A Ca\(^{2+}\)/CaM-regulated transcriptional switch modulates stomatal development in response to water deficit

Chan Yul Yoo\(^{1,6}\), Noel Mano\(^{2}\), Aliza Finkler\(^{3}\), Hua Weng\(^{1}\), Irene S. Day\(^{4}\), Anireddy S. N. Reddy\(^{4}\), B. W. Poovaiah\(^{5}\), Hillel Fromm\(^{3}\), Paul M. Hasegawa\(^{3}\) & Michael V. Mickelbart\(^{1,2}\)

Calcium (Ca\(^{2+}\)) signals are decoded by the Ca\(^{2+}\)-sensor protein calmodulin (CaM) and are transduced to Ca\(^{2+}\)/CaM-binding transcription factors to directly regulate gene expression necessary for acclimation responses in plants. The molecular mechanisms of Ca\(^{2+}\)/CaM signal transduction processes and their functional significance remains enigmatic. Here we report a novel Ca\(^{2+}\)/CaM signal transduction mechanism that allosterically regulates DNA-binding activity of GT2-LIKE 1 (GTL1), a transrepressor of STOMATAL DENSITY AND DISTRIBUTION 1 (SDD1), to repress stomatal development in response to water stress. We demonstrated that Ca\(^{2+}\)/CaM interaction with the second helix of the GTL1 N-terminal trihelix DNA-binding domain (GTL1N) destabilizes a hydrophobic core of GTL1N and allosterically inhibits third helix docking to the SDD1 promoter, leading to osmotic stress-induced Ca\(^{2+}\)/CaM-dependent activation (de-repression) of SDD1 expression. This resulted in GTL1-dependent repression of stomatal development in response to water-deficit stress. Together, our results demonstrate that a Ca\(^{2+}\)/CaM-regulated transcriptional switch on a trihelix transrepressor directly transduces osmotic stress to repress stomatal development to improve plant water-use efficiency as an acclimation response.

Plants sense and respond to external stimuli to acclimate and adapt to diverse environmental niches. In response to water deficit, plants limit transpirational water loss via reduced stomatal aperture as a rapid plant response\(^1\). As a longer-term response, plants may repress stomatal development, the result of which is reduced water loss to improve water-use efficiency (WUE) and/or drought tolerance\(^2\). Hypersmotic stress induces signaling pathways that activate or repress genes necessary for water-deficit acclimation responses\(^3\) and transcriptional regulation is a key regulatory mechanism that governs stomatal development\(^4\). The abundance of SPEECHLESS, the master transcriptional regulator of stomatal cell identity, is decreased by osmotic stress via a mitogen-activated protein kinase (MAPK) cascade\(^5\). However, the earliest signal transduction mechanism from sensing water-deficit to the direct regulation of gene expression responsible for repression of stomatal development remains enigmatic.

Calcium ions (Ca\(^{2+}\)) are versatile second messengers regulating plant growth and development. Environmental stimuli, and specifically hypersmotic stress, induce rapid and transient changes in cytosolic and nuclear Ca\(^{2+}\) concentrations that differ in amplitude and duration (Ca\(^{2+}\) signatures)\(^6\)-\(^12\). These stimuli-specific Ca\(^{2+}\) changes are decoded by Ca\(^{2+}\)-sensor proteins, including the ubiquitous Ca\(^{2+}\)-binding protein calmodulin (CaM). Ca\(^{2+}\) binding to CaM induces conformational change from a closed to an open state, facilitating hydrophobic and electrostatic interactions with basic amphipathic helices of target peptides to activate or repress the molecular activity of target proteins\(^8\)-\(^11\). Among various CaM-binding targets, Ca\(^{2+}\)/CaM-binding transcription

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1Department of Horticulture and Landscape Architecture, Purdue University, West Lafayette, IN, 47907, USA. 2Department of Botany and Plant Pathology, Purdue University, West Lafayette, IN, 47907, USA. 3School of Plant Sciences and Food Security, Faculty of Life Sciences, Tel Aviv University, Tel Aviv, 6997801, Israel. 4Department of Biology and Program in Cell and Molecular Biology, Colorado State University, Fort Collins, CO, 80523, USA. 5Department of Horticulture, Washington State University, Pullman, WA, 99164, USA. 6Present address: Department of Botany and Plant Sciences, Institute of Integrative Genome Biology, University of California Riverside, Riverside, CA, 92521, USA. Correspondence and requests for materials should be addressed to M.V.M. (email: mickelbart@purdue.edu)
Ca2+/CaM directly binds to the GTL1 N-terminal trihelical DNA-binding domain. We hypothesized that Ca2+/CaM directly binds to AtGT1, a negative regulator of stomatal development, to mediate water-deficit stress acclimation. The domain structure of GTL1 includes two trihelical DNA-binding domains in the N- and C-termini (Fig. 1a). A prediction algorithm identifies the α2 helix within the N-terminal DNA-binding domain (GTL1N) as a putative CaM-binding domain, but not within the C-terminal DNA-binding domain (GTL1C) (Supplementary Fig. S1). The α2 helix contains a CaM-binding domain, the GTL1N trihelical domain by NMR spectroscopy revealed that the trihelical tertiary structure is stabilized by hydrophobic residues forming a hydrophobic core with o3 docking to the CaM box in target gene promoters. GT2 factors have been implicated in developmental, light, and abiotic stress responses in Arabidopsis, poplar, soybean, and wheat. We previously reported that the Arabidopsis GT2-LIKE 1 (GTL1) is a transrepressor of STOMATAL DENSITY AND DISTRIBUTION 1 (SDD1) that negatively regulates stomatal development and transpiration. Ca2+/CaM-binding affinity has been demonstrated in AtGT2L and PtaGTL1. However, Ca2+/CaM signal transduction mechanisms of GTL1 and their biological implications are not known. Here, we show that Ca2+/CaM directly binds to the N-terminal DNA-binding domain of GTL1 and allosterically inhibits DNA-binding activity by disturbing the hydrophobic core. Hyperosmotic stress-induced Ca2+/CaM allosteric control on the GTL1 transrepressor leads to SDD1 derepression, both by inhibiting binding to and promoting release from the SDD1 promoter, leading to repression of stomatal development in response to water-deficit stress. We propose that this allosteric control of the GTL1 transcription factor by Ca2+/CaM is a transcriptional switch to modulate stomatal development, thereby conserving plant water loss as a long-term developmental adaptation during water stress.

**Results**

**Ca2+/CaM binds to the GTL1 N-terminal trihelical DNA-binding domain.** We hypothesized that Ca2+/CaM directly binds to AtGT1, a negative regulator of stomatal development, to mediate water-deficit stress acclimation. The domain structure of GTL1 includes two trihelical DNA-binding domains in the N- and C-termini (Fig. 1a). A prediction algorithm identifies the α2 helix within the N-terminal DNA-binding domain (GTL1N) as a putative CaM-binding domain, but not within the C-terminal DNA-binding domain (GTL1C) (Supplementary Fig. S1). The α2 helix contains a CaM-binding domain, the GTL1N trihelical domain by NMR spectroscopy revealed that the trihelical tertiary structure is stabilized by hydrophobic residues forming a hydrophobic core with o3 docking to the CaM box in target gene promoters. GT2 factors have been implicated in developmental, light, and abiotic stress responses in Arabidopsis, poplar, soybean, and wheat. We previously reported that the Arabidopsis GT2-LIKE 1 (GTL1) is a transrepressor of STOMATAL DENSITY AND DISTRIBUTION 1 (SDD1) that negatively regulates stomatal development and transpiration. Ca2+/CaM-binding affinity has been demonstrated in AtGT2L and PtaGTL1. However, Ca2+/CaM signal transduction mechanisms of GTL1 and their biological implications are not known. Here, we show that Ca2+/CaM directly binds to the N-terminal DNA-binding domain of GTL1 and allosterically inhibits DNA-binding activity by disturbing the hydrophobic core. Hyperosmotic stress-induced Ca2+/CaM allosteric control on the GTL1 transrepressor leads to SDD1 derepression, both by inhibiting binding to and promoting release from the SDD1 promoter, leading to repression of stomatal development in response to water-deficit stress. We propose that this allosteric control of the GTL1 transcription factor by Ca2+/CaM is a transcriptional switch to modulate stomatal development, thereby conserving plant water loss as a long-term developmental adaptation during water stress.

**The GTL1 N-terminal trihelical DNA-binding domain requires hydrophobic core formation for its DNA-binding activity.** Ca2+/CaM binding to the α2 helix within the GTL1 N-terminal trihelical DNA-binding domain suggests a potential link between Ca2+/CaM binding and the regulation of DNA-binding activity. To understand the functional characteristics of the trihelical domain, the structure of GTL1N was constructed by comparative homology modeling with the NMR structure of the AtGT1 trihelical DNA-binding domain as a template. The primary and secondary structural comparison indicates that the key residues required for hydrophobic core formation in GT-1 are conserved in the GTL1N trihelical DNA-binding domain with 35.94% sequence identity (Supplementary Fig. S3a), which is above 30%, a good standard for accuracy of homology modeling. The modeled GTL1N trihelical domain indicates that GTL1N α1 (residues 65–83) and α2 (residues 90–102) are oblique to each other, and that α3 (residues 107–125) is perpendicular to α1 (Fig. 3a). Ten residues (N terminus-W63; α1-L71, L72, and F82; α2-L91, W92, V95, and L99; and α3-F115 and V118) form a hydrophobic core to stabilize the GTL1N trihelical structure, which is conserved with hydrophobic residues important for the hydrophobic core of GT-1 (Fig. 3a and Supplementary Fig. S3b). Charged residues on the α3 helix, important for electrostatic interactions with the GT box cis-element, are also conserved between GT-1 and GTL1 (Supplementary Fig. S3c), suggesting that the functional structure of the trihelix DNA-binding domain is conserved between GT-1 and GTL1.
The GTL1N tertiary structure allows the α3 helix to interact with the major groove of the GT-3 box in the predicted docking structure between GTL1N and the SDD1 promoter (5′-gcttGGTAAAactt-3′) (Supplementary Fig. S4). The predicted model suggests that GTL1N-SDD1 promoter interactions occur via hydrogen bonds between α3 K109 (hydrogen donor) and the first guanine (Gua1) of the GT3 box (hydrogen acceptor), and α3 E116 (hydrogen acceptor) and the fourth adenine (Ade4) of the GT3 box (hydrogen donor) (Supplementary Fig. S4b). Other residues W63, α2-W92, and α3-K110, K112, K114, N117, Q119, and K120 interact with the negatively charged phosphate backbone of the GT3 box (Supplementary Fig. S4c), which stabilizes the GTL1N-SDD1 interaction.

Figure 1. CaM binds to the N-terminal DNA-binding domain of GTL1 in a Ca\(^{2+}\)-dependent manner. (a) Schematic illustration of GTL1 domain topology with N- and C-terminal trihelical domains. The primary and secondary structure of GTL1N (residue number 60–126) consists of three α-helices (α1 – grey, α2 – green, and α3 – orange) with α2 containing a predicted CaM-binding domain. (b) The helical wheel projection of the α2 helix shows a canonical amphipathic helix with hydrophobic (yellow) and hydrophilic (basic – blue and acidic – red) residues. (c) GTL1N interaction with Ca\(^{2+}\)/CaM was performed by in vitro pull-down assay using Escherichia coli expressed MBP-GTL1N or MBP to pull down in vitro-translated HA-AtCaM2 that was detected by immunoblots using anti-HA antibodies (upper panel). Immobilized MBP and MBP-GTL1N fusion proteins are shown in the Coomassie blue-stained SDS-PAGE gel (lower panel). (d) Schematic illustration of the bait and prey proteins used in the pull-down assays. Ca\(^{2+}\)-dependent CaM interaction with GTL1N or GTL1C were performed by pull-down assay using HA-AtCaM2. Bound and 10% of input HA-AtCaM2 fractions were detected by immunoblot using anti-HA antibodies. The MBP-GTL1N (Nt) and MBP-GTL1C (Ct) fusion protein bands are indicated by red arrows. Full-length blots and gels for c and d are presented in Supplementary Fig. S10.
promoter interaction. Based on the docking model of GTL1N-SDD1 promoter, we hypothesize that a hydrophobic core in the trihelical GTL1N DNA-binding domain is essential for its DNA-binding activity. To validate the importance of the hydrophobic core in tertiary structure for DNA-binding activity, hydrophobic residues (L91, W92, and V95) were substituted to acidic or basic amino acids to disrupt the formation of a hydrophobic core. Consistent with the homology modeling and docking prediction, site-directed mutations (L91R, W92R, and V95D) of hydrophobic residues to charged residues result in the complete loss of binding to the SDD1 promoter (Fig. 3b). These data support the model that the hydrophobic core formed by the hydrophobic residues in the GTL1N trihelical structure is required for native folding and its function as a DNA-binding transcription factor. However, the H94E mutation in the H94 residue required for Ca2+/CaM interaction did not result in the loss of DNA binding to the SDD1 promoter or rice PHYA promoter (Fig. 3c), suggesting that the α2 helix is not directly involved with DNA-recognition. Together, this result further supports the notion that the hydrophobic core formation results in a protein topology for surface exposure of the α2 (H94, R97, and K98) and α3 (K110, K112, K114, and K120) basic residues that are necessary for CaM and DNA interactions, respectively.

**Ca2+/CaM binding to α2 of GTL1N inhibits its association with and facilitates dissociation from the SDD1 promoter.** Ca2+/CaM binding to the α2 helix within the GTL1N N-terminal DNA-binding domain prompted us to postulate that Ca2+/CaM binding may allosterically inhibit GTL1N DNA binding to the SDD1 promoter. This hypothesis is based on the typical Ca2+/CaM-target peptide structure requiring hydrophobic residues9,34,

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**Figure 2.** Ca2+/CaM binds to the α2 helix of GTL1N. (a) Schematic illustration of two protein fragments (GTL1N and GTL1NΔdel) fused to maltose-binding protein (MBP). *In vitro* pull-down assay was performed using MBP-fusion proteins with Ca2+/HA-AtCaM2. Bound and 10% of input HA-AtCaM2 fractions were detected by immunoblot using anti-HA antibodies. The MBP-GTL1N and MBP- GTL1NΔdel fusion protein bands are indicated by red arrows. (b) Sequence alignment of the α2 helix of the N-terminal (Nt) or C-terminal (Ct) trihelical domain from GT2 family proteins, including GTL1. AtGTL1_Nt and AtGT2L_Ct have CaM-binding activity. *Indicates the residue that distinguishes CaM-binding and non-CaM-binding groups. (c) *An in vitro* pull-down assay was performed using MBP-GTL1N or MBP-GTL1N[H94E] fusion proteins with Ca2+/HA-AtCaM2. Bound and 10% of input HA-AtCaM2 fractions were detected by immunoblot using anti-HA antibodies. The MBP-GTL1N and MBP-GTL1N[H94E] fusion protein bands are indicated by red arrows. Full-length gels for a and c are presented in Supplementary Fig. S10.
which are important to form the hydrophobic core of GTL1N (Fig. 3a). We first tested this hypothesis with an in vitro electrophoretic mobility shift assay (EMSA) that was previously used to show GTL1 DNA binding to the SDD1 promoter27. GTL1N was able to interact with the SDD1 promoter fragment in the presence of CaM or CaM with EGTA (Fig. 4a). However, addition of Ca\(^{2+}\) substantially reduced the binding affinity with SDD1 (Fig. 4a). Interestingly, when GTL1N was pre-incubated with the SDD1 promoter fragment, the addition of CaM in the presence of Ca\(^{2+}\) was also able to dissociate GTL1N from the SDD1 promoter fragment (Fig. 4b). This result indicates that Ca\(^{2+}\)/CaM is able to dissociate GTL1N from the SDD1 promoter. To further confirm the effect of Ca\(^{2+}\)/CaM on GTL1N
DNA-binding activity, the EMSA was performed using GTL1N or GTL1N[H94E], and a biotin-labeled SDD1 promoter fragment that includes the GT3 box. (a) GTL1N (1 or 5 μg) was pre-incubated with human CaM (100 ng) without or with 2 mM CaCl₂ or 10 mM EGTA, and then incubated with the biotin-labeled SDD1 promoter fragment (250 ng). (b) GTL1N (2.5 or 5 μg) was pre-incubated with the SDD1 promoter fragment without or with 2 mM CaCl₂, and then with human CaM (100 ng). (c) GTL1N or GTL1N[H94E] was pre-incubated with the SDD1 promoter fragment without or with 2 mM CaCl₂ and then with human CaM (50 or 100 ng). (d) ChIP assays using anti-HA antibody were performed using protoplasts isolated from gtl1-4 plants, transformed with HA-GTL1 or HA-GTL1[H94E], and incubated with 0 or 200 mM mannitol for 1 h. Input is the total isolated chromatin before immunoprecipitation. Mouse IgG was the negative control for immunoprecipitation. The SDD1 promoter region including the GT3 box was amplified by SDD1 promoter-specific primers.

**Figure 4.** Hyperosmotic stress-induced Ca²⁺/CaM binding to the α2 helix inhibits DNA-binding activity of GTL1 to the SDD1 promoter. (a–c) EMSA was performed using GTL1N or GTL1N[H94E], and a biotin-labeled SDD1 promoter fragment that includes the GT3 box. (a) GTL1N (1 or 5 μg) was pre-incubated with human CaM (100 ng) without or with 2 mM CaCl₂ or 10 mM EGTA, and then incubated with the biotin-labeled SDD1 promoter fragment (250 ng). (b) GTL1N (2.5 or 5 μg) was pre-incubated with the SDD1 promoter fragment without or with 2 mM CaCl₂, and then with human CaM (100 ng). (c) GTL1N or GTL1N[H94E] was pre-incubated with the SDD1 promoter fragment without or with 2 mM CaCl₂ and then with human CaM (50 or 100 ng). (d) ChIP assays using anti-HA antibody were performed using protoplasts isolated from gtl1-4 plants, transformed with HA-GTL1 or HA-GTL1[H94E], and incubated with 0 or 200 mM mannitol for 1 h. Input is the total isolated chromatin before immunoprecipitation. Mouse IgG was the negative control for immunoprecipitation. The SDD1 promoter region including the GT3 box was amplified by SDD1 promoter-specific primers.
tobacco and Arabidopsis\textsuperscript{35–38}. Mannitol reduced GTL1 association with the SDD1 promoter, but did not substantially reduce GTL1[H94E] association with the SDD1 promoter (Fig. 4d). Together, both in vitro and in vivo results indicate that hyperosmotic stress-induced Ca\textsuperscript{2+}/CaM attenuates GTL1 docking to the SDD1 promoter.

**Hyperosmotic stress induces SDD1 expression through Ca\textsuperscript{2+}/CaM-dependent inhibition of the GTL1 transrepressor.** To understand the biological significance of Ca\textsuperscript{2+}/CaM interaction with GTL1 on the regulation of SDD1 promoter-binding activity, we hypothesized that Ca\textsuperscript{2+}/CaM-dependent inhibition of the GTL1/SDD1 promoter complex interaction is a key signal transduction mechanism to activate (derepress) SDD1 expression in response to water-deficit stress. Indeed, transcript abundance of SDD1 mRNA is up-regulated in response to water-deficit stress when DREB2A, a drought-induced marker gene, was also up-regulated (Fig. 5a).

Next, to determine if hyperosmotic stress regulates SDD1 transcription through GTL1 and Ca\textsuperscript{2+} and CaM signaling, we first established the 2 kb SDD1 promoter-reporter system (Fig. 5b). SDD1-LUC activity was higher in gtl1-4 than in wild-type (Col-0) protoplasts, whereas GTL1 promoter-driven GTL1-GFP expression in gtl1-4 (GT11/gtl1-4) protoplasts repressed SDD1 expression to a level comparable to wild-type protoplasts (Fig. 5b), which is consistent with previous results in mature leaves\textsuperscript{36,37}. The SDD1 promoter-reporter system established that the 2 kb promoter fragment of SDD1 including the GT3 box is sufficient for SDD1 expression and GTL1 transrepression.

We next determined the involvement of hyperosmotic stress-induced Ca\textsuperscript{2+} and CaM signaling in the transcriptional regulation of SDD1. SDD1 expression in wild type protoplasts was induced by hyperosmotic stress (200 mM mannitol addition to the protoplast incubation solution), which was partially inhibited by pretreatment with either the Ca\textsuperscript{2+} channel blocker gadolinium ion (Gd\textsuperscript{3+}) or the CaM antagonist W7 and abrogated when incubated with both (Fig. 5c). This result indicates that hyperosmotic stress-induced SDD1 expression is Ca\textsuperscript{2+}- and CaM-dependent. Constitutive SDD1 expression was evident in gtl1-4 protoplasts without hyperosmotic treatment (Fig. 5c), confirming that GTL1 transrepresses SDD1 expression\textsuperscript{37}. Hyperosmotic stress resulted in a slight increase in SDD1 expression in gtl1-4. W7, Gd\textsuperscript{3+}, or Gd\textsuperscript{3+} + W7 pretreatment only marginally reduced SDD1 expression in gtl1-4 plants compared to the response in Col-0 (Fig. 5c). This could be due to involvement of a GTL1-independent pathway or other members of the GT2 family in the regulation of SDD1 expression. A defect of Ca\textsuperscript{2+}-dependent hyperosmotic SDD1 expression in gtl1-4 was restored in GT11-GFP expressing lines (gtl1-4; GT11 promotergtl1-4GFP) (Supplementary Fig. S5). These results indicate that GTL1 is required for Ca\textsuperscript{2+}- and CaM-dependent hyperosmotic SDD1 expression. Together, these results indicate that hyperosmotic stress induces SDD1 expression through Ca\textsuperscript{2+}/CaM signaling and by attenuating GTL1 transrepressor activity.

To determine the in vivo direct outcome of the Ca\textsuperscript{2+}/CaM-binding function on GTL1 DNA-binding activity, SDD1 promoter-LUC reporter activity in response to hyperosmotic stress was compared by expressing wild-type GTL1 or GTL1[H94E]. Hyperosmolality increased SDD1 expression in wild-type and GTL1-expressing gtl1-4 protoplasts but not in GT11[H94E]-expressing gtl1-4 protoplasts (Fig. 5d). Moreover, GT11[H94E] expression suppressed constitutive SDD1 expression in gtl1-4 protoplasts (Fig. 5d). Together, these results indicate that SDD1 expression is activated or derepressed in response to hyperosmotic stress via allosteric inhibition of GTL1 docking to the SDD1 promoter by Ca\textsuperscript{2+}/CaM.

**Water-deficit stress represses stomatal development in a Ca\textsuperscript{2+}/CaM- and GTL1-dependent manner.** Hyperosmotic stress inhibits stomatal development in Arabidopsis\textsuperscript{39}. Overexpression of SDD1 represses stomatal development in species including Arabidopsis, tomato, and maize\textsuperscript{40–42}. We have shown that hyperosmotic stress results in up-regulated SDD1 expression in a Ca\textsuperscript{2+}/CaM- and GTL1-dependent manner (Fig. 5). This suggests that water-deficit stress may repress stomatal development through Ca\textsuperscript{2+}/CaM-dependent inhibition of the GTL1 transrepressor in plants. To test whether osmotic stress-induced repression of stomatal development is dependent on GTL1, we quantified changes in stomatal development in wild type and gtl1-4 plants in response to water-deficit stress (Supplementary Fig. S6). The presence of a significant water stress was confirmed by reductions in leaf area (Fig. 6a) and stomatal length (Fig. 6b), both of which are well-established indicators of water stress\textsuperscript{39}.

Stomatal index was reduced in gtl1-4 plants compared to wild type plants (Fig. 6c) as previously reported\textsuperscript{37}. We also observed a reduction in stomatal development in severely water-stressed Col-0 plants, evidenced by a decrease of stomatal index in both adaxial and abaxial leaves (Fig. 6c). Mild water-deficit stress (50% MWC) resulted in reduced leaf area, but no change in stomatal index, suggesting that inhibition of stomatal development depends on the severity of the stress. In addition, the number of stomatal precursor cells produced by asymmetric divisions (meristemoids and guard mother cells) was increased in response to severe water-deficit stress (Fig. 6d). In contrast to the water-deficit-induced reduction in stomatal index in Col-0 plants, stomatal development in gtl1-4 was insensitive to severe water-deficit stress (Fig. 6c), indicating that GTL1 is required for water-deficit stress-induced repression of stomatal development.

To further test the hypothesis that water-deficit stress-repressed stomatal development is dependent on Ca\textsuperscript{2+}/CaM signaling, plants were grown on media containing the CaM antagonist GdCl\textsubscript{3} with or without mannitol to induce osmotic stress. A lower concentration of W7 (50 and 100 \textmu M), relative to protoplast assays, was used to determine the effects on stomatal development in seedlings without major effects on growth. W7, whether alone or in combination with GdCl\textsubscript{3}, did not significantly affect stomatal index of Col-0 under water-sufficient conditions. However, W7 completely rescued the water-deficit-induced repression of stomatal development regardless of GdCl\textsubscript{3} treatment (Fig. 6e). There was no effect of W7 or mannitol on stomatal index of gtl1-4 plants (Fig. 6e). Leaf area was lower in both Col-0 and gtl1-4 when grown with mannitol, and this phenotype was not rescued by W7 (Supplementary Fig. S7). Together, these results indicate that water-deficit stress results in decreased stomatal development through Ca\textsuperscript{2+}/CaM signaling, and that GTL1 is required for transducing the osmotic stress signal to the repression of stomatal development.
Discussion

Hyperosmotic stress-induced Ca\(^{2+}\)/CaM signaling requires a rapid and efficient signal transduction mechanism to modulate global transcriptional regulation to cope with water deficit during drought stress. The widely accepted model of this signal transduction mechanism is that Ca\(^{2+}\)/CaM directly interacts with transcription factors (TFs). This interaction modulates the activity of TFs, leading to changes in gene expression and thereby affects the plant's response to drought. The model suggests that Ca\(^{2+}\)/CaM-dependent TF activation is crucial for plants to adapt to water stress, ensuring survival under drought conditions.
factors to modulate their activity and regulate transcription. However, molecular and structural mechanisms of this modulation specifically on a transcriptional repressor are largely unknown. In this set of experiments, we have demonstrated a mechanism of Ca\(^{2+}\)/CaM-dependent allosteric control of the GTL1 transcriptional repressor that directly controls its DNA-binding activity in response to hyperosmotic stress (Fig. 7). Under water-sufficient conditions, GTL1 binds to the SDD1 promoter and represses SDD1 expression, resulting in maintained stomatal development. Water-deficit induces a transient increase of intracellular Ca\(^{2+}\) level\(^{36,38,43,44}\), which is transduced by CaM that directly binds to GTL1, destabilizes the trihelical structure of the DNA-binding domain, and facilitates

Figure 6. Stomatal development is repressed under severe water-deficit conditions through Ca\(^{2+}\)/CaM and the GTL1 transrepressor. (a) Leaf area of Col-0 and gtl1-4 plants grown under well-watered (WW), mild water-deficit (50% media water content, MWC), and severe water-deficit (30% MWC) conditions. These leaves were used to quantify stomatal length (b), stomatal index (c), and stomatal precursor index (d) in adaxial and abaxial leaf surfaces of Col-0 and gtl1-4 plants. Adaxial (upper case) and abaxial (lower case) leaves were analyzed separately for the statistical comparisons. (e) Stomatal index of the abaxial surface was quantified in Col-0 and gtl1-4 plants grown under 0 and 200 mM mannitol conditions with or without W7 (50 and 100 \(\mu\)M) and GdCl\(_3\) (85 and 170 \(\mu\)M). Col-0 (left panel) and gtl1-4 (right panel) were analyzed separately for the statistical comparisons. Data shown are the means with SD for 8–12 replicates (a–d) and 7 replicates (e). Columns with the same letters above are not significantly different from each other based on Tukey's Honestly Significant Difference (HSD) test (\(P < 0.05\)) (Two-way ANOVA).
dissociation of GTL1 from the SDD1 promoter. This novel signal transduction mechanism leads to transcriptional derepression (activation) of SDD1, a negative regulator of stomatal development, and thus inhibits stomatal development to reduce transpirational water loss under water-deficit conditions as an acclimation response (Fig. 7).

A prototypical CaM-binding domain exists in the basic amphipathic region of GTL1Nα2 (Fig. 1b and Supplementary Fig. S8). It is well known that Ca2+/CaM binds to target peptides via hydrophobic and electrostatic interactions9,34,45. Ca2+ binding to CaM causes a conformational change from a dumbbell shape to a globular shape, which exposes hydrophobic pockets and negatively charged conserved glutamate residues that can interact with protein target regions such as the basic amphipathic α2 helix of GTL1N (Supplementary Fig. S8). The prediction of the GTL1Nα2-Ca2+/CaM complex suggests that α2 basic residues interact with conserved acidic glutamate residues of CaM through electrostatic interactions (Supplementary Fig. S8)9,11,45. The α2 hydrophobic residues interact with the hydrophobic pocket of Ca2+/CaM. Since α2 hydrophobic residues are also important for the formation of the hydrophobic core and GTL1N DNA-binding activity (Fig. 3), Ca2+/CaM binding to the α2 may interfere with the formation of the hydrophobic core and prevent binding to the SDD1 promoter. Our results demonstrate that Ca2+/CaM binds to the α2 inner-core hydrophobic residues, which destabilizes GTL1N tertiary structure by conformational change, preventing SDD1 promoter docking, which results in transcriptional derepression of SDD1 expression.

This allostery is in contrast to the Ca2+/CaM steric hindrance mechanism of E proteins in which Ca2+/CaM blocks promoter binding of E proteins but does not dissociate the E protein-promoter complex in animals12. Since GTL1 is associated with the SDD1 promoter, resulting in repression under well-watered conditions, this allosteric mechanism allows for dissociation of GTL1 from the promoter and the prevention of free GTL1 binding to the promoter, thereby maintaining SDD1 activation through both mechanisms. Allostery can involve large or subtle conformational change or can function without conformational change by redistributing electrostatic interactions necessary for binding interface46. Whether Ca2+/CaM-mediated allosteric regulation of GTL1 DNA-binding activity requires conformational change of trihelical tertiary structure is unknown without analyzing the solution structure of the Ca2+/CaM-GTL1 complex. It is, however, well-accepted that allostery is one of the fundamental molecular mechanisms in cellular signaling pathways of all living organisms47. We posit that the allosteric mechanism may fine-tune transcriptional regulation that is necessary for precise gene expression required when an organism must decode different signatures encoded by environmental changes necessary for adaptive fitness48.

Water deficit in soil results in decreased cell turgor pressure and imposes osmotic stress on plant cells, which activates various acclimation responses, including physiological and developmental changes49. Mannitol-induced hyperosmotic stress has been reported to induce cytosolic and nuclear Ca2+ transients in tobacco and Arabidopsis36,38,41,44. Hyperosmotic stress also results in decreased stomatal development in Arabidopsis36,38,41,44. Consistent with these reports, our study also showed that water-deficit stress repressed stomatal development as evidenced by a reduction of stomatal index (Fig. 7). However, in gtl1-4 plants, stomatal index was not further decreased by mannitol, indicating that GTL1 is required for transducing the signal from the osmotic stress to the modulation of stomatal development. We showed that the Ca2+/CaM-GTL1-SDD1 modular relay system allows for transduction of water-deficit (e.g. turgor reduction or low water potential) sensing. When water is sufficient, GTL1 transrepression of SDD1 (a negative regulator of guard cell lineage) is necessary to facilitate stomatal

Figure 7. A Ca2+/CaM-regulated GTL1 transrepressor is a transcriptional switch to control stomatal development, transpiration, and water-use efficiency in plants. Proposed model for Ca2+/CaM-regulated transcriptional switch to repress stomatal development and to improve water-use efficiency and drought tolerance through GTL1.
The Ca^{2+}/CaM-GTL1-SDD1 module appears to be a transcriptional switch mechanism through which plants can acclimate to reduced water availability via a repression of stomatal development (Fig. 7). Since YODA (a MAPKKK) is known to act downstream of SDD1, we posit that the Ca^{2+}/CaM-regulated transcriptional switch through the GTL1-SDD1 module is the most upstream signal transduction pathway responsible for this response identified to date. Whether the function of SDD1 is to send a signal in response to water deficit stress remains to be tested.

A Ca^{2+}-dependent transcriptional derepression mechanism can be considered to be an efficient signal transduction mechanism for rapid activation of acclimation genes^{17-19}. Recently, a similar signal transduction mechanism through the SR1/CAMTA3 transrepressor has been reported in Arabidopsis. SR1/CAMTA3 is a Ca^{2+}/CaM-binding transcriptional repressor that suppresses the expression of salicylic acid (SA)-related genes involved in plant immunity^{24}. Ca^{2+}/CaM binding to AtSR1/CAMTA3 is required for repression of the immune response. Low temperature induces expression of SA-related genes through de-repression of SR1/CAMTA3 activity by Ca^{2+}/CaM^{17,20}. However, the CaM-binding domain is located in the C-terminus, while the DNA-binding domain is located in the N-terminus. It is not known how Ca^{2+}/CaM activates CAMTA transcriptional activity or DNA-binding activity. This report together with our results suggest that Ca^{2+}/CaM-regulated transcriptional derepression is a common signal transduction mechanism in response to various environmental stresses.

The capacity for plants to sense water-deficit through the Ca^{2+}/CaM-GTL1-SDD1 module and to transduce this signal to a developmental program may be an important mechanism to modulate plant water use. The presence of GTL1 orthologs in a large number of diverse species suggests that this acclimation mechanism is conserved among higher plants^{22-24}. Supporting this notion, TaGT2L1D, a GTL1 ortholog in wheat, is also a transrepressor of SDD1 to negatively regulate stomatal development^{25}. We reported that the poplar GTL1 ortholog interacts with Ca^{2+}/CaM through the C-terminal DNA-binding domain that binds to the GT2 box of the poplar SDD1 promoter^{26}. It is interesting that the Ca^{2+}/CaM-binding domain of AHT2L2 is located in the α2 helix of the C-terminal trihelical domain, whereas in PtaGTL1 it is located in the α1 of the C-terminal trihelical domain. Some of the GT2 family do not have Ca^{2+}/CaM-binding domains within their trihelical domain due to a change of residue to an acidic residue (Fig. 2b). This suggests that the Ca^{2+}/CaM-binding property might have evolved in specific members of the GT2 family, such as GTL1. The Ca^{2+}/CaM-binding domains within their trihelical domain due to a change of residue to an acidic residue (Fig. 2b). This suggests that the Ca^{2+}/CaM-binding property might have evolved in specific members of the GT2 family, such as GTL1. The Ca^{2+}/CaM-binding domains within their trihelical domain due to a change of residue to an acidic residue (Fig. 2b). This suggests that the Ca^{2+}/CaM-binding property might have evolved in specific members of the GT2 family, such as GTL1. The Ca^{2+}/CaM-binding domains within their trihelical domain due to a change of residue to an acidic residue (Fig. 2b). 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Arabidopsis protoplast transient expression and luciferase reporter assay. Protoplasts were isolated from 4-week-old plants as described previously with modifications. Shoots from 150 seedlings were incubated in 30 mL of 1 M mannitol solution for 30 min and then incubated in 20 mL of enzyme solution (1% cellulose R-10, 0.25% macerozyme R-10, 500 mM mannitol, 1 mM CaCl₂, 10 mM MES, 20 mg BSA) for 12 h to digest cell walls. The protoplast mixture was filtered through 100 mm mesh and transferred onto 20 mL of 21% sucrose solution, which was centrifuged at 57  g for 4 min. Supernatant was removed, replaced by 20 mL of W5 solution, and incubated on ice for 6 h to stabilize protoplast.

SDD1promoter:LUC (29 mg), 35S promoter:LUC (1 mg), HA-GLT1L (5 mg), or HA-GLT1L[H94E] (5 mg) was transformed into protoplasts using the polyethylene glycol (PEG)-mediated transformation method. First, W5 solution was replaced by mannitol solution (400 mM mannitol, 30 mM MgCl₂, and 5 mM MES, pH 5.6). Plasmids (30 mL) were added to 300 mL of protoplast MaMg solution. Then, 300 mL of PEG solution (400 mM mannitol, 100 mM Ca(NO₃)₂, 40% PEG6000) was added, mixed by inverting for 1.5 min, and incubated at room temperature for 30 min. PEG protoplast solution was slowly diluted with 5 mL of W5 solution by adding 1 mL of solution every 2 min. The protoplast solution was centrifuged at 43  g for 4 min. Supernatant was removed, replaced by 2 mL of W5 solution, and incubated at room temperature for 12 h. To induce hypertonic stress, 200 mM mannitol was added and protoplasts were incubated for 1 h. To inhibit Ca²⁺ or CaM signaling, protoplasts were pre-incubated in 1 mM GdCl₃, 0.5 mM LaCl₃, or 600 mM W7 (N-(6-aminohexyl)-5-chloro-1-naphthylsulfonamide-hydrochloride) for 10 min prior to mannitol treatment. After incubation with mannitol, protoplasts were harvested by centrifugation at 57  g for 2 min, then frozen in liquid nitrogen. Frozen protoplasts were lysed with passive lysis buffer (Promega) and the SDD1-LUC and LUC activities were measured using a dual-luciferase assay kit according to the manufacturer’s instructions (Promega) using a luminometer (TD20/20). Relative SDD1-LUC activity was shown as 100 × (SDD1-LUC activity/Rrluc activity). Osmotic potential (ψₒ) of the osmotic stabilization solution (W5; (ψₒ = −1.19 MPa) and hypertonic solution (W5 with 200 mM mannitol; ψₒ = −1.73 MPa) was calculated by the equation (ψₒ = −CRT; C = osmolality, R = gas constant, T = Kelvin temperature). Osmolarity of the solution was measured using a vapor pressure osmometer (Wescor 5200).

Purification of recombinant proteins. MBP and MBP-fusion proteins (GT1N, GT1C, GT1Δdel, GT1N[L91R], GT1N[W92R], GT1N[H94E], and GT1N[V95D]) were purified by amylose resin (NEB) as described previously.

CaM-binding assays. CaM-binding assays were performed using bacteria lysates expressing MBP-fusion proteins and in vitro-translated HA-AtCaM2 as described previously with the following modifications. Briefly, MBP fusion proteins were expressed in Escherichia coli strain Rosetta (DE3) carrying pMAL-c2 vectors. After harvesting by centrifugation, the cells were lysed by French press in lysis buffer containing 50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 1% DMSO, 2 mM DTT, and protease inhibitor cocktail (Sigma-Aldrich). The cell extract was prepared by centrifugation at 10,000  g for 20 min at 4 °C and cleared by filtration. The clear lysates were incubated with amylose resin at 4 °C for 2 h. The beads with immobilized MBP fusion proteins were washed with wash buffer containing 50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 1% DMSO, 2 mM DTT, and 0.1% Nonidet P-40. AtCaM2 with an N-terminal HA tag was synthesized using plasmid pCMX-PL2 and the TNT T7 Coupled Reticulocyte Lysate System (Promega) according to the manufacturer’s protocol. The in vitro-translated HA-AtCaM2 was diluted with CaM-binding buffer (50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 1% DMSO, 2 mM DTT, and 2 mM CaCl₂) and incubated with the affinity-purified MBP fusion proteins immobilized on the beads at 4 °C for 2 h. The beads were then washed four times with wash buffer. Bound HA-AtCaM2 was eluted by boiling in 1x SDS sample buffer and subjected to 12% SDS-PAGE. Input and immunoprecipitated HA-AtCaM2 proteins were detected by immunoblots using rabbit polyclonal anti-HA antibodies (Abcam) and the Enhanced Chemiluminescence Plus protein gel blotting detection system (GE Healthcare). The amount of MBP fusion proteins used in each CaM-binding assay was separated by another 8% SDS-PAGE gel and visualized by staining with Coomassie Brilliant Blue.

Electrophoretic mobility shift assay (EMSA). Purified recombinant proteins were used to determine DNA-binding activity based on interaction with biotin-labeled Arabidopsis SDD1 promoter fragments that contain the GT3 box. EMSA procedures were as described previously, with the following modifications to determine the effect of Ca²⁺/CaM on DNA-binding activity of recombinant proteins. To determine the effect of CaM on the association between GT1N and the SDD1 promoter, human CaM protein (50 or 100 ng) was incubated with recombinant proteins with 2 mM CaCl₂ or 10 mM EGTA for 1 h. Then, biotin-labeled SDD1 promoter was added and incubated for 20 min. To determine the CaM effect on the dissociation of GT1N from the SDD1 promoter, biotin-labeled SDD1 promoter was incubated with GT1N or GT1N[H94E] for 20 min. Then, CaM protein (100 ng) was added to the GT1N-SDD1 promoter complex mixture with 2 mM CaCl₂ and incubated for 1 h. The mixture was separated in a 6% PAGE gel and transferred to a Biodyne B Nylon membrane (Thermo Fisher Scientific).
Chromatin immunoprecipitation. ChIP assays were performed using chromatin isolated from protoplasts of 4-week-old gtl1-4 plants using an EZ ChIP chromatin immunoprecipitation kit (Millipore) as described previously27 with the following modifications. Protoplasts were transformed with HA-GTL1 or HA-GTL1[H94E] and incubated for 12 h at room temperature, 200 mM mannitol was added to one group (each group includes 3 replicates) to induce hyperosmotic stress for 1 h. Then, formaldehyde was added for crosslinking of chromatin and 0.1 M glycine to quench crosslinking. After centrifuging briefly, protoplasts were resuspended in lysis buffer. Chromatin was sheared to 200–1000 bp fragments by sonication. Isolated and pre-cleared chromatin was incubated with mouse monoclonal anti-HA antibody or mouse IgG for 4 h at 4 °C to immunoprecipitate the chromatin associated with HA-GTL1 or HA-GTL1[H94E]. Immunoprecipitated chromatin was reverse-crosslinked and purified according to the manufacturer’s protocol. Input chromatin (immunoprecipitated by anti-HA antibody or IgG antibody) was used for PCR analysis by the SDD1 promoter-specific primers (SDD1ChIP-3F and SDD1ChIP-3R)27.

Stomatal development analysis. Following irrigation and drainage of excess water, containers were weighed to obtain media saturated weight (MSW). Plants were irrigated to saturation (well-watered) or water was withheld to 50 or 30% media water content (MWC), calculated as [media fresh weight (MFW) − media dry weight (MDW)]/[media saturated weight (MSW) − MDW] × 100. During the experiment, a MDW of 49 g was used to determine the appropriate amount of water to add to containers, based on prior experiments. After harvesting, containers were dried in a forced-air oven and MDW was obtained for each plant. Final average MWC of the 50 and 30% MWC treatments was final average MWC of the 50 and 30% MWC treatments was between 38% (prior to watering) and 53% (post-watering), and 25% (prior to watering) and 33% (post-watering), respectively (Supplementary Fig. S6). Once the treatment MWC was reached, containers were wrapped with plastic film to prevent water loss by evaporation. Containers were weighed every two days and re-watered to bring the MWC to target levels (Supplementary Fig. S6). This treatment was maintained for 30 days after the emergence of the target leaf, enabling the leaf to complete development under a given water deficit treatment.

Leaf appearance was tracked with photographs to determine leaf number. Once the target leaves had fully expanded, leaf adaxial and abaxial surfaces were pressed onto cyanoacrylate droplets (Henkel Manufacturing) on glass slides. Impressions were viewed under a BH-2 light microscope (Olympus) at 200× magnification, with a field of view of 0.113 mm². Four images were collected from each surface impression, and cell types were counted in ImageJ. Cell types were distinguished by shape; meristemoids were roughly triangular cells, while long oval cells were identified as guard mother cells (GMCs)55. Stomatal index was calculated as the number of stomata/(number of stomata + epidermal cells). A diagonal line was drawn through each image and the 5 stomata closest to this line were used for stomatal length measurements. Length was measured from end to end of the outer cell wall.

To test the necessity of calcium-CaM binding for the inhibition of GTL1-dependent stomatal development under osmotic stress conditions, a pharmacological disruption experiment was performed in vivo. Seedlings were grown for 6 weeks, at which point the first true leaf was collected and fixed in a 9:1 ethanol:acetic acid solution. Following clearing, leaf tissue was stored in 85% ethanol. Tissue was rehydrated by transferring to a 70% ethanol solution, then to 70, 50, and 30% methanol solutions, each for ca. one hour. The tissue was then stained with 0.2% toluidine blue and mounted for DIC microscopy in 50% glycerol. All microscopy was done at a 400× magnification. Leaf tissue used for microscopy was also photographed for measurements of leaf area in ImageJ.

Homology modeling, model validation, and computational docking. Homology modeling of the GTL1 N-terminal trihelix DNA-binding domain (residues 63 to 126) was modeled using the RaptorX web-based server56. Structure refinement of the predicted models was performed using ModRefiner57. The predicted model of GTL1N was validated by PROCHECK58 to check stereochemical quality of a native structure. Ramachandran plot (Supplementary Fig. S9a) and plot statistics (Supplementary Table S2) indicate 95.3% residues in most favored regions, showing a good quality of GTL1N structure. The GTL1N model was further evaluated using the ProSA web server59. ProSA calculates the Z-score (−4.13) for the model to show the overall model quality (Supplementary Fig. S9b) and knowledge-based energy plot of residue scores (Supplementary Fig. S9c).

The structure of the GTL1N-SDD1 promoter complex was determined using the High Ambiguity Driven bio-molecular DOCKing (HADDOCK) web server60. The active residues of GTL1N involved in docking are chosen based on the NMR structure of GT1 and GT-1 box element and are N61, R62, K109, E113, E116, N117, K120, and K126. The residues around active residues were automatically defined as passive residues. 3D structure of the SDD1 promoter fragment (54–gtcctGTTAaAactt-34) including a GT3-box was generated by 3D-DART, a DNA structure-modeling server61. The GT2 box residues in the SDD1 promoter fragment (Gua5, Gua6, Thy7, Ade8, Ade9, Ade10) and their complementary residues were considered active residues. A total of 20,000 structures were generated using rigid body energy minimization. The 200 lowest–energy structures were refined by semiflexible refinement and explicit solvent calculations. The model with the best (lowest) HADDOCK score (−175.6 ± 6.2) and root mean square deviation (2.0 ± 1.2) was selected and used for the analysis. All 3D structures were analyzed and visualized by 3D structure viewer UCSF Chimera62.

Statistical analysis. All column graphs were analyzed by Tukey’s honestly significant difference (HSD) test (one-way ANOVA) using SAS 9.2 software (SAS Institute).
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Author Contributions

C.Y.Y., P.M.H. and M.V.M. conceived and designed the experiments. C.Y.Y., N.M., A.F., H.W. and I.S.D. performed the experiments. C.Y.Y., N.M., A.F., H.W., H.I.F., P.M.H. and M.V.M. analyzed and discussed the data. A.S.N.R. and B.W.P. originally identified GTL1 as a Ca2+-calmodulin binding protein, analyzed and discussed data. C.Y.Y., P.M.H. and M.V.M., wrote the manuscript. All authors read and approve the final manuscript.

Additional Information

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