Elucidation of Thioredoxin Target Protein Networks in Mouse*

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Thioredoxin 1 (Trx1) is a key redox modulator that is functionally conserved across a wide range of species, including plants, bacteria, and mammals. Using a conserved CXXC motif, Trx1 catalyzes the reduction of cysteine disulfides and S-nitrosothiols. In contrast to small molecular reductants such as glutathione and cysteine that can reduce a wide range of oxidized proteins, Trx1 reduces only selected proteins via specific protein-protein interaction. Trx1 has been shown to regulate numerous signal transduction pathways, and its dysfunctions have been implicated in several diseases, including cancer, inflammation, and neurodegenerative and cardiovascular diseases. Identification of Trx1 target proteins may help to identify novel signaling mechanisms that are important for Trx1 antistress responses. In this study, we performed an iTRAQ proteomics study for the identification of Trx1 target proteins from the hearts of a cardiac specific Trx1-overexpressing transgenic mouse model (Tg-Trx1). Trx1-reduced proteins were distinguished from Trx1-induced proteins by comparison of the iTRAQ results with those obtained using a parallel iTRAQ (isobaric tags for relative and absolute quantitation) protein expression analysis. We were able to identify 78 putative Trx1 reductive sites in 55 proteins. Interestingly, we identified a few protein functional networks that had not been shown previously to be regulated by Trx1, including the creatine-phosphocreatine shuttle, the mitochondrial permeability transition pore complex, and the cardiac contractile apparatus. The results presented here suggest that in addition to a general antioxidant function, Trx1 may be involved in the coordination of a wide array of cellular functions for maintaining proper cardiac energy dynamics and facilitating muscle contraction. Molecular & Cellular Proteomics 8:1674–1687, 2009.

Thioredoxins (Trxs) are a class of antioxidant proteins that mediate the reduction of specific disulfide bonds and S-nitrosothiols within oxidized proteins. Trx and the NADPH-dependent thioredoxin reductase (TrxR) form a protein reductive system that plays essential roles in the repair of oxidatively damaged proteins and the restoration of cellular redox homeostasis. Two isoforms of Trx have been widely studied: Trx1 is mainly found in the cytosol and nucleus, whereas Trx2 is mitochondrial-specific. Each Trx isoform is coupled with its own thioredoxin reductase (TrxR) form, forming a protein reductive system. Trx1 has been shown to promote cell growth and proliferation and to inhibit apoptosis by modulating both caspase-dependent and -independent pathways. Trx1 has also been demonstrated to be a potent regulator of a wide variety of transcription factors and other gene expression regulators by preserving the reductive states of specific cysteines within transiently oxidized proteins such as nuclear factor xB (1), hypoxia inducible factor-1α (2), histone deacetylase 4 (3), and glucocorticoid receptor (4). A wide range of bioanalytical techniques has been applied to identify Trx1 targets, including two-dimensional electrophoresis-based differential thiol labeling (5, 6), Trx affinity chromatography (7, 8), and ICAT (9). The ICAT technique is of particular interest to redox analysis because of its capability to aid the identification of potential reduction sites within substrate proteins as well as the quantification of the extent of reduction (10). This technique was pioneered by Aebersold and co-workers (11) for the quantification of proteins by isotope tagging of cysteines. The advantage of this technique resides within its ability for the enrichment of cysteine-containing peptides for the reduction of sample complexity, resulting in more depth in stable isotope-based quantitative proteome coverage than conventional shotgun proteomics methods (11). Cohen and co-workers (12, 13) have tailored this technique to probe the redox status of protein cysteines and identified redox-sensitive cysteines in cardiac sarcoplasmic reticulum proteins including ion channels that may be important for modulating cardiac functions. Furthermore they have also elegantly mapped differential cysteine thiol redox sensitivities of p21ras GTPase toward peroxynitrite and oxidized glutathione (14). More recently, Svensson and co-workers (9) have used a...
modified ICAT method to discover over 100 in vitro Trx reduction targets in plants in which Trx-induced changes in protein disulfides were quantified. This report has provided the largest number of Trx-targeting plant disulfides to date, demonstrating the feasibility of using this technique for uncovering Trx targets in mammalian systems. In this study, we adopted a redox ICAT strategy to detect potential Trx1 reduction targets in rodent tissues.

Trx1 has been associated with a wide variety of diseases with oxidative imbalance, including cancer (15), human immunodeficiency virus infection (16), neurodegenerative diseases (17), and cardiovascular diseases (18). We were interested in understanding the roles of Trx1 in the protection of heart function in a rodent model of cardiac hypertrophy, the adaptive enlargement of the heart when under stress. However, prolonged pathological hypertrophy has been associated with metabolic disorder, inadequate ATP supply, contractile dysfunction, and gradual development into heart failure (19). We have shown previously that overexpression of Trx1 (in Tg-Trx1 animals) plays a central role in the activation of cardioprotective signal transduction pathways within hypertrophic hearts (3, 20). Mechanistically Trx1 may also exert its function through the regulation of gene expression, translation, and post-translational modifications. In a previous RNA microarray study (21), we reported that a wide range of genes is significantly altered in the hearts of Tg-Trx1 mice, including the up-regulation of genes involved in oxidative phosphorylation and the tricarboxylic acid cycle and several stress-related transcriptional factors. Given its known function as a protein reductant, a significant aspect of Trx1 function is expected to be exerted through the selective reduction of target proteins. For example, recently we have reported that histone deacetylase 4 is an important target of Trx1 reduction in heart (3). Trx1 facilitates the formation of a histone deacetylase 4-containing multiprotein complex resulting in histone deacetylase 4 nuclear translocation and thus regulates the expression of antihypertrophic genes. It is likely that in addition to regulating the function of individual proteins Trx1 may also exert its cardiac protective function by coordinately regulating a series of protein networks. To identify such protein networks, we conducted an ICAT-based proteomics study to identify proteins whose cysteine thiols became more reduced in the hearts of Tg-Trx1 animals compared with the control animals. The ICAT results were compared with data obtained from an in vitro study using a Trx1 overexpression construct and a Trx1-induced protein expression analysis to reveal genuine Trx1 reduction target proteins as opposed to Trx1-induced proteins. We discovered that protein networks associated with energy production and utilization processes, such as glycolysis, the tricarboxylic acid cycle, β-oxidation, the mitochondrial permeability transition pore (MPTP) complex, and the contractile apparatus of the myofibrils, were affected by Trx1 overexpression, suggesting a role for Trx1 in maintaining heart energetics.

MATERIALS AND METHODS

Chemicals and Reagents

HPLC grade ACN and water were purchased from J. T. Baker Inc. Sequencing grade modified trypsin was from Promega (Madison, WI). Tris, α-cyano-4-hydroxycinnamic acid, catalase, protease inhibitor mixture (P8340), and other chemicals were purchased from Sigma-Aldrich unless stated otherwise. MS calibration standard peptides, Glu-fibrinopeptide, and human adenosinecorticotropic hormone 18–39 were bought from AnaSpec (San Jose, CA). Cleavable ICAT reagents and iTRAQ reagents were obtained from Applied Biosystems (ABI, Foster City, CA). Recombinant human DJ-1 (ab651198, Abcam, Cambridge, MA), human Trx1 (T8690, Sigma), rat TrxR1 (American Diagnostica, Greenwich, CT), anti-Trx1 antibody (ab16835, Abcam), anti-adenine-nucleotide translocase 1 (ANT1; sc-9299, Santa Cruz Biotechnology, Santa Cruz, CA), and anti-biotin M antibody (MB-9100, Vector Laboratories, Burlingame, CA) were used in this study.

Transgenic Mouse Generation and Transverse Aortic Constriction (TAC) Surgery

Mice with cardiac specific overexpression of Trx1 (Tg-Trx1) were generated on an FVB background using the α-myosin heavy chain promoter as described previously (21, 22). Induction of cardiac hypertrophy was accomplished by surgical constriction of the transverse thoracic aorta and was performed on both the control and Tg-Trx1 mice as reported earlier (23). Briefly mice were anesthetized with pentobarbital sodium solution (60 mg/kg, intraperitoneal) and ventilated using a rodent ventilator. The left chest was entered through the second intercostal space, and the aorta was isolated. A Prolene suture was placed around the aorta between the innominate artery and the left carotid artery. A 27-gauge needle was tied onto the aorta and later removed. The chest was then closed in layers. The animals were kept warm, and the rectal temperature was maintained at 37 °C. After weaning from the ventilator, the mice were kept in an incubator with humidified oxygen and returned to cages after recovering from anesthesia. All protocols regarding the use of animals were in compliance with the regulations of the Institutional Animal Care and Use Committee at the University of Medicine and Dentistry of New Jersey.

Protein Extraction

Protein extracts were prepared from ~100 mg of dissected left ventricular tissues after three cold PBS washes and were homogenized with a tissue homogenizer (Omi International, Marietta, GA) in 500 μl of lysis buffer. For the ICAT proteomics study, 6 μl urea, 2% CHAPS, 1% Triton X-100, and 30 mM Tris-HCl at pH 7.5 was used. For the iTRAQ proteomics study, 25 μl triethylammonium bicarbonate and 20 mM Na2CO3 at pH 8.0 was used. Protease inhibitors (10 μl/ml), 2.5 mM sodium pyrophosphate, 1 mM β-glycerophosphate, and 1 mM Na3VO4 were added to both lysis buffers. Protein concentrations were measured with the BCA protein assay (Pierce). Protein extractions and sample preparation were quickly carried out on ice and with minimum exposure to air. Protein extracts were immediately precipitated with cold acetone (−20 °C) to remove unwanted interferences such as metal ions and other small antioxidant molecules that may induce non-biological redox variations of proteins. Furthermore both control and Tg-Trx1 samples were processed in parallel so the impact of artificial oxidation, if any, would be comparable across all the samples.

ICAT Labeling and Multidimensional Chromatography

The ICAT labeling procedures used for this study have been described previously (10) and are illustrated in Fig. 1. In brief, 100 μg of
each of the three independent control extracts was labeled with the light ICAT reagents, and the same amounts of Tg-Trx1 samples (three animals) were labeled with the heavy ICAT reagents at 37 °C for 2 h. No reduction and alkylation steps were performed prior to ICAT labeling to preserve the native protein thiol redox states. Excess ICAT reagents were removed with the addition of 50 mM DTT. Newly generated free thiols were alkylated with 50 mM iodoacetamide. The light (L) and heavy (H) ICAT labeled proteins were mixed and subjected to tryptic digestions at a 50:1 protein/enzyme ratio. Tryptic digested peptides were acidified and fractionated using strong cation exchange chromatography (SCX). SCX was run on an ABI Biocad Sprint System equipped with a PolySulfoethyl A column (200 mm × 4.6 mm; PolyLC Inc., Columbia, MD). The gradient profile of SCX consisted of 10 min of 100% mobile phase A (10 mM KH2PO4 and 20% ACN, pH 2.7) and subjected to SCX fractionation using a LC Packings capillary HPLC system (Dionex, Sunnyvale, CA). Peptides were captured with an in-line trapping column (5 μm, 0.3 × 5 mm) and resolved on a PepMapTM C18 column (5 μm, 0.075 × 150 mm) with a 70-min linear gradient of 0–30% mobile phase B (95% ACN and 0.1% TFA) followed by 30 min of 30–90% mobile phase B at a flow rate of 400 nl/min. The RPLC eluents were mixed in line at a 1:2 ratio with a MALDI matrix (4 mg/ml α-cyano-4-hydroxycinnamic acid, 60% ACN, 0.1% TFA, 20 mM ammonium phosphate, and the internal mass calibrants) through a microtome and deposited onto the MALDI plates with a Probot (Dionex).

**iTRAQ Labeling and Multidimensional Chromatography**

Fifty micrograms of protein from each of the four samples (two controls and two Tg-Trx1 animals) was digested by trypsin and labeled with one of the four iTRAQ reagents according to the manufacturer’s protocol. Peptides derived from the two control samples were labeled with iTRAQ tags 114 and 115, and the two Tg-Trx1 samples were labeled with iTRAQ tags 116 and 117 (see Fig. 1). The iTRAQ-labeled peptides were first combined; dried in a SpeedVac; then resuspended in 500 μl of Buffer A, which contained 10 mM KH2PO4 and 20% ACN, pH 2.7; and subjected to SCX fractionation using a PolySulfoethyl A column (4.6 × 200 mm, 5 μm, 300 Å; PolyLC Inc.) on an ABI Biocad Sprint System. A 60-min gradient program consisting of 10 min of mobile phase A and 40 min of linear gradient from 0 to 50% mobile phase B (10 mM KH2PO4, 20% ACN, and 600 mM KCl at pH 2.7) followed by a 10-min linear gradient from 50 to 100% B was performed at a constant flow rate of 1 ml/min. Thirty-five fractions were collected and desalted using PepClean™ C18 spin columns (Pierce) according to manufacturer’s protocols. RPLC separation of peptides within SCX fractions was identical to the procedures used for ICAT peptide analysis described above.

**ICAT Analysis on Recombinant Human DJ-1**

Recombinant human DJ-1 (0.5 μg/μl) was first oxidized with 200 μM H2O2 for 30 min in the dark. The reaction was quenched by addition of catalase (0.1 μg/ml). The oxidized DJ-1 solution (100 μl) was split in half: one half was treated with the Trx1 reduction system, and the other half was treated with buffer only. The Trx1 reduction system was prepared as follows. Five micrograms of Trx1 was incubated in 2 μl of activation buffer (10 mM Tris and 2 mM EDTA, pH 7.5) for 15 min, and this solution was then mixed with 14 μl of reaction solution (10 mM Tris, 2 mM EDTA, 0.2 mM NADPH, and 1 μg of human Trx1 reductase, pH 7.5). Twenty-five micrograms of oxidized DJ-1 protein was added to the Trx1 reduction solution and then incubated at room temperature for 20 min. Trx1-reduced DJ-1 was then labeled with the ICAT H reagent, and the buffer-treated control was labeled with the ICAT L reagent. The remaining procedures followed the ICAT protocols described above except that the SCX step was carried out with an SCX cartridge provided in the ICAT labeling kit (ABI) according to the manufacturer’s protocol.

**MS and Database Search**

ICAT—RPLC-purified peptides were analyzed on a 4800 MALDI-TOF/TOF mass spectrometer (ABI) operating in the reflectron mode. First, 2,000 shots were accumulated for each MS spectrum with a mass range of m/z 800–3,800. Internal calibration standards were used for achieving a mass accuracy better than 50 ppm. Following MS analysis, the isotope ion cluster intensities of ICAT pair ions (heavy and light ions with a mass difference of 9.03 ± 0.03 Da) were extracted and quantified from the parent spectra with the GPS Explorer software (v3.5, ABI). Only ICAT pairs with at least 20% intensity differences and a signal-to-noise ratio over 50 were submitted for MS/MS analysis; 3,500 shots were accumulated for each MS/MS spectrum. The peak lists were generated with 4000 Series Explorer (v3.5, ABI). For peak detection, the signal-to-noise ratio threshold was set to 10, local noise window width was 250 m/z, and minimum peak width bin size was 2.9; resolution was set at 22,000 at m/z 2,400 for MS and 8,000 at m/z 2,000 for MS/MS. Peptide identification was performed with MASCOT (v1.9.1) integrated in the GPS Explorer software against a non-redundant Swiss-Prot database (release 54 containing 13,561 unique mouse sequences). The search parameters included one missed tryptic cleavage and 50-ppm MS and 0.3 Da MS/MS error tolerance. Variable modifications included ICAT L/H modifications, carbamidomethylation of cysteines, and methionine oxidation. Only peptides identified with confidence interval (C.I.) values at or above 95% were considered as confident identifications. Each MS/MS spectrum was subjected to manual inspection for the confirmation of positive identifications. False discovery rates were estimated to be less than 4% for all three experiments (calculation according to Peng et al. (24)). Typical ICAT quantification accuracy with this work flow on our instruments was evaluated by using BSA tryptic peptides of set mixing ratios. Analytical coefficient of variation values of less than 10% were observed (10). For the data presented in Table I, we decided to include only peptides with ICAT changes beyond 20% of the population median so that future biological validations are more likely to be successful.

iTRAQ—RPLC-separated iTRAQ-labeled peptides were analyzed on a 4800 MALDI-TOF/TOF MS instrument. MS spectra (m/z 800–3,500) were acquired in positive ion reflector mode, averaging 500 laser shots per spot. Data-dependent selection of the top 10 most intense ions in each MALDI spot were submitted for subsequent MS/MS analysis using 2-keV collision energy and 5 × 10−7-torr collision gas pressure with an accumulation of 2,000 shots per precursor mass. MS/MS ion peak lists were generated as described for the ICAT experiments. Protein database search by MASCOT (v1.9) was performed against the same Swiss-Prot database as that used for ICAT analysis. The following search parameters were used: trypsin with one missed cleavage; mass tolerance of 50 ppm for the precursor ions and 0.3 Da for the MS/MS fragment ions; fixed modifications including N-terminal iTRAQ labeling, iTRAQ-labeled lysines, and MMTS-modified cysteines; and variable modifications including methionine oxidation and iTRAQ-labeled tyrosines. Peptides identified with C.I. values no less than 95% were used for protein identification.
and quantitation. Protein expression ratios (Tg-Trx1/control) were calculated as described previously (25, 26). Briefly the iTRAQ reporter ion cluster areas were extracted using the GPS Explorer software (ABI), and only ion counts greater than 5,000 were used for quantification analysis. The individual reporter ion peak areas for each iTRAQ channel were normalized by the population median. For confident protein identification (average of at least two unique peptides per protein), the relative protein expression ratio distribution and the standard deviations between the animal groups were determined accordingly. In this study, two biological replicates of the iTRAQ-labeled sample were analyzed, and a Student’s t test was performed to determine differentially expressed proteins. Proteins quantified with p values no greater than 0.05 and a protein expression ratio beyond 20% of the population median were considered as differentially expressed (26).

**ANT1 Immunoprecipitation and Western Blotting**

Heart proteins were extracted with a buffer containing 20 mM Tris-HCl, 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, 1% Triton X-100, 1 mM sodium orthovanadate, 2.5 mM sodium pyrophosphate, and 1 mM β-glycerophosphate, pH 7.4, supplemented with protease inhibitor mixtures (10 μl/ml Cell extracts (500 μg of proteins) were first treated with 200 μM H2O2 for 30 min followed by the addition of 0.1 μg/ml catalase to remove the excess H2O2. Different protein reduction conditions were applied separately to the oxidized proteins for 30 min at room temperature: (1) reaction buffer (2.0 mM EDTA and 10 mM Tris-HCl, pH 7.5) only; (2) 5 μg of Trx1 in the reaction buffer; (3) 5 μg of Trx1, 1 μg of TrxR, and 0.2 mM NADPH in the reaction buffer; and (4) 5 μM tris(2-carboxyethyl)phosphine (TCEP). The treated proteins were precipitated with cold acetone (−20°C) and the pellets were washed twice with cold acetone. The pellets were resuspended in radioimmune precipitation assay buffer, and the protein cysteine thiols were alkylated with the addition of 0.2 μM N-(2-(6-(biotinamido)hexyl)-3-(2'-pyridylthio)-propionamide for 1 h. The biotinylated proteins were immunoprecipitated with an anti-ANT1 antibody. For Western blot analysis, the proteins were separated by 11% SDS-PAGE and transferred onto a nitrocellulose membrane. The membrane was probed with either an anti-biotin M antibody or an anti-ANT1 antibody, respectively. Horseradish peroxidase-conjugated secondary antibody (Santa Cruz Biotechnology) was used for detection and visualization by chemiluminescence.

**Statistical Analyses**

iCAT data are expressed as ICAT pair ratios (H/L: Tg-Trx1/control) from three independent experiments (Table I). Statistical significance was determined by performing a one-tailed Student’s t test with Microsoft Excel, and a p value ≤ 0.05 was considered significant.

**Trx1 Reductive Target Sequence Motif Search**

An iterative statistical program (27), motif-x, was used to extract potential Trx1 reductive motifs among the identified target sequences. Cysteine was assigned as the central amino acid, and foreground sequences were extended to 15 amino acids on each side of the cysteine. The significance threshold was set at 0.000001.

**RESULTS**

**Identification of Putative Trx1 Target Cysteines in Cardiac Proteins—Hypertrophy and increased oxidative stress in hearts were introduced with TAC surgery in both control and Tg-Trx1 mice. Western analysis confirmed the significant overexpression of Trx1 in transgenic animals (supplemental Fig. 1). TAC induced a significant increase in left ventricle weight/body weight (4.0 ± 0.2 mg/g), an index of cardiac hypertrophy in control mice, whereas the increases in left ventricle weight/body weight were significantly attenuated in Tg-Trx1 mice (3.5 ± 0.1 mg/g; p < 0.01), consistent with our previous observation (22). Both Tg-Trx1 and control animals, when subjected to TAC surgery, incurred elevated oxidation of many proteins (21, 22). However, the elevated Trx1 levels in the transgenic mice facilitate the reduction of specific cysteines in target proteins. We used a forward iCAT labeling strategy as described previously (10) to identify redox-sensitive cysteines that are putative targets of Trx1 in TAC-stressed mouse hearts (Fig. 1). ICAT H reagent was used to label Tg-Trx1 samples, whereas the ICAT L reagent was used to label the control samples. In such a scheme, Trx1-induced protein thiol reduction would be manifested as H/L ratios larger than 1.0. Within three independent ICAT experiments, we observed ~2,000 ICAT ion pairs with a mean ICAT ratio of 1.15 (ICAT ratios presented in Table I; ratios were normalized by the population median of the individual experiment); ~700 precursors were selected for MS/MS identifications. We identified 152, 166, and 195 unique peptides with C.I. value ≥ 95% in the three experiments (supplemental Table 1). According to our previous analysis of the ICAT redox proteomics work flow on our MS instrument (10), the typical quantification coefficient of variation is less than 10%. Therefore, a more stringent 20% ICAT quantification ratio was used as a cutoff value for determining significant reduction of protein thiol levels between control and Tg-Trx1 animals. Only peptides fulfilled the following criteria were included in Table I: (i) C.I. value ≥ 95%, (ii) identification in all three ICAT experiments, (iii) p value ≤ 0.05, and (iv) ICAT ratio changes beyond 20% of the population median.

We found that 78 cysteines within 55 proteins were significantly reduced by Trx1 overexpression in all three experiments (Table I; categorized by gene ontology functional groups), and an additional 70 peptides were found in at least two experiments (supplemental Table 1). Potential Trx1 target proteins demonstrated various degrees of sensitivity to Trx1 reduction, ranging from a 20 to ~300% increase in free cysteine thiol levels. The majority of the Trx1-responsive proteins included structural proteins, ion channels, stress response proteins, and metabolic enzymes in the glycolysis pathway, the tricarboxylic acid cycle, oxidative phosphorylation, and β-oxidation of fatty acids.

We previously showed that TAC treatment resulted in increased oxidative stress in heart, likely resulting in elevated protein oxidation (22). A large number of proteins were found to contain cysteines highly sensitive to Trx1 reduction, including glyceraldehyde-3-phosphate dehydrogenase (3.8), pyruvate dehydrogenase E1 component subunit α (2.0), long-chain-specific acyl-CoA dehydrogenase (3.3), aldose reductase (2.5), ADP/ATP translocase 1 (2.5), and aspartate aminotransferase (3.4) (numbers in parentheses indicate -fold...
changes of free thiol levels, Tg-Trx1/control). Many proteins possessed more than one cysteine that was sensitive to Trx1 reduction but to different extents. For example, four different creatine kinase (Q6P8J7) peptides were found to have five cysteines: Cys63, Cys67, Cys90, Cys180, and Cys317. Cys63 and Cys67 were non-responsive to Trx1 as no significant differences were found between the light and heavy ICAT-labeled peptides (supplemental Table 1). Cys90, Cys180, and Cys317 responded positively to the Trx1-mediated reduction leading to an increase in heavy ICAT-labeled peptide ions by 20, 60, and 70%, respectively (Table I). Protein levels of creatine kinase were comparable between the control and Tg-Trx1 animals as evidenced by an iTRAQ ratio of 0.9.

To understand whether the observed increase of free thiol content was exclusively due to Trx1-mediated protein reduction or also due to alterations in protein expression, we compared the results of the ICAT analysis with those of the iTRAQ analysis of similar samples. Because our ICAT work flow did not contain the initial reduction step prior to ICAT labeling, increases in H/L ratios could be due to increases in protein amounts and/or their free thiol levels. The rationale for the comparison of ICAT and iTRAQ results was that if the protein levels were constant according to the iTRAQ method different ICAT ratios should be attributed to reversible thiol modifications. In the iTRAQ analysis, 10 proteins (including Trx1 with iTRAQ ratio of 1.5; \( p < 0.01 \)) of 769 unique proteins (\( \sim 2\% \)) were found to be significantly changed in the Tg-Trx1 animals. Changed proteins have been validated by Western blotting analysis with more animal numbers. Some proteins were identified with average iTRAQ ratios beyond 20% of the population median but were not deemed significant (\( p \) values \( >0.05 \)), for example myosin-binding protein C. Myosin-binding protein C was found with an average iTRAQ ratio of 0.8 (\( p \) value of 0.26 from 41 peptides) and with average ICAT peptide ratios for three distinct peptides of 1.7, 2.1, and 1.7, respectively (Table I). It is likely that these peptides would have been detected as more reduced in Tg-Trx1 animals if this protein were not also down-regulated.

Validation of ANT1 as a Direct Trx1 Reduction Target—Following the ICAT proteomics study, we confirmed that one of the proteins, ANT1, contained an oxidized cysteine that can

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TABLE I
Putative Trx1-mediated reduction protein network

| Protein name (Swiss-Prot accession number) | Peptide sequence | ICAT ratio (H/L) | Avg<sup>2</sup> | ICAT ratio | Protein ITRAQ ratio | Reported Trx1 Targets (Refs.) |
|-------------------------------------------|-----------------|-----------------|---------------|------------|---------------------|-----------------------------|
| **Metabolic enzymes**                     |                 |                 |               |            |                     |                             |
| Carbohydrate metabolism                   |                 |                 |               |            |                     |                             |
| Acetyl-coenzyme A synthetase 2-like (O96981) | ZGGLGLTVEGP     | 1.0 (12)        |               |            |                     |                             |
| Glyceraldehyde-3-phosphate dehydrogenase  | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| (P16858)                                  |                 |                 |               |            |                     |                             |
| L-Lactate dehydrogenase B chain (P16125)   | TVEGP           | 1.0 (12)        |               |            |                     |                             |
| Phosphoglucomutase-1 (O90089)              |                 |                 |               |            |                     |                             |
| (P35486)                                  |                 |                 |               |            |                     |                             |
| Phosphoglycerate kinase 1 (P09411)         |                 |                 |               |            |                     |                             |
| Pyruvate dehydrogenase E1 component subunit α (P35486) |                 | 1.0 (12)        |               |            |                     |                             |
| Pyruvate dehydrogenase E1 component subunit β (Q90051) | PEBVTR          | 1.0 (12)        |               |            |                     |                             |
| Triose-phosphate isomerase (P17751)        |                 |                 |               |            |                     |                             |
| Tricarboxylic acid cycle                   |                 |                 |               |            |                     |                             |
| Citrate synthase (O9Cu6)                   |                 |                 |               |            |                     |                             |
| Isocitrate dehydrogenase (P54071)          |                 |                 |               |            |                     |                             |
| Malate dehydrogenase, mitochondrial (P0249) |                 |                 |               |            |                     |                             |
| **Fatty acid metabolism**                  |                 |                 |               |            |                     |                             |
| Carnitine O-palmitoyltransferase I (Q924X2) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Δ<sub>9</sub>-Δ<sub>9</sub>-Dienoyl-CoA isomerase (O35495) |                 | 1.0 (12)        |               |            |                     |                             |
| Enoyl-CoA hydratase (Q8B895)               |                 |                 |               |            |                     |                             |
| Fatty acid-binding protein (P04117)        |                 |                 |               |            |                     |                             |
| Long-chain-specific acyl-CoA dehydrogenase (P51174) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Long-chain-fatty-acid-CoA ligase 1 (P14216) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Very-long-chain-specific acyl-CoA dehydrogenase (P58545) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| **Oxidative phosphorylation**              |                 |                 |               |            |                     |                             |
| ATP synthase ε chain (P56382)              | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Electron transfer flavoprotein-ubiquinone oxidoreductase (Q921G7) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| NADH dehydrogenase α subcomplex subunit 10 (O96981) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| NADH dehydrogenase protein 2 (Q91905)      | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| NADH dehydrogenase protein 6 (P52503)      | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Succinate dehydrogenase flavoprotein subunit (O9Cu6) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| **Other metabolic proteins**               |                 |                 |               |            |                     |                             |
| 2-Oxoglutarate dehydrogenase E1 component (P65697) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Aldose reductase (P45375)                  | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Aspartate aminotransferase (P65202)        | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Creatine kinase M-type (P07310)            | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Creatine kinase, sarcomeric mitochondrial (Q6P87J) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Fumarilacetate hydrolase (Q8B898)          | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Malate dehydrogenase, cytoplasmic (P14152) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Nucleoside-diphosphate kinase B (Q01668)   | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| **Muscle and structural proteins**         |                 |                 |               |            |                     |                             |
| Actin, α cardiac muscle 1 (P68033)         | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Filamin-C (Q8VHH6)                        | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
| Four and a half LM domains protein 2 (O74033) | PLLVTR          | 1.0 (12)        |               |            |                     |                             |
be directly reduced by Trx1 (Fig. 2). First the fragmentations of a heavy ICAT-labeled peptide (m/z 1432.72) generated an MS/MS ion profile that matched to ANT1 153EFNGLGD/-pyridyldithio)-propionamide, immuno precipitated with an anti-ANT1 antibody, and then blotted with different antibodies. We observed that ANT1 protein levels were constant across all four samples (Fig. 2, C and E).

Western blot analysis with biotin showed that only trace levels of biotinylated ANT1 (a marker for free thiol levels) was detected in H2O2-treated samples, indicating that most of the ANT1 cysteines were oxidized (Fig. 2D, lane 1). Trx1 treatment led to a ~30% increase in free thiols (Fig. 2D, lane 2). Treatment of the cell lysate with the Trx1/TrxR/NADPH reduction system increased protein thiols by ~130% (Fig. 2D, lane 3), suggesting that Trx1 regeneration was important to its reduction of ANT1. Lastly a strong reductant, TCEP, restored ~210% of the oxidized cysteines to the reduced states (Fig. 2D, lane 4). This analysis suggested that ANT1 contained redox-sensitive protein thiols that were sensitive to 200 μM H2O2 oxidation, and these oxidized cysteines could in turn be reduced by TCEP nonspecifically reduced most of the redox-sensitive protein thiols that were sensitive to 200 μM H2O2 oxidation, and these oxidized cysteines could in turn be reduced by TCEP.

### Table I—continued

| Protein name (Swiss-Prot accession number) | Peptide sequence | ICAT ratio (H/L) | Avg. ICAT ratio | Protein TRX targets (Refs.) |
|------------------------------------------|-----------------|----------------|---------------|---------------------------|
| Myosin light polypeptide 3 (P09542)      | 181LMAGQEDSNQGNYEAFVK199 | 1.5 1.4 1.1 0.033 | 1.3 1.0 (13) |
| Myosin-6 (Q02566)                        | 202TEGFPVDKEREVK | 2.0 2.5 4.2 0.016 | 2.9 1.0 (83) |
| Myosin-binding protein C (O70468)        | 129ATNLQGCAOCCER1563 | 1.4 1.8 1.8 0.013 | 1.7 0.8 (41) |
| Obscurin (A2AAJ9)                        | 561CEVSNDENPR790 | 1.7 3.0 1.7 0.026 | 21.1 |
| Tropomyosin α-1 chain (P58771)           | 712LCETEGR290 | 1.5 1.8 1.7 0.005 | 1.79 |
| Transport and channel protein             | 926ADAGEYSCAEQGQK451 | 3.0 3.9 2.6 0.033 | 3.2 0.9 (10) |
| ADP/ATP translocase 1 (P48962)           | 297CGOAGGASSAETVEYRN1688 | 1.6 2.6 2.3 0.016 | 2.22 |
| Chloride intracellular channel protein 4 (Q02QY1B) | 196CAEELLEK458 | 1.5 2.8 1.5 0.043 | 1.9 0.9 (16) |
| Sarcoplasmic/endoplasmic reticulum calcium ATPase 3 (Q64518) | 1578EFNLGQLCLTK163 | 2.0 3.5 2.1 0.014 | 2.5 1.0 (14) |
| Other proteome                          | 220DEFTNTPSD238 | 1.7 1.7 2.3 0.011 | 1.9 1.0 (3) |
| Rab GDP dissociation inhibitor β (Q61598) | 433VGEATETALLC361 | 1.6 1.6 1.5 0.001 | 1.6 N/A |

a C represents an ICAT-labeled cysteine, C* indicates a carbamidomethylated cysteine; M indicates an oxidized methionine.

b One-tailed Student’s t test was performed on the percentages of free thiol levels (converted from the ICAT ratios (R) with equations 1/(1 + R) × 100% for control animals and R/(1 + R) × 100% for Tg-Trx1 animals).

c Average.

d Numbers in parentheses indicate the number of unique peptides used for iTRAQ quantification calculation. N/A indicates that the protein was not found in the iTRAQ analysis.
sponded to the addition of a cleaved heavy ICAT label to Cys53. To confirm that the reduction of this specific cysteine was Trx1-dependent, we simulated oxidative stress by treating recombinant human DJ-1 protein with H2O2 (200 μM) for 30 min followed by removal of excess H2O2 with catalase (at a final concentration of 0.1 g/ml). The oxidized DJ-1 samples

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A

Mouse DJ-1

Heavy ICAT labeled cysteine

y13 y12 y11 y10 y9 y7 y6 y5 y4

B

Human DJ-1

Heavy ICAT labeled cysteine

y13 y12 y11 y10 y9 y8 y7 y6 y5 y4

MCP
were then incubated with either the activated Trx1/TrxR/NADPH system or blank reaction buffer for 20 min at room temperature and subjected to ICAT labeling. We observed only trace levels of available protein thiols labeled with the light ICAT reagent; however, the Trx1/TrxR/NADPH system reduced a significant amount of the oxidized Cys\textsuperscript{S3} in peptide \textsuperscript{49}DVICPDASLED\textsuperscript{62} (Fig. 3\textit{B}, inset). Although the tryptic peptide sequences are slightly different between mouse and human DJ-1, Cys\textsuperscript{S3} is highly conserved in both species. In conclusion, both \textit{in vivo} and \textit{in vitro} experiments confirmed that Cys\textsuperscript{S3} is a redox-sensitive cysteine, and its redox state can be regulated by the Trx1 reduction system.

\textit{Identification of Potential Trx1-interactive Sequence Motifs}—We also examined all the putative Trx1 target sequences found in this study for the possible presence of consensus sequence motifs. Interestingly we discovered two potential CXXC motifs (where either one or both cysteines could be reduced by Trx1) resembling the catalytic site of Trx1 (supplementary Fig. 2). Interestingly three peptides from this study (NADH dehydrogenase protein 6 (\textsuperscript{101}TGTCGYS\textsuperscript{C}GLOFK\textsuperscript{112} where \textsuperscript{C} represents an ICAT-labeled cysteine), DnajA3 (\textsuperscript{282}GS\textsuperscript{293}ITNPC\textsuperscript{C}VGR\textsuperscript{295}), and four and a half LIM domains protein 2 (\textsuperscript{157}Q\textsuperscript{166}ALQ\textsuperscript{C}V\textsuperscript{168}CK\textsuperscript{169}) contained a CXXC motif where both cysteines were more reduced in Tg-Trx1 animals (Table I). Four and a half LIM domains protein 2 also had two additional peptides (\textsuperscript{84}EEQLL\textsuperscript{C}TDY\textsuperscript{C}SNEYSSK\textsuperscript{100} and \textsuperscript{148}EN\textsuperscript{149}NC\textsuperscript{C}TPYE\textsuperscript{C}K\textsuperscript{156}) where \textsuperscript{C} indicates a carbamidomethylated cysteine) matched to the CXXC motif. However, because of the limited sample size reported here, future studies including proteomics and molecular modeling approaches may be needed to validate the predictive value of such motifs.

\section*{DISCUSSION}

\textit{Heart Oxidative Stress and Trx1 Function}—Understanding the components of cellular antistress pathways may provide clues on how to develop specific therapeutic agents for stress-related heart diseases. Trx1 and the affiliated redox regulatory system are important mechanism by which cardiac myocytes counter the detrimental effects of oxidative stress. Our earlier studies suggested that Trx1 is protective in cardiac myocytes and elevated ATP content in Tg-Trx1 hearts (21). In this study, through a comparative ICAT proteomics study of Tg-Trx1 and control animals, we identified 55 potential Trx1 reduction targets in the heart, including several previously reported Trx1 targets (Table I), which validated our approach. In addition, we discovered novel Trx1 reduction protein networks for maintaining energy conductance from mitochondria to the cardiac contractile apparatus, which will be discussed below (Fig. 4).

\textit{Comparison of ICAT and iTRAQ Analysis Results}—The importance of Trx1 has spurred enormous interest in identifying its targets, an endeavor that has been aided by the developments of innovative techniques such as fluorescent diagonal electrophoresis (5) and affinity chromatography (28, 29). However, these approaches have not been able to provide quantitative information regarding the degree or the specific sites of reduction; both are critical for relating protein redox status and the associated biological functions. ICAT is a cysteine-specific isotope labeling method for peptide quantification at the MS levels, whereas iTRAQ reagents label primary amines for isobaric peptide quantification at the MS/MS levels. Both methods have been widely used for large scale quantification of proteins (9, 10, 13, 25). To capture cysteine thiol redox status, the reduction/alkylation steps were omitted for this ICAT analysis. This strategy allowed the detection of the increase of specific protein thiols in Tg-Trx1 animals. The increase of the ICAT ratios could be due to Trx1-mediated gene/protein expression differences, protein reduction, or a combination of these elements. To differentiate these possibilities, we compared the ICAT proteomics results with those obtained from the iTRAQ expression analysis in which the TCEP reduction/MMTS alkylation steps were done to determine the relative protein expression levels of the same samples. Interestingly the expression levels of the 55 significant Trx1 reduction targets (Table I) were not significantly changed in the iTRAQ study, indicating that the increase in cysteine thiol levels in Trx1 mouse hearts was largely due to Trx1-mediated reduction of redox-sensitive thiols.

There were also indications that some proteins (peptides) are more oxidized in Tg-Trx1 animals (catalase (ICAT ratio of 0.48) and ubiquinol-cytochrome-c reductase complex core protein 1 (ICAT ratio of 0.65); supplemental Table 1). However, these proteins did not satisfy our stringent criteria for target selection (e.g. observed in all three independent experiments). The number of oxidized proteins in Tg-Trx1 animals was...
relatively small, and they are included in supplemental Table 1. We hypothesize that the reason why some proteins were oxidized more significantly in Tg-Trx1 animals may be due to Trx1-induced activation of oxidoreductases (Ero1/H2O2 (30), for example) that can in turn oxidize downstream targets. Validation of Trx1 Targets—Many of the proteins revealed from this study have been reported previously to be direct Trx1 reduction targets (Table I), validating the reliability of the current approach. However, Trx1 may also contribute to the reduction of oxidized cysteines indirectly, e.g. via the peroxiredoxins (31) or glutathione system (32). We conducted additional experiments to validate selected proteins as direct Trx1 reduction targets. One was ANT1, a key component of the mitochondrial permeability transition pore complex, which is a multiprotein complex containing ANT1, voltage-dependent anion channel, cyclophilin-D, creatine kinase, and Bcl/Bax family proteins (33). The MPTP exerts its function as a mitochondrial intermembrane conduit for ATP/ADP exchange to supply the high energy phosphates for muscle contraction. Dysregulation of MPTP is characterized by the depolarization of the mitochondrial inner membrane, disruption of the crossmembrane potential, mitochondrial swelling, and eventual release of mitochondrial apoptogens, including cytochrome c (34, 35). ANT1 has been reported to be vulnerable to oxidative modifications leading to aberrant MPTP activity that results in reduced ATP production and utilization and increased apoptosis (35). Mice deficient in ANT1 exhibit severe cardiomyopathy (36). There are four cysteines within mouse ANT1, several of which have been shown to be prone to oxidation (37, 38). Oxidation of Cys159 in human ANT1 (corresponding to the conserved Cys160 in mouse) within the binding site of adenine nucleotides has been shown to be detrimental to pore function and nucleotide transport (38). In this study, although ANT1 protein levels were comparable for all animals, Cys160 was found to be more reduced in Tg-Trx1 than wild type hearts (Tg-Trx1/control ratio of 2.5; Table I and Fig. 2, A and B). We also found that ANT1 was oxidized by H2O2 (Fig. 2D, lane 1), and plain Trx1 was able to partially reverse the

![Cardiac Trx1 Targets](image-url)
oxidation of free thiols (Fig. 2, C and D, lane 2). The reduction effect was more pronounced when TrxR and NADPH were included for Trx1 regeneration (Fig. 2, C and D, lane 3), suggesting that ANT1 could be a direct target of Trx1 reduction. More importantly, Cys160 was likely to be a reactive cysteine within ANT1 that was reduced by Trx1 given that TCEP was more thorough at potentially reducing all oxidized cysteine thiols nonspecifically in ANT1 than was Trx1 (Fig. 2, C and D, lane 4). In this study, we also observed that other MPTP components were modulated by Trx1, including creatine kinase, voltage-dependent anion channel 1, and cyclophilin-D (supplemental Table 1), suggesting MPTP may be directly regulated by Trx1 in a redox-dependent manner to sustain the supply of ATP for the heart under TAC-induced oxidative stress.

In addition to validating Trx1 targets at the protein levels, we also performed experiments to confirm that such regulation is site-specific. DJ-1 is a redox-sensitive protein associated with the development of Parkinson disease (39). Although it is expressed in the heart, its cardiac function is largely unexplored. DJ-1 deficiency in neuronal cells has been shown to sensitize cells to oxidative stress and may result in cell death (40); overexpression of DJ-1 is cytoprotective (41). There are four cysteines in mouse DJ-1, several of which have been shown to be very sensitive to oxidative stress (39, 42). No reports have shown that Trx1 is able to repair DJ-1 oxidative damages. Our in vitro study showed that Cys33, a highly conserved residue among mammalian species, may be a Trx1 reductive target (Table I and Fig. 3A); this is significant given previous reports of cysteinylations on Cys33 upon oxidative stress (39). In a follow-up experiment, we were able to show that Cys33 in human DJ-1 is sensitive to H2O2 oxidation (Fig. 3B), and its oxidation can be almost completely reduced by direct interaction with Trx1 (Fig. 3B).

**Trx1 Targets within Energy Pathways**—Not much is known of the impact of Trx1 on metabolic protein functions in mammalian cells, especially in heart tissues, where the demand for metabolic fuel is high. Typically when diseased hearts become stressed, they become adaptively hypertrophied, exhibiting significant metabolic dysfunction (43). For example, the disruption of β-oxidation of fatty acids has been reported in hypertrophied hearts, resulting in both the reduction of acetyl-CoA supply for the tricarboxylic acid cycle and accumulation of toxic lipid derivatives (44). Deficiency of very-long-chain-acyl-CoA dehydrogenase is known to induce heart hypertrophy (45). Here we found that both long-chain- and very-long-chain-acyl-CoA dehydrogenases are potential Trx1 reduction targets (Table I), suggesting that within Tg-Trx1 animals the lipid metabolic profiles may be protected from oxidative stress to maintain proper heart functions (Fig. 4).

Creatine kinase plays an important role in shuttling ATP from the mitochondria to myofibrils, supporting robust heart muscle contraction. There are two forms of creatine kinases in cardiomyocytes, cytosolic creatine kinase and muscular creatine kinase, which work in tandem to transfer the high energy phosphates from ATP to the contractile apparatus (Fig. 4). Our ICAT analysis identified that both cytosolic creatine kinase and muscular creatine kinase are potential substrates for Trx1 reduction. Cys317 in the 311LGYILTCPSNLGTLGR326 peptide within the catalytic site of cytosolic creatine kinase has been shown to be sensitive to oxidation, resulting in the inhibition of its enzymatic activity (12, 46). About 60% more free thiols in this peptide were found in Tg-Trx1, suggesting that Trx1 may help to maintain the creatine kinase activity, facilitating energy conductance from mitochondria to the heart contractile machinery.

Oxidation of cardiac contractile proteins has been shown to impair the plasticity and contractility of muscles and could be reduced by antioxidants in vitro (47, 48). We observed that a wide spectrum of sarcomeric proteins were reduced by Trx1 overexpression (Table I), suggesting Trx1 may play a direct role in maintaining cardiac contractile apparatus functionality and energy utilization (Fig. 4).

**Limitation of the Current Redox ICAT Method**—Although the redox ICAT method is effective for finding potential Trx1 target proteins, many well characterized Trx1 targets such as peroxiredoxin 1 (Prx1), Prx2, and ribonucleotide reductase were missing from this study. There are several potential limitations of the current method for redox proteomics studies. 1) The peptides containing the Trx1-reactive cysteines may not be suitable for this analytical work flow. For example, the Prx1 tryptic peptide 53VYFFYPLDFTVCPTEIIAFSDFR62 (an underlined C represents a known Trx1 reduction target cysteine) contains a known Trx1 reductive site at Cys52 (49). This peptide is highly hydrophobic and may not be eluted from the RPLC column. Furthermore, its mass of m/z 3376.5 may be too large to be fragmented efficiently by MALDI tandem MS. A similar issue may also be applicable to the Prx2 tryptic peptide 35VVYFLFYPLDFTVCPTEIIAFSDFHAEDFR56. On the other hand, the Trx1 reductive site in peroxiredoxin 5, 51GVLFGVGAFTPGCSK58, was found in this study because this tryptic peptide was amenable to the LC/MS/MS analysis. 2) Tryptic peptides containing the Trx1 reductive sites from proteins including ribonucleotide reductase are amenable for LC/MS/MS analysis but may be relatively low in abundance to be detected. 3) Some peptides may show significant ICAT ratio changes in MS levels, but their MS/MS fragmentation may be poor due to their amino acid compositions or the attachment of a bulky ICAT tag. 4) Because the protein disulfide bond reduction step was omitted from this redox ICAT study, it is possible that some of the free cysteines buried inside non-linearized protein domains may not be labeled efficiently by the bulky ICAT reagents. 5) Redox-induced post-translational modifications on non-cysteine residues, such as phosphorylation, may occur non-uniformly in light and heavy ICAT-labeled peptides. Therefore quantification accuracy of ICAT may be affected. 6) Some proteins bound to Trx1 with high affinity may still be reduced in the control animals, leading...
Cardiac Trx1 Targets

...ing to similar reduction levels compared with Tg-Trx1 animals regardless of the different Trx1 expression levels between the animal groups. Therefore, they may not be found in this ICAT difference-based study. Alternative approaches, including “hooking” to C3SS Trx1 mutant, may be used to identify some of these Trx1 targets (7, 8).

Conclusions—In the present study, we performed a proteomics identification of Trx1 reduction target proteins from the hearts of cardiac specific Tg-Trx1 mice. Using the comparative analysis of both iTRAQ and ICAT results, we were able to reveal many putative Trx1 reduction substrates, several of which were previously unknown. We identified several protein networks whose functions may be regulated by Trx1, including the creatine-phosphocreatine shuttle, the MPTP complex, and the sarcomeric contractile apparatus.

The results presented here suggest that in addition to its antioxidant function Trx1 may be involved in the coordination of a wide array of cellular signaling pathways to maintain cardiac function.

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