The NAC-type transcription factor CaNAC46 regulates the salt and drought tolerance of transgenic Arabidopsis thaliana

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Abstract

Background: The NAC (NAM, ATAF1/ATAF2, and CUC2) transcription factors belong to a large family of plant-specific transcription factors in monocot and dicot species. These transcription factors regulate the expression of stress tolerance-related genes that protect plants from various abiotic stresses, including drought, salinity, and low temperatures.

Results: In this study, we identified the CaNAC46 transcription factor gene in Capsicum annuum. Its open reading frame was revealed to comprise 921 bp, encoding a protein consisting of 306 amino acids, with an isoelectric point of 6.96. A phylogenetic analysis indicated that CaNAC46 belongs to the ATAF subfamily. The expression of CaNAC46 was induced by heat, cold, high salt, drought, abscisic acid, salicylic acid, and methyl jasmonate treatments. Thus, CaNAC46 may be important for the resistance of dry pepper to abiotic stresses. A subcellular localization analysis confirmed that CaNAC46 is localized in the nucleus. The overexpression of CaNAC46 improved the tolerance of transgenic Arabidopsis thaliana plants to drought and salt stresses. The CaNAC46-overexpressing lines had longer roots and more lateral roots than wild-type lines under prolonged drought and high salt stress conditions. Additionally, CaNAC46 affected the accumulation of reactive oxygen species (ROS). Moreover, CaNAC46 promoted the expression of SOD, POD, RD29B, RD20, LDB18, ABI, IAA4, and P5CS. The malondialdehyde contents were higher in TRV2-CaNAC46 lines than in wild-type plants in response to drought and salt stresses. Furthermore, the expression levels of stress-responsive genes, such as ABA2, P5CS, DREB, RD22, CAT, and POD, were down-regulated in TRV2-CaNAC46 plants.

Conclusions: Under saline and drought conditions, CaNAC46 is a positive regulator that activates ROS-scavenging enzymes and enhances root formation. The results of our study indicate CaNAC46 is a transcriptional regulator responsible for salinity and drought tolerance and suggest the abiotic stress-related gene regulatory mechanisms controlling this NAC transcription factor are conserved between A. thaliana and pepper.

Keywords: Pepper, NAC transcription factor, Stress tolerance, Transgenic plants
Background

Pepper (Capsicum annuum; family: Solanaceae) is a popular vegetable crop cultivated worldwide. China is the largest producer and consumer of chili peppers, with approximately 40% of the global chili cultivation area [1]. In addition to being widely used as a condiment, dry peppers are also used in the medical and military industries [2]. However, there are some environmental conditions, such as salinization and drought, that adversely affect the main dry pepper-producing areas.

Abiotic stresses, including drought, high-salinity, and temperature extremes, are the main factors responsible for decreased dry pepper quality [3]. To protect against the detrimental effects of environmental stresses, plants have gradually evolved appropriate defense mechanisms, in which the regulation of gene expression plays an important role. Transcription factors are critical regulators of abiotic stress tolerance [4]. In plants, the large NAC (NAM, ATAF1/ATAF2, and CUC2) transcription factor family is involved in abiotic stress responses [5]. The N-terminal of NAC transcription factors is highly conserved, in contrast to the variable C-terminal [6]. This transcription factor family has recently been extensively studied in Arabidopsis thaliana, rice, tobacco, and soybean [7–10]. There has also been increasing interest in these transcription factors in pepper crops [11, 12]. A genome-wide analysis of 104 NAC transcription factors in hot pepper and an expression analysis of 22 NAC transcription factor genes revealed diverse responses to various stresses [13]. As a transcriptional activator of CaPLD4, CaNAC1 helps degrade phospholipids in green pepper in response to low temperatures [11]. The pepper CaNAC2 transcription level is reportedly considerably up-regulated by an abiotic stress treatment, but is down-regulated by osmotic stress and a salicylic acid (SA) treatment [14].

The CaNAC46 gene was identified during a search of a pepper cDNA library with A. thaliana AtNAC2/ATAF1/ANAC002 (At1g01720) as the query [14]. The CaNAC46 transcription factor helps regulate responses to abiotic stresses (e.g., drought, salinity, and low temperatures) and phytohormones [e.g., abscisic acid (ABA), methyl jasmonate (MeJA), and SA]. We demonstrated that CaNAC46 positively regulates plant tolerance to drought and salinity stresses, and developed a model for the role of CaNAC46 in the abiotic stress response system [15]. The results may provide the basis for a comprehensive analysis of the function and regulatory network of CaNAC46 related to plant stress resistance, with potential implications for the application of CaNAC46 in future genetic engineering experiments.

Results

Identification of the ATAF subfamily gene CaNAC46 in pepper

The NAC transcription factors are involved in mediating plant responses to abiotic stresses [16, 17]. The CaNAC46 gene was isolated from C. annuum var. ‘Qingnong No. 2’. The encoded protein comprised 306 amino acids, with a molecular weight of 34.94 kDa and a pI of 6.96 (Additional file 1: Table S1). Additionally, the proportions of basic and acidic amino acids were similar between CaNAC46 and the homologous NAC proteins in other model plants. There were also similarities in the other examined physical and chemical characteristics between CaNAC46 and its homologs in other model plants. These results suggest that NAC transcription factors have a conserved secondary structure.

In this study, CaNAC46 was clustered with ATAF1 [18], OsNAC5 [19], and OsNAC6 [20], which belong to the ATAF subfamily (Fig. 1A). A multiple sequence alignment revealed a sequence identity of 71.19%, implying that the NAC transcription factor sequences were highly conserved. The CaNAC46 C-terminal contains the EVQS [E/x] PK [W/I] sequence, which is consistent with the sequences of typical ATAF subfamily members [3]. A phylogenetic analysis revealed that CaNAC46 is most closely related to OsNAC5, OsNAC6, and ANAC2 of the ATAF subgroup (Fig. 1B).

CaNAC46 is localized in the nucleus

An analysis of the fluorescent signals for the green fluorescent protein (GFP) alone (control) and the CaNAC46-GFP fusion protein in tobacco cells with a confocal microscope indicated CaNAC46 is a nuclear protein (Fig. 2A).

CaNAC46 expression is induced by abiotic stresses and phytohormones

We examined the CaNAC46 expression level in response to various abiotic stresses over a 12-h treatment period. The expression of CaNAC46 was significantly up-regulated following drought, salt, heat, and cold stress treatments, especially from the 6-h time-point, with peak expression levels at 12 h (Fig. 2B). Additionally, CaNAC46 expression was also substantially induced by ABA, which is a phytohormone involved in salt and drought stress responses.

Overexpression of CaNAC46 enhances the drought and salinity tolerance of transgenic Arabidopsis thaliana

The effect of CaNAC46 overexpression on drought stress tolerance was investigated using transgenic plants treated with polyethylene glycol (PEG 6000). To assess the effect of CaNAC46 overexpression on salt stress tolerance, wild-type (WT) and transgenic seedlings were
grown on 1/2 MS agar medium supplemented with 200 mM NaCl (Fig. 3A). Under normal conditions, the germination rates of the WT and transgenic plants were approximately 95%. However, the germination rates under the simulated drought and salt stress conditions differed substantially between the WT and CaNAC46-overexpressing plants (Additional file 2: Figure S1).

The salinity and drought tolerance of the CaNAC46-overexpressing lines was further evaluated at the vegetative growth stage. Following the drought and high salt treatments, the leaves of the WT plants were severely wilted, but the leaves of the CaNAC46-overexpressing plants were not (Fig. 3B). And after dehydration treatments, plants were re-watered 3 days, the leaves of WT remained wilted, while the OE lines grown normally (Fig. 3B).

CaNAC46 overexpression promotes the root growth of transgenic Arabidopsis thaliana

Plant root and shoot systems modify signaling and metabolic pathways to adapt to complex environmental changes and maintain normal growth and development [21]. A developed root system is important for improving plant stress resistance. In the current study, the NaCl treatment inhibited the lateral root development of WT seedlings. In contrast, the root elongation of two analyzed CaNAC46-overexpressing lines was only slightly inhibited, ultimately resulting in more lateral roots on
the transgenic plants than on the WT plants (Fig. 3A and C). These observations implied that the overexpression of CaNAC46 may enhance abiotic stress resistance by promoting root growth (Fig. 3). Moreover, the expression levels of LBD18 and IAA4, which are involved in lateral root development, were significantly upregulated in the CaNAC46-overexpressing plants under drought and saline conditions (Fig. 4B).

**CaNAC46 overexpression decreases reactive oxygen species accumulation**

High CaNAC46 expression levels in the CaNAC46-overexpressing A. thaliana lines were confirmed by quantitative real-time PCR (qRT-PCR) (Additional file 3: Figure S2). Drought and salinity stresses usually induce the excessive accumulation of reactive oxygen species (ROS), which eventually leads to irreversible damages. We measured the SOD, POD, and CAT activities in the WT and CaNAC46-overexpressing plants under normal and stress conditions (Fig. 4A). There were no severe differences in the enzyme activities between the WT and transgenic plants under normal conditions. However, exposures to drought and salt stresses resulted in considerable increases in the SOD, POD, and CAT activities in both the WT and CaNAC46-overexpressing plants, but the increases were more pronounced in the CaNAC46-overexpressing plants.

**CaNAC46 overexpression activates the expression of stress-related genes**

To clarify the putative regulatory effect of CaNAC46 on abiotic stress tolerance, qRT-PCR analyses were completed to examine the expression patterns of stress-responsive genes (e.g., ABI, P5CS1, RD29B, and RD20) in the WT and CaNAC46-overexpressing plants before and after the abiotic stress treatments (Fig. 4B). We observed that a relatively low CaNAC46 expression level impaired the water-deficit response of whole plants and detached leaves. In contrast, a relatively high CaNAC46 expression level protected plant from the damages due to abiotic stresses.

**Silencing of CaNAC46 decreases plant tolerance to high salt and drought conditions**

An analysis of the CaNAC46 expression levels in the CaNAC46-silenced (TRV2-CaNAC46) and WT plants before the stress treatments confirmed that CaNAC46 expression was significantly down-regulated in the TRV2-CaNAC46 lines (Additional file 4: Figure S3). The TRV2-CaNAC46 and WT plants were then subjected to drought stress (i.e., dehydrated) and salinity stress (i.e., treated with a 200 mM NaCl solution). The TRV2-CaNAC46 lines showed a more weaker growth state than that of none-silenced lines under both normal and stress conditions (Fig. 5A). The cellular H$_2$O$_2$ level determined by diaminobenzidine (DAB) staining was higher in the TRV2-CaNAC46 leaves than in the WT leaves after 3-week salt and drought treatments (Fig. 5B). Additionally, the malondialdehyde (MDA) content, which may be used as an indicator of lipid peroxidation, increased considerably in the TRV2-CaNAC46 plants, but not in the WT plants, during the later stages of the drought and salinity treatments (Fig. 5C). Moreover, in the TRV2-CaNAC46 plants, CAT6 (encoding a catalase important for plant cell responses to stress), PR1, and RD22 expression levels were down-regulated and the ROS-scavenging activity was low, which increased the sensitivity of these plants to saline and drought conditions (Fig. 5D).

**Discussion**

Diverse stresses adversely affect the growth and productivity of dry pepper and represent the major yield-limiting factors. In plants, specific transcription factors enhance the resistance to external stresses [22]. We previously identified seven stress-responsive CaNAC46 transcription factors in Capsicum species. In the current
study, we proved that the overexpression of \textit{CaNAC46} increases the tolerance to various abiotic stresses, including drought and salinity.

The \textit{CaNAC46} gene cloned from \textit{C. annuum} 'Qingnong No. 2' consists of 921 bp that encode 306 amino acid residues. An analysis of the physicochemical properties revealed differences between \textit{CaNAC46} and NAC transcription factors from other plant species. This diversity may be relevant for future functional characterizations of NAC transcription factors. Additionally, \textit{CaNAC46} has three disordered amino acid regions, but their effects on the functions of the transcription factor remain to be determined. Ooka [8] classified NAC proteins into Groups I and II, with Group I divided into 14 subgroups. The NAC transcription factors in subgroups ATA \\ [23], AtNAC3 [24], and OsNAC3 [25] are involved in plant stress responses. The conserved sequences and phylogenetic relationships have been analyzed. The NAC transcription factors consist of a NAM subdomain with 160 amino acid residues as well as five other conserved subdomains (A, B, C, E, and D). In previous studies, the highly conserved subdomains A, C, and D were revealed to be important for NAC transcription factor functions. Subdomains C and D are mainly involved in DNA binding. The significance of subdomain A results from the insertion of the \textit{dTph1} transposable element to form the \textit{NAM} mutant [8]. Additionally, subdomains B and E vary among NAC proteins, suggesting they may be associated with the functional diversity of the NAC transcription factors [8]. Sequence comparisons with NAC family members confirmed that \textit{CaNAC46} belongs to the ATA subfamily, and is most closely related to \textit{A. thaliana} ATA1.

Plant root system development is a relatively complex process, enabling plants to adapt to various environmental stresses. Salinity stress inhibits root meristem growth by decreasing auxin levels [26], whereas drought stress limits root growth and inhibits cytokinin production [27]. Root size and architecture influence the uptake of
water and nutrients from the soil. In this study, CaNAC46-overexpressing plants produced longer roots and more lateral roots than WT plants following an exposure to drought and salinity stresses. Orthologs of LBD18 in nonleguminous plants are required for lateral root development [28]. In A. thaliana, lateral root formation is regulated by indole-3-acetic acid (IAA) [29]. Our data indicated that LBD18 and IAA4 expression levels were up-regulated in the CaNAC46-overexpressing plants in response to drought and high salt conditions. Accordingly, CaNAC46 may promote lateral root formation and root elongation by up-regulating LBD18 and IAA4 expression to enhance the tolerance to drought and salinity stresses. The germination rate of CaNAC46-overexpressing lines was significantly lower than that of WT plants following the salt and drought treatments. These treatments also resulted in higher chlorophyll contents in the transgenic plants than in the WT controls.

In plants, proline is necessary for growth and development. Additionally, proline protects cells from damage by acting as a radical scavenger and an osmotic agent [15]. In the TRV2-CaNAC46 lines, the down-regulated expression of P5CS increased the sensitivity of the plants to salt and drought stress conditions. A previous study indicated that RD26 is a transcriptional activator mediating ABA-inducible gene expression in plants subjected to abiotic stress [30]. The rice NAC family transcription factors OsNAP and OsNAC2 respond to abiotic stresses via the ABA pathway [31, 32]. The NAC transcription factors may minimize the accumulation of ROS and regulate the expression of stress-related genes under drought and high salt conditions [33]. We speculate that the observed enhanced tolerance to abiotic stresses is mainly due to the up-regulated expression of stress-responsive genes, including RD20, RD29B, DREB, and RD22. Following drought and salt treatments, RD20 and RD29B expression levels were up-regulated in the

![Fig. 4](image_url)

Fig. 4 The physiological and stress-responsive gene expression analysis under stress treatments. A The activities of SOD, POD, and CAT were measured in the WT and CaNAC46-overexpressing plants under normal and stress conditions. B The expression levels of stress-responsive genes were examined by qRT-PCR analyses the WT and CaNAC46-overexpressing plants before and after the abiotic stress treatments. Three independent biological experiments were performed. Superscript letters indicate significant differences (P < 0.05).
Fig. 5 Suppression of CaNAC46 leads to salt and drought sensitivity in pepper. The role of CaNAC46 for salt and drought sensitivity was assessed by VIGS. A The TRV2-CaNAC46 and WT plants were subjected to drought stress by dehydration and salinity stress by watering with a 200 mM NaCl solution for 3 weeks. B DAB staining was in TRV2-CaNAC46 and WT leaves than in WT leaves after for 3 weeks of the salt and drought treatments. C The MDA content of TRV2-CaNAC46 and WT plants was measured under drought and salinity stresses. D The expression levels of stress-responsive genes were examined by qRT-PCR analyses the WT and TRV2-CaNAC46 plants before and after the abiotic stress treatments. Three independent biological experiments were performed. Superscript letters indicate significant differences ($P < 0.05$).
CaNAC46-overexpressing plants and DREB and RD22 expression levels were down-regulated in the TRV2-CaNAC46 lines. The SOD, POD, and CAT activities were higher in the CaNAC46-overexpressing plants than in the control plants. Moreover, more MDA accumulated in the TRV2-CaNAC46 plants than in the WT plants. The DAB staining results proved that ROS accumulation was greater in the TRV2-CaNAC46 plants than in the control plants. The silencing of CaNAC46 expression via virus-induced gene silencing (VIGS) resulted in oxidative damage, as indicated by the accumulation of \( \text{H}_2\text{O}_2 \). These results suggested that CaNAC46 can activate the expression of stress signaling-related genes in response to abiotic stresses. In the CaNAC46-overexpressing plants, the expression levels of ABI, POD3, SOD2, P5CS1, and other genes involved in the ROS-scavenging system were up-regulated to promote ROS scavenging and decrease stress-induced damages. The TRV2-CaNAC46 plants had relatively weak ROS activities, making them more sensitive to salt and drought stresses than the WT plants. Furthermore, CaNAC46 expression was correlated with CAT, POD, and SOD activities as well as with the transcription of most SOD and POD genes. Thus, silencing CaNAC46 expression may be associated with increasing ROS accumulation and sensitivity to salt and drought stresses.

**Conclusion**

Our results suggest that CaNAC46 localized in the nucleus helps regulate abiotic stress responses. In this study, CaNAC46 expression was induced by high salinity, drought, heat, cold, ABA, SA, and MeJA treatments. The ectopic expression of CaNAC46 in A. thaliana resulted in enhanced root formation and stronger development following drought and salt stress treatments. Moreover, the expression levels of several stress-responsive genes related to ROS scavenging and lateral root development were significantly up-regulated by drought and salt stress. Additionally, CaNAC46-silenced lines exhibited relatively weak growth and increased ROS production (i.e., more intense DAB staining) in response to salt and drought stress treatments. Furthermore, the expression of ROS-related genes was down-regulated in the CaNAC46-silenced lines. These findings imply that CaNAC46 may regulate salt and drought tolerance in pepper via its effects on ROS scavenging.

**Methods**

**Plant materials, growth conditions, and treatments**

*Capsicum annuum* L. var. ‘Qingnong No. 2’ was used as the plant material for various treatments and RNA isolation because of its tolerance to abiotic stresses. This cultivar was selected by researchers at Qingdao Agricultural University (Qingdao, China), the Qingdao Seed Station, and Dezhou Academy of Agricultural Sciences. Additionally, it was approved by the Shandong Variety Examination and Approval Committee in 2015 (deposition number: 2015–057-1). Pepper seeds were preserved at Qingdao Agricultural University (Qingdao, China). After a 3-day germination, seedlings were grown in a growth chamber at 28 °C with a 16-h light; 8-h dark cycle. All treatments were completed using 2-week-old seedlings. Regarding the simulated drought and saline conditions, seedling roots were treated with solutions containing 20% (w/v) PEG 6000 and 200 mM NaCl, respectively. For the cold and heat treatments, plants were transferred to growth chambers set to 4 and 37 °C, respectively. The phytohormone treatments involved immersing roots in an aqueous solution of 100 μM ABA, MeJA, or SA. Leaves were collected at 0, 2, 4, 8, 12, and 24 h after each treatment, immediately frozen in liquid nitrogen, ground to a powder, and then stored at −80°C.

**Isolation and sequence analysis of CaNAC46**

Total RNA was extracted from leaves using the Plant Total RNA Kit (Yuanpinghao, Beijing, China). The RNA was then used as the template to synthesize cDNA with the PrimeScript RT Reagent Kit with gDNA Eraser (TAKARA, Dalian, China). The cDNA was diluted 20-fold for the subsequent TA cloning and qRT-PCR. The CaNAC46 open reading frame was amplified by PCR using the following primers: forward: 5′-ATGATCAA AGGAATCGTTGGAAA-3′; reverse: 5′-CTAAGGT TTTGATCTGTAAGA-3′. A multiple sequence alignment was completed with DNAMAN 6.0. The hydrophobicity, composition, and physicochemical properties of protein sequences were determined with the ExPASy online tools (http://www.expasy.org/). Additionally, a phylogenetic tree was constructed with MEGA 6.0.

**Quantitative real-time PCR**

Gene expression levels were analyzed by qRT-PCR, which was completed with the reverse transcribed cDNA and the LightCycler® 480 SYBR Green I Master kit (TAKARA). The reaction solution comprised 7 μL ddH2O, 10 μL SYBR Green I Master mix, 1 μL Forward Primer (10 μmol/L), 1 μL Reverse Primer (10 μmol/L), and 1 μL cDNA. The PCR program was as follows: 95 °C for 3 min; 40 cycles of 95 °C for 5 s, 60 °C for 20 s, and 72 °C for 20 s. This program was followed by a melting curve analysis. All reactions were performed in triplicate. Relative gene expression levels were determined based on the 2ΔΔCt method. Primer sequences are listed in Table 1.
Subcellular localization of CaNAC46

The CaNAC46 coding sequence was amplified by PCR and inserted into the SpeI and Ascl sites of the modified pMDC83 vector with the ClonExpress II One Step Cloning Kit (Vazyme, Nanjing, China). After sequencing to confirm the accuracy of the inserted fragment, the recombinant plasmid and the empty pMDC83 vector (i.e., GFP alone) were introduced into tobacco cells according to an Agrobacterium tumefaciens-mediated method [34]. After a 24-h incubation in darkness, GFP fluorescence in transformed tobacco cells was observed with the FluoView™ FV300 confocal microscope (Olympus, Japan).

Generation of CaNAC46-overexpressing transgenic Arabidopsis thaliana plants

The CaNAC46 coding sequence was amplified by PCR and cloned into the XbaI sites of the pBI121 vector for the subsequent expression under the control of the cauliflower mosaic virus (CaMV) 35S promoter. The accuracy of the resulting construct was confirmed by sequencing, after which the pBI121-CaNAC46 recombinant plasmid was inserted into A. thaliana plants according to a floral dip method involving A. tumefaciens strain GV3101. Putative transgenic plants were screened on MS medium containing kanamycin and identified by PCR. Four T3 homozygous CaNAC46-overexpressing transgenic plants were selected for further analyses.

Table 1 Primer sequences used for real-time qRT-PCR amplification

| Primer name | Primer sequence |
|-------------|----------------|
| At-acting-F  | TTCCTCATGAGCTACATCTCCGTCTT |
| At-acting-R  | CACGAGATCCACGAGAATACTAGTG |
| AtABI1-F     | GGATTCTCACTGGCATGAGAGG |
| AtABI1-R     | GTCACCGGAGTTAGCGACGAA |
| AtRd20-F     | TTAAGTCCGGTACACCAATCA |
| AtRd20-R     | TGTGTTCACTTGACGGAG |
| AtPscSI-F    | GTTTCCCTGCACCGCCGATTTTA |
| AtPscSI-R    | GAAAGCGCCGACGTTGGAAC |
| AtRd29F-R    | TTAATGCTCACTGGTCTTCATC |
| AtRd29B-R    | CAAATGCCGGAACAGAGATCC |
| AtPd3-F      | TCTCGACTATCTGTCGAG |
| AtPOD3-R     | GAAAGACCTTGCGCCTCCAG |
| AtSod4-R     | GATTTGGCATCTGTTGCAAC |
| AtSod4-R     | TGAAGAATCTTGCCCTCCAG |
| Attd14-R     | GACATCCGATGAATTGAG |
| Atlbd18-F    | GTGCCTACATCTTGTCTTC |
| Atlbd18-R    | AGTAGTCTGCTGTAGTGCAAAT |
| Calnac46-F   | ACTAAGTACACTGCAACC |
| Calnac46-R   | GAGCAGGAGAGAATGAGT |
| Capi-f       | ACTCGAATTATGGATCACCC |
| Capi-R       | GACATCCGATGAATTGAG |
| CarD22-F     | AGTAAACTCTGCAAGTG |
| CarD22-R     | GTAGTCTGCGCGCTTCTT |
| Ca-actin-F   | GGTGACGAGGCTCAATC |
| Ca-actin-R   | CTCTGGACGCAACAGAA |
| Cat6-F       | TCCTGGATATCCGGG |
| Cat6-R       | TGAGGGGCAAAACCAAC |

Malondialdehyde content and antioxidant enzyme activity measurements

The MDA contents as well as the SOD, POD, and CAT activities were quantified with commercial detection kits (Jiancheng Bioengineering Institute, Nanjing, China). The chlorophyll content was measured with a hand-held chlorophyll meter.

Virus-induced gene silencing

A VIGS experiment was performed with pTRV1 and pTRV2 vectors. The CaNAC46 sequence was inserted into the pTRV2 vector. The sequence to be silenced was determined using the SGN VIGS Tool (https://vigs.solgensemics.net/), and was amplified with the following primers (TRV-CaNAC46): forward: 5′-TTGGTCGGCATTATATGTC-3′; reverse: 5′-ATTTTTTGTGTTTTATTTATCACATCTCA-3′. Agrobacterium tumefaciens cells containing pTRV1 were mixed with A. tumefaciens cells transformed with pTRV2-CaNAC46. The combined bacterial solution was infiltrated into the lower leaves of pepper plants. At 3 weeks post-infiltration, the silencing of CaNAC46 was assessed via physiological measurements and histochemical assays.
Statistical analysis
Bioassay results were evaluated with Origin and DPS. The significance of the differences in the data was determined with Student’s t-test (P < 0.05).

Supplementary Information
The online version contains supplementary material available at https://doi.org/10.1186/s12870-020-02764-y.

Additional file 1: Table S1. A comparison of physical and chemical characterization of CaNAC6 transcription factors with its homolog NAC in other model plants.

Additional file 2: Figure S1. The germination rates of WT and CaNAC6-overexpressing plants under the simulated drought and salt conditions.

Additional file 3: Figure S2. The expression of CaNAC6 gene in WT and OE plants by RT-PCR (1–4 represent WT plants; 5–8 represent OE plants).

Additional file 4: Figure S3. The expression of CaNAC6 gene in WT and TRV-CaNAC46 plants by RT-PCR (A, 1–4 represent WT plants, 5–8 represent TRV-CaNAC6 plants) and by qRT-PCR (B, 1 represent WT, 2 represent TRV-CaNAC6 plants).

Abbreviations
ABA: Abscisic acid; ACC: 1-aminoacyclopropane-1-carboxylic acid; CaMV: Cauliflower mosaic virus; DIAB: Diamino-benzidine; IAA: Indole-3-acetic acid; LSD: Lateral organ boundaries domain; MAL: Malondialdehyde; MEJA: Methyl Jasmonate; NAC: Nam; ABAF1/ATAF2, and CUC2; PSCS: Δ1-pyrroline-5-carboxylate synthetase; PEG: Polyethylene glycol; PLD: Phospholipase D; PR: Pathogenesis related; qRT-PCR: Quantitative reverse transcription-polymerase chain reaction; RD: Response to dehydration; ROS: Reactive oxygen species; SA: Salicylic acid; VIGS: Virus-induced gene silencing; WT: Wild type

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Authors’ contributions
JIM, DL and LWY conceived, designed and wrote the manuscript. LWY analyzed data. LWY, JXD and YW performed all experiments. Both authors read and approved the final manuscript.

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Availability of data and materials
All data generated or analyzed during this study are included in this article (and its supplementary information files) or are available from the corresponding author on reasonable request. Resource data from this study can be found at Arabidopsis Information Resource (https://www.arabidopsis.org/) under the following accession numbers: ATAF1 (At1g01720), ATAF2 (At5g08790), ANAC036 (At2g17040), ANAC098 (At1g59550). The sequences of OsNAC005 (LOC_Os11g82110), OsNAC006 (LOC_Os11g66120), OsNAC072 (LOC_Os09g33260) and ONAC022 (LOC_Os03g04070) are available in Genome Annotation Batch Download of Rice Genome Annotation Project (RGAP: http://rice.plantbiology.msu.edu/downloads_gad.shtml). In addition, the sequences of SINAC053 (Solyc06g060230), SINAC048 (Solyc05g055470), SINAC067 (Solyc07g0535590) and SINAC091 (Solyc06g0018000) are available in locus search of Sol Genomics Network (SGN: https://solgenomics.net/search/locus) database, and CaNAC52 (Capan02g000477), CaNAC71 (Capan07g002159), CaNAC101 (Capan12g002456), CaNAC20 (Capan02g000502), CaNAC46 (Capan02g000569) can be downloaded from the Pepper Genome Platform (PGP: http://peppergenomesnucau.de/download.php). SINAC33 (CAC42087.1) is available in National Center for Biotechnology Information (NCBI, https://www.ncbi.nlm.nih.gov/).

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that there are no competing interests.

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