(SS) was very slow (Figure 11d, dashed line). However, addition of [Cu(Bcs)]$^{3+}$ returned the rate of reaction to that observed previously (Figures 11d (solid line) vs 11c). It can be concluded that binding of Cu(I) to the high affinity site Cys-xx-Cys of reduced Atox1 increased the reduction potential for the protein disulfide/thiol couple considerably and that this provided the thermodynamic gradient needed to drive the reduction. However, Atox1-(SS) was not reduced completely under these conditions due to increasing rates of the reverse processes shown in the inset scheme of Figure 8. The equilibrium concentration of [Cu'(Bcs)]$^{3+}$ predicted for full reduction is indicated by the black dashed line in Figure 11 (see also Figure S6).

Additional experiments varying the relative concentrations of ligand Bcs$^{2-}$ or GSH further demonstrated that the reduction rate and equilibrium concentrations of Atox1-(SS) are regulated by both GSH concentration and the solution p[Cu$^{+}$] value (Figure 12). Apparently, multiple processes and equilibria control the overall reduction process including the differences in reduction potentials between the redox couples Atox1-(SS)/Atox1, hGrx1-(SS)/hGrx1 and GSSG/GSH and competition for Cu(I) between hGrx1, Atox1, probe ligand Bcs$^{2-}$ and GSH.

In summary, the above experiments demonstrate that: (i) GSH provides a shallow thermodynamic gradient only for reduction of Atox1-(SS). Binding of Cu(I) to the reduced form of Atox1 increases the thermodynamic gradient for reduction. (ii) hGrx1-wt and its variants with the catalytic residue Cys23 in place can readily catalyse reduction of Atox1-(SS) by GSH to Cu$^{1-}$Atox1 but not to apo-Atox1 (dithiol form).

The origins for these observations lie in the differences in reduction potentials of the relevant species and the impact of the binding of Cu(I) on such differences. The proposed catalytic cycle is summarised in Scheme III. Catalysis is initiated by nucleophilic attack of the Cys23 thiolate of hGrx1 on the Atox1 disulfide bond to yield an intermediate complex featuring a mixed disulfide bond (step i). Reduction of this bond by a monothiol (step ii) and/or dithiol mechanism (step iii) releases fully reduced Atox1 for high affinity Cu(I) binding. The
fully reduced Atox1 for high affinity Cu(I) binding. The oxidised forms of hGrx1 are regenerated by GSH via mono- or di-thiol mechanisms (steps iv, v) to complete the catalytic cycle.

**Direct observation of the proposed Atox1-S-S-Grx1 intermediate**

Evidence for step (i) of Scheme III is provided by ESI-MS analysis of a reaction mixture containing an equimolar ratio of Atox1-(SS) and fully reduced hGrx1-tm (each 10 µM). While more than 95% of the reactants remained intact, the proposed protein complex was detected unambiguously in the gas phase (1-5%; Figure 13). An accurate molar mass estimated for the complex shown in Scheme III excludes the possibility of the single Cys41 in Atox1-(SS) being involved in complex formation (Table S1).

Notably, step (i) of Scheme III is not favoured under equilibrium conditions but is the potential rate-determining step. The underlying thermodynamic reason is that the reduction potential of the couple hGrx1-(SS)/Grx1-(SH)2 is more positive than that of Atox1-(SS)/Atox1-(SH)2 (see Figure 6) and that the overall catalytic cycle is driven to completion by a combination of the high affinity of the Atox1 protein for Cu(I) (steps (ii, iii)) and regeneration of active hGrx1 enzyme by GSH (steps (iv, v)). However, when Cuṁ⁺ is buffered at relatively high concentrations such as [Cu⁺] = 14.8-14.9 in buffer containing lower GSH concentrations (Figure 12a,b), the catalytic cycle may be inhibited partially by binding of Cu(I) to the catalyst hGrx1. The impact will be different for the different forms of hGrx1: the affinity of hGrx1-C26S for Cu(I) is lower than that of hGrx1-tm (Table 1) and so it is inhibited to a lesser extent (Figure 12a,b). On the other hand, with GSH at higher concentrations such as in Figure 12c, access of Cu(I) to either forms of hGrx1 is restricted and furthermore, the reduction is promoted via the monothiol mechanism (step (ii)) and suppressed via the dithiol mechanism (step (iii)). Consequently, inhibition of hGrx1 activity via formation of an internal disulfide bond as observed at high GSSG/GSH ratios in Scheme II does not operate in the cases of low GSSG/GSH ratios where both hGrx1-wt and hGrx1-C26S catalyse the reduction cycle via monothiol mechanism with comparable activity (Figure 12c).

It is widely held that the oxidation and reduction of protein thiols are associated with catalytic glutathionylation and deglutathionylation. However, the experimental evidence presented here suggests that this is not always the case. Oxidation of the protein thiols in Atox1 to Atox1-(SS) with hGrx1 as a catalyst involves a two-step process of glutathionylation and deglutathionylation on both the substrate and the catalyst (Scheme II). The reverse process seems to involve those steps for the catalyst hGrx1 only (Scheme III). The main reason is that the dominant oxidised form of Atox1 (and likely of all other protein molecules featuring Cys-xx-Cys motifs) features an internal disulfide bond and not a glutathionylated Cys-S-SG group.

**Concluding remarks and perspective**

This study demonstrates an *in vitro* correlation between redox sulfur chemistry and copper binding involving hGrx1, Atox1 and GSH (Schemes II, III). hGrx1 is a thiol-disulfide oxidoreductase enzyme in the cytosol and regulates the redox status of protein thiols in biological cells using GSSG/GSH as cofactors. The present work demonstrates that the affinity
of hGrx1 for Cu(I) at pH 7.0 is 100 times weaker than that of Atox1 whereas its reduction potential is 70 mV more positive. These differences, plus a very low pKₐ of Cys23 of hGrx1, allow rationalisation of the reactions proposed in Schemes II and III. The more positive reduction potential of hGrx1 (Figure 6) is consistent with the enzyme being a robust catalyst for oxidation of apo-Atox1 in Scheme II (Figure 8c) but not for the opposite reaction involving reduction of Atox1(−SS) in Scheme III (Figure 11d, dashed trace). On the other hand, high affinity binding of Cu(I) to Atox1, but not to hGrx1, is expected to invert the order of reduction potentials and to reverse the catalytic preferences of hGrx1 in Schemes II and III (c.f., Figure 8a versus 11d, solid line trace). It is apparent that the direction of sulfur redox catalysis of Atox1 by hGrx1 may be regulated by both the overall reduction potential and by the availability of Cu(I).

The formal oxidant and reductant in Schemes II and III are GSSG and GSH, respectively. The reduction potential of the couple GSSG/2GSH changes sensitively with both ratio and total concentration. For example, at a fixed total concentration of [GSH] + 2[GSSG] = 5.0 mM at pH 7.0, eqn 3 dictates that the reduction potential changes from -200 mV to -100 mV when the ratio [GSSG]/[GSH] changes from 0.1 to 10. On the other hand, when the ratio is fixed at, say, ~ 1% GSSG with total concentration varying between the range 1-10 mM (typical cellular conditions), the reduction potential will vary between -210 mV and -240 mV (Figure 6), as demonstrated recently by in-cell NMR spectroscopy for yeast cells. Under these conditions, the dominant forms of both Atox1 and hGrx1 in the cytosol will be their reduced forms. This prediction was confirmed recently.²³

However, the potential of the GSSG/2GSH couple changes considerably with the biological status of the cell: ~ -240 mV for proliferation, ~ -200 mV for differentiation and ~ -170 mV for apoptosis.²⁶ A reduction potential of E°' = -188 mV for Atox1 suggests that its oxidation level and thus its cellular function will be sensitive to cellular conditions and consequently, in addition to its classic role as a copper chaperone, Atox1 may assume an additional role as a cellular redox regulator. Indeed, its yeast homologue Atx1 was identified originally as an antioxidant (Atx) in yeast cells lacking a functional superoxide dismutase 1 (Sod1).²⁸ Further evidence is accumulating for the antioxidant role of Atox1.⁴⁵ Intriguingly, this activity relies on supply of copper from Ctrl.²⁸,⁴⁵ The present study supports the view that Atox1 may work at the intersection between copper homeostasis and cellular redox balance. Such versatility may be extended to hGrx1 as well: it will allow Atox1 to bind Cu(I) even at cell potentials (> -188 mV) that favour its disulfide form (Scheme III; Figure 6). This aspect may be relevant to the requirement for copper in Atox1’s antioxidant role: a reduced form is required under oxidising conditions.

Reactive oxygen species (ROS) such as HO₂ and H₂O₂ are produced during normal metabolism as by-products of the four-electron reduction of O₂ to H₂O. In healthy cells, these species are scavenged by the antioxidant enzymes Sod1 and catalase. However, their flux increases in cells under oxidative stress and, if the homeostases of copper and iron are perturbed, the Haber-Weiss reaction may occur (catalysed by couples Cu¹¹/Cu¹ and/or Fe³⁻/Fe²⁺ from ‘free’ metal ions):

\[
\text{HO}_2 + \text{H}_2\text{O}_2 \rightarrow \text{O}_2 + \text{H}_2\text{O} + \text{HO}^* \quad (5)
\]

The ROS species (HO₂, H₂O₂, HO*) can attack amino acid sidechains (Met, Cys, Trp), nucleic acid bases and lipids. They will also affect the GSSG/GSH ratio of cells via oxidation of GSH and move the cell potential in a positive direction, favouring conversion of Atox1 to its oxidised form Atox1-(SS). However, hGrx1 will always favour its reduced form hGrx1-(SH)₂ (Figure 6).

In ROS-stressed cells and in the presence of available Cu⁴⁺, hGrx1 can catalyse the production of Cu¹⁻Atox1 (Figure 11; Scheme III). The latter is capable of supplying up to three reducing equivalents (Cu⁹⁰/Cu¹ and Atox1-(SS)/Atox1-(SH)₂) to safely reduce strong oxidants such as HO₂, H₂O₂ and HO* to H₂O (e.g., Scheme IV). This process is related to the redox sulfur chemistry observed in mitochondrial copper transfer. Metallo-chaperone Cu⁻¹-Cox17 (with Cu²⁺ bound in the reduced Cys-xx-Cys site) can transfer both the metal ion and two electrons to the oxidised form of apo-Sco1 (containing a disulfide bond).⁴⁶

Copper imported by Ctrl is delivered to various cellular destinations via a number of copper chaperones in the cytosol. However, many aspects of the processes remain unknown. For example, although Atox1 has been proven to be an important copper delivery vehicle for the Cu-ATPases, this chaperone is not required absolutely for these ATPases to bind copper nor to transfer it across membranes.⁴⁵ In the case of a photosynthetic bacterium, the copper chaperone Atx1 was not essential for copper delivery but ensured safe delivery without damage to other metallo-protein sites.⁴⁷ While this observation suggests that ‘free’ Cu⁴⁺ can reach the cellular targets, other molecules may have evolved to deliver copper to the Cu-ATPases as well.
Recent work in human cell lines has revealed that hGrx1 interacts with copper transporters ATP7A/7B whose MBDS feature Cys-xx-Cys motifs. This could indicate that the availability of the MBDS and copper ions is crucial for the interactions. The present study suggests that these interactions are likely to involve multiple events including reductive binding of Cu(I) to P-(SS) sites in the MBDS that is promoted by hGrx1 and GSH (Scheme III with Atox1 replaced by MBD) and also possible formation of (transient) Cu(I)-bridged complexes between MBD and hGrx1. Consequently, hGrx1 is a feasible candidate to supplement Atox1 in copper delivery.

The mechanism by which copper chaperones acquire copper from Ctr1 is another puzzle. Recent evidence suggests that GSH may take up that key role. Intracellular 'free' Cu\(_{aq}\) is under tight control and is buffered at sub-femtomolar concentrations. Although GSH binds Cu(I) weakly, its cellular concentration is usually at millimolar range. Under such conditions, GSH may buffer free Cu\(_{aq}\) concentrations in the femtomolar range as well (p[Cu\(^+\)] \approx 15.4-16). It may form a partnership with Atox1 and other copper chaperones to ensure tight buffering.

In this context, hGrx1 can also buffer free Cu\(_{aq}\) concentrations in the femtomolar range. Cellular concentrations of hGrx1 have been estimated to be in the µM range in human red blood cells. It regulates the redox status of a broad spectrum of protein thiols and is likely to interact directly with many different protein partners. Therefore, while Cys23 is responsible for catalysis, Cys26 may be recruited for Cu(I) binding and hGrx1 has evolved the properties necessary to function in redox regulation, Cu(I) buffering and Cu(I) transport. It is likely to play a key role in the vital copper transport pathway from the plasma membrane to the trans-Golgi network and beyond. The comparative molecular properties of Atox1 and hGrx1 uncovered by the present work are consistent with a sophisticated mechanism of regulation of redox environments and copper homeostasis in cells.

### Experimental Section

**Materials and general procedures**

Na\(_3\)Bca, Na\(_2\)Bcs, GSH, GSSG were purchased from Sigma-Aldrich. Dtt was purchased from Astral Scientific and Tcep from Fluka. These materials were used as received. Concentrations of Bca and Bcs solutions were standardized routinely as described previously. Stock solutions of Dtt and GSH were prepared in deoxygenated Milli-Q water and stored in an anaerobic glove box. Their concentrations based on qualitative dissolution were confirmed and calibrated with the Ellman assay. Copper solutions were prepared from CuSO\(_4\)·5H\(_2\)O purchased from Sigma-Aldrich.

**Protein Expression and Purification**

Protein Atox1 was expressed and isolated as described previously. The plasmid for expression of hGrx1 was constructed by inserting the hGrx1 gene into pET24d expression vector between NcoI and EcoRI restriction sites and was donated kindly by Dr. Caryn Outten, the University of South Carolina, USA. Plasmids for expression of its protein variants C23S, C26S, C23,26S and C8,79,83S were generated via the overlapping PCR approach using wild type hGrx1 plasmid as template. Each expression plasmid was transformed into E. coli BL21(DE3) CodonPlus cell lines for protein expression. The expression cells were grown in 2YT medium supplemented with antibiotics kanamycin (10 mg/L) and chloroamphetamine (34 mg/L). At an optical density OD\(_{600}\) ≈ 1.0, protein expression was induced with IPTG (0.5 mM) overnight at room temperature. Expressed hGrx1 and its protein variants were purified by gradient elution (0 – 0.3 M NaCl) from a cation-exchange CM-52 column equilibrated in acetate buffer (20 mM; pH 5.4) containing β-mercaptoethanol (5.0 mM). The final step of purification was elution from a Superdex-75 FPLC gel filtration column in KPi buffer (20 mM; pH 7.0; 150 mM NaCl, 0.5 mM Tcep).

The purified hGrx1 proteins were confirmed to be > 95% pure by SDS-PAGE analysis and their identities confirmed by ESI-MS (Table S1). As reported previously, two components (7410.7 and 7270.5 Da) were detected in the Atox1 preparation, corresponding to molecules with and without the first methionine residue, respectively. Their ratio varied from batch to batch. All proteins were isolated in apo-forms with no detectable copper content. Prior to the copper binding studies, apo-proteins were reduced fully by incubation overnight with excess Tcep in a glove box under nitrogen. Excess reductant was removed via a Bio-Gel P-6 DG gel-desalting column (Bio-Rad).

**Concentration Assays**

UV-visible spectra were recorded on a Varian Cary 300 spectrophotometer in dual beam mode with quartz cuvettes of 1.0 cm path length. The concentrations of the apo-protein solutions were estimated by absorbance at 280 nm. The molar absorptivities \(\varepsilon_{280}\) derived from protein primary sequences were
2,980 M⁻¹ cm⁻¹ for both Atox1 and hGrx1 (and its protein variants). Protein concentrations based on this value match those determined from the thiol assay of the fully-reduced apo-forms using Ellman’s reagent DTNB (5,5'-dithiobis-(2-nitrobenzoic acid)). All protein samples used for the metal affinity studies were confirmed to be fully reduced by analysis of the free thiol content. Experiments were carried out on samples prepared anaerobically in KPi buffer (40 mM; pH 7.0; NaCl, 100 mM) in the glove-box.

**Electrospray Ionization Mass Spectrometry**

All experiments were conducted on an Agilent time-of-flight mass spectrometer (TOF-MS) (model 6220, Palo Alto, CA) coupled to an Agilent 1200 LC system. All data were acquired and reference mass corrected via a dual-spray electrospray ionisation (ESI) source. Mass spectra were created by averaging the scans across each peak and background subtracted against the first 10 seconds of the TIC. Acquisition was performed using the Agilent Mass Hunter Acquisition software version B.02.01 (B2116.30) and analysis was performed using Mass Hunter version B.05.00 (B5.0.519.0).

Mass spectrometer conditions: Ionisation mode: Electrospray ionisation; drying gas flow: 7 L/min; nebuliser: 35 psi; drying gas temperature: 325°C; capillary voltage (Vcap): 4000 V; fragmentor: 250 V; skimmer: 65 V; OCT RFV: 250 V; scan range acquired: 100–3200 m/z; internal reference ions: positive ion mode = m/z = 121.050873 & 922.009798.

Chromatographic separation was performed using an Agilent Zorbax Poroshell SB300-C18 2.1 x 12 mm, 5 µm column (Agilent Technologies, Palo Alto, CA) using an acetonitrile gradient (5% (v/v; 0.1% formic acid) to 75% (v/v; 0.1% formic acid)) over 8 min at 0.25 mL/min.

**Quantification of Cu(I) binding**

Two Cu(I) probe ligands, Bca and Bcs were employed for the quantification. They each react with Cu(I) quantitatively to yield well-defined 1:2 complex \([CuL_2]^{2+}\) \((L = Bca or Bcs)\) with characteristic solution spectrum \((ε = 7,900 M^⁻¹ cm⁻¹ at λ_max = 562 nm for L = Bca and ε = 13,000 M^⁻¹ cm⁻¹ at λ_max = 483 nm for L = Bcs)\) and different formation constant \((log β_2 = 17.2 and 19.8 for L = Bca and Bcs, respectively)\). They were used to define Cu(I) binding stoichiometry under non-competitive condition and Cu(I) binding affinity of (expressed as dissociation constant \(K_D\)) under competitive condition according to eqns 1 and 2.

The experiments were conducted in an anaerobic glove-box by reaction of apo proteins with \([CuL_2]^{2+}\) \((L = Bca, Bcs)\) in deoxygenated KPi buffer (40 mM; pH 7.0; 100 mM NaCl) as described previously. Briefly, apo protein was titrated, in various quantities, into a series of \([CuL_2]^{2+}\) solutions of defined molar ratio \(L:Cu^I ≥ 2.5\) (to ensure the presence of the 1:2 complex \([CuL_2]^{2+}\) with negligible contribution from the 1:1 complex \([CuL]^{+}\)). All solutions were diluted to a fixed volume to provide a series of solutions with constant total concentrations of Cu(I) and ligand L but varying concentrations of protein P. Transfer of Cu(I) from \([CuL_2]^{2+}\) to protein P was established by the change in solution absorbance. By selection of probe ligands with different Cu(I) affinities and/or by systematic variation of their concentrations, conditions were searched and set that favored either non-competitive or competitive reaction. The metal binding stoichiometry was derived from the noncompetitive reaction whereas the dissociation constant \(K_D\) was estimated from the competitive reaction via eqns 1, 2.

The \(K_D\) values for Atox1 and hGrx1-am were also determined and compared at different pH within the range 5-7 in buffers Na-Mes (pH 5.0, 5.5, 6.0) and KPi (6.0, 6.5, 7.0) using Cu(I)-Bca as probe and under denaturated condition in KPi buffer (pH 7.0) containing urea (7 M) using Cu(I)-Bcs as probe.

**Catalytic reduction of model substrate mercaptoethanol disulfide by GSH**

The isolated hGrx1 and its protein variants were evaluated for their catalytic functions for several reactions under various conditions. The first is a classic assay based on two coupled enzymatic reactions as shown in Figure 7a. The assay was conducted at room temperature in KPi buffer (100 mM, pH 7.5) with following solution compositions: \((HOC_2H_5S)_2\) (2.5 mM), NADPH (0.25 mM), Grase (1.2 units/mL), GSH (0.5 mM) and hGrx1 at various concentrations (0-25 nM) or different protein variants of hGrx1 at a same concentration of 12.5 nM. The impact of Cu(I) on the reaction was tested by control experiments in the presence and absence of added Cu (3.0 eq relative to hGrx1). Nonenzymatic background reaction(s) without hGrx1 were subtracted from each assay under the otherwise identical conditions.

**Catalytic oxidation of protein thiols by GSSG**

The second assay was based on catalytic oxidation of Cu(I)-binding protein Atox1 with GSSG as an oxidant (see Scheme in Figure 8). The oxidation was followed by monitoring a loss of the high affinity Cu(I) binding affinity of the target protein Atox1 upon its oxidation. The probes used were chromophoric Cu(I) complexes \([Cu(Bca)_3]^{3+}\) and \([Cu(Bcs)_3]^{3+}\) which were prepared in situ from reaction of Cu(II) and excess ligand Bca or Bcs in reduced Mops buffer (50 mM, pH 7.0, NH₄OH, 1.0 mM). In the assays given in Figure 8a,b, the probe complex was included in the pre-assay mixture to make up an assay solution which contained typically Atox1 (35 µM), hGrx1 enzyme (2.0 µM), Cu(I) probe \([CuL_2]^{2+}\) (35 µM) with excess free \(L^2\)-ligand at various concentrations and GSSG (400 µM). The catalysis was started by last addition of the oxidant GSSG and monitored, in real time, by change in solution absorbance for \([CuL_2]^{2+}\) with background control without the enzyme hGrx1 (Figures 8a,b; S3a,b). In the assays given in Figure 8, the probe complex used was the weaker affinity probe \([Cu(Bca)_2]^{2+}\) with \([Bca]_{tot}/[Cu]_{tot} < 3\), but was not added to the catalytic reaction until quantification. Such probe contained relative high concentration free \(Cu_{apo}^{2+}\) and was able to saturate the high affinity Cu(I) site in Atox1 rapidly and consequently to quench and quantify the oxidation quantitatively upon its addition. Therefore, these later procedures separated the catalysis and...
detection into two sequential processes. In practice, the catalysis was started by mixing the fully reduced Atox1 (35 µM) and GSSG (400 µM) in the presence of catalyst hGrx1 (2.0 µM) in KPi (20 mM, pH 7.0) and a small fraction of the assay mixture was withdrawn at various reaction time points and added to an aliquot of probe solution [CuI(Bca)2]2+ with Cu : Bca ~ 1:3. The absorbance at 560 nm for [CuI(Bca)2]2+ was recorded for each solution and was found to increase with reaction time due to the Atox1 oxidation (Figures 8c; S3c). The fraction of the Atox1 oxidation was calculated by dividing the absorbance change observed by the total change expected when the Atox1 was fully oxidised.

An alternative approach of detection is by alkylation and ESI-MS analysis of the protein oxidation products. During the course of the reaction, a small fraction (~ 20 µL) of the above assay mixture was removed at various reaction time points and added to an alkylation solution of iodoacetamide (~ 1 µL 100 mM; > 50 folds excess). The oxidation levels were quantified by ESI-MS and were found to be consistent with the assay based on Cu(I)-Bca probe (Figure 8c). This approach also allowed a direct identification of the oxidised protein forms of Atox1 and hGrx1 (Figures 9, 10, S3).

Catalytic reduction of protein disulphide bond by GSH

The third assay was based on reduction of a disulphide bond in Atox1 (i.e., Atox1-SS) by GSH with hGrx1 as catalyst. It was an assay related to the second one but in the reversed direction (c.f., Schemes in Figures 8, 11). Atox1-SS was prepared under anaerobic condition by incubation of Atox1 (10 - 20 mg/mL) in KPi (20 mM, pH 7.0) with 2-3 equivalents of K2[Fe(CN)6] for ~ 1 h, followed by separation over a desalting column. The catalytic reaction was conducted under anaerobic condition in a mixing cuvette with Atox1-SS (100 µM) being loaded in one half-cell and the rest reaction components (GSH, hGrx1, probe Cu(I)-L complex at specified concentrations) in the other half-cell. After recording the solution absorbance before mixing at 483 nm for L = Bcs or 562 nm for L = Bca, the reaction was started by a rapid mixing and followed by change in the absorbance (note: the initial concentration of each reaction component was halved after the mixing). For control reactions without Cu(I)-L probe, the concentration of each component was increased by 5% and after certain reaction time (~ 20 min), a 20-time concentrated Cu(I)-L probe solution was added into the reaction mixture to estimate and follow the reactions before and after the Cu(I) probe addition (Figure 11d).

Determination of reduction potentials for Atox1 and hGrx1

They were determined by equilibration overnight of each protein (~ 10 µM) in a series of redox buffers of GSSG/2GSH at mM total concentration under anaerobic condition in Mops buffer (50 mM, pH 7.0). The reduction potential of each buffer was controlled by the ratio of GSSG/2GSH and calculated via eqn 3 with $E_{GSH}^{0} = -240$ mV at pH 7.0. The reaction was quenched by alkylation of free thiols with excess iodoacetamide (~ 10 folds excess). After incubation for > 1 h, the oxidation level of each protein was estimated by ESI-MS analysis. Control experiments with extra GSSG being included in the iodoacetamide solution detected essentially the same level of oxidation for either protein, confirming that the alkylation procedure was fast enough to reflect the original redox equilibrium.

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Notes and references

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e Electronic supplementary information (ESI) available: Experiments for determination of Cu(I) KD for variants hGrx1,C23S and -C26S; Figures S1-S6; Table S1. See DOI: 10.1039/b000000x/


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35 Control ESI-MS experiments with samples prepared by mixing various molar ratios of pure fully reduced and fully oxidized Atox1 or hGrx1 after pre-alkylation demonstrate that the relative intensities detected in the gas phase match their compositions in the solutions (see Figure S5).


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Redox Sulfur Chemistry of the Copper Chaperone Atox1 as Catalyzed by the Enzyme Glutaredoxin 1 is Regulated by Both the Reduction Potential of the Glutathione Couple GSSG/2GSH and the Availability of Cu(I)

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Supporting Information
Determination of $K_D$ for variants hGrx1-C23S and -C26S

In a buffer containing $[\text{Cu}^I(Bca)_2]^3-$ with $p[\text{Cu}^+] \sim 13$, both variants hGrx1-C23S and hGrx1-C26S removed Cu(I) from the probe complex but the protein products precipitated. In the same metal buffer with the free $\text{Cu}^{\text{aq}}^+$ buffered at a lower concentration ($p[\text{Cu}^+] \sim 14$), both variants competed for Cu(I) effectively and precipitation was slow. This allowed an estimation of their Cu(I) affinities via eqns 1 and 2 (Figure S1):\(^1\)

\[
[\text{Cu}^I\text{L}_2]^3- + P \rightleftharpoons \text{Cu}^I\text{P} + 2 \text{L}^2- \quad (L = \text{Bca}) \quad (1)
\]

\[
\frac{[P]_{\text{total}}}{[\text{Cu}]_{\text{total}}} = K_D \beta_2 \left( \frac{[\text{L}]_{\text{total}}}{[\text{Cu}^I\text{L}_2]} - 2 \right) \left( \frac{[\text{Cu}^I\text{L}_2]}{[\text{Cu}]_{\text{total}}} \right)^2 \left( 1 - \frac{[\text{Cu}^I\text{L}_2]}{[\text{Cu}]_{\text{total}}} \right) + 1 - \frac{[\text{Cu}^I\text{L}_2]}{[\text{Cu}]_{\text{total}}} \quad (2)
\]

where the term $[\text{Cu}^I\text{L}_2]$ is the equilibrium concentration of probe complex $[\text{Cu}^I\text{L}_2]^3-$ in eqn 1 and may be determined directly from the solution absorbance at equilibrium under the condition that this complex is the only absorbing species. The other terms in eqn 2 are the known total concentrations of the relevant species. Curve fitting of the experimental data to eqn 2 based on the known $\beta_2$ for $[\text{Cu}^I(Bca)_2]^3-$ ($10^{17.2} \text{ M}^{-2})^2$ estimated $K_D = 10^{-13.5} \text{ M}$ and $10^{-13.3} \text{ M}$ for hGrx1-C23S and hGrx1-C26S, respectively (Figure S1). However, such weaker Cu(I) binding caused protein precipitation readily, likely due to considerable conformation change upon Cu(I) binding.

References


Figure S1. Determination of the Cu(I) affinities for hGrx1-C23S (empty circle) and hGrx1-C26S (solid circle). The probe complex anion $[\text{Cu}^I\text{Bca}_2]^3$ was prepared with following compositions: $[\text{Cu}]_{\text{tot}} = 33 \ \mu\text{M}; [\text{Bca}]_{\text{tot}} = 200 \ \mu\text{M}$ and $[\text{NH}_2\text{OH}]_{\text{tot}} = 1.0 \ \text{mM}$ in KPi buffer (50 mM, pH 7.0). The solid traces are the fitting curves of the experimental data points to eqn (2) which generate Cu(I) $\log K_D = -13.5$ and -13.3 for hGrx1-C23S and hGrx1-C26S, respectively. Shown in empty triangles are the experimental data points with hGrex1-wt under the same conditions.
Figure S2. Determination of Cu(I) binding affinity in KPi buffer (25 mM, pH 7.0, 100 mM NaCl). Decrease in $[\text{Cu}^{(Bcs)}_2]^3$ concentration (total composition: $[\text{Cu}]_{\text{tot}} = 31.4$ µM; $[\text{Bcs}]_{\text{tot}} = 80$ µM; $[\text{NH}_2\text{OH}]_{\text{tot}} = 1.0$ mM) with increasing concentrations of: (a) hGrx1-wt; (b) hGrx1-tm. The experimental data points shown in empty circles in (ii) were obtained from a 1:1 dilution of each solution in (i). The solid traces are the fitting curves of the experimental data points to eqn (2) and these curve-fittings derive a consistent Cu(I) $\log K_D = -15.6 \pm 0.1$ for both hGrx1-wt and hGrx1-tm proteins. The dash trace in (ii) is the simple 1:1 dilution curve of data set (i).
Figure S3. Comparison of oxidation rate of Atox1 (35 µM) by GSSG (400 µM) with catalyst (each 2.0 µM) of hGrx1-wt (shown in blue trace), hGrx1-C26S (shown in green trace) or no catalyst (shown in red trace) in KPi buffer (20 mM; pH 7.0; NH₄OH, 1.0 mM) with free [Cu<sup>aq</sup>⁺] buffered to: (a) ≥ 10⁻¹⁶ M by Cu<sup>aq</sup>⁺-buffer of [Cu(I)]<sub>tot</sub> = 35 µM and [Bca]<sub>tot</sub> = 500 µM; (b) ≥ 10⁻¹⁷ M by Cu<sup>aq</sup>⁺-buffer of [Cu(I)]<sub>tot</sub> = 35 µM and [Bcs]<sub>tot</sub> = 140 µM; (c) no Cu<sup>aq</sup>⁺-buffer. The rates in (a,b) were followed in real time by increase in concentration of the Cu(I) probe [Cu<sup>I</sup>(Bca)<sub>2</sub>]<sup>3⁺</sup> whereas the rate in (c) by quenching the oxidation at various reaction time points with the same probe [Cu<sup>I</sup>(Bca)<sub>2</sub>]<sup>3⁺</sup> but less [Bca]<sub>tot</sub> = 100 µM (see Experimental section for details). Inset: scheme for the assay that consists of the catalysis (framed in green dots) and the associated Cu<sup>aq</sup>⁺-buffering and Cu(I)-transferring detection (framed in red dots).
Figure S4. ESI-MS analysis of the oxidized forms of hGrx1 (10 µM) upon oxidation by GSSG in the following redox buffer (potential $E^*$): (a) GSSG (0.06 mM)/GSH (3.88 mM) (-222 mV); (b) GSSG (0.68 mM)/GSH (0.64 mM) (-145 mV); (c) GSSG (0.39 mM)/GSH (0.03 mM) (-69 mV). Each reaction mixture was incubated overnight under anaerobic condition in KPi (50 mM, pH 7.0), followed by alkylation with iodoacetamide (~10 mM) and ESI-MS analysis.
Figure S5. Control experiments for comparison of ESI-MS peak intensities from a solution mixture containing equal molar concentrations of reduced and oxidised forms (both pre-alkylated and quantified separately before mixing) of either Atox1 (a) or hGrx1-tm (b).
Figure S6. Solution spectra in KPi buffer (50 mM, pH 7.0; 100 µM NH₂OH) with following further composition:

(a) \([\text{Cu}^+(\text{Bcs})_2]^{3-}\) generated from \([\text{Cu}]_{\text{tot}} = 37 \mu\text{M}\) and \([\text{Bcs}] = 100 \mu\text{M}\);

(b) as (a) but with extra GSH (800 µM) (no change with addition of extra 50 µM hGrx1-D26S);

(c) as (b) but with extra hGrx1-tm (50 µM)

(d) as (b) but with extra fully reduced Atox1 (50 µM).

The red dashed line indicated the solution absorbance when Atox1-(SS) (50 µM) reached a reduction equilibrium in solution (b).
Table S1. ESI-MS data of protein species $^a$

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$^a$ -SA: acetamidated cysteine thiol; -SSG: glutathionylated cysteine thiol; ND: not detected.