Detection of the Inflammasome in the Transplanted Heart
A Chloroplast-Localized Rubredoxin Family Protein Gene from *Puccinellia tenuiflora* (*PutRUB*) Increases NaCl and NaHCO₃ Tolerance by Decreasing H₂O₂ Accumulation

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Abstract: Rubredoxin is one of the simplest iron–sulfur (Fe–S) proteins. It is found primarily in strict anaerobic bacteria and acts as a mediator of electron transfer participation in different biochemical reactions. The *PutRUB* gene encoding a chloroplast-localized rubredoxin family protein was screened from a yeast full-length cDNA library of *Puccinellia tenuiflora* under NaCl and NaHCO₃ stress. We found that *PutRUB* expression was induced by abiotic stresses such as NaCl, NaHCO₃, CuCl₂ and H₂O₂. These findings suggested that *PutRUB* might be involved in plant responses to adversity. In order to study the function of this gene, we analyzed the phenotypic and physiological characteristics of *PutRUB* transgenic plants treated with NaCl and NaHCO₃. The results showed that *PutRUB* overexpression inhibited H₂O₂ accumulation, and enhanced transgenic plant adaptability to NaCl and NaHCO₃ stresses. This indicated *PutRUB* might be involved in maintaining normal electron transfer to reduce reactive oxygen species (ROS) accumulation.

Keywords: rubredoxin family protein; *Puccinellia tenuiflora*; ROS

1. Introduction

Nowadays, soil alkalinization is very severe. Salinity and alkalinity are major environmental stresses on soil that limit crop distributions and yields worldwide [1-3]. In Northeast China, the total area of alkaline saline soil is about 3.73 × 10⁶ ha and is expanding at a rate of 1.4% annually, making it one of the three largest sodic-saline areas in the world [4,5]. In such soil, plants suffer adverse stress factors, such as Na⁺ and HCO₃⁻/CO₃²⁻ ions, and high pH. These factors can directly affect nutrient uptake, organic acid balance, and ion homeostasis at a whole-plant level [6-9]. Research shows that only a small number of plants can complete their entire life cycle in highly alkaline areas. *Puccinellia tenuiflora*, a graminaceous and alkali-tolerant halophyte species, is one such plant. It can complete its life cycle in highly alkaline soil (pH 10). *P. tenuiflora* has evolved various strategies to adapt to saline or alkaline stress, such as maintaining ion balance, osmotic homeostasis adjustment, and removal of reactive oxygen species (ROS) [10–15]. Thus, it is considered a saline-alkali soil pioneer plant. To date, several stress-responsive genes in *P. tenuiflora* have been isolated and their biological functions have been tested [16–21]. *P. tenuiflora* is also considered an ideal model plant for discovering resistance genes and studying the resistance mechanism of halophytes.
Rubredoxin is a kind of non-heme iron protein that contains a \([\text{Fe(SCys)}_4]\) center. It is one of the simplest iron–sulfur (Fe–S) proteins, containing a single Fe atom and four S atoms [22]. Rubredoxin is mostly found in strict anaerobic bacteria, including bacteria, archaeabacteria and a few microaerobic bacteria in nature [23–27]. Relatively few of these proteins have been found in plants. Research suggests rubredoxin proteins act as mediators of electron transfer for various enzymes in anaerobic bacteria [24,28–30]. Recently, a new pathway called the Super Oxide Reductase reaction (SOR) was identified that provides an alternative superoxide reduction pathway in anaerobic microorganisms. Rubredoxins of various anaerobes appear to act as an electron donor in the SOR reaction to reduce the production of \(O_2\) [31–33].

In our previous work, we identified some candidate salt-responsive genes in \(P.\) tenuiflora using the Full-length cDNA Over-eXpressor gene (FOX)-hunting system. This system is a very effective tool for plant functional gene research that does not require any knowledge of the genome of interest or genetic mapping. It has opened up novel approaches for elucidating the functions of genes that control metabolic pathways and determine plant morphological characteristics [34,35]. \(PutRUB\) was one of the genes we identified. In this study, we analyzed its expression under different stress conditions. The function of \(PutRUB\) in NaCl and NaHCO\(_3\)-stress conditions was investigated in Arabidopsis transgenic the \(PutRUB\) gene. We hope these results will provide a theoretical basis for future exploration of the adaptation mechanism of \(P.\) tenuiflora.

2. Results and Discussion

2.1. Characteristics of \(PutRUB\)

The full-length cDNA of \(PutRUB\) was 1005 bp, with a 5′-UTR of 52 bp, an open reading frame (ORF) of 774 bp, and a 3′-UTR of 179 bp. From the ORF, we deduced that \(PutRUB\) encoded a 225-amino acid protein with a predicted molecular mass of 27.38 kDa and a theoretical pI of 9.85. The sequence contained an N-terminal signaling peptide with the most likely cleavage site between positions 22 and 23 (SHC-AD).

To study the phylogenetic relationships of \(PutRUB\) and other candidate rubredoxin proteins, we selected 31 complete protein sequences from 19 species to construct a phylogenetic tree. These included 13 candidate rubredoxin protein sequences from monocotyledons (3 from \(Zea\) \(mays\): NP_001183375, AFW57239, AFW57238; 1 from \(Triticum\) \(urartu\): EMS68403; 2 from \(Hordeum\) \(vulgare\) subsp. \(vulgare\): BAJ94117, BAK07366; 3 from \(Brachypodium\) \(distachyon\): XP_003573446, XP_010234369, XP_014756405; 2 from \(Oryza\) \(sativa\) Japonica Group: XP_015648775, XP_015615806; 2 from \(Zostera\) \(marina\): KMZ75006, KMZ59555), 15 candidate proteins from other plants (2 from \(Medicago\) \(truncatula\): XP_003610879, XP_013453352; 3 from \(Arabidopsis\) \(thaliana\): NP_568342, NP_001078598, NP_568749; 2 from \(Glycine\) \(soja\): KHN35119, KHN25601; 1 from \(Glycine\) \(max\): NP_001235582; 1 from \(Eutrema\) \(salsugineum\): XP_006400247; 1 from \(Arabidopsis\) \(lyrata\) subsp. \(lyrata\): XP_002871751; 2 from \(Theobroma\) \(cacao\): XP_007029491, XP_007038169; 1 from \(Phaseolus\) \(vulgaris\): AGV54433; 1 from \(Populus\) \(trichocarpa\): XP_002321763; 1 from \(Arabis\) \(alpina\): KFK26723) and two from green algae (\(Synechococcus\) sp. PCC 7002: AAL78082, \(Ostreococcus\) \(tauri\): XP_003080003) (Figure 1a). The results showed that \(PutRUB\) had the closest phylogenetic relationship with \(Z.\) \(mays\) NP_001183375, and had close relationships with most monocotyledonal plants.

Alignment of the amino acid sequence of PutRUB with sequences from \(T.\) \(urartu\) (EMS68403), \(H.\) \(vulgare\) subsp. \(vulgare\) (BAJ94117), \(O.\) \(sativa\) Japonica Group XP_015648775, \(Z.\) \(mays\) (NP_001183375), \(A.\) \(thaliana\) (AED92391.1) and \(G.\) \(soja\) (KHN35119) (Figure 1b) suggested PutRUB possessed characteristics common to rubredoxin family proteins in these plants and contained two domains. In the N-terminal region, there is a putative PDZ domain; in the C-terminus, there is a rubredoxin domain. The PDZ domain is potentially involved in interactions of protein to protein [36,37]. The conserved regions of PutRUB and the phylogenetic tree suggested that \(PutRUB\) belonged to the rubredoxin family, but its function have yet to be elucidated.
Figure 1. Phylogenetic analysis of PutRUB. (a) Amino acid sequence of PutRUB and phylogenetic trees analysis of 30 rubredoxin family protein sequences in plants; (b) The analysis of the amino acid sequence of PutRUB with rubredoxin family protein in Triticum urartu, Hordeum vulgare, Oryza sativa Japonica Group, zea mays, Arabidopsis thaliana and Glycine soja.
2.2. Subcellular Localization of the PutRUB: GFP Fusion Protein

To determine the accurate subcellular localization of the PutRUB protein in plant cells, we used transgenic plants containing pBI121-PutRUB-GFP. The GFP signal was stably accumulated in the chloroplast (Figure 2).

![Figure 2](image)

**Figure 2.** Subcellular localization of PutRUB in Arabidopsis protoplasts. (a) PutRUB: GFP within Arabidopsis protoplasts; (b) GFP within Arabidopsis protoplasts. The protoplasts were stained with MitoTracker Red. Scale bar = 10 µm.

2.3. Expression of the PutRUB Gene Is Induced by Abiotic Stresses

The steady-state mRNA levels of PutRUB in different tissues and under different stresses (NaCl, NaHCO₃, CuCl₂ and H₂O₂ treatments) were assayed with Semi-quantitative Polymerase Chain Reaction (RT-PCR) (Figure 3). PutRUB was accumulated in all tissues that we studied (Figure 3a). The highest expression levels were in leaves, while the lowest levels were in roots, indicating that PutRUB expression has tissue specificity. PutRUB was localized in chloroplasts, similarly to a rubredoxin in Synechococcus sp. PCC 7002, and had the highest expression in leaves. The Synechococcus sp. PCC 7002 rubredoxin is localized in chloroplasts and was confirmed to be involved in the building of the interpolypeptide (4Fe–4S) cluster Fx of PSI [38,39]. The above results suggested that PutRUB expression may be associated with photosynthesis.

Previous studies used the PutRUB gene to enhance the resistance of yeast to biotic and abiotic stresses [34]. In this paper, we analyzed the pattern of expression of the PutRUB gene under different abiotic stresses (Figure 3a). All of the stresses induced PutRUB expression, but the highest expression was observed under NaCl and NaHCO₃ stresses. Subsequently, the expression of PutRUB was monitored over a time course under NaCl, NaHCO₃, CuCl₂ and H₂O₂ stresses (Figure 3b). Figure 3b shows that PutRUB was up-regulated in both roots and leaves by exposure to 200 mM NaCl, 200 mM NaHCO₃, 150 µM CuCl₂ and 6 mM H₂O₂, indicating that PutRUB may be involved in responses to these stresses.

2.4. Response to NaCl and NaHCO₃ Stress in PutRUB Transgenic Plants

To analyze the function of PutRUB under NaCl and NaHCO₃ stresses, we constructed transgenic Arabidopsis plants overexpressing PutRUB under the control of the strong constitutive CaMV 35S promoter. Three independent T₁ and T₃ generation transgenic A. thaliana lines were identified by PCR amplification and northern blot analysis (Figure 4). The results showed that each of these lines expressed PutRUB. Thus, independent transgenic plants overexpressing PutRUB (#1, #2, #3) were used for assays of the root length and fresh weight.
We analyze the phenotypes (fresh weights and root lengths) of WT and transgenic seedlings in the presence and absence of NaCl and NaHCO₃ (Figure 5a–f). Under 1/2MS medium, no differences were observed between the WT and PutRUB transgenic seedlings. When the NaCl and NaHCO₃ concentration was increased, the growth of all seedlings was gradually retarded. Under stress treatments (100 and 125 mM NaCl; 1.5 and 3 mM NaHCO₃), the root lengths and fresh seedling weights of the transgenic seedlings were significantly higher than those of the WT.
Figure 5. Phenotypic analyses of PutRUB transgenic plants treated with NaCl, NaHCO$_3$, and H$_2$O$_2$. (a) Phenotypes of WT and PutRUB transgenic seedlings treated with 150 and 125 mM NaCl; (b) Effects of NaCl on the root length of wild-type (WT) and PutRUB transgenic plants; (c) Effects of NaCl on fresh weight of wild-type (WT) and PutRUB transgenic plants; (d) Phenotypes of WT and PutRUB transgenic seedlings treated with 1.5 and 13 mM NaHCO$_3$; (e) Effects of NaHCO$_3$ on the root length of wild-type (WT) and PutRUB transgenic plants; (f) Effects of NaHCO$_3$ on fresh weight of wild-type (WT) and PutRUB transgenic plants; (g) Phenotypes of WT and PutRUB transgenic seedlings treated with 2 and 14 mM H$_2$O$_2$; (h) Effects of H$_2$O$_2$ on the root length of wild-type (WT) and PutRUB transgenic plants; (i) Effects of H$_2$O$_2$ on fresh weight of wild-type (WT) and PutRUB transgenic plants. Single and double asterisks represent significant differences from WT at $p < 0.05$ and $p < 0.01$, respectively.
Salt and alkali stress are major abiotic stresses. To adapt to these stresses, plants have developed some sophisticated mechanisms to sense external pressure signals. Plants can change their physiological and morphological characteristics to adapt to adversity [9,40]. Multiple stresses often induce the same cell signal transduction pathways. All of these stresses will lead to a common adverse effect that is oxidative stress [41–44]. When transgenic lines were exposed to H₂O₂, they showed much better root and leaf growth than the WT, and their fresh weights and root lengths were higher than those of the WT (Figure 5g–i). Thus, we suspect PutRUB may play a vital role in reducing the damage from H₂O₂.

2.5. PutRUB Is Involved in Reducing of Reactive Oxygen Species (ROS) Detoxification

Staining with nitroblue tetrazolium (NBT) specifically tests superoxide radicals. NBT staining demonstrated that dealing with NaCl and NaHCO₃ induced O₂⁻ accumulation in the leaves of PutRUB transgenic lines and WT plants (Figure 6a). In the presence of 100 and 125 mM NaCl, and 1.5 and 3 mM NaHCO₃, less O₂⁻ was accumulated in the leaves of PutRUB plants than in the WT. We obtained similar results for measurements of H₂O₂ content. As shown in Figure 6b, under NaCl and NaHCO₃ stresses, H₂O₂ accumulation in PutRUB transgenic lines was lower than in WT plants. These results suggest that PutRUB is involved in decreasing H₂O₂ accumulation in plants and plays important roles in response to NaCl stress and NaHCO₃ stress.

![Figure 6](image-url)
Previous reports have shown that when plants are exposed to NaCl stress or oxidative stress, H$_2$O$_2$ and superoxide anions increase, injuring the plant cells [45]. Rubredoxins from *Pyrococcus furiosus* and *Desulfovibrio vulgaris* play an important role in sharp contrast to the role of superoxide dismutase, effectively reducing damage from O$_2$ in superoxide detoxification [31–33]. *AtRUB* (At5g17170) plays an important role in the control of ROS detoxification in response to NaCl stress [37]. These results all show that rubredoxins are key players in plant tolerance and responses to oxidative stress. Rubredoxins are electron carriers for various enzymes that reduce the damage from oxidative stress and protect the normal microbial life cycle [30,31,46]. Therefore, we assume that *PutRUB* maintains normal electron transfer to enhance transgenic plant tolerance and reduce ROS accumulation under NaCl and NaHCO$_3$ stresses.

3. Materials and Methods

3.1. Plant Materials and Stress Treatments

Wild-type of *P. tenuiflora* plants, organs/tissues (roots, stems, leaves, flowers and seeds) and mature seeds were collected from the AnDA experimental base of the Alkali Soil Natural Environmental Science Center (ASNESC), Northeast Forestry University (Harbin, China) in Songnen Plain in Northeast China. Plants were germinated in water for 3 weeks before stress treatments. Plants were harvested at 6, 12, 24 and 48 h after stress treatments and preserved at −80 °C for real-time PCR.

3.2. Phylogenetic Analysis

All protein sequences used for multiple alignments and phylogenetic analysis were extracted from the NCBI non-redundant protein sequences (nr) database using Blastp searches with an e-value cutoff of 1.0 × 10$^{-6}$. The results were used to retrieve proteins with high similarity to the *PutRUB* protein. Then, we used CD search (PMID: 25414356) to analyze the conserved domains within these proteins [47]. Proteins with PDZ/PDZ signaling superfamily or rubredoxin-like superfamily domains were used for further analysis. Alignments of the protein sequences were created with ClustalW (ref) (European Molecular Biology Laboratory, Heidelberg, Germany) using the following parameters: gap opening penalty 10, gap extension penalty 0.2. An unrooted phylogenetic tree was constructed with MEGA 6 (Molecular Evolutionary Genetics Analysis version 6.0, Research Center for Genomics and Bioinformatics, Tokyo Metropolitan University, Hachioji, Tokyo, Japan.) using the maximum likelihood method, and analyses were analysis using the Poisson correction model [48]. We used Gamma distribution to model rate variation sites with shape parameter = 1 distribution to model rate variation sites (shape parameter = 1).

3.3. Constructs

The ORF of *PutRUB* was amplified from the *P. tenuiflora* cDNA library using the primers *PutRUB*-F and *PutRUB*-R. The gene was cloned using *PutRUB*-YE, PB-F and *PutRUB*-YE, PB-R (Table 1). The amplified product was digested with *BamHI* and *XhoI*, cloned into the plant expression vector pBI121 then sequenced, which was used for subsequent analysis.

To construct GFP fusion proteins, *PutRUB* was amplified with the primers *PutRUB*-GFP-FW and *PutRUB*-GFP-RV (Table 1). The amplified product without the stop codon was digested with *BamHI* and *KpnI*, and cloned into the pEFP vector (Invitrogen, Carlsbad, CA, USA). The constructed plasmid pEFP-*PutRUB*-GFP was digested with *BamHI* and *EcoRI*, and cloned into the pYES2 vector to obtain the plasmid pYES2-*PutRUB*-GFP. Then the product was digested with *BamHI* and *SacI*, and cloned into the pBI121 vector to obtain the plasmid pBI121-*PutRUB*-GFP.

The constructs pBI121-*PutRUB*, pBI121-*PutRUB*-GFP and pBI121-GFP (control) were used to generate transgenic Arabidopsis by the floral dip method [49].
### Table 1. All primers used for PCR analysis.

<table>
<thead>
<tr>
<th>Primers Name</th>
<th>Sequences (5’→3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PutRUB-F</td>
<td>ATGGCTGCCAGCCTCTCCTCTGT</td>
</tr>
<tr>
<td>PutRUB-R</td>
<td>CTATTGCAGGCCATATACAAGCAG</td>
</tr>
<tr>
<td>PutRUB-YE,PB-F</td>
<td>GGATCCATGGCTGCCAGCCTCTCCTGT</td>
</tr>
<tr>
<td>PutRUB-YE,PB-R</td>
<td>CTCGAGCTATTGCAGGCCATATACAAGCAG</td>
</tr>
<tr>
<td>PutRUB-GFP-FW</td>
<td>GGTACCATGGCTGCCAGCCTCTCCTGT</td>
</tr>
<tr>
<td>PutRUB-GFP-RV</td>
<td>GGATCCTTGACGGCCATATACAAGCAG</td>
</tr>
<tr>
<td>PutRUB-QPCR-F</td>
<td>GTTCACCAAGACAGCCATCCAG</td>
</tr>
<tr>
<td>PutRUB-QPCR-R</td>
<td>CAGTCAGCTTCCGCTAAAAATCG</td>
</tr>
<tr>
<td>Put-Tubulin-F</td>
<td>GTGTCAGCCATACTGTGCCAATTC</td>
</tr>
<tr>
<td>Put-Tubulin-R</td>
<td>TTGCTCATCGGTCAGCAATACC</td>
</tr>
</tbody>
</table>

### 3.4. Subcellular Localization Assay

pBI121-GFP and pBI121-PutRUB-GFP were stably transformed into Arabidopsis. Protoplasts were extracted from transgenic leaves as described in [50]. An Axio Vision fluorescent microscopy system (Axio Imager Z2, Zeiss, Germany) was used to observe fluorescence.

### 3.5. Quantitative Real-time PCR

Total RNA was extracted by TRIzol reagent (Invitrogen) and cDNA was synthesized with PrimeScript Reverse Transcriptase (Takara Bio, Shiga, Japan) used oligo dT as primer, referring to the manufacturer’s instructions. The cDNA were diluted 5 times with double distilled water for quantitative real-time PCR (qRT-PCR) as templates, with the primers: PutRUB-QPCR-F and PutRUB-QPCR-R; Put-Tubulin-F and Put-Tubulin-R. The cDNA was amplified using TransStart Top Green qPCR SuperMix (TransGen Biotech) on a Stratagene Mx3000P QCR system (Agilent Technologies, Cold Spring, NY, USA). The comparative CT method was used to calculated relative expression levels, with the PutTubulin gene as an internal control.

### 3.6. Stress Response Analysis in Transgenic Arabidopsis

The plasmid pBI121-PutRUB was transformed into Arabidopsis by A. tumefaciens-mediated floral dipping. We isolated T₀-generation seeds of A. thaliana on 1/2MS medium containing 50 mg·L⁻¹ kanamycin, then collected T₃ transgenic lines (T₃ #1, #2, and #3). Transgenic seedlines were identified by PCR and northern blotting. The CTAB method were used to extract plant genomic DNA [51].

For abiotic stress treatment, WT and T₃ generation transgenic Arabidopsis seeds were grown on 1/2MS agar plates for 7 days. The seedlings were then transferred to 1/2MS medium supplemented with different chemicals for stress treatments (100 and 125 mM NaCl, 1.5 and 3 mM NaHCO₃, 2 and 4 mM H₂O₂); 1/2MS medium was used as a control. The seedlings were allowed to grow for 10 days (vertical culture, 16/8 h light/dark, temperature 22/18 °C), after which the fresh weights and root growth of the seedlings were measured. Statistical analyses were carried out using Student’s t-test.

### 3.7. ROS Detection

Three-week-old transgenic T₂ generation seedlings of the PutRUB were cultivated on 1/2MS medium containing 125 mM NaCl and 3 mM NaHCO₃ for 12 h at 22 °C, and the in situ accumulation of O₂⁻ was determined by histochemical staining with NBT. Briefly, in an amber-colored bottle, 0.1 g NBT was dissolved in 50 mM sodium phosphate buffer (pH 7.5) and the volume was increased up to 50 mL to get a 0.2% solution. The solution was mixed thoroughly using a magnetic stirrer. The NBT staining solution was prepared fresh before use. The seedlings were placed in test tubes and immersed in NBT staining solution to detect H₂O₂. The tubes were wrapped with aluminum foil and keep overnight at room temperature. Then, the chlorophyll was removed for proper visualization of the stain. This was
done by immersing the seedlings in absolute ethanol and heating in a boiling water-bath for 10 min (or more if necessary, with intermittent shaking [52].

H$_2$O$_2$ levels were quantified by the absorbance at a wavelength of 415 nm using titanium sulfate precipitation as described in [53].

4. Conclusions

Here, we revealed that a chloroplast-localized rubredoxin family protein from _P. tenuiflora_ might be involved in the plant response to environmental stresses. _PutRUB_ overexpression increased growth and inhibited H$_2$O$_2$ accumulation under NaCl and NaHCO$_3$ treatments. Our results indicate that _PutRUB_ might be involved in maintaining normal electron transfer to enhance transgenic plant adaptability to adversity and reduce ROS accumulation under NaCl and NaHCO$_3$ stresses.

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Author Contributions: Ying Li and Shenkui Liu conceived and designed the experiments; Ying Li performed the experiments; Ying Li analyzed the data; Panpan Liu contributed reagents/materials/analysis tools; Ying Li wrote the manuscript. Shenkui Liu, Tetsuo Takano revised and final approval of the article.

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

References


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