

Replacement of threonine-55 with glycine decreases the reduction rate of OsTrx20 by glutathione

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ABSTRACT

Thioredoxins (Trxs) are small ubiquitous oxidoreductase proteins with two redox-active Cys residues in a conserved active site (WCG/PPC) that regulate numerous target proteins via thiol/disulfide exchanges in the cells of prokaryotes and eukaryotes. The isoforms OsTrx23 with a typical active site (WCGPC) and OsTrx20 with an atypical active site (WCTPC) are two Trx h- type isoforms in rice that were previously found to be reduced by NADPH-dependent thioredoxin reductase and GSH/Grx system, respectively. In the present work the reduction of mutants G41T_{OsTrx23}, T55G_{OsTrx20}, K48D_{OsTrx20} and T55G-K48D_{OsTrx20} as well as wild types OsTrx23 and OsTrx20 were tested in the reaction containing either NADPH/NTR or glutathione (GSH). The results revealed that reduction rate of T55G_{OsTrx20} was remarkably decreased by GSH as compared to WtOsTrx20 highlighting the critical role of Thr-55 in interaction of OsTrx20 with GSH. On the other hand a significant decrease in the reduction rate of G41T_{OsTrx23} was observed in reaction containing NADPH-dependent thioredoxin reductase as compared with reduction rate of WtOsTrx23. These results suggest that first residue after N-terminal active site Cys is one of the critical residue in determination of system that Trxs can be reduced in.

Keywords: Glutathione; NADPH-dependent thioredoxin reductase; Rice; Thioredoxin

INTRODUCTION

Thioredoxins (Trxs) are small proteins (~12-14 kDa) with the highly conserved active site sequence motif, WCG(A/P)PC, and protein disulfide reductase activity [1]. Trxs act as electron donors to enzymes of metabolism such as ribonucleotide reductase, and play a critical role in the maintenance of reduced environment inside the cell [2]. In plants, Trxs are present in multiple forms based on primary structures and subcellular localization. Whereas Trxs f, m, x, and y are found in the chloroplast, Trxs o and h are

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localized to the cytoplasm or mitochondrion [3]. Trx f is reduced by ferredoxin/thioredoxin reductase and in turn reduced by ferredoxin generated during photosynthetic electron transport [4]. In contrast, reduction of Trx o and two large Trx h subgroups (subgroups I and II) depends on NADPH and involves NADPH-dependent thioredoxin reductase (NTR) [3, 5]. In subgroup III, there are Trx h isoforms belonging to poplar and *Medicago truncatula* that have been previously reported to be reduced by a GSH-dependent pathway [6,7].

NTRs in bacteria, archaea, fungi and plants are homodimers of ~35 kDa subunits, each containing a NADP-binding domain and a FAD-binding domain [8, 9]. The NADP-binding domain of NTR contains two redox-active cysteines in a conserved active site sequence motif, CAV(T)C [10]. The cysteines form a disulfide bond in the oxidized NTR and receive electrons from NADPH via the coenzyme FAD. For the reduction of Trx, one of the two cysteine thiols in the reduced NTR attacks the active-site disulfide bond in the oxidized Trx, and an intermolecular disulfide bond is formed as a reaction intermediate [11]. Structural information of this intermediate is available in an engineered NTR-Trx complex from barley obtained at a resolution of 3.0 Å [12].

NTR-Trx interaction is highly specific, and commonly NTR from different species prefer Trx from the same species as substrate [13-15]. For instance Arabidopsis NTR (AtNTR) shows 80-fold higher affinity towards AtTrx than *E. coli* Trx (EcTrx) despite the high structural conservation between NTRs and Trxs from the two organisms [13]. Moreover, the affinity of HvNTR isoforms is 20 fold higher than that from AtNTR towards HvTrxh isoforms [16]. These incompatibilities between NTR and Trx from different sources show the high specificity in the interaction between NTR and Trx [16].

We have previously cloned and produced the recombinant form of two cytoplasmic/mitochondrial type NTR (OsNTRA and HvNTRB) [17]. and three rice Trx h isoforms (OsTrx1, OsTrx20 and OsTrx23) [18]. An extensive analysis on the reaction kinetic of the rice NTR/Trx system showed OsTrx1 and OsTrx23 were reduced by both OsNTR isoforms [19]. However, the isoform OsTrx20, a member of subgroup III of *h-type*, with an atypical active site ${}_{53}\text{WCTPC}_{57}$ was not reduced with NTR isoforms but instead it was reactivated by GSH/Grx system. Similarly MtTrx8 with WCSPC was not reduced by MtNTR isoforms [6, 19]. Another obvious difference between OsTrx20 with other plant Trx h isoforms was the presence of Lys-48 as a basic amino acid that has been occupied with Asp or Asn in the most of other plant Trx h isoforms.

In the present work, to understand the role of Thr55_{OsTrx20}, Lys48_{OsTrx20} and Gly41_{OsTrx23} in reduction of Trxs by either NADPH/NTR system or GSH/glutaredoxin system the previously made mutants T55G_{OsTrx20}, K48D_{OsTrx20}, T55G-K48D_{OsTrx20} and G41T_{OsTrx23} [20]. as well as the wild types WtOsTrx20 and WtOsTrx23 were heterologously expressed and purified as His.tag fusion protein and were used in reactions that either NADPH or GSH served as electron donor.

MATERIALS AND METHODS

Heterogenous expression and purification of proteins: The previously *E. coli* strains producing OsNTRB [17], WtOsTrx20, WtOsTrx23 [18], G41T_{OsTrx23}, T55G_{OsTrx20}, K48D_{OsTrx20} and T55G-K48D_{OsTrx20} [20]. were grown at 37°C in LB medium supplemented with ampicillin (100 µg/ml) and chloramphenicol (5 µg/ml) to

an OD₆₀₀ of 0.6. Cultures were induced with 100 μM isopropyl β-d-thiogalactopyranoside (IPTG) for 4 h. The Cells were harvested by centrifugation and frozen at -80°C until use. For extraction of soluble proteins, cells were resuspended in Tris-HCl 10 mM, pH 8. The cells were lysed by sonication with 70 % cycle and 100 % amplitude for 45 min at 4°C. After centrifugation (12,000 rpm for 20 min at 4°C), the extracted soluble proteins in the supernatant were transferred in new tubes. The purification of His-tagged recombinant protein was performed using HisTrap HP columns (GE Healthcare). HisTrap HP columns are prepacked with Ni Sepharose High Performance and designed for simple, high-resolution purification of histidine-tagged proteins by immobilized metal ion affinity chromatography (IMAC). The extracted soluble proteins was applied onto His-Trap HP columns preequilibrated with loading buffer (10 mM imidazole, 500 mM NaCl, 30 mM Tris-HCl [pH 8.0]) and eluted in a gradient of 10–50 % imidazole. Eluted fractions were analyzed by the SDS-PAGE procedure. Purified proteins were desalted in 10 mM Tris-HCl buffer (pH 8.0) at 4°C using 12 kDa molecular weight cut-off cellulose acetate membranes (Sigma-Aldrich). The concentration of proteins were determined by A₂₈₀ and Beer-Lambert law using molar extinction coefficient 17085 M⁻¹ cm⁻¹ for WtOsTrx20, T55G_{OsTrx20}, K48D_{OsTrx20} and T55G-K48D_{OsTrx20}, 8542.5 M⁻¹ cm⁻¹ for WtOsTrx23 and G41T_{OsTrx23} and 28545 M⁻¹ cm⁻¹ for OsNTRB.

DTNB assay: Reduction of WtOsTrx20 and mutants with OsNTR1 was assayed in reaction containing NADPH as electron donor and DTNB (5, 5- dithiobis (2-nitrobenzoic acid) as final substrate [21]. The assay mixture contained 100 mM potassium phosphate pH 7.5, 10 mM EDTA, 0.1 mg/ml bovine serum albumin (BSA), 200 μM DTNB, 200 μM NADPH and 5 μM Trx. The reaction was initiated by addition of 80 nm NTR and the reduction of DTNB was measured by use of a spectrophotometer at 412 nm. The results reflected the formation of 2-nitro-5-thiobenzoic acid (TNB) ($\epsilon_{412\text{nm}} = 13,600 \text{ M}^{-1} \text{ cm}^{-1}$) [22]. The reaction mixture without OsTrx was used as a control. All reactions were repeated three times.

GSH- dependent insulin assay: The reduction of Trxs by GSH was tested in a reaction containing 300 μl reaction mixture containing GSH as electron donor and insulin as final substrate. The reaction mixture contained 100 mM potassium phosphate buffer, pH 7, 10 mM EDTA and 1mg/ml insulin and 5 μM OsTrx. The reactions were initiated by addition of 20 mM GSH and monitored spectrophotometrically at 650 nm for 100 min. A reaction mixture without Trx was run as control. All reactions were repeated three times.

Sequence Analysis: Multiple alignment between different rice Trx h isoforms was performed using the ClustalW software [<http://www.ebi.ac.uk/Tools/msa/clustalw2/>].

RESULTS

A multiple alignment among the amino acid sequences of rice Trx h isoforms demonstrated that the active site sequences of OsTrx1, OsTrx10, OsTrx15, OsTrx18, OsTrx23, OsTrx24, and OsTrx26 corresponds to classical Trx-active site motif (Trp-

Cys-Gly -Pro-Cys). In the active site sequence of OsTrx20, however, Gly has been replaced with Thr. Moreover, in other plant Trx h isoforms a conserved Asn/Asp is observed before active site. This amino acid has been replaced by a Lys in OsTrx20 (Fig. 1).

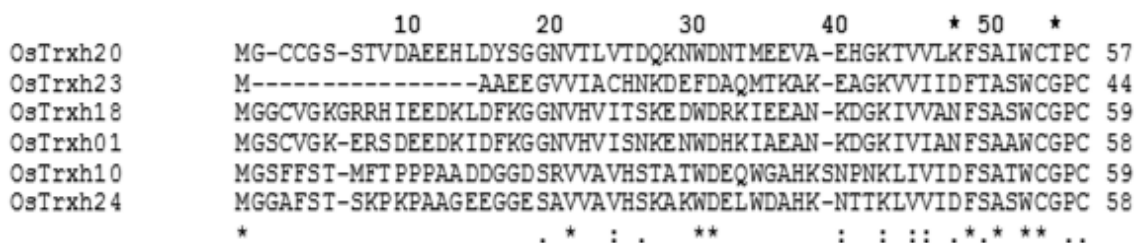


Figure1: Multiple alignment between different rice Trx h isoforms using the ClustalW software (<http://www.ebi.ac.uk/Tools/msa/clustalw2/>)

The proteins Wt_{OsTrx20}, Wt_{OsTrx23}, OsNTRB and mutants K48D_{OsTrx20}, T55G_{OsTrx20}, T55G-K48D_{OsTrx20} and G41T_{OsTrx23} with a His.tag at the N-terminus were found in the soluble fraction of the *E. coli* transformant culture after induction with IPTG. The theoretical molecular mass of Wt_{OsTrx20}, Wt_{OsTrx23}, OsNTRB, K48D_{OsTrx20}, T55G_{OsTrx20}, T55G-K48D_{OsTrx20} and G41T_{OsTrx23} were 17.5, 15.4, 39.9, 17.5, 17.5, 17.5 and 15.4 KDa respectively. SDS-PAGE of cell extracts revealed a prominent polypeptide band of the expected molecular mass (Fig. 2, lanes 2–7 and 14). The recombinant proteins were purified from the crude extracts by nickel affinity chromatography (Fig. 2, lanes 8-13 and 15).

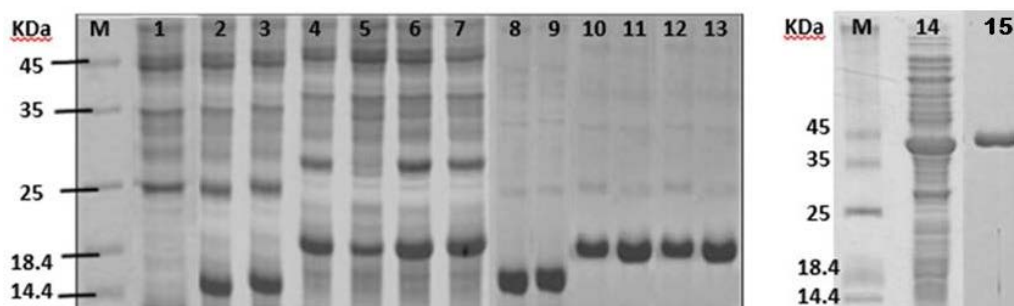


Figure 2: Overexpression of wild types and mutants in *E. coli*. SDS-PAGE analysis for verification of expression and purification. Total soluble protein extracted from *E. coli* harboring pET-15b (lane 1), pET15b-Wt_{OsTrx23} (lane 2), pET15b-G41T_{OsTrx23} (lane 3), pET15b-Wt_{OsTrx20} (lane 4), pET15b-K48D_{OsTrx20} (lanes 5), pET15b-T55G_{OsTrx20} (lane 6), pET15b-T55G- K48D_{OsTrx20} (lane 7) and OsNTRB (lane 14) 4 h after addition of IPTG. Purified His-Wt_{OsTrx23} (lane 8), His-G41T_{OsTrx23} (lane 9), His-Wt_{OsTrx20} (lane 10), His-T55G_{OsTrx20} (lane 11), His-K48D_{OsTrx20} (lane 12), His-(T55G)(K48D)_{OsTrx20} (lane 13) and OsNTRB (lane 15).

The insulin precipitation was monitored by the increase in turbidity at 650 nm (ΔOD_{650}) in insulin-disulfide reduction assays containing GSH in the presence of Wt_{OsTrx20}, T55G_{OsTrx20}, K48D_{OsTrx20} or T55G-K48D_{OsTrx23} (Fig. 3). The results showed that GSH actively reduced Wt_{OsTrx20}. However, in comparison to Wt_{OsTrx20} the rate of insulin reduction remarkably decreased when T55G_{OsTrx20} and T55G-K48D_{OsTrx20} was used in reaction. In addition, the rate of reaction slightly decreased in

the presence of K48D_{OsTrx20}. These results suggest that Thr55 has critical role in reduction of OsTrx20 by GSH.

The results of DTNB assay showed that recombinant form of OsNTRB was able to actively reduce WtOsTrx23 (Fig. 4). However the initial velocity of reaction was remarkably decreased when the mutant G41T_{OsTrx23} was used instead of WtOsTrx23. In contrast to OsTrx23, the isoform OsTrx20 was not reduced by OsNTRB. However the substitution of Thr 55 with Gly and Lys48 with Asp slightly increased the reaction of this isoform with OsNTRB. These results suggest that the residue Gly41 which is located in the first position after N-terminal active site Cys is critical for NTR-Trx interaction.

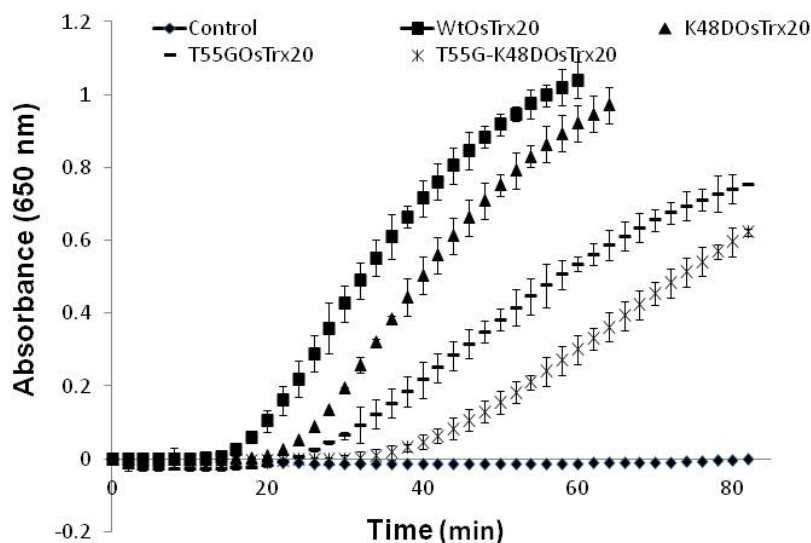


Figure 3: Insulin reduction WtOsTrx20 and mutants in the presence of GSH. The control shows reaction containing GSH and insulin without addition of Trx.

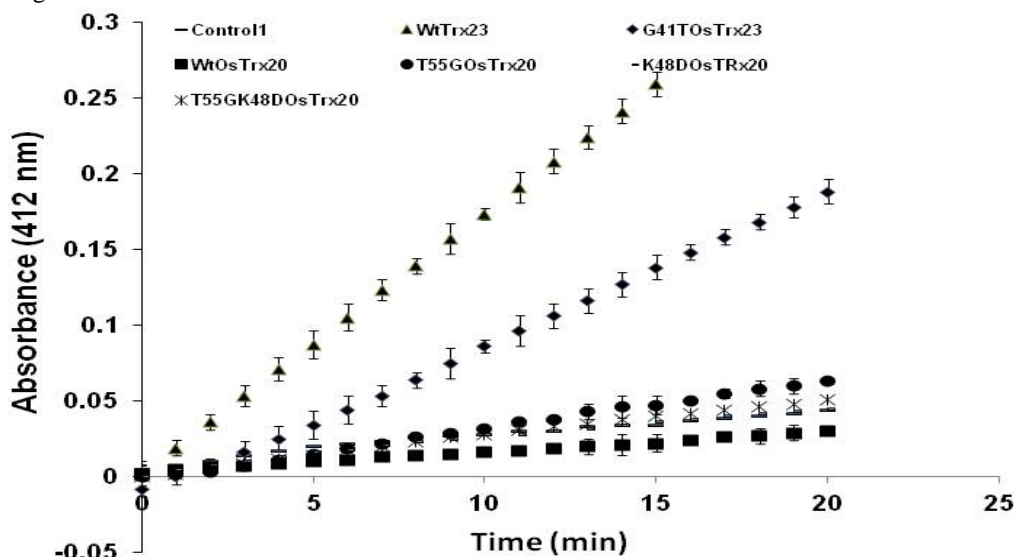


Figure 4: Time course of reduction of Trxs by OsNTRB as monitored by reduction of DTNB. Control shows the time course of DTNB reduction by OsNTRB without addition of OsTrx.

DISCUSSION

It has been very well documented that some members of the Trx h subgroup III characterized in plants with an atypical active site (WCTPC or WCSPC) are not reduced by NADPH/NTR system but they can be reactivated by GSH/Grx system [6, 7]. In contrast to Subgroup III, the members of subgroup I and II with typical WCGPC active site are reduced by cytoplasmic/mitochondrial NTRs [16, 19]. In rice Trx h isoforms can be divided in three subgroups on the basis of their amino acid sequences [19]. Previously we show that OsTrx23, a member of subgroup I is actively reduced by both OsNTRB and OsNTRA, but cannot be reduced by GSH [19]. In contrast, OsTrx20 which belongs to subgroup III and carry an atypical active site can be reactivated by GSH/Grx system [19]. The isoform OsTrx23 and OsTrx20 are predicted to be dominantly localized in cytoplasm and chloroplast, respectively [23]. The isoform OsTrx20 is expressed under light in the leaves of rice seedlings [23]. The activity of OsTrx20 for reduction of target proteins was demonstrated to be pH-dependent and the residues Thr55 and Lys48 were shown as key residues for instability of its activity under pH changes [20].

In this work, we showed that the rate of reduction of G41T_{OsTrx23} by OsNTRB decreases in comparison to WtOsTrx23 suggesting the presence of Thr in this position impairs the interaction of OsTrx23 with OsNTRB. In addition we showed that the isoform WtOsTrx20 was actively reduced by GSH and substitution of Thr55 with Gly in OsTrx20 resulted in remarkable decrease of rate of reduction by GSH. These results supports the critical role of residues situated between two active site Cys in redox activity of Trxs [24]. and highlight the role of first residue after N-terminal active site Cys in determination of the system that Trxs can be reduced in.

Taken together the presence of atypical active site for some h-type Trx isoforms, with Thr or Ser substitution for the largely conserved Gly in the first position after N-terminal active site Cys might be an evolution change that enables these isoforms to be reduced by GSH and adapt them in the cell organelles lacking cytoplasmic/mitochondrial NTR isoforms.

Conflict of Interest: The authors declare no conflict of interest.

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