Unglycosylated recombinant human glutathione peroxidase 3 mutant from *Escherichia coli* is active as a monomer

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Glutathione peroxidase 3 (GPx3) is a glycosylated member of GPx family and can catalyze the reaction of different types of peroxides with GSH to form their corresponding alcohols in vitro. The active center of GPx3 is selenocysteine (Sec), which is incorporated into proteins by a specific mechanism. In this study, we prepared a recombinant human GPx3 (rhGPx3) mutant with all Cys changed to Ser from a Cys auxotrophic strain of *E. coli*, BL21(DE3)cys. Although lacking post-translational modification, rhGPx3 mutant still retained the ability to reduce H$_2$O$_2$ and PLPC-OOH. Study on the quaternary structure suggested that rhGPx3 mutant existed as a monomer in solution, which is different from native tetrameric GPx3. Loss of the catalytic activity was considered to be attributed to both the absence of glycosylation and the failure of the tetramer. Further analysis was performed to compare the structures of rhGPx3 and GPx4 mutant, which were quite similar except for oligomerization loop. The differences of amino acid composition and electrostatic potentials on the oligomerization loop may affect the binding of large substrates to rhGPx3 mutant. This research provides an important foundation for biosynthesis of functionally selenium-containing GPx3 mutant in *E. coli*.

Glutathione peroxidases (GPxs) are a family of enzymes that catalyze the reduction of hydroperoxides to their corresponding alcohols. Thus far eight isoforms of GPx enzymes (GPx1–GPx8) have been found in mammals, of which five are selenoenzymes (GPx1–GPx4 and GPx6). GPx3, also known as plasma glutathione peroxidase or extracellular glutathione peroxidase, is the only known glycosylated extracellular isoform, which makes it unique among the selenium-containing members of GPx family. The human GPx3 gene is located at region q32 of chromosome 5 and divided into five exons spanning about 10 kb, the first of which encodes the signal peptide for secretion. The kidney is considered to be the main source for the GPx3 in plasma, which is primarily synthesized in the kidney proximal tubular cells. GPx3 mRNA is also detected in other tissues, such as liver, heart, breast, lung, skeletal muscle, pancreas, brain, mature absorptive epithelial cells and thyroid, from where GPx3 is secreted into the surrounding extracellular environment. Although GPx3 can reduce different types of peroxides using GSH as the reducing agent in vitro, the function of this enzyme is still a mystery due to the low concentration of GSH in plasma.

GPx3 exists as a tetramer in its native state with a subunit molecular weight of approximately 23 kDa. The crystal structure of human GPx3 revealed that each subunit contained a selenocysteine (Sec) residue and the active Sec-53 was located in a pocket on the protein surface with Gln-87, Trp-161 and Asn-162 nearby, which formed a catalytic tetrad highly conserved in the selenium-dependent GPx family. Sec, the 21st amino acid, is encoded by UGA, which is normally regarded as a stop signal. The decoding of UGA as Sec depends on a cis-acting element, known as the selenocysteine insertion sequence, and several trans-acting factors. The mechanism of Sec incorporation is quite different between prokaryotes and eukaryotes, thus very little work has been done in the area of preparation of recombinant mammalian GPx in *E. coli* for the last decades. But recently our group have prepared a GPx1 and GPx4 mutant with all Cys converted to Ser from an *E. coli* BL21(DE3)cys, which showed significant activity comparable to the native enzymes.

The aim of this paper is to explore the possibility of preparing an active selenium-containing GPx3 mutant from *E. coli*. After obtaining this GPx3 mutant, we discussed the effect of glycosylation on enzymatic activity.
Furthermore, the reason for the difference of quaternary structure between native GPx3 and its mutant was also explored.

Results

Preparation of rhGPx3 mutant. The hGPx3 mutant gene was cloned into expression vector pCold I for preparation of selenium-containing rhGPx3 mutant in E. coli BL21(DE3)cyS using a single protein production system (Takara). The expression plasmid pCGPx3M in fusion with translation enhancing element sequence for improving the expression level, the hexahistidine tag for purification and factor Xa cleavage site was constructed successfully. Sequencing analysis confirmed that the Sec and Cys codon(s) of hGPx3 gene was changed to Cys and Ser codon(s), respectively, and no undesired mutation was introduced during the cloning process. SDS-PAGE analysis of rhGPx3 mutant linked with excess amino acid sequence at N-termini showed a single band with a molecular mass of about 25.6 kDa. After being treated with factor Xa, the recombinant protein showed a mass decrease because of the removal of the excess amino acid sequence. Further analysis by Western blot showed that rhGPx3 mutant with excess amino acid sequence could be recognized by the monoclonal antibody against the hexahistidine tag, but not in the case of the protein treated with factor Xa (Figure 1a and 1b). The results indicated that rhGPx3 mutant was successfully expressed and purified from E.coli BL21(DE3)cyS. The yield of the recombinant selenoprotein was approximately 2.6 mg/L of culture. The concentration of selenium was found to be about 2.6 μg per mg protein, indicating that substitution ratio of Sec for Cys residues was approximately 84%.

Analysis of the quaternary structure of rhGPx3 mutant. The purified rhGPx3 mutant was treated with loading buffer with or without β-mercaptoethanol and boiling, and the resulting product was separated by SDS-PAGE. We used a recombinant hGPx1 mutant prepared from E. coli BL21(DE3)cyS as a control, which had been determined to exist, at least partially, as a tetramer. The tetrameric and monomeric forms of rhGPx1 mutant could be detected under non-reducing condition (Figure 1c). Unlike rhGPx1 mutant, the rhGPx3 mutant was found to migrate as a single band under both reducing and non-reducing conditions. The results revealed that the rhGPx3 mutant existed as a monomer, which was different from native human GPx3.

Assay of enzyme activity. The rhGPx3 mutant showed activity of 25.0 U/mg and 8.5 U/mg using H2O2 and 1-palmitoyl-2-(13-hydroperoxy-cis-9, trans-11-octadecadienoyl)- L-3-phosphatidylcholine (PLPC-OOH) as oxidizing substrate, respectively (Figure 2). We also determined the activity of the rhGPx3 mutant with the excess amino acid sequence. Equal amounts of proteins were added for spectrophotometric activity assays as described in Methods. The results revealed that there was no significant difference in the activity between the rhGPx3 mutant with or without the excess amino acid sequence. The results suggested that the additional amino acids had little effect on the enzyme activity.

Determination of optimum temperature and pH. The rhGPx3 mutant showed the highest activity at 45°C and pH 8.9 (Figure 3), which was similar to that of native GPx1 (42°C and pH 8.8)17. Enzymatic activity decreased rapidly when temperature was raised above 50°C and the mutant was almost completely inactivated at 55°C. The enzyme was relatively stable over a pH range of 7.4–8.9, but it lost 93% of the original activity if the pH was raised to 9.8.

Molecular modeling. The profile-3D score of rhGPx3 mutant was determined to be 72.2, compared with the expected high score of 86.6 and the expected low score of 39.0. Ramachandran plot analysis showed 88.7% of residues in the core region, 11.3% in the allowed region, and no residues in the disallowed region. These results suggested that the conformation of rhGPx3 mutant was reliable. Figure 4a showed that rhGPx3 mutant was superimposed on U46C GPx4 mutant (PDB entry 20B1) with a root mean square deviation of 2.87 Å, for 156 C-Alpha atoms.

Discussion

GPx, one of the most efficient antioxidants, can catalyze the reduction of hydroperoxides and protect cells against oxidative stress. Although GPx is a potential drug target, its therapeutic usage is restricted because of the limited sources. As the catalytic active site of selenium-containing GPx, Sec is irreplaceable with respect to activity. However, it was difficult to prepare mammalian selenium-containing GPx from E. coli due to the unique mechanism of Sec incorporation into protein. Recently, our group found that substitution of Ser for Cys would not cause dramatic loss of GPx activity compared to changing those Cys to Sec15,16. By using this method, we produced a GPx3 mutant, which showed significant activity in reducing both H2O2 and PLPC-OOH. Compared with native human GPx3, less than 10% of the activity was retained when the rhGPx3
mutant catalyzed the reaction of H₂O₂ to form the corresponding H₂O, while 15.6% of the activity was retained using PLPC-OOH as the oxidizing substrate (Figure 2). The number was the lowest compared with the GPx1 mutant (28%) and the GPx4 mutant (40%) produced in our previous work 15,16. GPx3 has been reported to be a glycoprotein12, but rhGPx3 mutant produced in E. coli lacked such post-translational glycosylation because of the difference between prokaryotic and eukaryotic cells. Considering the fact that the activity of recombinant GPx3 from mammalian cells was similar to that of partially purified human plasma GPx3 18, the significant drop in activity should be related to the absence of the post-translational modification. In addition, the substitution ratio of Sec for Cys residues using E. coli has been proved to be 84% in this study, which would also lower the activity of the mutant.

The tetrameric GPx3 was constituted by two asymmetric dimers13, but the results above showed that rhGPx3 mutant did not even form such dimer. Unlike rhGPx3 mutant, the monomeric form of the GPx1 mutant seemed to undergo “self-assembly” to form tetramer, at least partially, although the E. coli cells could not provide the same conditions as mammalian cells. Taking into account that rhGPx3 mutant was prepared using the same method, the failure to form tetramer should be mainly due to the structure factor. Aside from lacking post-translational glycosylation, rhGPx3 mutant have a different amino acid composition compared to native GPx3. In E. coli BL21(DE3)cys, the selenoprotein was produced via tRNA Cys misloading, suggesting that all Cys in the protein would be inevitably replaced by Sec. And previous studies in our laboratory have shown that conversion of Cys residues into Sec residues in GPx1 and GPx4 could result in dramatic loss of the catalytic activity15,16. To avoid the adverse effects caused by the introduction of multi-Sec residues, all Cys in GPx3 were changed to Ser in this study. However, analysis on native GPx3 showed a different mobility of reduced and nonreduced GPx3 monomer on SDS-PAGE, indicating the possible presence of disulfide bridges in native enzyme 19. The results of the crystal structure of human GPx3 suggested the disulfide bridge was probably formed by Cys-12 and Cys-136 13. Therefore, it can not be excluded that the existence of the disulfide bridge may contribute to the formation of tetramer, while no such disulfide bridge is present in rhGPx3 mutant. The absence of the disulfide bridge could cause structural change, which is another reason for the failure of formation of tetramer.

It has been reported that native GPx3 has a similar structure to that of the bovine GPx111. Study on bovine GPx1 revealed that only two GSH molecules could bind to each tetramer, indicating that the GSH binding site were formed by residues at least from two subunits20. Therefore, it is tempting to speculate that a similar situation might occur in the case of native GPx3, which would show the highest activity in tetrameric form. In comparison, rhGPx3 mutant seems more likely to catalyze the reaction just like GPx4, which is the only monomeric member among the GPx family, due to the fact that the mutant exists as a monomer in its native state. Actually the structures of U46C GPx4 mutant and rhGPx3 mutant are quite similar (Figure 4a) although their sequence identity is only 34%. The main difference between the two structures is the region opposite the active

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**Figure 3 | Effects of temperature and pH on activity of rhGPx3 mutant.**

(a) Plot of GPx activity versus temperature. The activities were measured at the temperature range of 15–55 °C at pH 7.4. The enzyme activity was assumed as 100% at 45 °C. (b) Plot of GPx activity versus pH. The activities were measured in the pH range of 4.7–9.8 at 37 °C. The enzyme activity was assumed as 100% at pH 8.9. The concentration of GSH and H₂O₂ used for assay was 1 mM and 0.5 mM, respectively. The scale of relative activity (%) indicates the percentage of the activity relative to the highest activity (100%) within each experiment.

**Figure 4 | Comparison between rhGPx3 mutant and U46C GPx4 mutant (PDB entry 2OBI).** (a) The superposition of rhGPx3 mutant (orange) and U46C GPx4 mutant (green). The catalytic tetrad consisting of Sec(Cys), Gln, Trp and Asn is labeled. (b) The electrostatic surface potential of rhGPx3 mutant. The red denotes negative electrostatic potential and the blue denotes positive electrostatic potential. Potential isocontours are shown at + 3 kT/e (blue) and −3 kT/e (red). The circles in panels B and C indicate oligomerization loop.
site, which is termed as oligomerization loop. The type of oligomerization loop in rhGPx3 mutant is mainly lined by hydrophilic residues (Thr-165, Ser-166, Asp-167, Arg-168, Trp-171 and Glu-172) and the surface of oligomerization loop is negatively charged (Figure 4b). The oligomerization loop of rhGPx3 protrudes from an area near the catalytic tetrad. In contrast, this area is a positively charged flat surface in GPx4 (Figure 4c) and has been considered to be involved in the efficient binding of complex lipid molecules at the active site. The type of oligomerization loop in GPx3 was supposed to be related to the formation of the tetrameric complex. But in rhGPx3 mutant, the hydrophilic environment and the negatively charged surface of oligomerization loop may limit the access of large, complex lipid substrates to the active site. This could explain why the activity of rhGPx3 mutant using H$_2$O$_2$ as oxidizing substrate was in the same order of magnitude as that of human native GPx4 (34.8 U/mg) and GPx4 mutant with all Cys converted to Ser (14.0 U/mg), while it was much lower than that of native GPx4 (64.6 U/mg) and the GPx4 mutant (24.6 U/mg) in the case of using PLPC-OOH as the oxidizing substrate.

In conclusion, a rhGPx3 mutant with Cys converted to Ser was obtained from E. coli BL21(DE3) cells. Although the activity of this mutant was about one order of magnitude lower than that of native GPx3, the expression level of rhGPx3 mutant in E. coli was significantly higher than that of recombinant GPx3 in eukaryotic cells. There is good reason to believe that the rhGPx3 mutant would exhibit the same function as native enzyme if the tetramer is formed. Future studies are needed to explore the mechanism of tetramer formation.

**Methods**

**Construction of hexahistidine-tagged recombinant human GPx3 (rhGPx3) mutant expression vector.** A GPx3 mutant gene without signal peptide sequence (nucleic acids 1–60) was designed on the basis of the cDNA sequence of human GPx3 (GenBank accession No. NM_002084.3), in which the codons for Cys-12 (TGC), Cys-136 (TGT) were replaced with the codons for Ser (TCG) and the absence of post translational modifications. To our knowledge, this is the first time that a potential substitute for human GPx3 has been prepared from E. coli. Mass preparation of this enzyme is of great significance to explore the role of GPx3. There is good reason to believe that the rhGPx3 mutant would exhibit the same function as native enzyme if the tetramer is formed. Future studies are needed to explore the mechanism of tetramer formation.

**Overexpression and purification of rhGPx3 mutant.** The hGPx3 mutant was expressed in an E. coli BL21(DE3) auxotrophic strain following the previously described protocol with slight modifications. In detail, E. coli BL21(DE3) cells cotransformed with plasmids pCPG3M and pMalF2 (TaKaRa) were grown at 37°C in 400 mL of MM medium (modified M9 minimal medium) supplemented with ampicillin (100 μg/mL), kanamycin (25 μg/mL), chloramphenicol (25 μg/mL) and Cys (50 μg/mL) on a shaker. At OD$_{600}$ of 0.5, the culture was chilled in an ice-water bath for 5 min and incubated at 15°C for 45 min on a shaker. The cells were collected by centrifugation (5000 g for 5 min at 15°C) and washed twice with 0.9% NaCl at 15°C. The pellet was resuspended in 10 mL of MM medium supplemented with ampicillin (100 μg/mL), kanamycin (25 μg/mL), chloramphenicol (25 μg/mL), isopropyl-β-D-1-thiogalactopyranoside (1 mM) and Sec (600 μg/mL). Then the culture was incubated at 15°C for 16 more hours on a shaker.

The cells resuspended in binding buffer (50 mM sodium phosphate, 300 mM NaCl, and 20 mM imidazole, pH 7.4) were sonicated for 10 min with middle pulse and then filtered using 0.45 μm filter. The supernatant was loaded on an immobilized metal affinity chromatography column previously equilibrated with binding buffer. The rhGPx3 mutant with the hexahistidine tag was bound on the Nde I/III fragment from pUC57-GPx3M into the PCold I vector (TaKaRa). The construct was verified by DNA sequencing.

**SDS-PAGE and Western blot analysis.** Twenty μL of sample containing 5 mg of purified rhGPx3 mutant was mixed with 20 μL of loading buffer (20 mM Tris-HCl, 8% SDS, 10% β-mercaptoethanol, 30% glycerol and 0.004% bromophenol blue at pH 6.8) and heated in boiling water for 5 min. The samples were subjected to standard SDS-PAGE on 12% polyacrylamide gels with a 5% stacking gel. The gel was stained with Coomassie brilliant blue R-250 to visualize protein. To prepare non-reduced samples, β-mercaptoethanol was omitted from loading buffer and samples were loaded without boiling. Western blot analysis was performed using an anti-hexahistidine monoclonal antibody as described above. Samples were separated on a 12% SDS-PAGE gel and transferred onto a nitrocellulose membrane. The membrane was incubated for 1 h in blocking buffer composed of Tween-20 buffered saline (TTBS, 100 mM Tris, pH 7.5, 150 mM NaCl, 0.2% Tween 20) and 5% nonfat dry milk at 37°C. Then the membrane was incubated with mouse monoclonal anti-hexahistidine monoclonal antibody (Sigma) diluted (1:3000) in TTBS at 37°C for 1 h. The membrane was washed three times with TTBS for 5 min each and incubated with horseradish peroxidase conjugated goat anti-mouse IgG (Sigma) diluted (1:3000) in TTBS at 37°C for 1 h. After three washes with TTBS, immunoreactive protein band was identified by DAB staining. Total protein content was estimated by the Bradford method using bovine serum albumin as a standard. The selenium content of rhGPx3 mutant was determined by hydride generation atomic fluorescence spectrometer as described previously.

**Assay of enzyme activity.** GPx activities of rhGPx3 mutant were determined using a previously described method. Sodium phosphate (50 mM pH 7.4), EDTA (1 mM), GSH (2 mM), NADPH (0.25 mM), glutathione reductase (1 U) and samples (2.5 μL) were mixed in a cuvette at 37°C. The reagent mixture was incubated for 3 min at 37°C and the reaction was initiated by addition of (final concentration) 60 μM H$_2$O$_2$, 60 μM GSH and 50 μM PLPC-OOH. The peroxidase activity was determined by measuring the decrease of NADPH absorption at 340 nm per min. Activity units (U) are defined as the amount of enzyme necessary to oxidize 1 μmol of NADPH per min at 37°C. The specific activity is expressed in U/mg. Samples were run in triplicate, and the values were averaged.

**Determination of optimum temperature and pH.** The optimum temperature and pH for GPx activity of rhGPx3 mutant was determined by performing enzymatic assays at different temperatures (15 to 55°C) at pH 7.4. The optimum pH was determined by performing enzymatic assays at different pH levels (4.7 to 9.6) at 37°C. GPx activities were measured using H$_2$O$_2$ as an oxidizing substrate with the same method as described above. Samples were run in triplicate, and the values were averaged.

**Molecular modeling.** Molecular modeling was performed with the Insight II package, version 2000 (Accelrys, San Diego, CA), using the X-ray crystal structure of GPx3 with Sec-53 mutated to Gly (Protein Data Bank ID: 2R37) was used for the starting structure. The 3D structure of the GPx3 mutant with all Cys converted to Ser was refined by MD simulations and analyzed with Profile-3D and Procheck following the method as described previously. The electrostatic potential was calculated using the DelPhi module.

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