Isolation of Salt Stress-Related Genes from *Aspergillus glaucus* CCHA by Random Overexpression in *Escherichia coli*

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The halotolerant fungus *Aspergillus glaucus* CCHA was isolated from the surface of wild vegetation around a saltern with the salinity range being 0–31%. Here, a full-length cDNA library of *A. glaucus* under salt stress was constructed to identify genes related to salt tolerance, and one hundred clones were randomly selected for sequencing and bioinformatics analysis. Among these, 82 putative sequences were functionally annotated as being involved in signal transduction, osmolyte synthesis and transport, or regulation of transcription. Subsequently, the cDNA library was transformed into *E. coli* cells to screen for putative salt stress-related clones. Five putative positive clones were obtained from *E. coli* cells grown on LB agar containing 1 M NaCl, on which they showed rapid growth compared to the empty vector control line. Analysis of transgenic *Arabidopsis thaliana* lines overexpressing CCHA-2142 demonstrated that the gene conferred increased salt tolerance to plants as well by protecting the cellular membranes, suppressing the inhibition of chlorophyll biosynthesis. These results highlight the utility of this *A. glaucus* cDNA library as a tool for isolating and characterizing genes related to salt tolerance. Furthermore, the identified genes can be used for the study of the underlying biology of halotolerance.

1. Introduction

Halotolerant or halophilic microorganisms, able to live in saline environments, offer a large number of potential applications in various fields of biotechnology [1]. In order to survive high salinity, such organisms have developed adaptive features to function under extreme conditions [2]. Salinity stress is one of the major factors that reduce crop plant growth and productivity, resulting in crucial economic losses worldwide. Therefore, it is vital to isolate and functionally characterize salinity stress-related genes to elucidate the mechanisms underlying halotolerance and develop salinity stress-tolerant plants [3].

As the fundamental cellular adaptive mechanisms are conserved and play precise roles in salinity stress [4], it can be supposed that stress-related genes from various species can be screened efficiently via overexpression in simple organisms. Indeed, there are several reports of such functional screening of stress-related genes by overexpression in yeast [5–7] or *E. coli* [8–10]. The gene for a catalytic subunit of the *Beta vulgaris* (sugar beet) protein kinase CK2 (BvCKA2) related to salinity was cloned by functional expression in yeast of a NaCl-induced *B. vulgaris* cDNA library [6]. In addition, a plant RelA/SpoT homolog has been found to confer salt tolerance to *E. coli* [10]. It is thought that the cellular response to environmental stress is highly conserved among all three superkingdoms, namely, the eukaryotes, the eubacteria, and the archaea [4]. Although plants may have unique adaptive mechanisms to accommodate various stresses, the basic cellular stress response (CSR) mechanism, which is activated in response to stress, is similar in prokaryotes, lower plants, and eukaryotes [4, 10]. Accordingly, stress-resistance genes of halotolerant fungal species have the potential to improve the stress tolerance of microorganisms and plants [11–14].
The Aspergillus glaucus (A. glaucus) CCHA strain is one of the salt-tolerant fungi that are found to thrive in salt mines. Its optimum salt concentration is 50 g L\(^{-1}\) and it can survive from 0 g L\(^{-1}\) to 310 g L\(^{-1}\) salt [15]. Therefore, A. glaucus might serve as a valuable resource from which salt tolerance related genes are identified. The isolation and identification of salinity stress-resistance genes will promote effort to develop plants tolerant of salinity stress. In this study, an A. glaucus CCHA cDNA library was constructed and functionally screened for halotolerance-related genes. A number of clones were identified, and five positive clones were analyzed for their roles in stress tolerance. The results demonstrate that this cDNA library represents an efficient tool for isolating and characterizing genes related to salt tolerance.

2. Materials and Methods

2.1. Media, Strains, and Growth Conditions. A. glaucus (strain CCHA), which was isolated and characterized at Jilin University, China [15], was kindly provided by Professor Shihong Zhang. It was cultivated using potato dextrose agar (PDA) and potato dextrose broth (PDB) with 0.85 M NaCl. Liquid cultures were grown at 37°C of antibiotics in accordance with the requirements of the cultures.

2.2. RNA Isolation. For RNA isolation, A. glaucus (strain CCHA) was grown in PDB liquid medium with 3 M NaCl and harvested by centrifugation after 4 days. The pellet was frozen in liquid nitrogen and homogenized using a mortar and pestle. Total RNA was isolated from 200 mg homogenized tissue using a Norgen RNA Purification Kit (Norgen Biotek Corp., Ontario, Canada). The quality and quantity of total RNA were analyzed using a NanoDrop 2000 UV-Vis spectrophotometer (Thermo, Wilmington, USA) and gel electrophoresis.

2.3. Library Construction in E. coli DH5α. A cDNA library was constructed using the SMART cDNA Library Construction Kit (Clontech, Mountain View, USA) according to the manufacturer’s instructions with minor modifications. Briefly, the first strand cDNA was synthesized from 5 μg total RNA using SuperScript III (Clontech, Mountain View, USA) with SMART IV oligonucleotide primer and CDS III/3’ PCR primer (Table 1). The products were used for the synthesis of second strand cDNA catalyzed by 50× Advantage 2 Polymerase Mix. The resulting double strand cDNA (ds cDNA) (2-3 μg in 50 μL) was digested by proteinase K (20 μg/μL) to inactivate the DNA polymerase. The purified ds cDNA was digested with SfiI at 50°C for 4 h, 2 μL 1% xylene cyanol dye was added, and cDNA size fractionation was performed using CHROMA SPIN-400 (Clontech, Mountain View, USA). The concentration and yield of the ds cDNA fractions were estimated using a NanoDrop 2000 UV-Vis spectrophotometer (Thermo, Wilmington, USA) and gel electrophoresis. The ds cDNA longer than 700 bp was pooled, precipitated, and used for ligation reactions with modified vector pYES2G digested by SfiI [16]. Subsequently, 5 μL products of the reaction were electroporated into 100 μL E. coli Electro-Cells DH5α (TAKARA, Dalian, China), at 1500 V with a Gene Pulser (BIO-RAD, Hercules, USA). The titer of the library was determined by serial dilution in LB medium (0, 10\(^{-1}\), 10\(^{-2}\), and 10\(^{-3}\)). In order to estimate the integrity and annotation of the inserts, 100 clones were sequenced randomly. The library was mixed with the freezing medium (15% glycerol, 85% LB medium), frozen, and stored at −80°C.

2.4. Library Transfer to E. coli BL21 Star (DE3). An aliquot of the library that contained approximately 4.8 × 10\(^{9}\) transformants was inoculated into 50 mL liquid LB medium with ampicillin and grown on a rotary shaker at 180 rpm and 37°C. After overnight incubation, total plasmid DNA was isolated with the AxyPrep Plasmid Miniprep Kit (Axygen Scientific, Tewksbury, USA). The plasmid DNA was transformed into E. coli BL21 Star (DE3) cells by electroporation.

2.5. Identification of Clones Conferring Salt Tolerance in E. coli BL21 Star (DE3). To characterize the growth of transformants under salt stress, 100 μL E. coli BL21 Star (DE3) library aliquots were plated on LB plates (100 mg L\(^{-1}\) ampicillin) containing a range of concentrations of NaCl (0.7, 0.75, and 0.85 and 1 M). The screening plates were then incubated at 37°C for 48 h and transformants with increased salt tolerance were selected for further study. PCR was performed to assess the insert in the pYES2G vector in a 20 μL reaction volume, with 10 pmol vector-specific primers (T7 and pYES2G-R). The thermal profile of the reaction was as follows: 5 min denaturation at 95°C, followed by 35 cycles of 30 s at 95°C, 30 s at 53°C, and 2 min and 30 s at 72°C, with the final elongation step of 7 min at 72°C. The PCR products were analyzed by agarose electrophoresis and the colonies with the desired length were sequenced by Sangon Biotech Co., Ltd. (Sangon, Shanghai, China). The nucleotide sequences were analyzed and their putative functions were identified by comparison with GenBank database entries using BLASTX. All colonies showing tolerance to salt stress were mixed with the freezing medium (15% glycerol, 85% LB medium), frozen, and stored at −80°C until further analysis.

2.6. Salt-Tolerance Assays. Salt tolerance of the positive E. coli BL21 Star (DE3) transformants was compared to that of transformants carrying the empty vector. Spot assays were carried out to ascertain the functions of the screened genes from A. glaucus (strain CCHA) in E. coli BL21 Star (DE3) cells. Cells were grown in LB medium until OD 600 reached 0.6. Then, the cultures were serially diluted to 10\(^{-5}\), 10\(^{-4}\), and 10\(^{-3}\), and 5 μL of each dilution was spotted on LB plates supplemented with 0.7 M NaCl or 1 M NaCl and cultured for 3 d.

Functional analyses of five clones showing tolerance to 1 M NaCl were also performed in LB liquid culture supplemented with 1 M NaCl. E. coli BL21 Star (DE3) cells with recombinant plasmid or empty vector were grown in LB liquid culture, diluted to OD 600 = 0.6, and 1 mL cells were inoculated in 100 mL LB medium containing 1 M NaCl and
Table 1: Primers used in this study.

| Primer          | Primer sequence                          | Description                                    |
|-----------------|------------------------------------------|-----------------------------------------------|
| SMART IV oligonucleotide | 5'-AAGCAGTGGTATCAACGCAGAGTGCGGATTACGGCCGGG-3' | First strand synthesis, forward               |
| CDS III/3 PCR primer | 5'-ATTCTAGGGCGGAGCCGCGCGACATG-d(T(N=3)N=1-N=3, N=A,G,C; T; N=1=A,G,C) | First strand synthesis, reverse               |
| T7              | 5'-CGCAACAATCCCACTATCTCTCGCAAG-3'        | Transgenic screening primer, forward          |
| PYES2G-R        | 5'-GATAACATCGCAAGACCGGCAACAGG-3'         | Transgenic screening primer, reverse          |
| CCHA-42-F       | 5'-CGGACGAAATCCAACCCTTC-3'               | Real-time PCR primer, forward                 |
| CCHA-42-S       | 5'-TTACCAGGCACATAGTCA-3'                 | Real-time PCR primer, reverse                 |
| CCHA-14-F       | 5'-CCGGGCAAAAGGAGCATAAA-3'               | Real-time PCR primer, forward                 |
| CCHA-14-S       | 5'-AACCGCTGCTTAGTCTGGAACA-3'             | Real-time PCR primer, reverse                 |
| CCHA-29-F       | 5'-ATTTCGCGCCGGGATATGGG-3'               | Real-time PCR primer, forward                 |
| CCHA-29-S       | 5'-ACGGGCCCATTACCCGAA-3'                 | Real-time PCR primer, reverse                 |
| CCHA-21-F       | 5'-GGTGAATTCGACCCAATAAC-3'               | Real-time PCR primer, forward                 |
| CCHA-21-S       | 5'-CCAGCCGCTGATGTCTATTTT-3'              | Real-time PCR primer, reverse                 |
| CCHA-47-F       | 5'-CTTGATGCCCCCHACGTACA-3'               | Real-time PCR primer, forward                 |
| CCHA-47-S       | 5'-TGGGCGCTCAAATGAACAAAT-3'              | Real-time PCR primer, forward                 |
| Actin2-F        | 5'-GCACCCCTGTTCTTCTTACCG-3'              | Standard control primer, forward              |
| Actin2-R        | 5'-AACCTCTGTAATGGCAGAACA-3'              | Standard control primer, forward              |
| Tubulin1-F      | 5'-GAGTTGACCACAGCAGATG-3'                | Standard control primer, forward              |
| Tubulin1-R      | 5'-CAGTTTCTTTCTTGGAC-3'                  | Standard control primer, reverse              |

incubated at 37°C, 200 rpm. The bacterial suspensions were harvested every 2 h through 14 h and OD 600 was measured.

2.7. Real-Time Quantitative PCR Analysis. A. glaucus samples were grown in PDB medium with 1 M NaCl at 35°C and 180 rpm. After 2 d, mycelium pellets were harvested and incubated in PDB medium with 3 M NaCl for 0 (control), 0.5, 1, 6, 24, 48, and 72 h. Mycelium pellets were harvested and used for the extraction of RNA. The cDNA was prepared using 1 μg total RNA with the PrimeScript RT reagent kit (TAKARA, Dalian, China). The expression profiles of five representative genes, CCHA-2142, CCHA-2114, CCHA-2227, CCHA-2229, and CCHA-2247, in response to salt stress were determined with tubulin as the internal reference. The primers were designed using primer 5 (Table I). The reaction mixture in a total reaction volume of 20 μL included 2 μL cDNA, 10 μL SYBR Premix Ex Tag (2x) (TAKARA, Dalian, China), and 0.4 μL gene-specific primers. The cycling conditions were as recommended by the manufacturer (10 s at 95°C, 40 cycles of 95°C for 5 s, and 60°C for 31 s). Specificity of amplification was verified by melting curve analysis (60 to 95°C) after 40 PCR cycles. The experiments were repeated three times independently. The relative expression was calculated according to the method of Livak and Schmittgen [17].

2.8. Verifying the Salt Stress Tolerance Activity of Genes in Arabidopsis thaliana. CCHA-2142 cDNA was digested with SfiI and then inserted into the pBI121G vector, which is remodeled as previously described [18]. The recombinant vector was transformed into Agrobacterium tumefaciens EHA105 by electroporation at 2500 V. Arabidopsis thaliana ecotype Columbia plants were transformed by the floral dip method [19]. T₀ seeds were collected and sown on sterile medium containing 50 μg mL⁻¹ kanamycin and then confirmed by PCR and RT-PCR using the primers described above and those for Actin2 (Table I) as the internal control. Wild-type plants (ecotype Columbia) and T₃ CCHA-2142-overexpressing Arabidopsis thaliana plants were sown in soil under a long-day cycle (16 h light, 8 h dark). After 30 d, transgenic and control wild-type plants were soaked in Hoagland's medium with, for 1 d, 3 d, and 5 d, or without 200 mM NaCl. Then 0.5 g leaf tissue was ground in liquid nitrogen and homogenized in 10 volumes of extraction buffer (45.8 mM Na₂HPO₄·12H₂O, 4.25 mM
NaH₂PO₄·2H₂O, pH 7.8) per unit fresh weight. After centrifugation at 12000×g for 30 min at 4°C, the supernatant was used for measurements of total superoxide dismutase (SOD) activity and malondialdehyde (MDA) content.

Total SOD activity was assayed by the method described in Giannopolitis and Ries [20]. MDA, a biomarker for lipid peroxidation, was assayed by the method described in Heath and Packer [21]. Leaf tissue (0.2 g) and 20 mL ultrapure water were mixed, and 3 hours later, the electrical conductivity was measured. Then the mixtures were heated in boiling water for 20 min and then cooled to 25°C to determine the total electrical conductivity. The degree of ion leakage was calculated as electrical conductivity (before boiling)/electrical conductivity (after boiling) × 100%. Leaf tissue (0.1 g) was used for chlorophyll content measurements. Total chlorophyll content, chlorophyll a content, and chlorophyll b content were measured after extraction with 80% acetone [22]. All the experiments were carried out independently at least thrice with four lines (wild type and 3 transgenic plant lines).

3. Results

3.1. Identification of A. glaucus Genes Expressed in the Presence of Salt. We began by constructing a cDNA library from A. glaucus (strain CCHA) grown on medium containing 3 M NaCl. To gain an overview of genes expressed in A. glaucus, one hundred clones were randomly selected for sequencing and bioinformatics analysis. The sizes of the inserted cDNA fragments in the library were ~500–1500 bp. The sequences were annotated based on alignment with sequences in various databases including the National Center for Biotechnology Information (NCBI) nonredundant protein (Nr) database, the NCBI nonredundant nucleotide sequence (Nt) database, and UniProt/Swiss-Prot. Of the 100 unigenes analyzed, 71 had significant matches in the Nr database, Nt database, and Swiss-Prot database. Based on their similarity to known proteins, the annotated genes appear to be functionally involved in signal transduction, osmolyte synthesis and transport, and regulation of transcription (Table 2).

3.2. Screening for A. glaucus Genes Conferring Salt Tolerance. The A. glaucus cDNA library was then transferred to E. coli BL21 Star (DE3) cells for expression. We performed a salt screen assay, in which the E. coli BL21 Star (DE3) library was plated on LB containing 0.7, 0.75, and 0.85 M NaCl. Whereas there was a lawn of colonies on LB plates (containing 0.7 M NaCl), many fewer colonies survived on LB plates containing higher concentrations of NaCl. Sixty transformants that could grow on LB with 0.75 M NaCl were sequenced, and the sizes of the inserts ranged from 300 to 1500 bp. In addition, a total of 22 unique sequences were identified based on supporting growth on LB containing 0.85 M NaCl. Putative functions were identified for 20 of the genes by comparison with deposited sequences in the GenBank database (Table 3). Notably, A. glaucus CCHA-1408 was most similar to RNase PH-related exoribonuclease, which appears to be an important salt stress protein [23]. Two other genes, CCHA-2227 and CCHA-2114, belonged to the DJ-I/Pfpl family, which may be related to salt stress [24]. Based on their putative functional annotations, 14 transformants were selected for further analyses.

3.3. Analysis of Salt Tolerance Confferred by Selected Clones from A. glaucus in E. coli BL21 Star (DE3). Spotting assays of selected transformants on LB plates (Figure 1(a)) did not reveal any differences in growth at 0.5 M NaCl. By contrast, while the control strain carrying an empty plasmid did not grow on LB containing 0.7 or 1 M NaCl, all of the selected A. glaucus clone transformants grew at 0.7 M NaCl. Five of them (CCHA-2142, CCHA-2114, CCHA-2221, CCHA-2229, and CCHA-2247) could support growth at 1 M NaCl, and their accession numbers are KJ934998, KJ935001, KJ935002, KJ934999, and KJ935000, respectively (Table 4). The growth of these five transformants was also analyzed in LB liquid medium containing 1 M NaCl. All five showed increased growth compared to the control strain with empty vector alone (Figure 1(b)).

3.4. Bioinformatics Analysis of A. glaucus CCHA-2247, CCHA-2229, and CCHA-2142. The three genes conferring the strongest resistance to salt were chosen for detailed bioinformatics analysis. The CCHA-2247 cDNA contains a predicted gene with a full-length coding sequence of 408 bp encoding a 135 aa protein of 15.6 kDa. The predicted pI is 10.53 and there is no signal peptide. According to the CELLO program (http://cello.life.nctu.edu.tw/), this protein may be located in the periplasm. CCHA-2247 showed 93% identity to ribosomal L27 protein of Aspergillus flavus (XP_002381915.1), 93% identity to ribosomal L27 protein of Aspergillus terreus (XP_00121815.1), and 92% identity to ribosomal L27 protein of Aspergillus clavatus (XP_001269541.1) at the amino acid level. Phylogenetic analysis of CCHA-2247 revealed that it clustered with a ribosomal protein from Aspergillus flavus (Figure S1b, see Supplementary Material available online at http://dx.doi.org/10.1155/2014/620959).

The CCHA-2229 cDNA includes a gene with a predicted full-length coding sequence of 1038 bp. The deduced amino acid sequence revealed a protein consisting of 345 aa with a molecular mass of 37.9 kDa. As predicted, the pI is 5.54 and it has no signal peptide. Using the TMHMM program (http://www.cbs.dtu.dk/services/TMHMM/), we predicted that it is an inner membrane protein with 8 transmembrane domains. The BLASTP analysis results suggested that it is a putative plasma membrane low affinity zinc ion transporter. CCHA-2229 shares 94% identity with proteins from Aspergillus ruber (EYE97956.1) and 76% identity with one from Aspergillus clavatus (XP_00127799.1). Phylogenetic analysis placed CCHA-2229 at the base of the putative plasma membrane low affinity zinc ion transporter family (Figure S1d), which indicates that it may be an ancient form of a plasma membrane low affinity zinc ion transporter.

The cDNA of CCHA-2142 encodes a predicted protein consisting of 347 amino acids with a molecular mass of 39.5 kDa. There is a signal peptide in this sequence and its pI is 6.06. Preliminary analysis result of subcellular localization showed that it is likely to be a periplasmic...
Table 2: *A. glaucus* coding sequences expressed during growth on 3 M NaCl. One hundred cDNA clones were randomly selected and sequenced.

| A. glaucus unigenes | Annotation (based on similarity to database sequences) |
|---------------------|--------------------------------------------------------|
| 0307, 23171         | RRM superfamily, nuclear and cytoplasmic polyadenylated RNA-binding protein publ |
| 2229, 2243, 23119, and 0501 | ZIP zinc/iron transport family, plasma membrane low affinity zinc ion transporter |
| 0508, 2247, 2372, 2390, 0209, 2302, 2341, 2380, 2382, 23144, 23151, 23159, and 2378 | Ribosomal_L27e superfamily, 60S ribosomal protein L27e |
| 0509, 23136, and 2018 | Similar to alpha/beta hydrolase fold-3 domain containing protein |
| 0314 | Six-hairpin glycosidase |
| 1402, 2037, and 2113 | 4HTB_3 (beta-hydroxydecanoyl-acyl carrier protein-(ACP-) dehydratase) structure and subsequently in 4HBT (4-hydroxybenzoyl-CoA thioesterase) |
| 1408, 23140, and 23134 | RNase_PH superfamily, exosome complex endonuclease 2/ribosomal RNA processing protein |
| 2250 | MFS general substrate transporter (secondary transporters that include uniporters, symporters, and antiporters) |
| 2330 | Casein kinase II beta 2 subunit |
| 2334 | Putative intracellular protease/amidase |
| 2354, 23179, 2042, 2048, 2142, and 2263 | DUF3431 superfamily |
| 2362 | WD repeat protein (adaptor/regulatory modules in signal transduction, Pre-mRNA processing and cytoskeleton assembly; typically contains a GH dipeptide II–24 residues) |
| 23131 | p450 superfamily, cytochrome P450 [secondary metabolites biosynthesis, transport, and catabolism/methylsterigmatocystin oxidoreductase] |
| 23149 | PF03619 domain protein (organic solute transporter Ostalpha) |
| 23207 | WD40 superfamily. Nuclear pore complex subunit (adaptor/regulatory modules in signal transduction) |
| 2265 | Spore germination lipase lipC |
| 2251 | Family 2 glycosyl transferase |
| 2258 | Phosphoprotein |
| 0203 | 7 transmembrane receptors |
| 2301 | Exodeoxyribonuclease V subunit beta |
| 2302, 2304 | Protein CRE_17637 |
| 2305 | Aspartic proteinase nepenthesin-1-like |
| 2306 | Carnitine acetyl transferase |
| 2339 | TIM_phosphate_binding superfamily, nucleoporin SONB |
| 2377 | PDI_b_family, PDI_a_family (thioredoxin_like superfamily), disulfide-isomerase precursor |
| 2383, 23163 | MDR superfamily, delta-aminolevulinic acid dehydratase |
| 23103 | PING superfamily, regulator of gluconeogenesis Rmd5 |
| 23104 | Fasciclin superfamily, fasciclin domain family protein |
| 23120 | Lactamase_B superfamily, Zn-dependent hydrolases of the beta-lactamase fold |
| 23135 | NADH dehydrogenase subunit 1 |
| 23187 | Rho_GDI superfamily (rho-gdp dissociation inhibitor) |
| 23189 | P-loop NTPase domain superfamily (ras-like GTP-binding protein) |
| 2397 | SocE |
| 2253, 2227, 2214, 2242, 2369, 23156, 23197, 0215, 0217, and 2346 | DJ-1/Pfi family protein (Type 1 glutamine amidotransferase- (GATase-)- like domain) |
| 2252 | Isonitrile hydratase |
| 22651, 2375, 23102, 23143, 23160, 23168, 23191, and 2248 | 4-Methyl-5-(B-hydroxyethyl)thiazole monophosphate biosynthesis protein |
| 2243 | Zinc/iron permease |
| 22652, 2389, and 2246 | Unknown; no similar proteins found |
Table 3: *A. glaucus* coding sequences that were identified from the cDNA expression library as putatively conferring increased salt-resistance.

| *A. glaucus* unigenes | Annotation (based on similarity to database sequences) |
|-----------------------|------------------------------------------------------|
| 2229                  | ZIP zinc/iron transport family, plasma membrane low affinity zinc ion transporter |
| 2247                  | Ribosomal L27e superfamily, 60S ribosomal protein L27e |
| 1402                  | 4HTB3 (beta-hydroxydecanoyl-acyl carrier protein- (ACP-) dehydratase) structure and subsequently in 4HBT (4-hydroxybenzoyl-CoA thioesterase) |
| 1408                  | RNase PH superfamily, exosome complex endonuclease 2/ribosomal RNA processing protein |
| 2227, 2114            | DJ-1/Pfpl family protein (Type I glutamine amidotransferase- (GATase1-) like domain) |
| 2246, 22652, and 2221 | Unknown; no similar proteins found |
| 2250                  | MFS general substrate transporter (secondary transporters that include uniporters, symporters, and antiporters) |
| 2258                  | Phosphoprotein |
| 2142                  | DUF3431 superfamily |
| 22651, 2248           | 4-Methyl-5 (B-hydroxyethyl)thiazole monophosphate biosynthesis protein |
| 22654                 | Spore germination lipase lipC |
| 0203                  | 7 transmembrane receptors |

Figure 1: *A. glaucus* genes confer salt tolerance to *E. coli*. (a) Spot assay of BL21/pYES2G (empty plasmid), BL21/CCHA-2142, BL21/CCHA-2229, BL21/CCHA-2114, BL21/CCHA-2247, BL21/CCHA-2243, BL21/CCHA-2248, BL21/CCHA-2258, BL21/CCHA-2264, BL21/CCHA-2246, BL21/CCHA-2227, BL21/CCHA-2250, BL21/CCHA-1402, BL21/CCHA-22652, and BL21/CCHA-22654 on LB plates with 0.7 M NaCl and 1 M NaCl. (b) Growth analysis of CCHA-2142, CCHA-2114, CCHA-2229, CCHA-2227, and CCHA-2247 in LB liquid medium with 1 M NaCl. OD$_{600}$ was recorded every 2 h.
Table 4: *A. glaucus* coding sequences that were obtained from the cDNA expression library selected with increased salt-resistance.

| GenBank accession number | Gene       | Putative gene function                                      | Selection |
|--------------------------|------------|-------------------------------------------------------------|-----------|
| KJ934998                 | CCHA-2142  | DUF3431 superfamily                                        | 1 M NaCl  |
| KJ935001                 | CCHA-2114  | DJ-1/Pip1 family protein (Type 1 glutamine amidotransferase- (GATase-like) domain) | 1 M NaCl  |
| KJ934999                 | CCHA-2229  | COGO428 superfamily (plasma membrane low affinity zinc ion transporter) | 1 M NaCl  |
| KJ935002                 | CCHA-2221  | Unknown; no similar proteins found                         | 1 M NaCl  |
| KJ935000                 | CCHA-2247  | 60S ribosomal protein L27e                                  | 1 M NaCl  |

The amino acid sequence alignment of CCHA-2142 is shown in Figure S1f. In addition, sequence analysis showed that this gene belongs to DUF3431 superfamily. CCHA-2142 shares 66% identity with an *Aspergillus terreus* protein (XP_001213981.1), 62% identity with a *Neosartorya fischeri* protein (XP_001262501.1), and 61% identity with an *Aspergillus fumigatus* protein (XP_7464951.1). Phylogenetic analysis of CCHA-2142 indicated that it clustered with one hypothetical protein of *Neosartorya fischeri* NRRL (XP_001262501) (Figure S1f).

3.5. Expression Profiles of CCHA-2142, CCHA-2114, CCHA-2227, CCHA-2229, and CCHA-2247 in *A. glaucus* under Salt Stress. Quantitative real-time PCR was performed to analyze the expression patterns of selected candidate halotolerance-conferring genes from *A. glaucus*. Overall, all five genes showed upregulation in response to salt stress, albeit to various degrees. The expression profiles of CCHA-2114 (Figure 2(a)) and CCHA-2142 (Figure 2(d)) in *A. glaucus* under salt stress were similar to each other, with an immediate response at 0.5 h, reaching a peak at 6 h or 24 h and then gradually decreasing to the original level at 72 h. The expression of CCHA-2229 showed no obvious upregulation until treatment had lasted for 72 h, at which point expression was increased 5.17-fold (Figure 2(c)). CCHA-2221 (Figure 2(b)) and CCHA-2247 (Figure 2(e)) were upregulated 2.57-fold and 2.22-fold, respectively, at 24 h of salt stress, with no significant change at other treatment durations.

3.6. *A. glaucus* CCHA-2142 Confers Salt Tolerance to *Arabidopsis thaliana* in Transgenic Plants. To examine the potential ability of CCHA-2142 to confer salt tolerance to other species, we generated transgenic *Arabidopsis thaliana* lines constitutively expressing *A. glaucus* CCHA-2142 which were detected by PCR (Figure 3(a)) and RT-PCR (Figure 3(b)). Three transgenic lines and a control line (wild type, WT) were used for physiological studies under salt stress. In total, more serious damage was observed on the leaves of the wild-type *Arabidopsis* than that of the transgenic lines when treated with 200 mM NaCl. After 72 hours, both the wild-type and transgenic plants began to wilt, while injurious spots gradually appeared on the leaves of wild-type plants, which was not observed on transgenic lines. And the growth of the wild type was much more inhibited compared to the transgenic lines. After 7-day stress, all the wild-type plants permanently died, while the transgenic lines could present recovery of vegetative growth and reproductive growth partially after being rewatered for 14 days (Figure 4(a)). Without salt treatment, the SOD activities of CK and transgenic lines were 159.5, 158.5, 154.1, and 167.5 U/g FW, respectively, and there was no significant difference between them. Following salt stress treatment for 5 d, the SOD activities were 225.4, 298.9, 327.0, and 270.2 U/g FW. At all durations of salt stress, the SOD activity in CCHA-2142-overexpressing plants was significantly higher than that in control plants (Figure 4(b)).

Lipid peroxidation is a marker of oxidative damage under high salinity, and we measured MDA content as an indicator of lipid peroxidation (Figure 4(c)). Without salt stress treatment, the MDA content of the control and transgenic plants was similar. With salt stress treatment of 1, 3, and 5 d, the MDA content changed by only a small amount in transgenic plants but increased dramatically in control plants. These results indicate that CCHA-2142 overexpression led to reduced lipid peroxidation and thus likely increased tolerance to salt stress.

The membrane integrity of leaves was assessed by measuring electrical conductivity, which reveals the degree of ion leakage from cellular membranes. Although electrical conductivity increased in both transgenic and control plants in response to salt stress, the degree of ion leakage increased less in the transgenic lines at 3 and 5 d than in the control plants at those time points (Figure 4(d)). This suggests that overexpressing CCHA-2142 helps to maintain membrane permeability under high salinity conditions, directly or indirectly.

Without salt stress, no significant differences in the contents of total chlorophyll, chlorophyll a, and chlorophyll b were detected between the control and CCHA-2142-overexpressing plants. However, after salt stress for 1 d, 3 d, and 5 d, the total chlorophyll content was reduced more dramatically in the transgenic plants than in the control plants (Figure 4(e)). In addition, the chlorophyll a content and chlorophyll b content have the same solution (Figures 4(f) and 4(g)). These results suggest that the photosynthetic capacity of the transgenic plants overexpressing CCHA-2142 remained higher during salt treatment than that of the control plants.

4. Discussion

The cellular mechanisms of stress tolerance are similar from bacteria to higher plants, and halotolerant fungi are
Figure 2: Expression patterns of candidate *A. glaucus* salt-tolerance genes. Real-time PCR analysis of *CCHA-2114* (a), *CCHA-2221* (b), *CCHA-2229* (c), *CCHA-2142* (d), and *CCHA-2247* (e) in *A. glaucus* CCHA under 3 M NaCl stress for 0, 0.5, 1, 6, 24, 48, and 72 h.

Good potential sources of biotechnologically important genes. As abiotic stress could affect the cellular gene-expression machinery, it is evident that a lot of genes are up- or downregulated. For instance, heterologous expression of selected *Debaryomyces hansenii* and *Hortaea werneckii* genes markedly improved the salt tolerance of *Saccharomyces cerevisiae* [25, 26]. Lots of cellular responses to salt stress are conserved between the prokaryotes and the eukaryotes. Due to the similarity of the general cellular stress responses, these significant genes related to salinity tolerance have been studied by their overexpression in simple organisms, such as bacteria or yeast [7, 27].

*A. glaucus* (strain CCHA), which thrives on salt mines, represents another rich resource to mine for genes involved in
Figure 3: The identification of CCHA-2142-overexpressing Arabidopsis thaliana. (a) PCR analysis of transgenic plants with CCHA-2142: M means DM 2000 marker; WT means wild type; 1, 2, 3, 4, and 5 mean transgenic lines; and plasmid means positive control (plasmid pBI121G-CCHA-2142). (b) Confirmation of CCHA-2142 was expressed in transgenic lines by RT-PCR; WT means wild type; and 1, 2, and 3 mean transgenic lines.

Salt tolerance. To isolate genes of potential biotechnological utility, we constructed a full-length cDNA library from A. glaucus under salt stress. Given that some of the protective mechanisms are common to both prokaryotes and eukaryotes under stress conditions [10, 28], we expressed the A. glaucus cDNA library in the model prokaryote E. coli, taking advantage of its short life cycle and ease of manipulation. The identification of over 100 transformants with increased halotolerance demonstrates the efficiency of this screening system for identification of candidate genes involved in basic cellular stress responses.

Characterization and identification of genes in halotolerant or halophilic microorganisms provide an important basis to elucidate the mechanisms underlying stress tolerance [29]. About 60% of the candidate genes identified in this study had not been characterized previously, and we confirmed their potential roles in salt stress using spot assays. All five clones selected for more detailed analysis and their putative functions were identified by comparison with GenBank database entries. PsRPL30E (ribosomal protein-L30E), one of the components of the large (60 S) ribosomal subunit from Pisum sativum, confers high salinity resistance to E. coli, and the expression of PsRPL30E is induced by high salt stress and repressed by cold stress in Pisum sativum pea [3]. The ribosomal structure between eukaryotes and prokaryotes is different; however, it may confer tolerance via a pathway yet to be illuminated. A yeast ribosomal 60 S subunit homologue was induced more than 2.0-fold under salt stress [30]. Here, we found that CCHA-2247, encoding a ribosomal 60 S subunit homologue, improved the salinity tolerance of the E. coli. This, along with its expression profile, suggests that CCHA-2247 may be involved in salinity stress tolerance and thus may represent a new salt-tolerance candidate gene from A. glaucus (CCHA).

Another A. glaucus gene conferring salt tolerance to E. coli, CCHA-2229, encodes a protein that appears to be a member of the COGO428 superfamily, which includes divalent heavy-metal cations transporters playing an important role in zinc homeostasis at the cellular level [31]. The low affinity system has a lower affinity for substrate, and it is temperature, time, and concentration dependent. In Saccharomyces cerevisiae, the ZRT2 gene encodes the low affinity zinc transporter which could increase low affinity uptake [32]. Considering CCHA-2229 with 8 transmembrane domains, the gene works as an ion transporter when suffering from abiotic stress. In addition, CCHA-2114, which was induced more than 3.2-fold under salt stress in this study, belongs to the DJ-1/Pfpl family. The Saccharomyces cerevisiae HSP31 gene, also a member of the DJ-1/Pfpl family, protects cells against oxidative stress and complements other stress protection systems within the cell [24]. Although no similar sequences were found in the database, CCHA-2221 also improved the salinity tolerance of E. coli cells. CCHA-2221 was induced more than 2.57-fold under salt stress, supporting the idea that it plays a role in halotolerance. Finally, we found that to be induced more than 4.4-fold under salt stress, CCHA-2142 encodes a protein containing a DUF3431 domain.

In order to assess the function of these candidate halotolerance genes, CCHA-2142 was chosen as a representative gene to express in Arabidopsis thaliana. As is known to all, chlorophyll content, MDA content, and the degree of membrane damage are important indicators of injury and aging [33, 34]. Under salt stress, plant cells produce scavengers (mainly SOD) to remove free radicals induced by environmental stress. SOD can eliminate the accumulation of reactive oxygen species to reduce the damage to cellular membrane structures [35]. Abiotic stress can decrease chlorophyll content by inhibiting chloroplast formation and biosynthesis rates [36]. In our study, the levels of SOD in CCHA-2142-expressing plants under salt stress were higher than those in control plants, suggesting that CCHA-2142 overexpression may enhance the tolerance of transgenic plants through influencing the levels of reactive oxygen scavengers. The CCHA-2142-expressing plants appeared to suffer less stress in the presence of salt, as evidenced by less reduction in chlorophyll content, less MDA accumulation, and less membrane damage compared to control plants. These data further confirmed and highlighted the functions of A. glaucus CCHA-2142 in enhancing plant salt tolerance. However, the functional mechanism of this gene still needs further research. In view of the constitutive CaMV 35S promoter is
Before treatment Treatment for 3 d After treatment for 7 d rewater for 14 d

Figure 4: Physiological comparison of control and CCHA-2142-overexpressing Arabidopsis thaliana. (a) Salt tolerance of transgenic plants and WT. Delayed senescence was observed for the transgenic line as compared with WT upon stress with 200 mM NaCl for 3 d. Comparison of SOD activity (b), malondialdehyde (MDA) content (c) and electrical conductivity (d), total chlorophyll content (e), chlorophyll a content (f), and chlorophyll b content (g) between CCHA-2142-transformed and wild-type Arabidopsis thaliana plants under 200 mM NaCl stress for 1 d, 3 d, and 5 d. 0 d: without stress. Lane CK: negative control (wild-type Arabidopsis thaliana); Lines 2142-1, 2142-6, and 2142-12: transgenic Arabidopsis thaliana lines. Data represent means ± standard deviations (SD) of three biological replicates. One asterisk and two asterisks indicate significant differences at $P \leq 0.05$ and $P \leq 0.01$ compared with WT by Student's $t$-test, respectively.
active in all tissues and growth phases, the stress-inducible promoter rd29A is considered in the further experiment. This kind of promoter could direct gene expression only in induced osmotic stress and minimize the negative effective effects on plant growth [37, 38]. Also as there may be codon preference in different species, it would be considerable to improve the genetic code of these target genes due to the codon usages in our donor and recipient strains.

In this work, we identified a number of candidate genes related to salt tolerance in A. glaucus, which may shed light on novel mechanism of salinity tolerance. Overall, this study provided an important contribution for our better understanding of stress tolerance and showed that intensified studies of this halophilic eukaryotic microorganism are warranted to promote understanding of the targets, processes, and networks involved in salt tolerance, to increase our understanding of fundamental stress responses, and to identify genes that might enhance the properties of plants. However, It remains to identify these salinity stress induced genes to reveal a clear pathway for further experimentation.

5. Conclusion

In our study, a cDNA library from A. glaucus under salt stress was constructed by random overexpression, and a lot of genes related to salt tolerance were functionally annotated. Subsequently, five putative clones were selected from E. coli cells using LB agar with 1 M NaCl. Meanwhile, the analysis of CCHA-2142-expression transgenic Arabidopsis thaliana confirmed this gene could increase salt stress tolerance. All these results showed the utility of this cDNA library as a tool for isolating genes related to salt tolerance.

Conflict of Interests

The authors have declared that no conflict of interests exists.

Authors’ Contribution

Jie Fang, Xiaojiao Han, and Lihua Xie contributed equally to the work.

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