Characterization of redox-active cysteine residues of persulfide-responsive transcriptional repressor SqrR

Takayuki Shimizu and Shinji Masuda

Graduate School of Bioscience & Biotechnology, Tokyo Institute of Technology, Kanagawa, Japan; Center for Biological Resources and Informatics, Tokyo Institute of Technology, Kanagawa, Japan; Earth-Life Science Institute, Tokyo Institute of Technology, Tokyo, Japan

ABSTRACT

We recently identified the novel persulfide sensor SqrR that functions as a master regulator of sulfide-dependent gene expression in the purple photosynthetic bacterium *Rhodobacter capsulatus*. SqrR binds to the promoter regions of target genes to repress their expression in the absence of sulfide, and the repressor activity is negated by sulfide treatment. SqrR has 3 cysteine residues, 2 of which are conserved in SqrR homologs from other bacteria: Cys41 and Cys107. SqrR forms an intramolecular tetrasulfide bond between Cys41 and Cys107 when exposed to persulfide, which results in loss of the DNA-binding activity in vitro. Here, we address the mechanism through which these cysteine residues are modified by persulfides. We show that the predicted pH, value of Cys107, as revealed by a putative SqrR structural model, is lower than that of Cys41. Furthermore, C41S SqrR in which Cys41 was changed to serine forms an intermolecular disulfide bond between Cys107 of 2 SqrRs, suggesting high nucleophilic reactivity of Cyste107. These data suggest that Cys107 and Cys41 function as attacking Cys and resolving Cys, respectively; this occurs during tetrasulfide bond formation of WT SqrR, when it is exposed to persulfide.

Hydrogen sulfide is a toxic chemical that inhibits cellular respiration, and also works as the general biological gasotransmitter or signaling molecule in both prokaryotes and eukaryotes. Although cysteine thiol modifications are likely involved in this sulfide signaling, sulfide is negatively charged and thus will not react with nucleophilic thiolates. Recent studies in mammalian cells suggest that more oxidized and highly reactive sulfur-containing species, termed reactive sulfur species (RSS), are the actual sulfide signaling species. Although persulfidated (S-sulfhydrated) proteins have been identified and a plausible repair pathway(s) elucidated in mammalian cells, the functional impact of persulfidation of proteins and enzymes is as of yet not generally clear.

We recently identified the novel persulfide-responsive transcriptional repressor SqrR from the purple photosynthetic bacterium *R. capsulatus*. SqrR forms an intramolecular tetrasulfide bond between Cys41 and Cys107 when incubated with the sulfane sulfur donor glutathione, and then loses the repressor activity. Cysteine-dependent enzymes, e.g. thioredoxin (TrxA) and peroxide sensor (OxyR), which form an intramolecular disulfide crosslink, generally have what are termed attacking cysteine and resolving cysteine. The attacking cysteine initiates the crosslinking catalysis by intramolecular nucleophilic attack, so that its reactivity is higher than that of the resolving Cys. Here, to reveal the functional roles of the 2 conserved Cys residues for tetrasulfide bond formation of SqrR, we estimated the pH values of Cys41 and Cys107 that were calculated from a putative SqrR structure. Obtained results are compared with oligomerization characteristics of C41S and C107S SqrRs under oxidized conditions.

We first obtained a SqrR structural model by using the SWISS-MODEL server homology modeling pipeline (http://swissmodel.expasy.org/). Homology modeling was performed with the crystal structure of the transcriptional repressor BigR, a SqrR homolog, from *Xylella fastidiosa* (PDB ID: 3PQJ) as a template. Because the obtained structure was dimer, we deleted one subunit, chain B, using PyMOL (https://www.pymol.org/) to obtain a monomer structure. Then, pH values of Cys41 and Cys107 at pH7.0 in the SqrR structure were estimated by the web server called PROPKA 3.1 (http://propka.org/). By the calculation, pH values of Cys41 and Cys107 were predicted to be 10.13 and 9.36, respectively. Many hydrophobic amino acids are located...
around Cys107 (e.g., Leucine, methionine, isoleucine and alanine) that may influence the lower pKₐ value of Cys107. These results suggest that the nucleophilicity of Cys107 is higher than that of Cys41.

Next, we tested the reactivity of the 2 Cys residues by biochemical analysis. Recombinant C41S and C107S SqrR were expressed by an E. coli overexpression system followed by affinity chromatography as described previously. C41S and C107S SqrR were treated with CuCl₂ to induce disulfide bond formation and then applied to size-exclusion chromatography (Fig. 1A, C). The chromatogram of C41S SqrR displayed a single peak (fraction 44–48) (Fig. 1A). On the other hand, C107S SqrR showed 2 peaks; one is similar to that of C41S SqrR (fraction 44–48) and the second peak (fraction 37–40) is unique for C107S (Fig. 1C). To determine the oligomerization status of the proteins, each peak fraction was subjected to SDS-PAGE under non-reducing and reducing conditions (Fig. 1B, D). In the case of C41S SqrR, a single band estimated to be ≈24 kDa was observed in the fraction 45 and 46 under non-reducing conditions (Fig. 1B). Given the molecular weight of SqrR is 12.1 kDa, calculated from its deduced amino-acid sequence, C41S SqrR exists as dimer. On the other hand, no signal was detected above fraction 38 under both conditions (Fig. 1B). In the case of C107S SqrR, dimer and tetramer were detected in fractions 45–46 and 38–39, respectively (Fig. 1D). Moreover, a double band was observed in each fraction, indicating that there are at least 2 patterns of intermolecular crosslink between cysteine residues (e.g., Cys9-Cys9, Cys41-Cys41 and/or Cys9-Cys41). The pKₐ value of Cys41 was predicted to be higher than that of Cys107, so that the reactivity of Cys41 might be weaker than that of Cys107. Therefore, Cys41 reactivity is not particularly high and could react with various cysteine residues like Cys9. On the

**Figure 1.** Redox state of C41S and C107S SqrR. The recombinant *R. capsulatus* C41S and C107S SqrR were purified from *E. coli* as described previously. Purified SqrRs were treated with 50 μM CuCl₂ for 1 hour at room temperature to induce intermolecular disulfide-bond formation and then were loaded onto a Superdex 75 gel filtration column (GE) in 20 mM Tris-HCl (pH 8.0), 500 mM NaCl, 6% glycerol. (A and C) The size-exclusion chromatogram of C41S and C107S SqrR. Expected elution fractions are as follows: dimer, fraction number 44–48; tetramer, fraction number 37–40. Protein elution is monitored by measuring absorbance at 280 nm using an AKTA prime. (B and D) SDS-PAGE analysis of C41S SqrR and C107S SqrR of each fraction indicated by arrows in A and C under oxidizing (–β-mercaptoethanol; β-ME) and reducing (+β-ME) conditions. The gels were stained with SYPRO Ruby stain (Molecular Probes) and imaged using UV-transillumination.
other hand, C41S SqrR only forms an intermolecular disulfide bond between Cys107 of the 2 proteins immediately after induction with CuCl₂, indicating that the reactivity of Cys107 is higher than those of Cys9 and Cys41. Indeed, in the case of thioredoxin (TrxA), a strong intermolecular crosslinking between attacking cysteine residues is observed when the resolving Cys was changed to Ser.¹⁹ A single band estimated to be ≈12 kDa was observed under reduced conditions in all fractions, indicating that dimeric and tetrameric SqrRs were due to intermolecular disulfide cross-link formation (Fig. 1B, D). Overall, these results suggest that Cys107 functions as the redox-active attacking cysteine residue for the disulfide-bond formation.

SqrR senses persulfide by tetrasulfide formation between Cys41 and Cys107 based on nucleophilic reaction of thiol of cysteine residues and persulfide.²⁰ This reaction may be undertaken by similar mechanisms of the disulfide formation. Our structural prediction analysis as well as biochemical characterization of Cys mutants strongly suggested that Cys41 and Cys107 function as resolving and attacking residues during tetrasulfide-bond formation; Cys107 is initially modified by persulfide and then forms a tetrasulfide bond by performing an intramolecular nucleophilic attack on Cys41. Further characterization of the redox-dependent reaction of cysteine residues of SqrR should be important for understanding the step-by-step molecular mechanism of persulfide sensing in cells.

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