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OPEN Phosphorylation of serine residue modulates cotton Di19-1 and Di19-2 activities for responding to high salinity stress and abscisic acid signaling

Li-Xia Qin^{1,2}, Xiao-Ying Nie¹, Rong Hu¹, Gang Li¹, Wen-Liang Xu¹ & Xue-Bao Li¹

Di19 (drought-induced protein 19) family is a novel type of Cys2/His2 zinc-finger proteins. In this study, we demonstrated that cotton Di19-1 and Di19-2 (GhDi19-1/-2) proteins could be phosphorylated in vitro by the calcium-dependent protein kinase (CDPK). Mutation of Ser to Ala in N-terminus of GhDi19-1/-2 led to the altered subcellular localization of the two proteins, but the constitutively activated form (Ser was mutated to Asp) of GhDi19-1/-2 still showed the nuclear localization. GhDi19-1/-2 overexpression transgenic Arabidopsis seedlings displayed the hypersensitivity to high salinity and abscisic acid (ABA). However, Ser site-mutated GhDi19-1(S116A) and GhDi19-2(S114A), and Ser and Thr double sitesmutated GhDi19-1(S/T-A/A) and GhDi19-2(S/T-A/A) transgenic Arabidopsis did not show the salt- and ABA-hypersensitive phenotypes. In contrast, overexpression of Thr site-mutated GhDi19-1(T114A) and GhDi19-2(T112A) in Arabidopsis still resulted in salt- and ABA-hypersensitivity phenotypes, like GhDi19-1/-2 transgenic lines. Overexpression of GhDi19-1/-2 and their constitutively activated forms in Atcpk11 background could recover the salt- and ABA-insensitive phenotype of the mutant. Thus, our results demonstrated that Ser phosphorylation (not Thr phosphorylation) is crucial for functionally activating GhDi19-1/-2 in response to salt stress and ABA signaling during early plant development, and GhDi19-1/-2 proteins may be downstream targets of CDPKs in ABA signal pathway.

Drought, high salinity and low temperature are the most common abiotic stresses that limit distribution of plants and affect crop productivity and quality^{1,2}. Plant adaptation to these stresses is dependent on the activation of cascades of molecular networks involving in stress perception, signal transduction and expression of specific stress-related genes^{3,4}. Abscisic acid (ABA) as an important phytohormone is required for plant adaptation to environmental stress by affecting some physiological processes in cells during plant growth and development⁵. Several signal molecules, such as calcium ion (Ca²⁺), reactive oxygen species (ROS) and protein kinases [including mitogen-activated protein kinase (MAPK), calcium-dependent protein kinase (CDPK), and calcium/ calmodulin-dependent protein kinase (CCaMK)] play important roles in ABA signal transduction for plant response to abiotic stresses⁶⁻¹⁰. For instance, overexpression of AtCPK4 or AtCPK11 in Arabidopsis enhanced plant ABA- and salt-sensitivity during seed germination and seedling growth, but loss-of-function mutations of CPK4 and CPK11 resulted in plant ABA- and salt-insensitive phenotype¹¹.

Transcriptional modulation is thought to be one of the most important ways for plants responding and adapting to stress conditions, and a number of genes are induced or repressed in plants under abiotic stresses¹² Previous studies revealed that plant transcription factors (TFs) (such as bZIP, HD-ZIP, NAC, MYB, MYC and AP2/ERF) are involved in plant stress response^{3,16}. These TFs regulate expression of the stress-related genes by specifically binding to the cis-elements in the target promoters¹⁷.

¹Hubei Key Laboratory of Genetic Regulation and Integrative Biology, School of Life Sciences, Central China Normal University, Wuhan 430079, China. ²Institute of Cotton, Shanxi Academy of Agricultural Sciences, Yuncheng 044000, China. Correspondence and requests for materials should be addressed to W.-L.X. (email: wenliangxu@mail.ccnu. edu.cn) or X.-B.L. (email: xbli@mail.ccnu.edu.cn)

Cys2/His2 (C2H2)-type zinc-finger proteins (ZFPs), also called classical TFIII-types zinc-finger proteins, represent a large family of eukaryotic transcription factors. The C2H2-type zinc finger domain is one of the well-characterized DNA-binding motifs involved in protein-DNA, protein-RNA and protein-protein interactions in plants^{18–20}. This domain consists of approximate 30 amino acid residues with the common sequence $CX_{2-4}CX_3FX_5LX_2HX_{3-5}H$, and a zinc ion is tetrahedrally coordinated by two cysteines and two histidines in order to maintain its stability²⁰. In *Arabidopsis* and rice, 176 and 189 zinc-finger proteins that contain one or more zinc-finger motifs have been identified, respectively^{21,22}. Some zinc-finger proteins play important roles in plant response to abiotic stresses. For example, *Zat10* (STZ) and *Zat12* promote plant tolerance to drought, salinity and oxidative stresses^{22–24}. ThZF1, a C2H2-type TF from salt cress (*Thellungiella halophila*), is involved in drought and salt stresses²⁵. Similarly, several C2H2-type ZFPs are related to rice response to drought and salinity stresses^{4,26,27}. Loss of function of a rice DST (drought and salt tolerance) protein leads to the enhanced plant drought- and salt-tolerance²⁸.

Di19 (drought-induced 19) proteins contain two unusual and evolutionarily conserved C2H2 zinc-finger domains²⁹. These proteins may share a common or closely related biological function based on the conserved C2H2 zinc finger-like motif. Arabidopsis Di19 family comprises of seven hydrophilic protein members. AtDi19-1 and AtDi19-3 are rapidly induced by dehydration, while transcript amounts of AtDi19-2 and AtDi19-4 are increased in response to high-salinity stress. However, most of AtDi19 genes are not transcriptionally induced by ABA³⁰. Recently, a study revealed that *Arabidopsis* Di19-1 as a transcription factor participates in plant response to drought stress by binding to the TACA(A/G)T element within the promoters of PR1 (pathogenesis-related 1), PR2 and PR5 genes31. Subsequently, we reported that AtDi19-3 as a transcriptional activator is involved in plant response to high salinity, drought, ABA and H₂O₂³². In addition, Arabidopsis Di19 proteins could be phosphorylated by CPK11 and CPK3 in vitro in a Ca²⁺-dependent manner^{30,33}. However, the function of most Di19 proteins in plants remains unknown as yet. Our previous study revealed that two cotton Di19 proteins (GhDi19-1 and GhDi19-2) are involved in plant response to salt stress and ABA signaling. Overexpression of GhDi19-1 and GhDi19-2 in Arabidopsis enhanced plant sensitivity to high salinity and exogenous ABA34. In this study, we further demonstrated that the serine (Ser) site in N-terminus of the GhDi19-1/-2 proteins is crucial for functionally activating GhDi19-1/-2 in response to salt stress and ABA signaling during early seedling development. The calcium-dependent protein kinase (CDPK) is able to phosphorylate these Di19 proteins at Ser116 of GhDi19-1 and at Ser114 of GhDi19-2.

Results

GhDi19-1/-2 is phosphorylated by calcium-dependent protein kinase (CDPK) in a Ca²⁺-dependent manner. Sequence analysis showed that both GhDi19-1 and GhDi19-2 contain Ser/Thr kinase phosphorylation sites in the conserved nuclear localization signal region next to the two zinc finger domains (GhDi19-1: Thr114 and Ser116; GhDi19-2: Thr112 and Ser114), and the predicted Ser phosphorylation is potentially stronger than the Thr phosphorylation (http://myhits.isb-sib.ch/cgi-bin/motif_scan). Then, we employed an *in vitro* kinase assay to determine whether GhDi19-1 and GhDi19-2 are phosphorylated by a kinase (such as AtCPK11). As shown in Fig. 1a, the S₁₁₆ of GhDi19-1 was mutated to A₁₁₆ [GhDi19-1(S116A)], and the S₁₁₄ of GhDi19-2 was mutated to A₁₁₄ [GhDi19-2(S114A)]. The *in vitro* kinase assay revealed that the recombinant AtCPK11 phosphorylated GhDi19-1 and GhDi19-2 in the presence of Ca²⁺. Additionally, weaker phosphorylation of GhDi19-1(S116A) was detected in the presence of Ca²⁺, and no phosphorylation of GhDi19-2(S114A) was found with or without Ca²⁺ (Fig. 1b). The above results indicated that GhDi19-1/-2 is phosphorylated *in vitro* by a kinase (such as AtCPK11) in a Ca²⁺-dependent manner, and the 116th Ser of GhDi19-1 or the 114th Ser of GhDi19-2 is required for the phosphorylation.

Ser site is crucial for normal subcellular localization of GhDi19-1 and GhDi19-2 proteins. Bioinformatics analysis predicted that both GhDi19-1 and GhDi19-2 contain a conserved nuclear localization signal (NLS) in its sequence. To confirm nuclear localization of the two GhDi19 proteins, the coding regions of the two genes were fused with an enhanced GFP (eGFP) reporter gene under the control of a cauliflower mosaic virus (CaMV) 35S promoter, respectively, and expressed constitutively in Arabidopsis. To determine whether Ser site and/or Thr site play an essential role in subcellular localization of GhDi19-1 and GhDi19-2 proteins, the S_{116} of GhDi19-1 was mutated to A_{116} [GhDi19-1(S116A)], the S_{114} of GhDi19-2 was $mutated \ to \ A_{114} \ [GhDi19-2(S114A)], \ the \ T_{114} \ of \ GhDi19-1 \ was \ mutated \ to \ A_{114} \ [GhDi19-1(T114A)], \ the \ T_{112} \ of \ GhDi19-1 \ was \ mutated \ to \ A_{114} \ [GhDi19-1(T114A)], \ the \ T_{112} \ of \ GhDi19-1 \ was \ mutated \ to \ A_{114} \ [GhDi19-1(T114A)], \ the \ T_{112} \ of \ GhDi19-1 \ was \ mutated \ to \ A_{114} \ [GhDi19-1(T114A)], \ the \ T_{112} \ of \ GhDi19-1 \ was \ mutated \ to \ A_{114} \ [GhDi19-1(T114A)], \ the \ T_{112} \ of \ GhDi19-1 \ was \ mutated \ to \ A_{114} \ [GhDi19-1(T114A)], \ the \ T_{112} \ of \ GhDi19-1 \ was \ mutated \ to \ A_{114} \ [GhDi19-1(T114A)], \ the \ T_{112} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ [GhD119-1(T114A)], \ the \ T_{112} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ [GhD119-1(T114A)], \ the \ T_{112} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ [GhD119-1(T114A)], \ the \ T_{112} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ [GhD119-1(T114A)], \ the \ T_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ [GhD119-1(T114A)], \ the \ T_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ [GhD119-1(T114A)], \ the \ T_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ [GhD119-1(T114A)], \ the \ T_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ [GhD119-1(T114A)], \ the \ T_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ [GhD119-1(T114A)], \ the \ T_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ [GhD119-1(T114A)], \ the \ T_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ of \ GhD119-1 \ was \ mutated \ to \ A_{114} \ of \ A_{114}$ of GhDi19-2 was mutated to A_{112} [GhDi19-2(T112A)], both the T_{114} and S_{116} of GhDi19-1 were mutated to A [GhDi19-1(S/T-A/A)], and both the T₁₁₂ and S₁₁₄ of GhDi19-2 were mutated to A [GhDi19-2(S/T-A/A)]. Furthermore, S₁₁₆ was mutated to D₁₁₆ in GhDi19-1 and S₁₁₄ was mutated to D₁₁₄ in GhDi19-2 for generating the constitutively activated GhDi19-1 and GhDi19-2 [named GhDi19-1(S116D) and GhDi19-2(S114D), respectively]. The coding sequences of the above mutated GhDi19s were also fused with an enhanced GFP (eGFP) reporter gene under the control of a cauliflower mosaic virus (CaMV) 35S promoter, respectively, and expressed constitutively in Arabidopsis. As shown in Fig. 2a,b, the GhDi19-1:eGFP and GhDi19-2:eGFP fusion proteins were mainly accumulated in the nuclei of root cells of the transgenic seedlings, demonstrating that GhDi19-1 and GhDi19-2 are two nuclear-localized proteins. Likewise, strong GFP fluorescence signals were detected in the nuclei of the transgenic root cells harboring the GhDi19-1(S116D):eGFP (Fig. 2c), GhDi19-2(S114D):eGFP (Fig. 2d), GhDi19-1(T114A):eGFP (Fig. 2e), and GhDi19-2(T112A):eGFP (Fig. 2f), respectively. On the contrary, GFP fluorescence was found in cytoplasm, cytomembrane and nuclei of root cells of the transgenic plants expressing Ser site-mutated GhDi19-1(S116A) (Fig. 2g) and GhDi19-2(S114A) (Fig. 2h), and Ser/Thr double sites-mutated GhDi19-1(S/T-A/A) (Fig. 2i) and GhDi19-2(S/T-A/A) (Fig. 2j), indicating that subcellular localization of both GhDi19-1 and GhDi19-2 proteins was altered as the Ser site (but not Thr site) was mutated to Ala in these proteins, while GFP fluorescence signals were observed in whole cells of the transgenic seedlings harboring

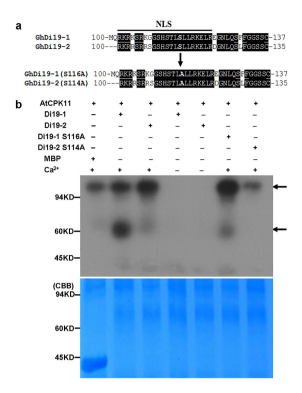


Figure 1. *In* vitro assay of GhDi19-1 and GhDi19-2 phosphorylation. (a) The map of mutation of specific amino acids in the phosphorylation of GhDi19-1 and GhDi19-2. NLS, nuclear localization signal domain; S, serine (Ser); A, alanine (Ala). The arrowhead indicated the mutated amino acids. (b) AtCPK11 phosphorylates GhDi19-1 and GhDi19-2 *in vitro*. Top: Autoradiograph showing AtCPK11 autophosphorylation and GhDi19-1/-2 phosphorylation; Bottom: the same SDS-PAGE gel with Coomasie Brilliant Blue (CBB)—stained AtCPK11 and GhDi19-1/-2 proteins. GhDi19-1 and GhDi19-2 proteins were purified as a MBP-fusion and incubated with AtCPK11-MBP. A control reaction with the MBP protein was also performed. The upper arrowhead indicated autophosphorylation of AtCPK11-MBP, while the lower arrowhead indicated phosphorylation of GhDi19-1-MBP and GhDi19-2-MBP.

only GFP vector (Fig. 2k). The above data implied that the Ser site (but not Thr site) is essential for normal subcellular localization of GhDi19-1 and GhDi19-2 proteins.

Ser site is crucial for activating GhDi19-1 and GhDi19-2 proteins in response to salt stress and ABA signaling. Our previous study indicated that overexpression of the *GhDi19-1* and *GhDi19-2* genes in *Arabidopsis* enhances plant sensitivity to salt stress in ABA-dependent manner³⁴. To further investigate the mechanism of GhDi19-1 and GhDi19-2 involving in response to salt stress and ABA signaling, the S₁₁₆ of GhDi19-1 was mutated to A₁₁₆, and the S₁₁₄ of GhDi19-2 was mutated to A₁₁₄. The coding sequences of *GhDi19-1(S116A)* and *GhDi19-2(S114A)* were inserted into plant expression vector pBI121 driven by CaMV 35S promoter, and introduced into *Arabidopsis*, respectively. Homozygotes of T3 generation of *GhDi19-1* overexpression transgenic lines (L4 and L19), *GhDi19-1(S116A)* overexpression transgenic lines (L2 and L8), *GhDi19-2* overexpression transgenic lines (L3 and L6) and *GhDi19-2(S114A)* overexpression transgenic lines (L1 and L4) were analyzed by PCR to ensure the presence of the respective transgene and by quantitative RT-PCR to confirm transgene expression (Fig. 3a).

Seeds of GhDi19-1, GhDi19-1(S116A), GhDi19-2 and GhDi19-2(S114A) overexpression transgenic lines and wild type were sowed on MS medium with or without NaCl and ABA, respectively. On MS medium, all the seeds from wild type and GhDi19-1/-2 variants were able to fully germinate (≤ 4 days) after sowing, and there was no significant difference in seed germination rate between each other. In the presence of 150 mM NaCl, however, the GhDi19-1 and GhDi19-2 transgenic seeds germinated much later than those of wild type. After 4 day, approximately 75% of wild type seeds germinated, but only about 35%–40% of GhDi19-1 and GhDi19-2 transgenic seeds germinated. After 6 days, germination rate of wild type seeds reached to about 90%, but both GhDi19-1 and GhDi19-2 transgenic seeds reached to only approximately 60%. In contrast, seed germination rates of GhDi19-1(S116A) and GhDi19-2(S114A) transgenic lines were similar to those of wild type with NaCl treatment. On MS medium supplemented with 0.8 μ M ABA, seed germination rate of both GhDi19-1 and GhDi19-2 transgenic lines was significantly lower than that of wild-type. About 60% of wild type seeds, but only 40% of GhDi19-1 and 30% of GhDi19-2 overexpression transgenic seeds germinated after 4 days. On the contrary, seed germination rate of GhDi19-1(S116A) and GhDi19-2(S114A) overexpression transgenic lines was as same as those of wild type under ABA treatment (Fig. 4a).

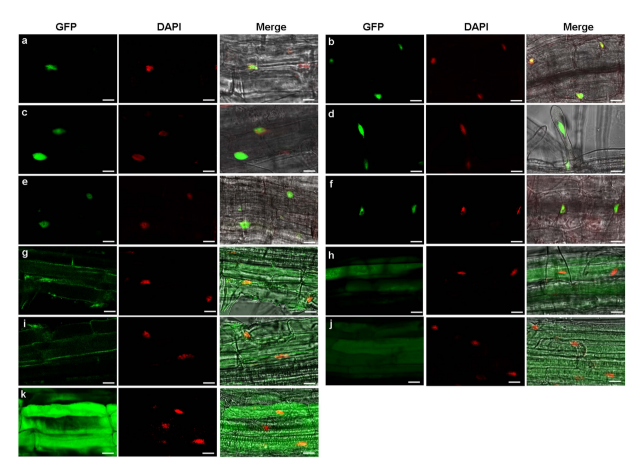


Figure 2. Subcellular localization of GhDi19-1/-2 and their mutated proteins in cells. (a-f) GFP fluorescence signals were mainly detected in nuclei of the transgenic Arabidopsis root cells harboring the GhDi19-1:eGFP, GhDi19-2:eGFP, GhDi19-1(S116D):eGFP, GhDi19-2(S114D):eGFP, GhDi19-1(T114A):eGFP and GhDi19-2(T112A):eGFP, respectively. (a,b) Normal subcellular localization of GhDi19-1 (a) and GhDi19-2 (b). (c,d) Subcellular localization of the Ser site-mutated (constitutively activated) GhDi19-1(S116D) (c) and GhDi19-2(S114D) (d). (e,f) Subcellular localization of the Thr site-mutated GhDi19-1(T114A) (e) and GhDi19-2 (T112A) (f). (g-j) The altered subcellular localization of the Ser site-mutated GhDi19-1(S116A) (g), GhDi19-2(S114A) (h), the Ser and Thr double sites-mutated GhDi19-1(S/T-A/A) (i) and GhDi19-2(S/T-A/A) (j). GFP fluorescence signals were detected in cytoplasm, cytomembrane and nuclei of the transgenic Arabidopsis root cells. (k) pBI121-eGFP construct was used as positive control. (Left) Confocal microsopy of GFP fluorescence. (Midst) Nuclear DAPI staining of the same cell on the left. (Right) Left and midst $superimposed over the bright-field image. \ Bars = 10 \ \mu m. \ S116D, the \ S_{116} \ of \ GhDi19-1 \ was \ mutated \ to \ D_{116}; \ begin{tabular}{l} Policy of the limit of the limit$ S114D, the S_{114} of GhDi19-2 was mutated to D_{114} ; T114A, the T_{114} of GhDi19-1 was mutated to A_{114} ; T112A, the T_{112} of GhDi19-2 was mutated to A_{112} ; S116A, the S_{116} of GhDi19-1 was mutated to A_{116} ; S114A, the S_{114} of GhDi19-2 was mutated to A₁₁₄; S/T-A/A, the S₁₁₆ and T₁₁₄ of GhDi19-1 were mutated to A₁₁₆ and A₁₁₄, and the S₁₁₄ and T₁₁₂ of GhDi19-2 were mutated to A₁₁₄ and A₁₁₂, respectively. S, serine (Ser); A, alanine (Ala); T, threonine (Thr); D, aspartic acid (Asp).

To assess effects of Ser site on GhDi19-1 and GhDi19-2 involving in response to salt and ABA during early seedling development, we used two approaches in the experiments. One way is that seeds were directly planted on MS medium containing 150 mM NaCl or $1\,\mu\text{M}$ ABA for determining growth status of the seedlings after germination (Fig. 4b). Another is that seeds were germinated on MS medium for 48 h after stratification and then transferred onto MS medium containing 150 mM NaCl and 5 μM ABA in the vertical position (Fig. 4d). The results obtained with these two approaches were similar. There was no significant difference among the GhDi19-1/-2 and their site-mutated transgenic lines and wild type when seedlings grew on MS medium. However, the GhDi19-1 and GhDi19-2 overexpression seedlings were more sensitive to salt and ABA than wild type, but growth status of GhDi19-1(S116A) and GhDi19-2(S114A) transgenic seedlings was almost as same as those of wild type (Fig. 4b). The cotyledon greening rates of GhDi19-1 overexpression lines (L4 and L19) and GhDi19-2 overexpression lines (L3 and L6) were significantly lower than those of wild type under NaCl and ABA treatments, while cotyledon greening rates of GhDi19-1(S116A) and GhDi19-2(S114A) transgenic lines (L2 and L8, L1 and L4, respectively) were similar to those of wild type (Fig. 4c).

When the seedlings were transferred onto MS plates in the vertical position without any stress treatments, root growth of all *GhDi19-1/-2* transgenic seedlings and their site-variants was almost as same as those of wild type

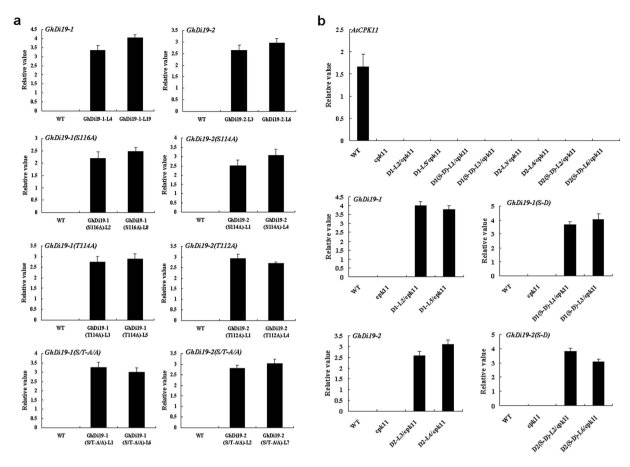


Figure 3. Quantitative RT-PCR analysis of expression of the related exogenous GhDi19-1/-2 genes in transgenic Arabidopsis. (a) Quantitative RT-PCR analysis of expression of the GhDi19-1/-2 and their mutant genes in wild type, and the GhDi19-1/-2 and their site-mutated transgenic lines. 10-day-old seedlings were used for RNA extraction. WT, wild type; Di19-1-L4 and -L19, the GhDi19-1 transgenic line 4 and 19; GhDi19-1(S116A)-L2 and -L8, the site-mutated GhDi19-1(S116A) transgenic line 2 and 8 (S116A, the S₁₁₆ of GhDi19-1 was mutated to A₁₁₆). GhDi19-1(T114A)-L3 and -L5, the site-mutated GhDi19-1(T114A) transgenic line 3 and 5 (T114A, the T₁₁₄ of GhDi19-1 was mutated to A₁₁₄). GhDi19-1(S/T-A/A)-L1 and -L6, the double sites-mutated GhDi19-1(S/T-A/A) transgenic line 1 and 6 (S/T-A/A, the S₁₁₆ and T₁₁₄ of GhDi19-1 were mutated to A₁₁₆ and A₁₁₄. respectively). GhDi19-2-L3 and -L6, the GhDi19-2 transgenic line 3 and 6; GhDi19-2(S114A)-L1 and -L4, the sitemutated GhDi19-2(S114A) transgenic line 1 and 4 (S114A, the S_{114} of GhDi19-2 was mutated to A_{114}); GhDi19-2(S114A) transgenic line 1 and 4 (S114A, the S_{114} of GhDi19-2 was mutated to A_{114}); GhDi19-2(S114A) transgenic line 1 and 4 (S114A, the S_{114} of GhDi19-2 was mutated to A_{114}); GhDi19-2(S114A) transgenic line 1 and 2 (S114A) transgenic line 1 and 3 (S114A) transgen 2(T112A)-L1 and -L4, the site-mutated GhDi19-2(T112A) transgenic line 1 and 4 (T112A, the T_{112} of GhDi19-2 was mutated to A₁₁₂); GhDi19-2(S/T-A/A)-L2 and -L7, the double sites-mutated GhDi19-2(S/T-A/A) transgenic line 2 and 7 (S/T-A/A, the S_{114} and T_{112} of GhDi19-2 were mutated to A_{114} and A_{112} , respectively). (b) Quantitative RT-PCR analysis of expression of the GhDi19-1/-2 and their mutant genes in wild type, Atcpk11 mutant, and GhDi19-1/2- and their mutated transgenic lines in Atcpk11 mutant. 10-day-old seedlings were used for RNA extraction. WT, wild type; cpk11, Atcpk11-2 mutant; D1-L2/cpk11 and D1-L5/cpk11, the GhDi19-1 overexpression transgenic line 2 and 5 in Atcpk11-2 mutant; D1(S-D)-L1/cpk11 and D1(S-D)-L3/cpk11, the site-mutated GhDi19-1(S-D) overexpression transgenic line 1 and 3 in Atcpk11-2 mutant; D2-L3/cpk11 and D2-L4/cpk11, the GhDi19-2 overexpression transgenic line 3 and 4 in Atcpk11-2 mutant; D2(S-D)-L2/cpk11 and D2(S-D)-L6/cpk11, the sitemutated GhDi19-2(S-D) overexpression transgenic line 2 and 6 in Atcpk11-2 mutant (S-D, the S₁₁₆ of GhDi19-1 was mutated to D_{116} and the S_{114} of GhDi19-2 was mutated to D_{114}). The assays were repeated three times along with three independent repetitions of the biological experiments. S, serine (Ser); A, alanine (Ala); T, threonine (Thr); D, aspartic acid (Asp).

(Fig. 4d). However, when the seedlings were transferred onto MS medium supplemented with 150 mM NaCl or 5μM ABA for several days, primary root growth of the GhDi19-1 and GhDi19-2 overexpression transgenic seedlings was significantly inhibited much more than that of wild type, but growth of *GhDi19-1(S116A)* and *GhDi19-2(S114A)* transgenic roots was similar to that of wild type (Fig. 4d). Under NaCl and ABA treatments, roots of GhDi19-1 and GhDi19-2 overexpression seedlings were shorter than those of wild type, whereas there was little significant difference in primary root length between *GhDi19-1(S116A)* and *GhDi19-2(S114A)* transgenic lines and wild type (Fig. 4e). In addition, we also introduced the "empty vector" (pBI121-eGFP) into *Arabidopsis* as a negative control in the above experiments, and found that the phenotypes of the transgenic plants harboring the empty vector were similar to those of wild type. Collectively, these data suggested that overexpression of *GhDi19-1*

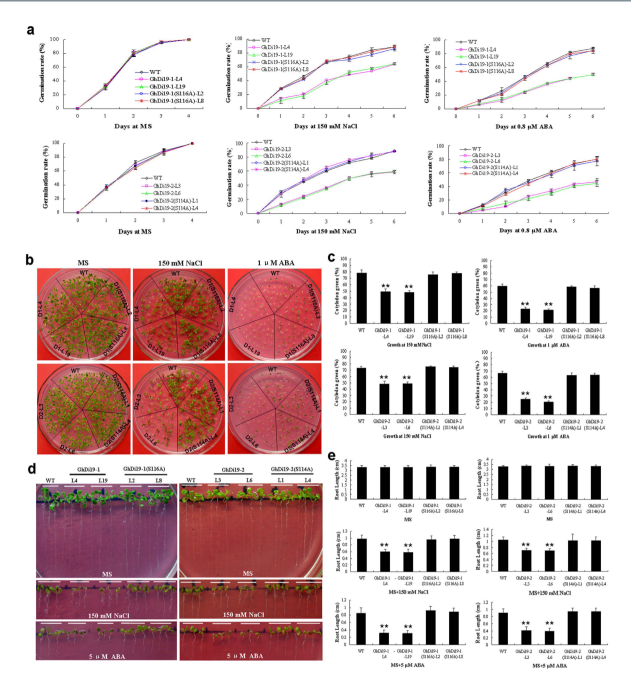


Figure 4. Effect of serine site-mutation on GhDi19-1 and GhDi19-2 responding to salt and abscisic acid (ABA) during seed germination and early seedling development. (a) Statistical analysis of seed germination rate. Seeds of the transgenic lines and wild type were germinated on MS medium without (control) or with 150 mM NaCl and 0.8 µM ABA, respectively. (b) Growth status of the transgenic seedlings and wild type grown on MS medium without or with 150 mM NaCl and 1 µM ABA for 10 days, respectively. (c) Statistical analysis of cotyledon expansion and greening of seedlings grown on MS medium with 150 mM NaCl and 1 μM ABA for 10 days, respectively. (d) Phenotypic analysis of GhDi19-1/-2(S-A) transgenic Arabidopsis under NaCl and ABA treatments. 10-day-old seedlings of wild type, GhDi19-1/-2 and GhDi19-1/-2(S-A)-overexpression transgenic lines growing on MS medium with or without 150 mM NaCl and 5 µM ABA, respectively. (e) Statistical analysis of root length of 10-day-old seedlings of wild type and GhDi19-1/2 and GhDi19-1/2(S-A) transgenic lines growing on MS medium without (control) or with 150 mM NaCl and 5 µM ABA, respectively. Mean values and standard errors (bar) were shown from three independent experiments (n > 60 seedlings per each line). Asterisk represents that there was (very) significant difference between the transgenic lines and wild type in independent t-tests (one asterisk: P value < 0.05; two asterisks: P value < 0.01). WT, wild type; GhDi19-1-L4 (D1-L4) and -L19 (D1-L19), the GhDi19-1 transgenic line 4 and 19; GhDi19-1(S116A)-L2 [D1(S116A)-L2] and -L8 [D1(S116A)-L8], the site-mutated GhDi19-1(S116A) transgenic line 2 and 8 (S116A, the S₁₁₆ of GhDi19-1 was mutated to A₁₁₆); Di19-2-L3 (D2-L3) and -L6 (D2-L6), the GhDi19-2 transgenic line 3 and 6. GhDi19-2(S114A)-L1 [D2(S114A)-L1] and -L4 [D2(S114A)-L4], the site-mutated GhDi19-2(S114A) transgenic line 1 and 4 (S114A, the S₁₁₄ of GhDi19-2 was mutated to A₁₁₄). S, serine (Ser); A, alanine (Ala).

and *GhDi19-2* in *Arabidopsis* enhances plant sensitivity to salt stress in ABA-dependent manner, and the Ser site is essential for GhDi19-1 and GhDi19-2 proteins involving in plant response to salt and ABA during seed germination and early seedling development.

Thr site is inessential for GhDi19-1 and GhDi19-2 in response to salt stress and ABA signaling. To understand whether Thr phosphorylation site also play an important role in GhDi19-1 and GhDi19-2 involving in response to salt and ABA signaling, the T_{114} of GhDi19-1 was mutated to A_{114} , the T_{112} of GhDi19-2 was mutated to A_{112} , both T_{114} and S_{116} of GhDi19-1 were mutated to A, and both T_{112} and S_{114} of GhDi19-2 were mutated to A. The coding sequences of GhDi19-1(T114A), GhDi19-1(S/T-A/A), GhDi19-2(T112A) and GhDi19-2(S/T-A/A) were inserted into pBI121 vector, and introduced into Arabidopsis, respectively. Homozygotes of T3 generation of the respective overexpression transgenic lines were analyzed by PCR to ensure confirm the presence of the respective transgene and by quantitative RT-PCR to confirm that the transgene was expressed (Fig. 3a). Two transgenic lines (L3 and L5) with higher GhDi19-1(S/T-A/A) expression, two lines (L1 and L6) with higher GhDi19-1(S/T-A/A) expression, two lines (L1 and L6) with higher GhDi19-2(S/T-A/A) expression were selected for analyzing their phenotypes under different stresses.

When seeds sowed on MS medium, there was no significant difference in seed germination between all the transgenic lines and wild type under normal conditions (Fig. 5a,d,g, j). When sowing on MS medium containing 150 mM NaCl or 0.8 µM ABA, however, the *GhDi19-1(T114A)* and *GhDi19-2(T112A)* transgenic seeds germinated much later than those of wild type, and showed salt- and ABA-sensitivity. In the presence of 150 mM NaCl, only about 35%–40% of the *GhDi19-1(T114A)* and *GhDi19-2(T112A)* transgenic seeds germinated, while approximately 65% of wild type germinated after 3 days. And germination rate of the transgenic seeds only reached to 60%, while wild type was about 80% after five days of germination (Fig. 5b,h). In addition, seed germination rate of the *GhDi19-1(T114A)* and *GhDi19-2(T112A)* transgenic lines was reduced much more than that of wild type under ABA treatment. When treated with 0.8 µM ABA, the seed germination rates of *GhDi19-1*, *GhDi19-2*, *GhDi19-1(T114A)* and *GhDi19-2(T112A)* transgenic lines were decreased to approximately 40%, whereas wild type retained 75% seed germination rate after 4 days (Fig. 5c,i). On the other hand, there was no significant difference in seed germination rate between *GhDi19-1(S/T-A/A)* and *GhDi19-2(S/T-A/A)* transgenic lines and wild type under NaCl and ABA treatments (Fig. 5e,k,f,l).

We also calculated cotyledon greening of the transgenic seedlings and wild type under salt stress. As shown in Fig. 6a, both GhDi19-1(T114A) and GhDi19-2(T112A) transgenic seedlings growing on MS containing 150 mM NaCl were more sensitive than wild type. Cotyledon greening of GhDi19-1(T114A) and GhDi19-2(T112A) transgenic seedlings was drastically inhibited by NaCl treatment, compared with that of wild type. Similarly, cotyledon greening of GhDi19-1(T114A) and GhDi19-2(T112A) transgenic plants was also inhibited more than that of wild type. In the presence of $1\,\mu$ M ABA, approximately 20% and 25% of the GhDi19-1(T114A) and GhDi19-2(T112A) transgenic seedlings with expanded and turned-green cotyledons was observed, respectively, while wild type reached to 60% cotyledon greening rate. In contrast, GhDi19-1(S/T-A/A) and GhDi19-2(S/T-A/A) transgenic plants showed the similar phenotype to wild-type under NaCl and ABA treatments (Fig. 6b).

Additionally, seeds of all the transgenic *Arabidopsis* lines and wild type germinated on MS medium for 48 h, and then were transferred onto NaCl- and ABA-containing MS medium for investigating growth of the seedlings responding to high salinity and ABA (Fig. 6c). When transferred onto MS plates, root growth of all the transgenic seedlings was almost as same as that of wild type. However, when the seedlings were transferred onto MS medium supplemented with 150 mM NaCl or 5 µM ABA for several days, primary roots growth of the *GhDi19-1(T114A)* and *GhDi19-2(T112A)* transgenic seedlings was more significantly inhibited than that of wild type by NaCl or ABA. The roots of *GhDi19-1(T114A)* and *GhDi19-2(T112A)* transgenic seedlings were shorter than those of wild type under NaCl and ABA treatments. However, there was no significant difference in root length between *GhDi19-1(S/T-A/A)* and *GhDi19-2(S/T-A/A)* transgenic lines and wild type under NaCl and ABA treatments (Fig. 6c). These results demonstrated that overexpression of the mutated *GhDi19-1(T114A)* and *GhDi19-2(T112A)* genes in *Arabidopsis* still resulted in the transgenic plants salt- and ABA-hypersensitivity, like *GhDi19-1/-2* transgenic lines, suggesting that the Thr site is inessential for GhDi19-1 and GhDi19-2 proteins involving in response to salt stress and ABA signaling during seed germination and early seedling development.

Phosphorylation of Ser site by calcium-dependent protein kinase (CDPK) is important for functionally activating GhDi19-1/-2 in response to salt stress and ABA signaling. To explore the functional relationship of GhDi19s and CDPK, loss-of-function mutation of CPK11 (Atcpk11-2, SALK_054495) was employed in this study. A single copy of T-DNA was inserted into the genome of the Atcpk11-2 mutant, generating a 39-bp deletion from 320 to 358 bp downstream of the translation start codon¹¹. Quantitative RT-PCR analysis demonstrated that AtCPK11 mRNA was undetectable in Atcpk11-2 mutant. Meanwhile, S₁₁₆ was mutated to D₁₁₆ in GhDi19-1 and S_{114} was mutated to D_{114} in GhDi19-2 for generating the constitutively activated GhDi19-1 and GhDi19-2 [named GhDi19-1(S116D) and GhDi19-2(S114D), respectively]. The coding sequences of GhDi19-1, GhDi19-2, GhDi19-1(S116D) and GhDi19-2(S114D) under the control of CaMV 35S promoter were introduced into Atcpk11-2 mutant. Homozygotes of T3 generation of these GhDi19 transgenic lines in Atcpk11-2 background were analyzed by quantitative RT-PCR to confirm the transgene expression (Fig. 3b). Seeds of wild type, Atcpk11-2 mutant and GhDi19-1, GhDi19-2, GhDi19-1(S116D) and GhDi19-2(S114D) overexpression transgenic lines in Atcpk11-2 background (GhDi19-1/cpk11, GhDi19-2/cpk11, GhDi19-1(S116D)/cpk11 and GhDi19-2(S114D)/cpk11) were sowed on MS medium with or without ABA and NaCl, respectively. Seed germination rate of wild type and the transgenic lines was monitored under different abiotic stresses. On MS medium, all the seeds from wild type and the transgenic lines were able to fully germinate (≤4 days) after sowing, and there was no significant difference in seed germination rate between the transgenic lines and wild type. However, in the

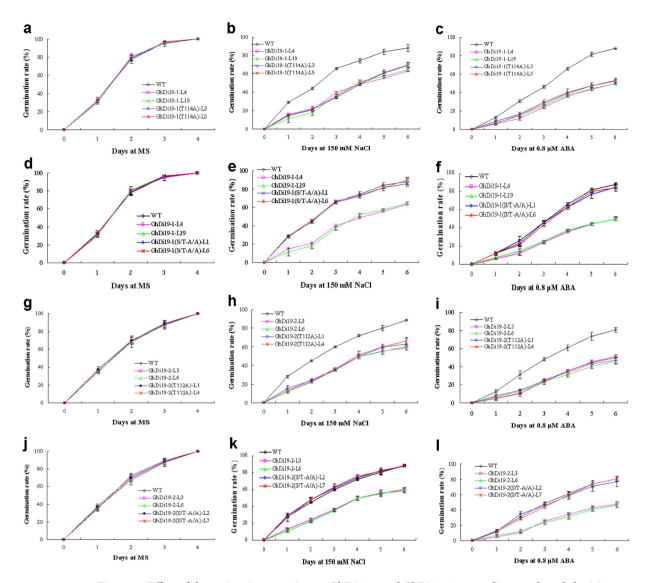


Figure 5. Effect of threonine site-mutation on GhDi19-1 and GhDi19-2 responding to salt and abscisic acid (ABA) during seed germination. Seeds of the transgenic lines and wild type were germinated on MS medium (a,d,g,j), MS medium with 150 mM NaCl (b,e,h,k) and MS medium with 0.8 μ M ABA (c,f,i,l), respectively. Statistical analysis of seed germination rate was determined. Mean values and standard errors (bar) were shown from three independent experiments (n > 200 seeds per each line). WT, wild type; Di19-1-L4 and -L19, the *GhDi19-1* transgenic line 4 and 19; GhDi19-1(T114A)-L3 and -L5, the site-mutated *GhDi19-1(T114A)* transgenic line 3 and 5; GhDi19-1(S/T-A/A)-L1 and -L6, the double sites-mutated *GhDi19-1(S/T-A/A)* transgenic line 1 and 6 (T114A, the T₁₁₄ of GhDi19-1 was mutated to A₁₁₄; S/T-A/A, the S₁₁₆ and T₁₁₄ of GhDi19-1 were mutated to A₁₁₆ and A₁₁₄). Di19-2-L3 and -L6, the *GhDi19-2* transgenic line 3 and 6; GhDi19-2(T112A)-L1 and -L4, the site-mutated *GhDi19-1(T112A)* transgenic line 1 and 4; GhDi19-1(S/T-A/A)-L2 and -L7, the double sites-mutated *GhDi19-2(S/T-A/A)* transgenic line 2 and 7 (T112A, the T₁₁₂ of GhDi19-2 was mutated to A₁₁₂; S/T-A/A, the S₁₁₄ and T₁₁₂ of GhDi19-2 were mutated to A₁₁₄ and A₁₁₂). The assays were repeated three times along with three independent repetitions of the biological experiments. S, serine (Ser); T, threonine (Thr); A, alanine (Ala).

presence of 0.8 µM ABA, seeds of *Atcpk11-2* mutant germinated much earlier and faster than those of wild type, and the germination rate of *GhDi19-1/cpk11* and *GhDi19-2/cpk11* seeds was similar to wild type, whereas seeds of *GhDi19-1(S116D)/cpk11* and *GhDi19-2(S114D)/cpk11* germinated much later than wild type. After 4 days, about 60% of wild type, *GhDi19-1/cpk11* and *GhDi19-2/cpk11* seeds germinated, and approximately 75% of *Atcpk11-2* mutant seeds germinated, but only about 35% of *GhDi19-1(S116D)/cpk11* and *GhDi19-2(S114D)/cpk11* seeds germinated. After 6 days, seed germination rate of *Atcpk11-2* mutant reached to about 90%, but those of *GhDi19-1(S116D)/cpk11* and *GhDi19-2(S114D)/cpk11* transgenic lines reached to only approximately 40–50%. In the presence of 150 mM NaCl, seed germination rate of *Atcpk11-2* mutant was significantly higher than that of wild type. On the contrary, germination of the *GhDi19-1(S116D)/cpk11* and *GhDi19-2(S114D)/cpk11* transgenic seeds was inhibited much more by NaCl, compared with wild type. Approximately 80% of *Atcpk11-2* mutant seeds and

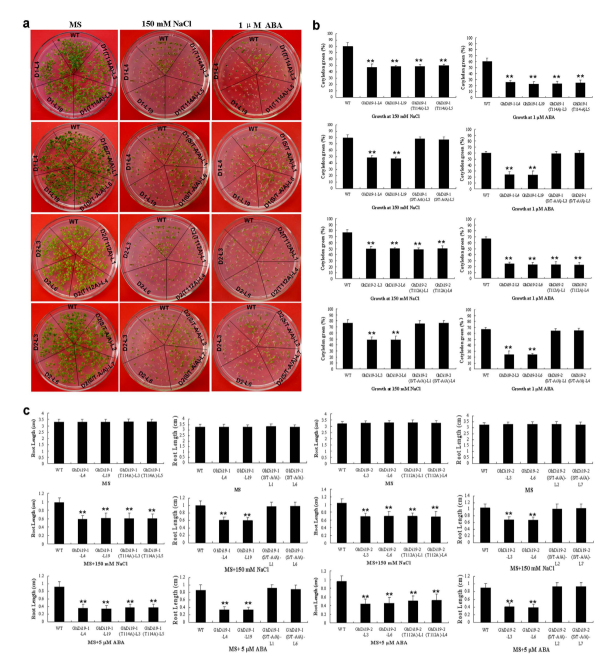


Figure 6. Effect of threonine site-mutation on GhDi19-1 and GhDi19-2 responding to salt and abscisic acid (ABA) during early seedling development. (a) Growth status of wild type, and the GhDi19-1/-2, GhDi19-1/-2(T-A) and GhDi19-1/-2(S/T-A/A) overexpression seedlings growing on MS medium with or without 150 mM NaCl and 1 µM ABA, respectively. (b) Statistical analysis of cotyledon expansion and greening of wild type, and the GhDi19-1/-2, GhDi19-1/-2(T-A) and GhDi19-1/-2(S/T-A/A) overexpression seedlings growing on MS medium with 150 mM NaCl and 1 μM ABA, respectively. (c) Statistical analysis of root length of 10-day-old seedlings of wild type, and the GhDi19-1/-2, GhDi19-1/-2(T-A) and GhDi19-1/-2(S/T-A/A) overexpression lines growing on MS medium without (control) or with 150 mM NaCl and 5 µM ABA. Mean values and standard errors (bar) were shown from three independent experiments (n > 60 seedlings per each line). Asterisk represents that there was (very) significant difference between the transgenic lines and wild type in independent t-tests (one asterisk: P value < 0.05; two asterisks: P value < 0.01). WT, wild type; GhDi19-1-L4 (D1-L4) and -L19 (D1-L19), the GhDi19-1 transgenic line 4 and 19; GhDi19-1(T114A)-L3 and -L5 [D1(T114A)-L3 and -L5], the site-mutated GhDi19-1(T114A) transgenic line 3 and 5; GhDi19-1(S/T-A/A)-L1 and -L6 [D1(S/T-A/A)-L1 and -L6], the double sites-mutated GhDi19-1(S/T-A/A) transgenic line 1 and 6 (T114A indicated the T₁₁₄ of GhDi19-1 was mutated to A_{114} , and S/T-A/A indicated both S_{116} and T_{114} of GhDi19-1 were mutated to A_{116} and A_{114}). GhDi19-2-L3 (D1-L3) and -L6 (D1-L6), the GhDi19-2 transgenic line 3 and 6; GhDi19-2(T112A)-L1 and -L4 [D2(T112A)-L1 and -L4], the site-mutated GhDi19-2(T112A) transgenic line 1 and 4; GhDi19-2(S/T-A/A)-L2 and -L7 [D2(S/T-A/A)-L2 and -L7], the double sites-mutated GhDi19-2(S/T-A/A) transgenic line 2 and 7 (T112A indicated the T_{112} of GhDi19-2 was mutated to A_{112} , and S/T-A/A indicated both S_{114} and T_{112} of $GhDi19-2\ were\ mutated\ to\ A_{114}\ and\ A_{112}).\ S,\ serine\ (Ser);\ T,\ threonine\ (Thr);\ A,\ alanine\ (Ala).$

70% of wild type, GhDi19-1/cpk11 and GhDi19-2/cpk11 seeds, but only 50% of GhDi19-1(S116D)/cpk11 and GhDi19-2(S114D)/cpk11 transgenic seeds germinated after 4 days (Fig. 7a).

During early seedling development, no significant difference was observed among the transgenic variants and wild type when seedlings grew on MS medium. However, the seedlings of *Atcpk11-2* mutant grew better than those of wild type on ABA- and NaCl-containing medium, but the *GhDi19-1(S116D)/cpk11* and *GhDi19-2(S114D)/cpk11* transgenic seedlings were more sensitive to ABA and salt than wild type, while growth status of *GhDi19-1/cpk11* and *GhDi19-2/cpk11* transgenic seedlings was similar to wild type under ABA and NaCl treatments (Fig. 7b). Under ABA and NaCl treatments, cotyledon greening rate of *Atcpk11-2* mutant was significantly higher than that of wild type, while cotyledon greening rate of *GhDi19-1/cpk11* and *GhDi19-2/cpk11* transgenic lines was similar to wild type. In contrast, cotyledon greening of *GhDi19-1(S116D)/cpk11* and *GhDi19-2(S114D)/cpk11* transgenic lines was inhibited much more by ABA and NaCl, compared with wild type (Fig. 7c).

When the seedlings were transferred onto MS plates in the vertical position without any stress treatments, the root growth of all transgenic variants was almost as same as those of wild type. However, when the seedlings were transferred onto MS medium supplemented with 5 µM ABA or 150 mM NaCl for several days, primary root growth of *Atcpk11-2* mutant was much better than that of wild type, and root growth of *GhDi19-1/cpk11* and *GhDi19-2/cpk11* transgenic lines was similar to wild type, whereas root growth of the *GhDi19-1(S116D)/cpk11* and *GhDi19-2(S114D)/cpk11* transgenic seedlings was inhibited much more than that of wild type (Fig. 7d). Roots of *Atcpk11-2* mutant were longer than those of wild type, and root length of the *GhDi19-1/cpk11* and *GhDi19-2/cpk11* transgenic lines was similar to wild type, but roots of *GhDi19-1(S116D)/cpk11* and *GhDi19-2(S114D)/cpk11* transgenic seedlings were shorter than those of wild type under ABA and NaCl treatments (Fig. 7e). These results indicated that overexpression of GhDi19-1/-2 in *Atcpk11* background could recover the salt- and ABA-insensitive phenotype of the mutant, implying the other CDPK (besides CPK11) may also activate GhDi19-1/-2 in the transgenic *Arabidopsis* plants, and Ser phosphorylation is important for functionally activating GhDi19-1/-2 in response to salt stress and ABA signaling during seed germination and early seedling development.

Discussion

A large number of genes encoding receptors, kinases, transcription factors and other signal molecules in plants are induced after exposure to various abiotic stresses^{35,36}. These genes function in various ways to confer plants stress tolerance^{37–39}. Due to much low similarity of the different Di19-related proteins outside the zinc finger domain, Di19 proteins may play diverse roles in plant development and in response to abiotic stresses³². Previous studies revealed that GhDi19-1 and GhDi19-2 are involved in plant response to salt stress and ABA signaling³⁴. In this study, we further investigated the mechanism of GhDi19-1 and GhDi19-2 involving in response to salt stress and ABA signaling, and found the Ser site is essential for GhDi19-1 and GhDi19-2 proteins involving in plant response to salt and ABA during seed germination and early seedling development.

Modification of phosphorylation by kinases and of dephosphorylation by phosphatases is an important physiological process as plants respond to abiotic stresses and ABA signaling. Some ABA/stress signaling regulators are modulated at the post-translational level by changing their phosphorylation states^{40–44}. Calcium-dependent protein kinases (CDPKs) are unique serine/threonine kinases in plants responding in abiotic stresses⁴⁵. Their multifunctionality and signaling specificity may be conferred by their ability to phosphorylate different substrates. Arabidopsis CPK4 and CPK11 positively regulate ABA signaling via phosphorylating downstream ABA-responsive transcription factors in plants¹¹. Furthermore, previous studies revealed that most of seven Arabidopsis AtDi19s are phosphorylated by CPK3 and CPK11 in vitro in a Ca⁺²-dependent manner³³. Likewise, both GhDi19-1 and GhDi19-2 contain a conserved nuclear localization signal (NLS), in which two putative kinase phosphorylation sites (Thr114 and Ser116 in GhDi19-1, and Thr112 and Ser114 in GhDi19-2, respectively) are located³⁴. In this study, our data indicated that GhDi19-1/-2 could be phosphorylated in vitro by CPK11 in a Ca²⁺-dependent manner, and the Ser site of GhDi19-1/-2 is required for phosphorylation and normal subcellular localization of GhDi19-1 and GhDi19-2 proteins. Previous study reported that CPK11 phosphorylates Ser104 and Ser107 sites within the AtDi19-1 bipartite NLS probably, but effects of Ser phosphorylation at different sites on Di19 proteins involving in plant response to salt and ABA are unknown so far³³. In this study, our data demonstrated the Ser site (but not Thr site) is crucial for functionally activation of cotton Di19-1/-2 in response to salt stress and ABA signaling.

AtCPK11 interacts with AtDi19-1 protein in the cell nucleus, and increases AtDi19-1 transactivation of *PR1*, *PR2*, and *PR5* expression³¹. Furthermore, AtCPK4 and AtCPK11 overexpression transgenic plants display the hypersensitivity to ABA during seeds germination and seedlings growth¹¹. Similar phenotype was also observed in the *GhDi19-1/-2* overexpression transgenic *Arabidopsis*³⁴. These findings indicated that there is a specific functional relationship of cotton Di19s and CDPKs. Additionally, overexpression of GhDi19-1/-2 in *Atcpk11* could recover the salt- and ABA-insensitive phenotype of the mutant, implying the other kinases (besides CPK11) may also activate GhDi19-1/-2 in the transgenic *Arabidopsis* plants.

Based on the data presented in this study, we thought that Di19 proteins may be potential downstream targets of CDPKs in ABA signal pathway during early plant growth and development. When plants are exposed to abiotic stress stimuli (such as drought and high salinity etc.), concentrations of intracellular ABA and Ca²⁺ may be increased in plants for responding the environmental stress signaling. Subsequently, Ser/Thr kinases (e.g. CDPK) are activated by ABA and in turn the activated kinases phosphorylate Di19 proteins at Ser site in the cell nucleus. Finally, the activated Di19 proteins transduce the signals to downstream ABA- and stress-responsive genes, thereby promoting plants response to abiotic stresses.

Methods

Plants materials and growth conditions. A T-DNA insertion mutant (named *Atcpk11-2*, SALK_054495) of *Arabidopsis* CPK11 (AT3g05700) was obtained from ABRC (www.arabidopsis.org/abrc). Seeds of *Arabidopsis*

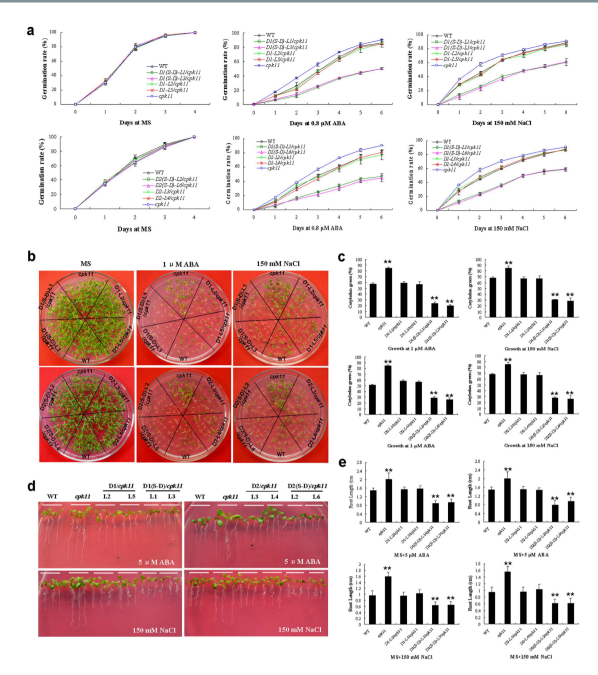


Figure 7. Phenotypic assay of GhDi19-1/-2 and their activated form GhDi19-1/-2 (S-D) transgenic seedlings in Atcpk 1 mutant under salt and abscisic acid (ABA) treatments. (a) Statistical analysis of seed germination rate. Seeds of wild type, Atcpk11 mutant, and GhDi19-1/-2 or GhDi19-1/-2(S-D) overexpression transgenic lines in Atcpk11 mutant were germinated on MS medium with or without 150 mM NaCl and 0.8 μM ABA, respectively. (b) Growth status of seedlings growing on MS medium with or without 1 μM ABA and 150 mM NaCl for 10 days, respectively. (c) Statistical analysis of cotyledon expansion and greening of seedlings growing on MS medium with 1 µM ABA and 150 mM NaCl for 10 days, respectively. (d) Phenotypes of seedlings under NaCl and ABA treatments. After germination on MS medium for 48 h, the wild type, Atcpk11 mutant, and GhDi19-1/-2 or GhDi19-1/-2(S-D) overexpression transgenic seedlings in Atcpk11 mutant were transferred and grew for 10 days on MS medium without or with 150 mM NaCl and $5\,\mu\text{M}$ ABA, respectively, in the vertical position. (e) Statistical analysis of root length of 10-day-old seedlings growing on MS medium with 5 µM ABA and 150 mM NaCl. Mean values and standard errors (bar) were shown from three independent experiments (n > 60 seedlings per each line). Asterisk represents that there was (very) significant difference between the transgenic lines and wild type in independent t-tests (one asterisk: P value < 0.05; two asterisks: P value < 0.01). WT, wild type; *cpk11*, the *Atcpk11-2* mutant; *D1-L2/cpk11* and *D1-L5/cpk11*, the *GhDi19-1* overexpression transgenic line 2 and 5 in Atcpk11-2 mutant; D1(S-D)-L1/cpk11 and D1(S-D)-L3/cpk11, the GhDi19-1(S-D) overexpression transgenic line 1 and 3 in Atcpk11-2 mutant; D2-L3/cpk11 and D2-L4/cpk11, the GhDi19-2 overexpression transgenic line 3 and 4 in Atcpk11-2 mutant; D2(S-D)-L2/cpk11 and D2(S-D)-L6/cpk11, the GhDi19-2(S-D) overexpression transgenic line 2 and 6 in Atcpk11-2 mutant. S-D, the S₁₁₆ of GhDi19-1 was mutated to D₁₁₆ and the S₁₁₄ of GhDi19-2 was mutated to D₁₁₄. S, serine (Ser); D, aspartic acid (Asp).

thaliana (Columbia ecotype), including wild type, mutant and transgenic lines, were surface-sterilized with 75% ethanol for 1 min and 10% NaClO for 5 min, followed by washing with sterile water. The sterilized Arabidopsis seeds were plated on Murashige and Skoog (MS) medium. After stratification at 4 °C for 2 days, the plates were transferred to a plant growth incubator (Sanyo, Osaka, Japan) for seed germination (16 h light/8 h dark at 22 °C) ten days later, seedlings were transplanted in soil and grown in a growth room under the conditions of 16 h light/8 h dark cycle at 22–24 °C. Tissues were derived from these seedlings for RNA extraction.

Generation of transgenic plant and mutant lines. The coding sequences of cotton *GhDi19-1* and *GhDi19-2* genes, amplified from its cDNA by PCR with the proofreading *pfu* DNA polymerase, were cloned into pBI121 vector with *BamH I/Sac I* sites to replace the *GUS* gene, respectively³⁴.

To introduce the Ser116-to-Asp (S116D) mutation of GhDi19-1 or the Ser114-to-Asp (S114D) mutation of GhDi19-2 for constitutively activated GhDi19-1 or GhDi19-2, primer-based site-directed mutations were performed, using primers as follows: *GhDi19-1(S116D)* 5′-CTCATTCCGACCTTGACCTTCTGAGGA -3′ and 5′- TCCTCAGAAGGTCAAGGTCGGAATGAG-3′; *GhDi19-2(S114D)* 5′-CATTCAGACCTT GACCTTCTGAGGAAGGTCAAGGTCAAGGTCTGAATG-3′. The coding sequences of *GhDi19-1(S-D)* and *GhDi19-2(S-D)* were cloned into pCAMBIA1301 vector. Mutant alleles of *Atcpk11-2* (SALK_054495) were used for experiments¹¹. Complementation experiments using *GhDi19-1/-2* and *GhDi19-1/-2* (S-D) transgenes were done in *Atcpk11-2* background.

All generated binary vectors were transformed into *Agrobacterium turmefaciens* strain GV3101. *Arabidopsis* transformation was performed by the floral dip method⁴⁶, and transformants were identified on selective medium with kanamycin or hygromycin.

Quantitative RT-PCR analysis. Total RNA was extracted from 10-day-old Arabidopsis seedlings of wild type, GhDi19-1-, GhDi19-1(S116A)-, GhDi19-1(T114A)- and GhDi19-1(S/T-A/ A)-overexpression transgenic lines, GhDi19-2-, GhDi19-2(S114A)-, GhDi19-2(T112A)- and GhDi19-2(S/T-A/A)-overexpression transgenic lines, Atcpk11 mutant, GhDi19-1(S-D)- and GhDi19-2(S-D)-overexpression transgenic lines in Atcpk11-2 mutant. Real-time quantitative RT-PCR (qRT-PCR) analysis was performed as described as previously⁴⁷. Primer pairs for qRT-PCR analysis are as follows: GhDi19-1 5'-ATGGATGCTGATTCATGGAGT-3' and 5'-TTATAAAATTTCATCAGGCAT-3'; GhDi19-1(S116A) 1(T114A) 5'-ATGGATGCTGATTCATGGAGT-3' and 5'-GGATCTCATTCCGCGCTTTCTCTTCTG -3'; GhDi19-1(S/T-A/A) 5'-ATGGATGCTGATTCATGGAGT-3' and 5'-TCATTCCGCGCTTGCT CTTCTGAGG-3'; GhDi19-1(S116D) 5'-ATGGATGCTGATTCATGGAGT-3' 5'-CTCATTCCGACCTTGACCTTCTGAGGA-3'; GhDi19-2 5'-ATGGATGCTGATCCATGGAC-3' and 5'-TCATAAAACATCATCAAGAAT-3'; GhDi19-2(S114A) 5'-ATGGATGCTGATCCATGGAC-3' and 5'-CTCATTCAACGCTTGCTCTTCTGAGGAAAG-3'; GhDi19-2(T112A) 5'-ATGGATGCTGATTC ATGGAGT-3' and 5'-GTGGATCTCATTCAGCGCTTTCTCTTC-3'; GhDi19-2(S/T-A/A) 5'-ATGGATGCTGATTCATGGAGT-3' and 5'-GATCTCATTCAGCGCTTGCTCTTCTGAGG-3'; GhDi19- $2 (S114D)\ 5' - ATGGATGCTGATCCATGGAC-3'\ and\ 5' - CATTCAGACCTTGACCTTCTGAGGAAAG-3';$ Atcpk11-2 5' - GAGAGAGTCAAAAAAATTGGAGAA-3' and 5' -AAACCAATTAGGCGATGAACC-3'. Expression level of Arabidopsis ACTIN2 was monitored with forward 5'-GAAATCACAGCACTTGCACC-3' and reverse 5'-AAGCCTTTGATCTTGAGA GC-3' primers to serve as an internal control. For all the above qRT-PCR reactions, the assays were repeated three times along with three independent repetitions of the biological experiments, and means of three biological experiments were calculated for estimating gene expression levels.

Phenotypic analysis of the transgenic *Arabidopsis* plants. Homozygous plants (T3 and T4 generations) of the *GhDi19-1/-2* transgenic lines and their site-mutations were used for phenotypic analysis, employing wild type, mutant and the transgenic line harboring the "empty vector" (i.e. pBI121-eGFP) as controls. We first tested a series of ABA and NaCl concentrations in the pre-experiments, and determined which concentration of abscisic acid (ABA) or NaCl is suitable to the respective experiments. Then, seeds of wild type and independent transgenic lines, mutants, or transgenic mutants were germinated on MS medium supplemented without or with 150 mM NaCl, 0.8 and l µM ABA, respectively. The seeds were incubated at 4 °C for 2 days and then transferred into a plant growth incubator at 22 °C conditions (16 h light/8 h dark). Seeds were considered successfully germinated when radicles completely penetrated the seed coats. Germination rate and proportion of seedlings with opened green cotyledons were expressed as a percentage of the total number of seeds plated.

The seedling growth experiments were performed as described previously 11 . Seeds were germinated after stratification on MS medium for 48 h and then transferred onto MS medium containing 150 mM NaCl and $5\,\mu\text{M}$ ABA in the vertical position. The growth status of seedling was recorded for 10 days and the length of seedling primary roots was measured at tenth day after the transfer. In addition, seedling growth was also assessed by directly planting the seeds on NaCl- or ABA-containing MS medium to investigate the response of seedling growth to salt or ABA for 10 days.

Statistical analysis was performed in all the experiments. At least 300 seeds of each line were used for analyzing seed germination rate and cotyledon greening rate, and 30–60 seedlings in vertical cultivation were used for measuring root length. The experiments were repeated at least three times with three technical replications.

Subcellular localization of GhDi19-1 and GhDi19-1 protein. The coding sequences of GhDi19-1, GhDi19-1(S116D), GhDi19-1(S116A), GhDi19-1(T114A), GhDi19-1(S/T-A/A), GhDi19-2, GhDi19-2(S114D), GhDi19-2(S114A), GhDi19-2(T112A) and GhDi19-2(S/T-A/A) were cloned into pBI121-eGFP vector to generate GhDi19-1:eGFP, GhDi19-1(S116D):eGFP, GhDi19-1(S116A):eGFP, GhDi19-1(T114A):eGFP, GhDi19-1(S/T-A/ A):eGFP, GhDi19-2:eGFP, GhDi19-2(S114D):eGFP, GhDi19-2(S114A):eGFP, GhDi19-2(T112A):eGFP and GhDi19-2(S/T-A/A):eGFP construct, respectively, and then introduced into Arabidopsis by the floral dip method. The harvested seeds were germinated on selective MS medium for selecting transgenic plants. Subsequently, GFP fluorescence in root cells of the transgenic seedlings was observed under a SP5 Meta confocal laser microscope (Leica, Germany) with a filter set ($488 \,\mathrm{nm}$ for excitation and $506 \,\sim 538 \,\mathrm{nm}$ for emission). SP5 software (Leica, Germany) was employed to record and process the digital images taken. Primers used in vector construction are as follows: GhDi19-15'-CTTGGATCCATGGATGCTGATTCATGGAG-3' and 5'-GGGTCTAGATTATAAAATTTCATCAGGC -3'; GhDi19-2 5'-CTTGGATCCATGGATGCTGATCCATGGAC-3' and 5'-GGGTCTAGATCATAAA ACATCATCAAGAA-3'; GhDi19-1(S116D) 5'-CTCATTCCGACCTTGACCTTCTGAGGA-3' and 5'-TCCTCAGAAGGTCAAGGTCGGAATGAG-3'; GhDi19-2(S114D) 5'-CATTCAGAC CTTGACCTTCTGAGGAAAG-3' and 5'-CTTTCCTCAGAAGGTCAAGGTCTGAATG-3'; GhDi19-1(S116A) 5'-TCCTCAGAAGAGCAAGCGTGGAATGAG-3' and 5'-CTCATTCCACGCTTGCTCTTCTGAGGA-3'; GhDi19-2(S114A) 5'-GAAGAGAAAGCGCTGAATGAGATCCAC-3' and 5'-GTGGATCTCATTCAG CGCTTTCTCTC-3'; GhDi19-1(T114A) 5'-CAGAAGAGAAAGCGCGGAATGAGATCC-3' and 5'-GGATCT CATTCCGCGCTTTCTCTG-3'; GhDi19-2(T112A) 5'-CTTTCCTCAGAAGAGCAAGCGTTGAATGAG -3' and 5'-CTCATTCAACGCTTGCTCTTCTGAGGAAAG-3'; GhDi19-1(S/T-A/A) 5'-CCTCAGAAGA GCAAGCGCGAATGA-3' and 5'-TCATTCCGCGCTTGCTCTTGAGG-3'; GhDi19-2(S/T-A/A) 5'-CCTCAGAAGAGCAAGCGCTGAATGAGATC-3' and 5'-GATCTCATTCAGCGCTTGCTCTTCTGAGG-3'.

In vitro phosphorylation assay. The coding sequences of AtCPK11, GhDi19-1, GhDi19-1(S116A), GhDi19-2 and GhDi19-2(S114A) were inserted downstream the malE gene, which encodes maltose-binding protein (MBP), in pMAL-c2X vector for expressing MBP-AtCPK11, MBP-GhDi19-1, MBP-GhDi19-1(S116A), MBP-GhDi19-2 and MBP-GhDi19-2(S114A) fusion proteins, respectively. In vitro phosphorylation assay was performed for the fusion proteins and MBP protein (control) purified from Escherichia coli strain BL21 by MBP's affinity for maltose (NEW ENGLAND), according to the method described previously^{48,49}. Primers used in vector construction are as follows: AtCPK11 5'-CTTGGATCCATGGAGACGAAGCCAAACCCTAG-3' and 5'-GGGGTCGACTCAGTCATCAGATTTTTCACCA-3'; GhDi19-1 5'-CTTGGATCCATGGA TGCTGATCCATGGAG-3' and 5'-GGGTCTAGATTATAAAATTTCATCAGGC-3'; GhDi19-25'-CTTGGATC CATGGATGCTGATCCATGGAC-3' and 5'-GGGTCTAGATCATAAAACATCATCAAGAA-3'; GhDi19-1(S116A) 5'-CTCATTCCACGCTTGCTCTTCTGAGGA-3' and 5'-TCCTCAGAAGAGCAAGCGTGGAATGAG -3'; GhDi19-2(S114A) 5'-CTCATTCAACGCTTGCTCTTCTGAGGA-3' and 5'-CTCATTCA ACGCTTGCTCTTCTGAGGAAAG-3' and 5'-CTCATTCA ACGCTTGCTCTTCTGAGGAAAG-3'.

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Author Contributions

X.B.L. and L.X.Q. conceived and designed the research. L.X.Q., X.Y.N., R.H., G.L. and W.L.X. performed the experiments. L.X.Q., W.L.X. and X.B.L. analyzed the data and wrote the article.

Additional Information

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