Potato NAC Transcription Factor StNAC053 Enhances Salt and Drought Tolerance in Transgenic Arabidopsis

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Abstract: The NAC (NAM, ATAF1/2, and CUC2) transcription factors comprise one of the largest transcription factor families in plants and play important roles in stress responses. However, little is known about the functions of potato NAC family members. Here we report the cloning of a potato NAC transcription factor gene StNAC053, which was significantly upregulated after salt, drought, and abscisic acid treatments. Furthermore, the StNAC053-GFP fusion protein was found to be located in the nucleus and had a C-terminal transactivation domain, implying that StNAC053 may function as a transcriptional activator in potato. Notably, Arabidopsis plants overexpressing StNAC053 displayed lower seed germination rates compared to wild-type under exogenous ABA treatment. In addition, the StNAC053 overexpression Arabidopsis lines displayed significantly increased tolerance to salt and drought stress treatments. Moreover, the StNAC053-OE lines were found to have higher activities of superoxide dismutase (SOD), catalase (CAT), and peroxidase (POD) under multiple stress treatments. Interestingly, the expression levels of several stress-related genes including COR15A, DREB1A, ERD11, RAB18, ERF5, and KAT2, were significantly upregulated in these StNAC053-overexpressing lines. Taken together, overexpression of the stress-inducible StNAC053 gene could enhance the tolerances to both salt and drought stress treatments in Arabidopsis, likely by upregulating stress-related genes.

Keywords: potato; NAC transcription factor; ABA; abiotic stress

1. Introduction

Abiotic stresses such as high salinity, drought, and extreme temperatures negatively affect plant growth and development, causing yield loss in crop plants [1]. In order to survive adverse environmental conditions, plants have developed various adaptive mechanisms that involve changes at both physiological and biochemical levels during evolution [2]. The molecular mechanisms that cope with abiotic stresses have been studied extensively in plants, and transcriptional regulation of gene expression exerts a significant role in this process. Members of several transcription factor families in plants, including NAC, WRKY, MYB, and bZIP, have been functionally characterized to be involved in transcription regulation of plant stress responses [2–5].

NAC proteins constitute one of the largest plant-specific transcription factor families, and have been shown to be involved in various developmental processes in plants, including leaf senescence [6,7], secondary wall formation [8,9], lateral root development [10–12], shoot apical meristem development [13,14], floral development [15], plant hormone signaling [16–18], and cell division [19]. Plant NAC proteins contain a highly conserved DNA-binding domain named NAC domain in the N-terminal region and a variable transcription regulatory domain in the C-terminal region. So far, a large number of NAC transcription
factors have been successively identified in different plant species, including 117 genes in Arabidopsis, 151 in rice (Oryza sativa) [20], 110 in potato (Solanum tuberosum) [21], 168 in durum wheat (Triticum turgidum L. ssp. Durum) [22], 152 in tobacco (Nicotiana tabacum) [23], 152 in soybean (Glycine max) [24], and 82 in melon (Cucumis melo) [25].

It has been reported that some NAC genes exert central roles in plants’ response to various abiotic stresses, including in the model plant Arabidopsis and crop plants such as rice (Oryza sativa), soybean (Glycine max), wheat (Triticum aestivum), and maize (Zea mays). ANAC019, ANAC055, and ANAC072 were demonstrated to significantly increase drought tolerance in transgenic Arabidopsis via binding to and activating the ERD1 gene, whereas ANAC069 and AtNAP function to negatively regulate salt stress tolerance through reducing ROS scavenging ability and repressing ARBE1, respectively [26–28]. In rice, the expression of SNAC3 was significantly induced by drought, salt, high temperature stress and ABA treatment, and its overexpression resulted in increased drought and high-temperature stress tolerance [29]. OsNAC10, an ABA-dependent NAC transcription factor, could enhance tolerance against drought stress and improve grain yield under field drought conditions in transgenic rice [30]. In rice, ONAC022-overexpressing transgenic plants showed an increased tolerance against salt and drought stresses via an ABA-mediated pathway [31]. Two soybean NAC proteins, GmNAC11 and GmNAC20, were identified to be able to strongly improve tolerance against high salinity stress in transgenic plants. Meanwhile, GmNAC20-overexpression also led to increased tolerance to freezing stress [32]. In wheat, overexpression of TaNAC2 conferred strong drought, salt and freezing stress tolerance to transgenic plants [33]. The stress-inducible TaNAC4 was found to function as a transcriptional activator involved in wheat responses to biotic and abiotic stresses [34,35]. Ectopic expression of TaSNAC11-4B in Arabidopsis promoted ROS accumulation and significantly accelerated drought and ABA-induced leaf senescence [36]. As a positive regulator in maize’s response to drought stress, ZmNAC111 expression was found to be higher in drought-tolerant lines and lower in maize plants that were more sensitive to water-stressed conditions [37]. When a maize drought-responsive NAC transcription factor, ZmNAC55, was used to transform Arabidopsis, the transgenic plants showed significantly enhanced drought tolerance [38]. ZmSNAC1 is another stress-responsive gene that enhanced drought tolerance in transgenic Arabidopsis [39]. Overall, results from these studies showed that NAC proteins play important roles in plants’ response to abiotic stresses, and changes in their expression can alter abiotic stress tolerance in plants.

Potato (Solanum tuberosum L.) is one of the most important horticultural crops and also the fourth principal food crop. The yield of potato is often affected by multiple abiotic stresses, especially drought and salt stresses. Higher tolerance to drought and salt stresses is thus among the major targets in potato breeding. In this study, StNAC053, an NAC transcription factor from potato, was cloned and functionally characterized. Its expression pattern analysis indicated that StNAC053 was induced by high salinity, drought and ABA treatments. StNAC053 protein was found to be localized in the cell nucleus and could function as a transcriptional activator. Arabidopsis plants overexpressing StNAC053 were generated in this study, which showed enhanced tolerances to drought and high-salinity stresses compared to WT. These results suggested that StNAC053 play an important role in potato’s response to drought and high salt stresses.

2. Results
2.1. Isolation and Characterization of the StNAC053 Gene

In the current study, an NAC gene was cloned from potato. According to its chromosomal localization, it has been designated as StNAC053 (GenBank: EU049847) in a previous study [21]. The full length CDS of StNAC053 is 891 bp that encodes a 296 amino acid protein. The molecular weight (MW) and theoretical isoelectric point (pI) of the protein are 33,898.5 Da and 6.929, respectively. Multiple sequence alignments showed that StNAC053 possesses a highly conserved NAC domain (8-132) at the N-terminus and a transcription activation region at the C-terminus. The NAC structure contained five subdomains.
(A, B, C, D, E) which have been predicted to function as a DNA-binding domain. Further, a nuclear localization signal (NLS) was identified in the subdomain D (Figure 1).

Figure 1. Multiple sequence alignment of StNAC053 with ten other NAC proteins from *Arabidopsis* and potato. The locations of nuclear location signal (NLS) and the five subdomains (A–E) are indicated by black lines.

2.2. Phylogenetic and Motif Analysis of the StNAC053 Protein

To investigate the evolutionary relationship between StNAC053 and other NAC proteins, we constructed a neighbor-joining tree of the StNAC053 protein sequence with 17 other plant NAC proteins from *Arabidopsis*, wheat, rice, soybean, and potato. As a result, StNAC053 was clustered together with ATAF1/ANAC002 and ATAF2/ANAC081 (Figure 2A), which are both abiotic stress-responsive genes [40]. Subsequently, the MEME online tool was used to analyze the conserved domains of StNAC053 and other plant NAC proteins. A total of 10 conserved functional domains were identified (Supplementary Figure S1). Among them, motif 2 represents the subdomain A and subdomain B. Motif 1 corresponds to subdomain C. Motif 3 corresponds to subdomain D. Motif 4 corresponds to subdomain E (Figure 2B). These results indicated that StNAC053 encodes a stress-responsive NAC transcription factor.
2.3. Expression Patterns and Promoter Analysis of StNAC053

To further investigate the gene expression pattern of StNAC053, qRT-PCR was conducted to detect the transcription levels of StNAC053 in different tissue types and in response to various abiotic stresses. The results showed that StNAC053 had relatively high expression levels in roots, root tips, stems, tubers, and the highest expression level was observed in senescent leaves. StNAC053 expression was relatively low in young leaves and stem tips (Figure 3A). Whole plants of treated potato seedlings were collected for qRT-PCR assays. Under salt stress conditions, the expression of StNAC053 was induced during 1–6 h after treatments and reached peak levels with a 32.6-fold increase at 3 h (Figure 3B). StNAC053 transcript was upregulated specifically at 1 and 3 h after drought stress and ABA treatments, and reached peak levels with 14.5- and 5.5-fold increases, respectively, at 1 h (Figure 3C,D). The cold stress treatments induced StNAC053 expression within 1 h, and then the expression declined until 6 h (Figure 3E). These results indicated that the expression of StNAC053 can be induced by salt and drought stresses. Promoter analysis showed that the StNAC053 promoter contains one MYB binding site (MBS) that is related to drought inducibility, one stress-responsive element (TC-rich repeats), and four elements related to ABA responsiveness (ABRE), suggesting that the StNAC053 might be involved in plants’ response to ABA-mediated abiotic stresses (Supplementary Figure S2).

2.4. Subcellular Location and Transactivation Activity Assay

In order to analyze the subcellular localization of the StNAC053 protein, the 35S::GFP-StNAC053 fusion vector and 35S::GFP empty vector (control) were transiently expressed in N. benthamiana and the subcellular localization of the GFP signals was observed by confocal microscopy. The green fluorescence of the StNAC053-GFP fusion proteins was only observed in the nucleus, whereas the signals of the control were distributed on plasma membranes and in the cytoplasm and nucleus (Figure 4). These results demonstrated that the StNAC053 protein was located in the nucleus.
Figure 3. The expression pattern of StNAC053 in potato. (A) The expression profiles of StNAC053 in seven tissues. R: root; RT: root tip; S: stem; ST: stem tip; YL: young leaf; SL: senescent leaf; T: tuber. (B–E) The expression pattern of StNAC053 under salt, drought, ABA, and cold treatments. Whole plants of treated potato seedlings were collected for qRT-PCR assays.

Figure 4. The subcellular localization of StNAC053 in N. benthamiana epidermal cells. GFP and StNAC053-GFP fusion construct under the control of CaMV-35S promoter were transiently expressed into N. benthamiana epidermal cells, respectively. Bar = 20 μm, DAPI, 4,6-diamidino-2-phenylindole for nuclear staining.

To investigate the transcriptional activation ability of StNAC053, the full length CDS, N-terminal region, and C-terminal region of the coding sequence were inserted into the pBridge vector, respectively (Figure 5A). The three constructs and pBridge empty vector (control) were then transformed into the yeast strain AH109, and all transformed yeast cells grew well on the SD/Trp medium. Subsequently, these well-grown yeast cells were transferred from SD/Trp medium to SD/Trp/gal medium. Activation of the reporter gene in the yeast cells was determined by assays of the β-galactosidase activity with x-gal as substrate. The yeast strain containing the full-length StNAC053 (GAL4BD-StNAC053) and the C-terminal of StNAC053 (GAL4BD-StNAC053C) were blue on the SD/Trp/gal medium, whereas the cells with the N-terminal of StNAC053 (GAL4BD-StNAC053N) and the pBridge empty vector were not blue (Figure 5B). The above described results showed that StNAC053 is a transcriptional activator, and its transactivation domain is in the C-terminal.
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![Diagram of yeast transformation](image)

Figure 5. Transactivation assay of StNAC053 in yeasts. (A) Different truncations of StNAC053 fused to the GAL4 DNA binding domain in pBridge vector. (B) Transactivation activity assay of StNAC053 in yeast strain AH109. β-galactosidase activity against x-gal was detected on the SD/-Trp medium.

### 2.5. StNAC053 Overexpression in Arabidopsis Confers ABA Hypersensitivity

In order to further study the roles of StNAC053 in response to salt and drought stresses, transgenic *Arabidopsis* plants overexpressing *StNAC053* were generated. Eight T0 transgenic lines were verified by normal PCR. The homozygous T3 transgenic lines were selected, and the expression level of *StNAC053* was analyzed using qRT-PCR. Two overexpression lines (OE2 and OE6) with higher expression levels of *StNAC053* were selected for further phenotypic analysis (Supplementary Figure S3).

As described above, expression of *StNAC053* in potato seedlings was upregulated within 3 h after ABA treatment and reached a peak at 1 h (Figure 3D). In order to find out whether StNAC053 is involved in ABA response, we conducted a germination assay to
study the sensitivity of \textit{StNAC053-OE} lines to ABA treatments. The results showed that without ABA treatment, the germination rates were almost identical between the \textit{StNAC053-OE} lines and WT. However, when grown on half-MS media containing 0.5 or 1.0 µM ABA, the germination rates of both \textit{StNAC053-OE} lines and WT reduced significantly, while germination rates of the \textit{StNAC053-OE} lines were significantly lower than that of WT (Figure 6A), indicating that \textit{StNAC053-OE} plants were more sensitive to exogenous ABA in the seed germination assay (Figure 6B).

![Figure 6. Comparison of the ABA sensitivity between \textit{StNAC053} overexpression and wild-type \textit{Arabidopsis}. (A) Seed germination assay between \textit{StNAC053} overexpression and wild-type \textit{Arabidopsis} under 0.5 and 1 µM ABA treatments. The seed germination rates were recorded six days after sowing. (B) The statistics of germination rates under normal condition, 0.5 and 1 µM ABA treatments. WT, wild-type. The data were means ± SD from three independent replications. ** $p < 0.01$ (t-tests).]
2.6. StNAC053 Overexpression Enhances Salt Tolerance in Transgenic Arabidopsis

Germination assays were also used to investigate the salt tolerance of StNAC053-OE lines. As a result, the StNAC053-OE lines displayed higher germination rates than WT on half-MS media containing 100 and 150 mM NaCl (Figure 7).

Figure 7. Comparison of seed germination between StNAC053 overexpressing and wild-type Arabidopsis under salt stress treatments. (A) Seed germination assay between StNAC053 overexpression and wild-type Arabidopsis under 100 and 150 mM NaCl treatments. The seed germination rates were recorded six days after sowing. (B) The statistics of seed germination under normal condition, 100 and 150 mM NaCl treatments. WT, wild-type. The data were means ± SD from three independent replications. * p < 0.05, ** p < 0.01 (t-tests).
In addition, a root elongation assay was conducted to further determine the response of the StNAC053-OE lines to salt stress. Seven days after growing on half-MS media containing 100 or 150 mM NaCl, primary roots of the StNAC053-OE lines were found to be significantly longer compared to that of the WT (Figure 8). Taken together, results from the salt treatment assays suggested that overexpression of StNAC053 improved the tolerance to salt stress in transgenic Arabidopsis plants.

Figure 8. Comparison of root growth between StNAC053 overexpressing and wild-type Arabidopsis under salt stress treatments. (A) The primary root length of StNAC053 overexpressing and wild-type Arabidopsis under 100 and 150 mM NaCl treatments. The primary root length were recorded seven days after growth. Bar = 1.5 cm. (B) The statistics of primary root length under normal condition, 100 and 150 mM NaCl treatments. WT, wild-type. The data were means ± SD from three independent replications. * p < 0.05 (t-tests).
2.7. StNAC053 Overexpression Improves Drought Tolerance in Transgenic Arabidopsis

To study the role of StNAC053 in drought response, one-week-old seedling of StNAC053-OE, StNAC053-OE6, and WT were transferred from half-MS media to water-saturated matrix and grown for four weeks. Subsequently, these plants were treated with drought stress via water withholding. After two weeks of drought treatments, while most of the WT plants wilted, the StNAC053-OE plants displayed slight leaf rolling (Figure 9A). After rewatering for three days, only 37% of the WT plants were recovered, whereas 82% and 78% of plants of the two StNAC053-OE lines survived, respectively (Figure 9B). Furthermore, the StNAC053-OE plants showed higher water content than WT plants during the 3 h dehydration (Figure 9C). These results suggested that the StNAC053 overexpression improved the tolerance of the Arabidopsis plants to drought stress.

Figure 9. Transgenic StNAC053 Arabidopsis showed higher drought tolerance. (A) Phenotypes of StNAC053 overexpressing and wild-type Arabidopsis under drought stress. The StNAC053 overexpressing and wild-type Arabidopsis were grown for four weeks under normal conditions. Watering was ceased for two weeks, followed by rewatering for recovery. (B) Statistical analysis of survival rates after three days for rewatering. (C) Water loss rates of detached rosette leaves of StNAC053 overexpressing and wild-type Arabidopsis. WT, wild-type. The data were means ± SD from three independent replications. *p < 0.05 (t-tests).

2.8. StNAC053 Overexpression Enhances ROS-Scavenging Capability in Transgenic Arabidopsis

Changes in reactive oxygen species (ROS) are often associated with plants’ response to abiotic stresses. In order to determine whether changes of antioxidant capacity are associated with increased tolerance to salt and drought stresses of the StNAC053-OE plants, we measured the stress-related physiological parameters of StNAC053-OE lines and WT plants with or without stress treatments for three days (Figure 10). The activities of the ROS-scavenging enzymes CAT (catalase), POD (peroxidase), and SOD (superoxide) were comparable between the StNAC053-OE lines and WT without drought or salt stress. When subjected to salt or drought stress, the activities of CAT, POD, and SOD in plants of StNAC053-OE lines and WT were all increased. However, significantly more increase of ROS-scavenging activities was observed in the StNAC053-OE lines than in WT. Moreover, there was no significant difference in MDA (malonic dialdehyde) contents between the StNAC053-OE lines and WT without stress treatments. However, significantly lower MDA levels were detected in StNAC053-OE plants when subjected to salt or drought stress.
These results suggested that StNAC053 transgenic Arabidopsis plants might have enhanced stress tolerance due to increased ROS-scavenging capability.

**Figure 10.** Changes of antioxidant enzyme activities and MDA contents in StNAC053 overexpressing and wild-type Arabidopsis after treatments with drought or salt stress. MDA, malonic dialdehyde; CAT, catalase; POD, peroxidase; SOD; superoxide. WT, wild-type. The data were means ± SD from three independent replications. *p < 0.05.

2.9. StNAC053 Regulates the Expression of Stress-Responsive Genes

qRT-PCR was performed to examine whether StNAC053 regulates salt and drought responses through regulating expression of stress-responsive genes. There was no significant difference in the expression of selected stress-responsive genes between StNAC053-OE and WT plants under normal growth conditions. When subjected to drought or salt stress, the expression of the stress-responsive genes was upregulated in both StNAC053-OE and WT plants. However, compared to the WT, the expression levels of the stress-responsive genes were significantly higher in the StNAC053-OE lines (Figure 11). These results suggested that StNAC053 transgenic plants might enhance drought or salt tolerance by activating the expression of stress-responsive genes.
3. Discussion

The NAC transcription factors comprise one of the largest transcription factor families in plants and play multiple roles in stress responses. In a previous study, a total of 110 potato NAC family members were identified and several of them were identified to be highly induced by various stress treatments [21]. A number of NAC genes has been characterized and analyzed in different plant species, including *Arabidopsis* and rice. The yield of potato could be severely affected by various abiotic stresses, especially salt and drought. However, little information is available on the biological functions of the potato NAC genes. In the present study, an NAC transcription factor was cloned from potato, which possessed a typical NAC structure and was classified in the ATAF subgroup (Figures 1 and 2A).

In plants, a number of NAC transcriptional factors have been identified to be involved in ABA-mediated abiotic stress responses [40,41]. In this study, we found that the promoter region of *StNAC053* possessed four ABRE motifs, which has been known as an important cis-element involved in ABA signaling (Supplementary Figure S2). The results from expression analysis revealed that *StNAC053* was significantly induced by ABA treatments (Figure 3D). Furthermore, seed germination rates of *StNAC053*-OE lines were significantly lower than those of WT plants, suggesting that overexpression of *StNAC053* caused transgenic plants to be hypersensitive to ABA (Figure 6).

Multiple NAC transcription factors have been reported to be involved in plants’ responses to abiotic stresses, including salt, drought, heat, and cold treatments [29,42]. In potato, *StNAC053* was found to be induced by various abiotic stress treatments, es-

![Figure 11. Expression of stress-responsive genes in *StNAC053* overexpressing and wild-type *Arabidopsis* after drought or salt stress treatments. WT, wild-type. The data were means ± SD from three independent replications. * p < 0.05, ** p < 0.01.](image-url)
pecialy drought and salt stresses, implying that StNAC053 may play important roles in potato’s response to drought and salt stresses (Figure 3). In this study, we focused on the role of StNAC053 in salt and drought stress responses. Under high salinity conditions, the germination rates and root lengths of StNAC053-OE plants were significantly higher than those of WT (Figures 7 and 8). In addition, StNAC053-OE plants had higher survival rates than WT plants under drought conditions (Figure 9). Thus, our results showed that StNAC053-OE Arabidopsis transgenic plants had significantly enhanced salt and drought tolerance. Similarly, overexpression of the StNAC053 homolog ATAF1 conferred drought tolerance in Arabidopsis [43]. Additionally, under drought or salt stress, a large amount of ROS were produced. Excessive ROS could damage the cellular membrane [44], resulting in the generation of massive secondary products such as MDA [45]. The antioxidant enzymes played an important role in coping with excess ROS to reduce oxidative stress. CAT, POD, and SOD are among the three most important antioxidant enzymes [44]. After drought or salt stress, the levels of MDA in StNAC053-OE lines were significantly decreased compared with WT, suggesting that StNAC053-OE lines suffered less oxidative damage (Figure 10). Meanwhile, the enzyme activities of CAT, POD, and SOD were significantly higher, suggesting that StNAC053-OE lines have higher ROS-scavenging capacity (Figure 10). Notably, although growth retardation is usually accompanied with the overexpression of NAC transcription factors [46,47], no significant developmental difference was observed between the transgenic plants and WT in this study.

Most of the NAC transcription factors were involved in stress responses function as transcription activators in regulating downstream stress-responsive genes [48]. AtJUB1 can directly activate the transcription of AtDREB2A in Arabidopsis [49]. Rice ONAC066 enhanced drought tolerance through regulating the expression of OsDREB2A [50]. In the current study, StNAC053 was identified to function as a transcription activator (Figures 4 and 5) and positively regulate the expression of multiple stress-responsive genes such as COR15A, DREB1A, KAT2, etc. (Figure 11). These genes encode stress-associated proteins that act as key mediators in protecting plants from oxidative or osmotic damages. The promoter regions of all these genes were found to harbor NAC binding sites (Supplementary Table S1) to which the StNAC053 transcription factor might bind when activating the expression of these stress-responsive genes under stress conditions. StNAC053 holds a promising utility as an excellent candidate in the genetic improvement of potato abiotic stress tolerance because it positively regulates plant stress tolerance but does not have any negative influence on plant growth.

4. Materials and Methods
4.1. Potato Plant Preparation and Treatments
Potato cultivar GN2 was used to analyze the expression of StNAC053 gene in this study. Potato sprouts were incubated on complete MS solid media by nodule cutting and cultivated in a growth chamber at 24 °C under continuous light. For stress treatments, seedlings were treated with dehydration, at 4 °C, with NaCl (200 mM), and ABA (50 µM) for 0, 1, 3, and 6 h, respectively. All of the harvested samples were immediately frozen in liquid nitrogen, and stored at −80 °C, prior to RNA extraction. Three biological replicates were used for each sample. The Arabidopsis thaliana Col-0 plants were used for transgenic study of StNAC053.

4.2. RNA Extraction and qRT-PCR
Total RNAs of the tested samples were extracted using the Ultrapure RNA Kit (cw-biotech, Beijing, China), and the first-strand complementary DNA (cDNA) was synthesized using the PrimeScript™ RT reagent Kit (TaKaRa, Dalian, China). Expression of six stress-responsive genes from Arabidopsis including COR15A (U01377.1), DREB1A (AB013815.1), ERD11 (D17672.1), RAB18 (X68042.1), ERF5 (NM_124094.3), and KAT2 (NM_001341273.1) was determined. The potato EF1α gene and Arabidopsis Actin 2 gene were used as internal controls [21,39], and quantitative real-time PCR (qRT-PCR) reactions were performed with
40 cycles in a Roche LightCycler 480 Real-Time PCR instrument. All expression data were obtained from three technical repeats, and calculated by the $2^{-\Delta\Delta CT}$ method. The primer sequences used in the current study are listed in Supplementary Table S2.

4.3. Cloning and Sequence Analysis of StNAC053

The coding sequence of StNAC053 was amplified using RT-PCR. The purified PCR products were inserted into the subclone vector pEASY-Blunt, and the constructed vector was then transformed into E.coli trans T1 competent cells (Trans Gen, Beijing, China). After sequencing, the recombinant plasmid was named Blunt-StNAC053. The ProtParam online tool (https://web.expasy.org/protparam/) (accessed on 20 January 2020) was used to analyze the molecular weight, isoelectric point of the StNAC053 protein. The NLS of NAC proteins was predicted via eNLS Mapper (http://nls-mapper.lab.keio.ac.jp) (accessed on 20 January 2020). Multiple sequence alignment was carried out using the DNAMAN tool. A neighbor-joining tree was generated using MEGA6 with 1000 replications. The NAC protein sequences were obtained from the TAIR (https://www.arabidopsis.org/) (accessed on 20 January 2020), SGN (http://solgenomics.net/) (accessed on 20 January 2020), and NCBI. The conserved motifs of NAC protein sequences were analyzed via MEME (http://meme-suite.org/) (accessed on 20 January 2020) with default parameters. To assess the promoter cis-acting elements of the StNAC053 gene, 2000 bp of promoter regions upstream of the start codon of the StNAC053 gene were extracted. PlantCARE (http://bioinformatics.psb.ugent.be/webtools/plantcare/html/) (accessed on 20 January 2020) was used for cis-acting regulatory element investigation.

4.4. Subcellular Localization

The coding sequence (without stop codon) of StNAC053 was amplified using PCR. The purified PCR products were inserted into the pCHF3-cGFP vector at the speI site by Infusion (Clontech), yielding plasmid 35S::StNAC053-GFP after sequencing. The construct was then transformed into Agrobacterium GV3101 competent cells, and transiently expressed in the leaves of Nicotiana benthamiana [51]. Simultaneously, the empty pCHF3-cGFP vector was used as a control. Three days after injection, the leaves were soaked in the DAPI (dye 4,6-diamidino-2-phenylindole) staining solution and observed under a confocal laser microscope (TCS-SP8 Leica, Wetzlar, Germany) to detect fluorescence signals of the fusion protein, as previously reported [52].

4.5. Transactivation Assay

For transactivation activity assay of StNAC053, the full length CDS and two truncated sequences of StNAC053 genes were amplified using PCR and inserted into the pBridge vector at the EcoRI site by Infusion (Clontech, Beijing, China) to fuse with a GAL4 DNA binding domain, yielding plasmid pBridge-StNAC053. Plasmid pBridge-StNAC053 and empty pBridge vector (control) were then transformed into the yeast strain AH109, respectively. The transformed yeasts were plated on SD media lacking tryptophan (SD/-Trp) and incubated at 30 °C for 3 days. Subsequently, the positive clones were transferred to SD/-Trp media supplemented with x-gal and incubated at 30 °C for 3 days. Transactivation activity of the fused proteins was determined based on the growth status (blue/white) of the transformants.

4.6. Generation of StNAC053 Transgenic Arabidopsis Plants

For overexpression analysis, the full-length CDS of StNAC053 was amplified using PCR and inserted into the pCHF3 vector at the SacI site by Infusion (Clontech, Beijing, China), yielding plasmid 35S::StNAC053 after sequencing. The construct was then transformed into Agrobacterium GV3101 competent cells. The positive colonies of Agrobacterium were then selected and used in transforming Arabidopsis Col-0 plants by the floral dip method [53]. The T0 generation seeds were screened by half-strength MS media with
50 mg/L kanamycin to obtain StNAC053 overexpression plants. T3 homozygous lines were selected for further phenotypic analysis.

4.7. Seedling Growth Assays

For seed germination assays, Arabidopsis seeds were firstly sterilized using 75% alcohol and then evenly sown on half-MS media containing ABA (0.5 and 1.0 µM) or NaCl (100 and 150 mM), respectively. Seeds were stratified at 4 °C for 2 days under darkness, and then moved to a growth chamber (23 °C, continuous light). After cultivation for 6 days, the germination rates were calculated with three replications.

For root elongation assays, Arabidopsis seedlings were grown normally on half-MS media for 7 days. Seedlings with similar growth status were then transferred to half-MS media containing 100 and 150 mM NaCl, respectively. The length of the primary root was measured after 7 days with three replications.

4.8. Drought Stress Tolerance Assays of StNAC053 Transgenic Arabidopsis Plants

For the drought tolerance assays, one-week-old seedlings germinated on half-MS media were transferred into pots containing soil mixture (peat moss and vermiculite, 3:1 v/v). Four-week-old plants grown under normal conditions were exposed to drought stress for 14 days. The plants were then rewatered for 3 days, and the survival rates were recorded. The detached leaves were air-dried for 3 h, and were weighed at 5 time points (0, 45, 90, 135, and 180 min). The rate of water loss was calculated based on the weight loss at each time point divided by the initial fresh weight.

4.9. Physiological Measurements

The physiological parameters were measured using fully expanded leaves obtained from well-watered, drought-stressed, or salt-stressed plants. The content of MDA and the enzymatic activities of superoxide (SOD), peroxidase (POD), and catalase (CAT) were determined as previously described [54]. Three biological replicates were performed.

4.10. Statistical Analysis

The statistical significance was analyzed using the GraphPad Prism 8 t test. p values less than 0.05 or 0.01 were regarded as significantly different from the control. All data were obtained from three replicates.

5. Conclusions

In this study, a typical NAC transcription factor StNAC053 was cloned and identified to be a transcription activator. The Arabidopsis plants overexpressing StNAC053 displayed significantly increased tolerance to salt and drought stress treatments, while the overexpression lines seem to be more sensitive to ABA treatments. Taken together, the transcription factor StNAC053 might be involved in drought and salt stress responses via ABA-mediated signaling in potato.

Supplementary Materials: The following are available online at https://www.mdpi.com/1422-0067/22/5/2568/s1, Figure S1: The putative conserved motifs in NAC proteins, Figure S2: Stress-related elements in the promoter regions of StNAC053 gene, Figure S3: The expression level of the StNAC053 gene in wild-type and two overexpression lines, the ratios of gene expression levels were calculated relative to the wild-type, ** p < 0.01, Table S1: Prediction of NAC binding cis-elements in the promoter of stress-responsive genes, Table S2: The primers used in this study.

Author Contributions: Q.W., and C.G., conducted the research and participated in drafting the manuscript. Z.L., J.S., Z.D., and L.W. assisted in data collection and analysis. X.L. and Y.G. conceived this research, designed the experiments and drafted the manuscript. All authors have read and agreed to the published version of the manuscript.

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