INTRODUCTION

Exposing plants to unfavorable environmental conditions such as low or elevated temperature, ultraviolet (UV) irradiation, or salt stress can induce the production of reactive oxygen species (ROS) such as the superoxide radical, hydrogen peroxide, and the hydroxyl radical. ROS cause serious damage to biological macromolecules. To protect against these toxic oxygen intermediates, plants use various enzymatic and non-enzymatic defense mechanisms that prevent or repair oxidative damage [1].

Peroxiredoxins (Prxs) constitute the most recently identified group of H\textsubscript{2}O\textsubscript{2}-decomposing antioxidant enzymes. Prxs are present in all organisms, from prokaryotes to eukaryotes, and they catalyze the reduction of various hydroperoxides into the corresponding alcohol or water [2-4]. Prxs are thiol-based peroxidases that possess at least one conserved cysteine in their N-terminus that reacts with peroxide substrates [5]. In addition to the reduction of H\textsubscript{2}O\textsubscript{2}, Prx proteins also detoxify alkyl hydroperoxides and peroxinitrite, despite the fact that significant differences exist in substrate specificity and kinetic properties [2]. Experimental evidence for Prx function in plants has revealed multiple mechanisms: it can serve as an antioxidant, a modulator of cell signaling pathways, and as a redox sensor [2]. Four Prx subgroups have been described in plants, and they are distinguished by the number and position of the conserved cysteine residues: 1-Cys Prx, 2-Cys Prx, type II Prx, and Prx Q [2,6]. 1-Cys Prx is a dormancy-related protein [7] that is localized in the nucleus, where it helps to protect macromolecules from oxidative damage. 2-Cys Prx is a...
nuclear-encoded protein targeted to the chloroplast, where it protects the photosynthetic membrane from oxidative damage [8]. Type II Prx was isolated as a target of cytoplasmic thioredoxin-h, and displays antioxidant activity in Arabidopsis thaliana [9]. Finally, Prx Q was identified initially in Sedum lineare as a homolog of the Escherichia coli bacterioferritin co-migratory protein [3]. Among the four groups, the type II Prxs are the least studied with respect to peroxidase activity, although it has recently been observed in yeast [9,10], Arabidopsis [6,9,11], poplar [4], and Chinese cabbage [12]. At least in Arabidopsis, the type II Prx is complex and composed of several different members. Six Prx II genes have been identified in Arabidopsis [6,11], one of them seems to be a pseudogene; the others are expressed in leaves. Three Prxs localized to the cytosol (AtPrx II B, C, and D), one to the chloroplast (AtPrx II E), and one to the mitochondria (AtPrx II F) [6,13,14].

Although the molecular and biochemical characterizations of Prx genes in plants are advancing rapidly, the functional role of the type II Prx remains poorly defined. To the best of our knowledge, no research to date has examined Prx in ginseng. Korean ginseng (Panax ginseng [P. ginseng] C. A. Meyer) is a perennial herb in the family Araliaceae. It has been cultivated for its highly valued root, which is used for medicinal purposes [15]. Despite its medicinal importance, little genomic analysis has been carried out on this plant. In the present study, the isolation and characterization of a peroxiredoxin cDNA isolated from P. ginseng, as well as its possible role in resisting various abiotic stresses, is reported.

MATERIALS AND METHODS

Plant materials and growth conditions

The specific type of ginseng used was P. ginseng cv. “Hwang-Sook (yellow berry variant)” (Ginseng Genetic Resource Bank, Suwon, Korea). Cultured 3-week-old plantlets were used for treatments and nucleic acid extractions, as described previously [16]. For analysis of gene expression in different organs, samples were collected from leaves, roots, and stems of 6-week-old P. ginseng plantlets. Plants were grown in a growth room at 25°C with a 16 h photoperiod.

RNA purification and construction of a cDNA library

Total RNA was isolated from leaves of 4-year-old ginseng plants, using the aqueous phenol extraction procedure described by Morris et al. [17]. Poly (A)+ RNA was isolated using an oligo (dT) cellulose column and the Poly (A) Quick mRNA isolation kit (Stratagene, La Jolla, CA, USA). A commercial cDNA synthesis kit was used to construct a cDNA library according to the manufacturer’s instructions (Clontech, Palo Alto, CA, USA). Size-selected cDNA was ligated into the λ TriplEx2 vector and packaged in vitro using a Gigapack III Gold Packaging Extract kit (Stratagene).

Nucleotide sequencing and sequence analysis

Homologous sequences of Prx expressed sequence tags (EST) were compared against the GenBank databases using a BLASTX algorithm. Nucleotide and amino acid sequence analyses were performed using the DNASIS program (Hitachi Software Engineering America Ltd., CA, USA). Deduced amino acid sequences were used to search for homologous proteins in the databases using the NCBI BLAST search engine. We used ClustalX with default gap penalties to perform multiple alignment of Prx isolated from ginseng with previously registered homologs in other species. Based on this alignment, a phylogenetic tree was constructed according to the neighbor-joining method using the MEGA3 program. Bootstrap analysis with 1,000 replicates was also conducted to obtain confidence levels for the branches. Secondary structure was determined by SOPMA program, and the hydropathy pattern was analyzed in a manner described by Kyte and Doolittle [18]. Protein properties were estimated using ProtParam [19].

Abiotic stress treatment

For chemical stress or plant hormone treatments, the plantlets were placed in Murashige and Skoog medium containing 100 mM NaCl for various periods. Chilling stress was applied by exposing the plantlets to a 4°C environment. For the UV treatment, the plantlets were irradiated under UV lamps at 1.35 μE m⁻² s⁻¹ (below 280 nm). Control and stress-exposed plant materials were immediately frozen in liquid nitrogen and stored at -70°C until used.

Real-time quantitative RT-PCR (qRT-PCR)

Total RNA was extracted from plantlets of P. ginseng using the RNeasy mini kit (Qiagen, Valencia, CA, USA). To perform RT-PCR, 5 μg total RNA was used as a template for reverse transcription using oligo(dT)₅ primer (0.2 mM) and AMV Reverse Transcriptase (10 U/μl; INTRON Biotechnology, Seongnam, Korea) according to the manufacturer’s instructions. qRT-PCR was performed using 10 ng cDNA in a 10 μL reaction volume with SYBR® Green SensimixPlus Master Mix (Quantrace, Watford, England). Specific primers for PgPrx, S'-TTCTAGCTGACGGATCAGG-3' and

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5’-GAATTCCCAACCAGTCTCAA-3’, were used. The thermal cycler conditions recommended by the manufacturer were used as follows: 10 min at 95°C, followed by 40 cycles of 95°C for 10 s, 58°C for 10 s, and 72°C for 20 s. The fluorescent product was detected in the last step of each cycle. Amplification, detection, and data analysis were performed using a Rotor-Gene 6000 real-time rotary analyzer (Corbett Life Science, Sydney, Australia). The threshold cycle (Ct) represents the number of cycles during which the fluorescence intensity is significantly higher than the background fluorescence during the initial exponential phase of PCR amplification. To determine the relative fold differences in template abundance for each sample, the Ct value for PgPrx was first normalized to the Ct value for β-actin, and relative differences were calculated using the formula 2^ΔΔCt. Three independent experiments were performed.

RESULTS

Isolation and amino acid sequence analysis of PgPrx

From our EST analysis of a cDNA library that was prepared from the leaves of 4-year-old field-grown *P. ginseng*, we identified a cDNA clone encoding a peroxiredoxin gene. This gene is named PgPrx (*P. ginseng* Prx), and its sequence has been deposited in GenBank. The obtained *PgPrx* cDNA is 716 nucleotides long and contains a putative open reading frame (ORF) of 489 base pairs. This ORF encodes a peroxiredoxin protein with 162 amino acids, beginning at the initiation codon ATG (position 98) and ending at the stop codon TAA (position 584). The aliphatic index, which is regarded as a positive factor that increases thermostability, was calculated to be 94.51, using ProtParam [19]. In the deduced amino acid sequence of the *PgPrx* protein, the total number of negatively charged residues (aspartic+glutamic) was 22, while the total number of positively charged residues (arginine+lysine) was 18. The instability index was computed to be 32.85, which classifies the protein as stable. The molecular mass of the mature protein was calculated to be approximately 17.4 kDa, with a predicted isoelectric point (pI) of 5.37.

Homology analysis

A GenBank BlastX search revealed that the deduced amino acid of *PgPrx* shares a high degree of homology with Prx of the Citrus hybrid cv. Shiranuhi (ABL67649; 83% identity, 93% similarity), *N. nucifera* (ABN46981; 83% identity, 91% similarity), *Plantago major* (AH58634; 82% identity, 90% similarity), and *Capsicum annuum* (AAL35363.2; 77% identity, 90% similarity). The comparison also showed strong homology with the type II Prx of *Lycopersicon esculentum* (*L. esculentum*; AAP34571; 76% identity, 90% similarity) and *Xerophyta viscosa* (ACA13182; 75% identity, 90% similarity) (Fig. 1A). A phylogenetic study comparing the *PgPrx* amino acid sequence to the sequences of the previously reported four types of proteins [17] led to the conclusion that *PgPrx* is a Prx II (Fig. 1B). In addition, in BlastP searches, *PgPrx* had high score hits when compared to the Prx5 family and PrxII; indeed, PrxII is homologous with mammalian Prx5 [17].

The hydropathy pattern of *PgPrx* is similar to that of other homologous Prxs (Fig. 2A). Secondary structure analysis and molecular modeling of *PgPrx* were performed using SOMPA. Secondary structure analysis revealed that *PgPrx* consists of 48 α-helices, 15 β-turns joined by 39 extended strands, and 60 random coils. The Citrus hybrid displays almost the same secondary structure as *PgPrx*: 48 α-helices, 11 β-turns joined by 42 extended strands, and 61 random coils. The other homologous plant Prxs including PrxII of *L. esculentum* also showed a similar structure (Fig. 2B).

Organ-specific expression patterns of *PgPrx*

The expression patterns of *PgPrx* in different *P. ginseng* organs were examined using real-time qRT-PCR. *PgPrx* was highly expressed in the leaf and root, with relative low levels of expression in the stem (Fig. 3).

Temporal expression of *PgPrx* in response to abiotic stresses

To elucidate the expression patterns of *PgPrx* in response to stressors such as UV, chilling, and salt, real-time qRT-PCR analysis was performed. Fig. 4A shows the expression pattern of *PgPrx* in response to salt stress (Fig. 4A). Salinity stress (100 mM NaCl) caused a marked (8-fold) and immediate (1 h) induction of *PgPrx*; however, transcription levels then decreased to 2-fold or remained at a similar level as controls for the subsequent 72 h. UV exposure also caused a dramatic and rapid induction of *PgPrx*, inducing transcript levels to increase by 64-fold 1 h post-treatment, and reaching a maximal induction (87-fold) in 4 h; thereafter, it decreased dramatically (Fig. 4B). Under chilling stress, *PgPrx* mRNA level seemed to have been slightly suppressed initially (0.67-fold 1 h post-treatment), but then increased by 10-fold within 4 h, and continued to be higher 8 h and 24 h later (15-fold and 64-fold, respectively). However, 48 hours after treatment, *PgPrx* expression was 10-fold lower; 72 hours post-treatment, it was 7-fold lower (Fig. 4C).
DISCUSSION

Here we provide the first report of the isolation and characterization of the peroxiredoxin gene in ginseng. BlastX database searches identified Prx sequences in other plant species as the best matches. Analyses of amino acid homology, hydropathy pattern, and secondary structure revealed that PgPrx is very similar to other homologous plant Prx genes (Fig. 1). The Prx genes can be divided into four subgroups: 1-Cys Prx, 2-Cys Prx, type II Prx, and Prx Q. Database searches suggested the PgPrx belongs to the type II family. All Prx molecules
have a similar basic protein structure containing a thioredoxin fold, and their molecular masses range between 17 and 22 kDa [2]. The $PgPrx$ cDNA encodes a Prx protein of 162 amino acids, with a predicted molecular weight of 17.4 kDa and a pl value of 5.37. Similarly, cytosolic AtPrxII B and C also have masses of 17.4 kDa with pl values of 5.17 and 5.33, respectively [11]. All of the Prx isoforms have conserved-catalytic cysteine (Cys) residues in common. The absolutely conserved Cys is the catalytic or peroxidatic Cys [20,21]. Catalytic Cys residues undergo oxidation during the peroxide-reduction reaction, and need to be reduced by electron donors. The active site [6], which neighbors Cys51, is well conserved among $PgPrx$ and homologous plant Prxs, whereas the second conserved Cys (Cys 76) is only present in some plants (Fig. 1A). The second Cys is also absent from cytosolic Prx from other plants such as N. mucifera and O. sativa, supporting the fact that this Cys is not absolutely necessary for the protein’s functional activity [22]. The transcripts of $PgPrx$ were well-detected in the leaf and root of ginseng plantlets. In the case of Arabidopsis, the expression of Prx II, Prx IIB, and E is most abundantly expressed in green tissue, and Prx IIC and D are highly expressed in the flower and root, whereas Prx IIF is ubiquitously expressed [21]. This suggests that Prx genes are expressed in most organs, but with different organ specificities.

Prx proteins from organelles may prevent DNA dam-

**Fig. 2.** Superimposed hydrophilicity profiles and secondary structure predictions for peroxiredoxin. (A) Hydrophathic index analysis. Hydrophobic domains are indicated by positive numbers; hydrophilic domains are above the line and hydrophilic domains are below it. Citrus hybrid (C. hybrid) cv. Shiranuhi, Capsicum annuum (C. annuum), Plantago major (P. major), and Lycopersicon esculentu (L. esculentu). The helix, sheet, turn, and coil structures are indicated in order from the longest to the shortest. (B) Comparison of the secondary structures of $PgPrx$, C. hybrid, C. annuum, P. major, and L. esculentu by SOPMA.
A few representatives of the type II Prx have been cloned from plant sources, but knowledge on their molecular functions is limited. Mitochondrial Prx II responds to oxidative stress in *Arabidopsis* [14], plays a role in plant fungal pathogen interactions in poplar [23], and accumulates in response to cold and heavy metal stress in pea plants [24]. Among six identified *AtPrx II* genes in *Arabidopsis*, the expression of cytosolic *AtPrx IIB* and *IIC* is induced by oxidative stress and salt stress. Chloroplastic *Prx IIE* is unregulated after bacterial exposure, and its expression decreases in response to salt stress and increases in response to bright light [6,11]. Mitochondrial *AtPrx IIF* expression is constitutive and is essential for root growth under stress conditions [14]. Mitochondrial *AtPrx IIF* expression increases in response to biotic rather than abiotic stress [23]. In peas, *Prx IIF* accumulates in cold- and heavy metal-treated plants [24].

When ginseng plantlets were exposed to stresses, *PgPrx* expression changed as a function of the exposure time. *PgPrx* expression seems to be up-regulated by salt, chilling, and UV treatment at early time points. *Arabidopsis* cytosolic *Prx II* is also up-regulated by salt, unlike chloroplast and mitochondrial *Prx II* [6,11,23]. *Prx IIB* mRNA levels increase slightly after 6–8 h and *Prx IIC* mRNA levels increase dramatically within 2 h in response to 50 mM NaCl treatment in *Arabidopsis* [6]. Salinity causes oxidative stress, and salt tolerance is related to the induction of antioxidant defenses [25]. UV irradiation can cause oxidative stress by generating ROS in plants [26], which leads to oxidative damage. Chilling stress is also involved in the ROS signaling pathway [27]. These conditions trigger oxidative stress and result in the rapid up-regulation of *PgPrx* gene expression within 1-4 h of exposure. These results may indicate a particular and important role of *PgPrx* in stress defense. Few studies have considered cytosolic Prx II in plants.
so further investigations are needed to better understand its mechanisms of action. The present study suggests that Prx II plays an important role in ginseng’s defense mechanism against cold, UV, and salinity stresses.

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REFERENCES