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Raf-like kinases and receptor-like (pseudo)kinase GHR1 are required for stomatal vapor pressure difference response

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Stomatal pores close rapidly in response to low-air-humidity-induced leaf-to-air vapor pressure difference (VPD) increases, thereby reducing excessive water loss. The hydroactive signal-transduction mechanisms mediating high VPD–induced stomatal closure remain largely unknown. The kinetics of stomatal high-VPD responses were investigated by using time-resolved gas-exchange analyses of higher-order mutants in guard-cell signal-transduction branches. We show that the slow-type anion channel SLAC1 plays a relatively more substantial role than the rapid-type anion channel ALMT12/QUAC1 in stomatal VPD signaling. VPD-induced stomatal closure is not affected in mpk12/mpk4GC double mutants that completely disrupt stomatal CO2 signaling, indicating that VPD signaling is independent of the early CO2 signal-transduction pathway. Calcium imaging shows that osmotic stress causes cytoplasmic Ca2+ transients in guard cells. Nevertheless, osca1-2/1.3/2.2/3.1/3.7 Ca2+-permeable channel quintuple, osca1.3/1.7-channel double, cngc5/6-channel double, cngc20-channel single, cnga19/20crispr-channel double, grf3/2.3-channel double, cpk-kinase quintuple, cb1/4/5/6/7 quintuple, and cb1/3 raf double mutants showed wild-type-like stomatal VPD responses. A B3-family Raf-like mitogen-activated protein (MAP)–kinase kinase kinase, M3K5/6/RAF6, activates the OST1/SnRK2.6 kinase in plant cells. Interestingly, B3 Raf-kinase m3k5/6 and m3k5/1/5/6/7 (raf3/6/5/4) quadruple mutants, but not a 14-generaf-kinase mutant including osmotic stress-linked B4-family Raf-kinases, exhibited slowed high-VPD responses, suggesting that B3-family Raf-kinases play an important role in stomatal VPD signaling. Moreover, high VPD–induced stomatal closure was impaired in receptor-like pseudokinase GUARD CELL HYDROPEROXIDE-RESISTANT1 (GHR1) mutant alleles. Notably, the classical transient “wrong-way” VPD response was absent in ghr1 mutant alleles. These findings reveal genes and signaling mechanisms in the elusive high VPD–induced stomatal closure response pathway.

Significance

With the continuing increase in global temperatures, plants transpire more water due to the increasing vapor pressure deficit. Stomatal pores in plants close rapidly in response to the rising vapor pressure deficit to counteract water loss. We demonstrate that mutations in the stomatal CO2-signal-transduction pathway do not impair the response to an increase in vapor pressure difference (VPD). Osmotic stress causes cytoplasmic Ca2+ transients in guard cells. Nevertheless, we show that diverse investigated higher-order calcium-signaling mutants do not affect the VPD response. We reveal that B3 family Raf-like protein kinases and a plasma membrane receptor-like protein GHR1 function in the elusive leaf-to-air VPD-mediated stomatal closure pathway. Notably, ghr1 mutant alleles disrupt the classical “wrong-way” stomatal VPD response.
Slow (S-type) anion channels (13) and rapid (R-type) anion channels (14) play important roles in mediating and regulating guard-cell ion efflux and stomatal closure in response to upstream signaling events. These anion channels mediate anion efflux and cause plasma-membrane depolarization in guard cells (13, 14). The S-type anion channel SLAC1 and the R-type anion channel ALMT12/QUAC1 function in stomatal closure (15–18). Mutants in the SLAC1 gene show impairment in stomatal closure in response to several environmental stimuli, including abscisic acid (ABA), CO2, ozone, and low relative humidity (17, 18).

The plant hormone ABA plays a central role in regulating stomatal closure (19–21). ABA-mediated stomatal regulation evolved 400 million years ago in moss species (22). ABA triggers stomatal closure (19–21). ABA-mediated stomatal regulation evolved in several ABA biosynthesis mutants and ABA signaling mutants in Arabidopsis. The resulting models have been, to a degree, debated. Early studies on ABA-deficient (abi1-1) and ABA-insensitive (abi1-1 and abi2-1) mutants suggested that ABA and ABA-signaling mutants are necessary for high VPD-induced stomatal closure (33). This is supported by the investigation of transgenic lines expressing a guard-cell-targeted dominant abi1-1 protein phosphatase, which showed a wild-type (WT)-like stomatal response to a rapid increase in VPD, including in time-resolved gas-exchange analyses (34). Overexpression of the Arabidopsis abil-1 mutant allele in gray poplar, Populus x canescens (Aiton) Sm., exhibited a defect in steady-state stomatal conductance in response to a VPD increase (35). In another study, an ABA biosynthesis mutant (aba2-13) and an OST1 protein kinase mutant (ost1-4) were identified by genetic screening for mutants defective in low-humidity-mediated steady-state leaf temperature regulation and thus were impaired in low-humidity-mediated stomatal closure (36). It has been proposed that high VPD/low humidity rapidly triggers ABA biosynthesis to induce stomatal closure (37, 38). However, whole-plant gas-exchange analyses showed that stomatal closure in QUAC1 and high VPD stimulation, in several ABA-deficient mutants defective in key steps of ABA biosynthesis, including the most severe nced3/nced5 double mutant (5, 39, 40), indicating that a direct increase in the ABA concentration of guard cells may not be essential for initiating high VPD-induced stomatal closure. Furthermore, a step toward low humidity (high VPD) caused a similar percentage degree of reduction in the guard-cell K+ content in WT and the ABA biosynthesis abi3-1 mutant, indicating a functional high VPD response in the ABA-deficient mutant (37). Moreover, an ABA receptor, pyl1/pyl2/pyp1/lpy2/lpy1/pyl5/pyl8 hextuple mutant, which disrupts ABA-induced stomatal closure (5, 41), exhibited a significantly reduced initial response rate and increased half-response time of stomatal closure during high VPD stimulation (5). These results indicate that ABA signaling may contribute to the stomatal closing rate of the high VPD response and steady-state stomatal apertures, but rapid enhancement of ABA signaling is not essential for high VPD-induced stomatal closing to be initiated and can proceed, albeit at a slower rate, in the above mutants (5).

It has been suggested that young leaves acquire stomatal ABA sensitivity by pre-exposure to ABA or low relative humidity for 1 d (42). Analyses of ABA concentration increases in response to drought or low humidity have shown that substantial rises in the ABA concentration appear to occur after 3 or more hours of stress treatment (43), whereas high VPD–induced stomatal closing typically occurs within 15 min. Studies have shown that guard cells in nonstressed leaves have measurable and higher ABA concentrations than mesophyll cells (44–47). Recent studies suggest that steady-state basal ABA signaling occurs in guard cells (46–48), and a model has been proposed that may explain many of the above findings, in which basal ABA signaling may contribute to the stomatal VPD response by amplifying the hydroactive stomatal closure pathway (20), similar to CO₂-triggered stomatal closure (46, 47). Furthermore, several studies suggest a key role for ABA in adjusting the steady-state stomatal conductance (34, 35). In the present study, we pursued analyses of other guard-cell signal-transduction pathways and higher-order mutants given the limited knowledge of early VPD sensing and signaling mechanisms.

In contrast to the findings of a functional high VPD response in ABA receptor hextuple mutants, the high VPD response was strongly impaired in the OST1 protein kinase mutant, ost1-3 (5), consistent with earlier research (36). It has been suggested that an ABA-independent pathway is involved in OST1/SnRK2 kinase activation during water deficiency (49, 50). However, apart from the requirement of the OST1/SnRK2.6 protein kinase, the molecular and genetic mechanisms that mediate VPD sensing and signal transduction in the rapid hydroactive response remain elusive.

Given the dearth of mutants and mechanisms known to affect the rapid hydroactive VPD-induced stomatal closing response, in the present study, we screened and investigated potential signaling mechanisms to determine mechanisms that mediate the hydroactive stomatal VPD response. We analyzed mutants in several guard-cell signal-transduction pathways to determine which genes, mechanisms, and pathways contribute to the VPD response and which gene combinations alone are not required for the VPD response. The time-dependent kinetics of stomatal VPD responses were investigated by intact leaf gas-exchange analyses in mutants in guard-cell signaling pathways. Differential contributions of the S-type anion channel SLAC1 and the R-type anion channel ALMT12 were found by analyzing the high VPD response in slac1-3, almt1-2, and slac1-3almt12-1 mutants. The involvement of osmotic/mechanical sensing and calcium signaling in the stomatal high VPD response was investigated in the putative osmotic/mechanical sensor osca1-2/13/2.2/3.3/3.1 quintuple and pathogen-associated molecular patterns-activated calcium channels oscal.1/3.1/7 double mutants, glutamate receptor-like channels gr3.2/3.3 double mutant, Ca++-permeable cyclic nucleotide-gated channels cneg5/6 double mutant, cngc20 and cngc19/20转型升级 double mutants, calcium-dependent protein kinase cpek3/5/6/11/23 and cpek3/4/5/6/11 quintuple mutants, and calcineurin-B like proteins cb1/14/5/8/9 quintuple and cb1/2/3/4/5 double mutants. The stomatal high VPD responses were also analyzed in strongly CO₂-insensitive mitogen-activated protein (MAP) kinase mpk12/mpk4GC double-mutant alleles showing that early CO₂ signaling does not participate in stomatal VPD signal transduction. Interestingly, mutant alleles in the receptor-like (pseudo)kinase GHR1 greatly impaired the stomatal VPD response and impaired the classical “wrong-way” response to stomata to VPD increases. We further investigated whether the recent identified B3-family and B4-family Raf-like MAP Kinase Kinase Kinases (M3Ks) (51), which function in OST1 activation (52–54), are required for stomatal closure in response to VPD elevation and show that B3-family, but not B4-family, Raf-like M3Ks function in the stomatal VPD response, with a central role for M3Kδ5/RAF6.

**Results**

**High VPD–Induced Stomatal Closure Is Slowed in ost1 Kinase Mutant and Guard-Cell Anion-Channel slac1 Single and slac1/ almt12 Double Mutants.** The OST1/SnRK2.6 protein kinase is an important positive transducer of stomatal closing and an activator of anion channels during stomatal closure. It has been shown that...
suggest that high VPD–induced stomatal closure in WT controls (5, 36, 40). Although a transition to elevated VPD (~2.0 kPa), caused a time-dependent reduction in stomatal conductance in both osl1-3 mutant and WT leaves, the final stomatal conductances were larger in osl1-3 mutant leaves, and the time course of the response was consistently slower in independent datasets (SI Appendix, Fig. S1A). The initial high-VPD stomatal closing responses were fitted with exponential one-phase decay functions. The average rate constant in osl1-3 mutant leaves was significantly smaller than in WT controls (SI Appendix, Fig. S1B; osl1-3: 0.10 ± 0.01 min⁻¹ vs. WT: 0.28 ± 0.06 min⁻¹, P < 0.03), and the average half-response times were significantly larger in osl1-3 mutant leaves than WT (SI Appendix, Fig. S1C; osl1-3: 8.9 ± 0.6 min vs. WT: 5.1 ± 0.4 min, P < 0.002). These results suggest that high VPD–induced stomatal closure in osl1-3 was slower than in WT, consistent with previous research (5, 36, 40). Similar results were also found in an independent set of experiments (SI Appendix, Fig. S1 D–F). These data show that altered stomatal response kinetics to VPD can be analyzed with our system by direct comparisons to WT plants grown in parallel and analyzed within the same datasets (Materials and Methods).

Activation of anion channels in guard cells plays an important role in stomatal closure. It has been shown that the slow-type anion channel SLAC1 is required for the high-VPD response (18). As a further test of our system, we analyzed the high-VPD response in slac1-3 mutant leaves. Similar to the findings of Vahisalu et al. (18), slac1-3 mutant leaves showed a slower response to high VPD (Fig. 1A and SI Appendix, Figs. S2 and 3). However, unlike osl1-3 mutant leaves, the steady-state stomatal conductances at low-VPD and high-VPD conditions in slac1-3 were comparable with WT under the imposed conditions (Fig. 1A). In another independent set of experiments, slac1-3 mutant leaves exhibited slightly increased steady-state stomatal conductances compared to parallel-grown WT controls (SI Appendix, Fig. S3A).

It has been shown that the rapid-activating R-type anion channel, ALMT12/QUAC1, mediates anion efflux in guard cells during ABA-triggered, darkness-triggered, and high-CO₂-triggered stomatal closure (15, 16, 55). To investigate whether ALMT12/QUAC1 is required for the high-VPD response, almt12-1 mutant leaves were analyzed. Unexpectedly, no obvious difference in the high-VPD response was found between almt12-1 mutant and WT leaves (Fig. 1B). We further analyzed the high-VPD response in slac1-3/almt12-1 double mutants. Interestingly, slac1-3/almt12-1 double-mutant leaves showed a higher steady-state stomatal conductance before and after high-VPD shifts compared to WT leaves and exhibited a slower stomatal closure in response to high-VPD stimulation, similar to slac1-3 mutant leaves (Fig. 1C and SI Appendix, Fig. S3). Average rate constants in slac1-3 and slac1-3/almt12-1 mutant leaves were significantly smaller than in WT controls (Fig. 1D; slac1-3: 0.13 ± 0.04 min⁻¹ vs. WT: 0.31 ± 0.04 min⁻¹, P < 0.05 and slac1-3/almt12-1: 0.10 ± 0.00 min⁻¹ vs. WT: 0.31 ± 0.04 min⁻¹, P < 0.009 and SI Appendix, Fig. S3 C and D; slac1-3: 0.11 ± 0.01 min⁻¹ vs. WT: 0.27 ± 0.03 min⁻¹, P < 0.001 and slac1-3/almt12-1: 0.10 ± 0.00 min⁻¹ vs. WT: 0.33 ± 0.03 min⁻¹, P < 0.001), and average half-response times were larger in slac1-3 and slac1-3/almt12-1 mutant leaves than in WT controls (Fig. 1E; slac1-3: 7.4 ± 1.0 min vs. WT: 4.6 ± 0.2 min, P < 0.002 and slac1-3/almt12-1: 9.0 ± 0.3 min vs. WT: 4.6 ± 0.2 min, P < 0.001 and SI Appendix, Fig. S3 E and F; slac1-3: 8.4 ± 1.0 min vs. WT: 4.7 ± 0.2 min, P < 0.006 and slac1-3/almt12-1: 8.7 ± 0.3 min vs. WT: 4.8 ± 0.2 min, P < 0.001). These experiments provide further evidence that SLAC1 and ALMT12/QUAC1 synergistically function in controlling the steady-state stomatal conductance. Furthermore, the finding that stomatal closing occurs in slac1-3/almt12-1 mutant leaves suggests that additional anion-channel genes could function in the VPD response.

Investigation of Potential Roles of Calcium-Linked Signaling Mechanisms in High-VPD-Triggered Stomatal Closure. It remains unclear how guard cells perceive and respond to increases in VPD to trigger stomatal closure. High VPD initially increases transpiration rates and leads to a rapid reduction of apoplastic water content in leaves (12). The alteration of water content and/or the initial transient increase in stomatal conductance, termed the wrong-way VPD response (56, 57) (e.g., Fig. 1), may trigger...

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osmotic/mechanical-induced cytosolic calcium concentration changes in guard cells to trigger stomatal closure. We tested the effect of 500 mM sorbitol (as osmoticum) on the cytosolic Ca\(^{2+}\) concentration on preopened stomata in leaf epidermal strips of 4- to 5-week-old Arabidopsis plants expressing the Cameleon YC3.60 sensor placed under the control of the pGC1 promoter (58). Both the addition of sorbitol (hyperosmotic stress) and removal of sorbitol (hypo-osmotic stress) caused clear Förster resonance energy transfer (FRET) of the Cameleon sensor, with a decrease of cyan fluorescent protein (CFP) fluorescence and an increase in cpVenus fluorescence (Fig. 2A). When representing the data as normalized ratios (ΔRR/RRo), which depict cytosolic Ca\(^{2+}\) concentration changes, rapid and transient cytosolic [Ca\(^{2+}\)] increases were observed after both the hyperosmotic and hypo-osmotic stimuli (Fig. 2B). Of note, the removal of sorbitol induced a higher maximum Ca\(^{2+}\) transient compared to the one observed after its addition (Fig. 2C).

Previous studies have indicated that OSCA1 and other OSCA family proteins are cation channels mediating calcium influx (59–64). Several studies have suggested that OSCA channels may be activated in response to osmotic or mechanical stimulation (59–63). We hypothesized that OSCA family proteins may be involved in transducing the high-VPD signal in guard cells. According to transcriptome datasets in the Arabidopsis Electronic Fluorescent Pictograph (eFP) browser, the transcripts of OSCA1, OSCA1.3, OSCA2.2, OSCA2.3, and OSCA3.1 are generally more highly expressed in guard cells (SI Appendix, Fig. S44) (58, 65). We therefore generated an osca-21.3.2.2.2/3.1 quintuple transfer DNA (T-DNA) insertion mutant, which included one reported intron allele in the OSCA1 gene (59) and four exon alleles in the OSCA1.3, 2.2, 2.3, and 3.1 genes (SI Appendix, Fig. S5 A and B).

To further investigate the potential roles of calcium-permeable channels in high-VPD-triggered stomatal closure, we focused on putative calcium channels with higher expression

![Fig. 2.](https://doi.org/10.1073/pnas.2107280118)

Both hyperosmotic and hypo-osmotic stress induce rapid cytoplasmic Ca\(^{2+}\) transients in guard cells, and no obvious stomatal VPD phenotype was found in the higher-order mutants of OSCA, GLR3, and CNGC Ca\(^{2+}\)-permeable channels. (A) Representative single cpVenus and CFP emissions (540 and 480 nm) measured from a single guard cell of Col-0 (WT) pGC1–YC2.60 plants initially bathed in a standard solution (5 mM KCl, 10 mM MES, and 50 μM Ca\(^{2+}\), pH 6.15) and then treated transiently with 500 mM sorbitol added to the same solution. Sorbitol was added and removed at the times indicated by arrows. (B) Representative normalized cpVenus/CFP ratio reported as ΔRR/RRo calculated with the single emissions shown in A. (C) Scatter dot plot of maximal cytosolic Ca\(^{2+}\) increases in response to sorbitol addition (I peak) and removal (II peak) of n = 34 guard cells from six independent experiments. In blue and green are the mean and SD for the I and II peaks, respectively. AUF, arbitrary unit of fluorescence. **p < 0.0001.** (D–I) Time-resolved stomatal conductance (Gs) responses to an increase in VPD in Col-0 (WT), osca-21.3.2.2/3.3, osca-3.1.7, glr3.2/3.3, cngc5.1/6.1, cngc20, and cngc19/20crispr mutant leaves under high light (450 μmol m\(^{-2}\)s\(^{-1}\) red light combined with 50 μmol m\(^{-2}\)s\(^{-1}\) blue light). VPDs are shown on top of the time axis in each panel. Data represent mean ± SEM. WT: n = 5 leaves in D–F, n = 4 leaves in G and I, and n = 3 leaves in H; osca-21.3.2.2/3.3: n = 5 leaves; osca-1.3/1.7: n = 5 leaves; glr3.2/3.3: n = 5 leaves; cngc5.1/6.1: n = 5 leaves; cngc20: n = 3 leaves; and cngc19/20crispr: n = 4 leaves.
levels in guard-cell protoplasts, including glutamate-like receptors genes GLR3.2 and GLR3.3 and cyclic nucleotide-gated channel genes CNGC5, CNGC6, and CNGC20 based on transcriptome analyses (SI Appendix, Fig. S4 B–D). Stomatal conductance responses to an increase in VPD were analyzed in gtr3.2/3.3, cngc5-1/6-1, and cngc20 T-DNA insertion mutants and a deletion mutant of two homologous tandem repeat genes CNGC19 and CNGC20, cngc19/20crispr (Fig. 2 F–I and SI Appendix, Fig. S5 C–F). Notably, no obvious differences in the VPD responses were observed compared to parallel-grown WT leaves in all of these mutants.

The high-VPD response was also investigated in cpk3/5/6/11/23 and cpk3/4/5/6/11 quintuple mutants that include mutant allelic in several genes previously found to contribute to stomatal Ca\(^{2+}\) responses and/or anion-channel regulation (67–71). Furthermore, we investigated VPD responses in plasma-membrane-targeted Ca\(^{2+}\) sensor cbl1/4/5/8/9 quintuple mutant leaves and in vacuolar Ca\(^{2+}\) sensor cbl2/3/5 (reduced function) double-mutant leaves (71–74). No major differences in stomatal VPD change responses were found in these higher-order cpk quintuple and cbl mutants compared to WT leaves (SI Appendix, Fig. S6).

**High VPD and High CO\(_2\) Use Different Pathways to Trigger Stomatal Closure.** Defects in several ABA biosynthesis and ABA signaling mutants in *Arabidopsis* increase steady-state stomatal conductances and thus shift VPD responses to higher stomatal conductance values (5, 37). Recent time-resolved stomatal conductance analyses have led to the model that the rapid stomatal conductance response to VPD elevation occurs in parallel to the ABA signaling pathway (5, 34). A signaling pathway parallel to ABA signaling is the CO\(_2\) signaling pathway (46). MAP kinases MPK4 and MPK12 are essential for high-CO\(_2\)-induced stomatal closure, but not for ABA-induced stomatal closure (75, 76). To investigate whether VPD and CO\(_2\) share a common signaling pathway upstream of anion-channel activation, the stomatal response to VPD elevation was analyzed in two independent *mpk12*4GC double-mutant plant lines, in which guard cell MPK4 was silenced by an artificial microRNA driven by the MPK12 promoter in the *mpk12-4* mutant background (75, 77). These *mpk12*4GC double-mutant alleles are the strongest presently known specific high-CO\(_2\)-response impaired mutants in *Arabidopsis* (75). These analyses showed that the *mpk12*4GC double-mutant leaves exhibited a much higher steady-state stomatal conductance than WT leaves (Fig. 3 A and B), consistent with previous findings (75). Previous research has shown a slightly lower stomatal index and no clear effect on stomatal density in *mpk12*4GC double mutants (75). Leaves of *mpk12*4GC double mutants responded rapidly to VPD elevations with response times similar to WT (Fig. 3 A and B). Relative changes in stomatal conductance at early time points in response to VPD increase (10 min) were even larger in these mutant leaves than in WT controls (Fig. 3 C and D). No difference in average rate constants was found between *mpk12*4GC double mutants and WT (Fig. 3E; *mpk12*4GC-1: 0.20 ± 0.03 min\(^{-1}\) vs. WT: 0.28 ± 0.05 min\(^{-1}\), P = 0.42; and *mpk12*4GC-2: 0.23 ± 0.02 min\(^{-1}\) vs. WT: 0.28 ± 0.05 min\(^{-1}\), P = 0.61).
B3-Family Raf-Like Kinases Are Involved in High VPD-Induced Stomatal Closure. Recent studies identified B3-family Raf-like M3Ks as an important mechanism for the activation of SnRK2.2/SnRK2.3/OST1 (SnRK2.6) protein kinases by ABA and osmotic stress (53, 54). Transcriptome datasets in the Arabidopsis eFP browser suggest that B3-family Raf-like M3Ks (M3Kδ1/RAF3, M3Kδ3/EDR1/RAF2, M3Kδ5/RAF6, M3Kδ6/SIS8/RAF5, M3Kδ7/RAF4, and CTR1/RAF1) are expressed in guard cells (SI Appendix, Fig. S4D). Our previous results showed that m3kδ1/δ7 double, m3kδ1/δ6-1/δ7 triple, and m3kδ1/crispr/δ6-2/δ7 crispr triple mutants exhibited partial degrees of ABA insensitivity in non-guard-cell tissues (54). Here, we examined whether M3Kδ1, M3Kδ6, and M3Kδ7 alone are required for the stomatal high-VPD response by analyzing these mutants. No obvious difference in stomatal VPD response was found between m3kδ1/δ7 double, m3kδ1/δ6-1/δ7 triple, and m3kδ1/crispr/δ6-2/δ7 crispr triple mutants compared to WT controls (Fig. 4 A–C). In addition to B3-family, B2- and B4-family Raf-like M3Ks are also involved in SnRK2 protein kinase activation in response to osmotic stress in plants (53, 78). We further investigated the stomatal high-VPD response in a B2-, B3-, and B4-family Raf-like kinase quatrourdecouple mutant, OK-quatdec, (raf16rafo/RAF24140/raf18raf20/raf25/RAF4/raf5-1/m3kδ1/raf3/m3kδ7raf4/m3kδ6raf5raf7raf9/raf10), in which the mutations in 14 Raf-like M3K kinase genes were confirmed, as described in Materials and Methods. However, no obvious difference in the stomatal VPD response was found between OK-quatdec mutant plants compared to WT controls (Fig. 4D).

The above mutants did not include mutations in the M3Kδ5 gene, which was the most highly expressed B3-family Raf-like M3K in guard cells (SI Appendix, Fig. S4D). We pursued in-gel kinase assays that showed that kinase domains of the B3-family member M3Kδ5 and M3Kδ6 kinase, but not the B2-family M3Kδ4 kinase, activated OST1/SnRK2.6 kinase in Arabidopsis mesophyll protoplasts (Fig. S4A). Stomatal VPD response assays showed that m3kδ5-1 and m3kδ5-2 single-mutant leaves exhibited increased steady-state stomatal conductances compared to WT controls (Fig. 5 B and C). Average rate constants in m3kδ5-1 and m3kδ5-2 single-mutant leaves were smaller than in WT controls (Fig. 5D; m3kδ5-1; 0.23 ± 0.02 min−1 vs. WT: 0.35 ± 0.03 min−1, P < 0.03; and m3kδ5-2; 0.23 ± 0.02 min−1 vs. WT: 0.35 ± 0.03 min−1, P < 0.03), and half-response times were larger in m3kδ5-1 and m3kδ5-2 mutant leaves than in WT controls (Fig. 5E; m3kδ5-1; 5.4 ± 0.2 min vs. WT: 4.2 ± 0.1 min, P < 0.001; and m3kδ5-2: 5.0 ± 0.1 min vs. WT: 4.2 ± 0.1 min, P < 0.001).

Based on findings with the two m3kδ5 mutant alleles, m3kδ1/δ5-1/δ6-1/δ7 and m3kδ1/δ5-1/δ6-1/δ7 quadruple mutants were generated to investigate their VPD responses. Both mutant alleles exhibited larger steady-state stomatal conductances under low-VPD control conditions in both high-light (Fig. 6; 500 μmol·m−2·s−1) and light (SI Appendix, Fig. S7; 150 μmol·m−2·s−1) conditions compared to WT controls (Fig. 6 A and B and SI Appendix, Fig. S7 A and B). Furthermore, average rate constants of high VPD–induced stomatal closing in m3kδ1/δ5-1/δ6-1/δ7 and m3kδ1/δ5-1/δ6-1/δ7 quadruple-mutant leaves were reproducibly and significantly smaller than in the WT controls grown in parallel (Fig. 6 C and D; m3kδ1/δ5-1/δ6-1/δ7: 0.21 ± 0.01 min−1 vs. WT: 0.44 ± 0.06 min−1, P < 0.009 and m3kδ1/δ5-2/δ6-1/δ7: 0.14 ± 0.02 min−1 vs. WT: 0.29 ± 0.05 min−1, P < 0.04 and SI Appendix, Fig. S7C; m3kδ1/δ5-1/δ6-1/δ7: 0.11 ± 0.03 min−1 vs. WT: 0.40 ± 0.05 min−1, P < 0.001 and m3kδ1/δ5-2/δ6-1/δ7: 0.13 ± 0.02 min−1 vs. WT: 0.40 ± 0.05 min−1, P < 0.001). Half-response times were significantly larger in m3kδ1/δ5-1/δ6-1/δ7 and m3kδ1/δ5-2/δ6-1/δ7 quadruple-mutant leaves than in the WT controls at both light intensities (Fig. 6 E and F; m3kδ1/δ5-1/δ6-1/δ7: 6.5 ± 0.2 min vs. WT: 4.3 ± 0.2 min, P < 0.001 and m3kδ1/δ5-2/δ6-1/δ7: 7.7 ± 0.2 min vs. WT: 5.0 ± 0.4 min, P < 0.002 and SI Appendix, Fig. S7D; m3kδ1/δ5-1/δ6-1/δ7: 8.3 ± 0.5 min vs. WT: 6.5 ± 0.2 min).
quadruple mutants were independently found in intact whole-plant gas-exchange analyses (SI Appendix, Fig. S8). These results suggest that B3-family Raf-like M3Ks play a role in stomatal closure induced by VPD elevation.

**GHR1 Is Required for High VPD–Induced Stomatal Closure.** GHR1 encodes a receptor-like (pseudo)kinase, which is involved in activating the S-type anion channel SLAC1 to trigger stomatal closure (77, 79, 80). The involvement of GHR1 in the stomatal high-VPD response was analyzed in ghr1-3 and ghr1-6 exon-insertion mutant leaves. Leaves of the strong ghr1-3 allele and the weak ghr1-6 allele mutants (80) showed higher steady-state stomatal

4.4 ± 0.3 min, P < 0.001 and m3kδ1/δ5/δ6-δ1/δ7: 7.9 ± 0.4 min vs. WT: 4.4 ± 0.3 min, P < 0.001). These data at high (Fig. 6) and low (SI Appendix, Fig. S7) light intensities provide evidence that the phenotypes found in the m3kδ1/δ5/δ6/δ7 quadruple-mutant alleles are not simply a result of the higher steady-state conductance in these mutants compared to WT plants.

As the slowing of the VPD response in these quadruple-mutant alleles was significantly, but only slightly, slower, we investigated these mutants in the independent laboratory of E.M. in genotype-blinded experiments using a distinct whole-plant gas-exchange system. Higher stomatal conductances and in part increased half-response times in m3kδ1/δ5/δ6/δ7 quadruple mutants were independently found in intact whole-plant gas-exchange analyses (SI Appendix, Fig. S8). These results suggest that B3-family Raf-like M3Ks play a role in stomatal closure induced by VPD elevation.

**GHR1 Is Required for High VPD–Induced Stomatal Closure.** GHR1 encodes a receptor-like (pseudo)kinase, which is involved in activating the S-type anion channel SLAC1 to trigger stomatal closure (77, 79, 80). The involvement of GHR1 in the stomatal high-VPD response was analyzed in ghr1-3 and ghr1-6 exon-insertion mutant leaves. Leaves of the strong ghr1-3 allele and the weak ghr1-6 allele mutants (80) showed higher steady-state stomatal

Fig. 5. Stomatal VPD response is slightly slower in B3-family Raf-like kinase m3kδ5 mutants. (A) The Raf-M3Kδ5 kinase domain activates OST1 in Arabidopsis mesophyll cells. The protein kinase domains of the Raf-like M3Ks were expressed transiently in mesophyll cell protoplasts isolated from pUB8Q10:OST1-HF Arabidopsis leaves. OST1 kinase activity was visualized by in-gel kinase assays (n = 3). GFP, green fluorescent protein. CBB, coomassie brilliant blue loading control. Anti-GFP, immunoblotting of GFP-fusion proteins. (B and C) Time-resolved stomatal conductance (Gs) in response to increases in VPD in Col-0 (WT) and Raf-like m3kδ5 mutant under high light (450 μmol m⁻² s⁻¹ red light combined with 50 μmol m⁻² s⁻¹ blue light). WT controls shown in B and C are the same because mutants and WT were grown in parallel and analyzed within the same experimental sets. (D and E) Rate constants and half-response times in response to VPD elevations were calculated and presented. Data represent mean ± SEM. n = 5 for each genotype. Different letters represent significant differences (P < 0.05) analyzed by one-way ANOVA followed by Holm–Sidak multiple comparisons.

Fig. 6. Stomatal VPD response is slower in B3-family Raf-like kinase m3kδ1/δ5/δ6/δ7 quadruple mutants. (A and B) Time-resolved stomatal conductance (Gs) in response to increases in VPD in Col-0 (WT), Raf-like m3kδ1/δ5/δ6-δ1/δ7 quadruple-mutant, and m3kδ1/δ5-2/δ6-1/δ7 quadruple-mutant leaves under high light (450 μmol m⁻² s⁻¹ red light combined with 50 μmol m⁻² s⁻¹ blue light). VPDs are shown on top of the time axis in each panel. (C–F) Average rate constants (C and D) and average half-response times (E and F) in response to VPD elevations were calculated and presented. Data represent mean ± SEM. n = 5 for each genotype in A. n = 3 for WT and n = 4 for the m3kδ1/δ5-2/δ6-1/δ7 quadruple mutant in B. *P < 0.05; **P < 0.01 (analyzed by Student’s t test compared to WT control). Data represent one of two independent sets of experiments in each mutant allele showing similar results.
conductions in both high-VPD and low-VPD conditions compared to WT controls (Fig. 7 A and B and SI Appendix, Fig. S9 A and B). Although high-VPD treatment triggered a greater degree of stomatal closure in ghr1-3 and ghr1-6 mutant leaves, ghr1 mutant leaves exhibited a clear difference in response kinetics compared to WT leaves under both high-light (500 μmol m⁻² s⁻¹; Fig. 7 A and B and SI Appendix, Fig. S9 A and B) and low-light (150 μmol m⁻² s⁻¹; SI Appendix, Fig. S9C) experimental conditions. Rate constants in the strong ghr1-3 mutant leaves were significantly smaller than in WT leaves (Fig. 7 C and D; ghr1-3: 0.14 ± 0.01 min⁻¹ vs. WT: 0.36 ± 0.06 min⁻¹, P < 0.003 and ghr1-6: 0.24 ± 0.02 min⁻¹ vs. WT: 0.29 ± 0.05 min⁻¹, P = 0.37 and SI Appendix, Fig. S9 D–F; ghr1-3: 0.15 ± 0.00 min⁻¹ vs. WT: 0.44 ± 0.07 min⁻¹, P < 0.003; ghr1-6: 0.25 ± 0.02 min⁻¹ vs. WT: 0.42 ± 0.04 min⁻¹, P < 0.03; ghr1-3: 0.15 ± 0.01 min⁻¹ vs. WT: 0.41 ± 0.08 min⁻¹, P < 0.03; and ghr1-6: 0.26 ± 0.05 min⁻¹ vs. WT: 0.41 ± 0.08 min⁻¹, P = 0.15). Half-response times were larger in strong ghr1-3 mutant leaves than in WT controls (Fig. 7 E and F; ghr1-3: 6.4 ± 0.3 min vs. WT: 4.5 ± 0.2 min, P < 0.002 and ghr1-6: 5.0 ± 0.3 min vs. WT: 4.9 ± 0.2 min, P = 0.83 and SI Appendix, Fig. S9 G–I; ghr1-3: 4.1 ± 0.3 min vs. WT: 6.3 ± 0.1 min, P < 0.001; ghr1-6: 4.6 ± 0.3 min vs. WT: 4.2 ± 0.2 min, P = 0.33; ghr1-3: 5.9 ± 0.1 min vs. WT: 4.2 ± 0.5 min, P < 0.03; and ghr1-6: 5.0 ± 0.3 min vs. WT: 4.2 ± 0.5 min, P = 0.17). Taken together, our results show that GHR1 is required for the rapid stomatal closure induced in response to VPD increases. Furthermore, in contrast to all other mutants investigated in the present study, ghr1 mutants showed no initial transient stomatal conductance increase (wrong-way response) as found under high-light conditions (Fig. 7 A and B).

Discussion

The continuing elevation in global temperature is suggested to lead to a trend of rising vapor pressure deficit, though VPD is variable depending on conditions (81). Increases in VPD affect transpiration and regulate stomatal conductance of terrestrial plants (82–85). The molecular mechanisms of VPD sensing and signal transduction remain to a large degree unknown. Previous studies have focused on the roles of ABA in the VPD response (introduction). In the present study, we have investigated mutants affected in several stomatal responses. Mutants that disrupt specific known signaling mechanisms or pathways were investigated.

We show here that mpkl2/mpk4GC mutant alleles that disrupt early stomatal CO₂ signal transduction (75) show robust stomatal closing responses to VPD elevation. Osmotic stress exposures caused rapid cytoplasmic Ca²⁺ transients in guard cells (Fig. 2 A–C). Several higher-order calcium-permeable channel mutants (oscal1-2/1.3li2/2.3/3.1 quintuple, oscal1.3/1.7 double, glr3.2/3.3 double, cnng5-1/6-1 double, cnng20 single, and cnng19/20crsir double) and calcium-signaling mutants (cpk3/5/6/11/23 quintuple, cpk3/4/5/6/11 quintuple, chl1/4/5/8/9 quintuple, and chl2/3rf double) were analyzed, but were not affected in their high-VPD-triggered stomatal closing responses. Interestingly, however, we uncover here that the recently identified B3 family members of M3K RAF-like kinases, particularly M3K5, but not B4 RAF-like M3Ks, are required for an intact rapid stomatal VPD response. Furthermore, we reveal that the transmembrane receptor-like pseudokinase GHR1 functions in the stomatal response to VPD elevation.

Analyses of the ost1-3 single mutant (SI Appendix, Fig. S1) confirmed that OST1/SnRK2.6 is a central regulator of hydroactive high VPD–induced stomatal closure (5, 36, 39) and provided an internal control for our analysis system. The OST1 protein kinase regulates both S-type anion channels and the R-type anion channel ALMT12/QUAC1 in guard cells or Xenopus oocytes in response to several stimuli (28, 86–88). In the present study, analyses of slac1-3 single-mutant leaves were used as an additional system control, confirming that the S-type anion channel SLAC1 contributes to high-VPD-mediated stomatal closing (Fig. 1 A, D, and E and SI Appendix, Fig. S3 A, C, and D) (18).

Notably, the single almt12-1 mutant exhibited high VPD–induced stomatal closing kinetics similar to WT plants (Fig. 1 B, D, and E). Interestingly, however, slac1-3/almt12-1 double-mutant leaves showed a dramatic increase in the steady-state stomatal conductance and a slowing of the high-VPD response, which was kinetically similar to the slac1-3 single mutant (Fig. 1 C, D, and E and SI Appendix, Fig. S3). These results indicate that S-type anion channels play a rate-limiting role, and R-type anion channels function together with S-type anion channels in setting the steady-state stomatal conductance.

![Fig. 7. Stomatal VPD responses, including initial transient increase in stomatal conductance, are impaired in ghr1 mutants. (A and B) Time-resolved stomatal conduc-
tance (Gs) in response to increases in VPD in Col-0 (WT) and ghr1-3, and ghr1-6 mutant leaves under high light (450 μmol m⁻² s⁻¹ red light combined with 50 μmol m⁻² s⁻¹ blue light). VPDS are shown on top of the time axis in each panel. (C–F) Average rate constants (C and D) and average half-
response times (E and F) in response to VPD elevations were calculated and presented as described. Data represent mean ± SEM. *P < 0.01 (as analyzed by Student’s t test compared to WT control). WT: n = 4 leaves in A and n = 5 leaves in B; ghr1-3: n = 5 leaves; and ghr1-6: n = 5 leaves. Data rep-
resent one of two independent sets of experiments. Similar results were found in an independent set of experiments in each mutant allele (SI Appendix, Fig. S8).](https://doi.org/10.1073/pnas.2107280118)
under the imposed conditions. Given redundancies and nonlinearities in signal-transduction networks (89, 90), our data do not unequivocally exclude a contribution of R-type anion channels to the high VPD–induced stomatal closing response. The slow, but continued, stomatal closing response in slac1slat1sl2 double-mutant leaves suggests that additional anion channels and mechanisms function in parallel to these channels. In this study, higher steady-state stomatal conductances in m3kδ5 single-mutant and m3kδ1s5/δ6/δ7 quadruple-mutant alleles were not attributable to increases in stomatal index (SI Appendix, Fig. S10). Furthermore, the slower response to an increase in VPD in m3kδ1s5/δ6/δ7 quadruple mutants was not simply caused by the increased initial steady-state stomatal conductance in these quadruple mutants for several reasons. First, we found that the larger half-response time in m3kδ1s5/δ6/δ7 quadruple mutants was consistently observed, even at low light, with reduced initial steady-state stomatal conductance levels (SI Appendix, Fig. S7). Second, unlike m3kδ1s5/δ6/δ7 quadruple-mutant alleles, no significant differences in half-response times were found in additional controls in WT plants exposed to high-light and low-light conditions under 400 ppm per million (ppm) CO2 or 150 ppm CO2 (SI Appendix, Fig. S11). Third, a rapid stomatal response to increases in VPD was observed in the early CO2 signaling mpk12/4GC mutant lines that also exhibited initial steady-state stomatal conductance (Fig. 3). Therefore, our findings suggest that B3 M3Ks are involved in accelerating the hydroscopic signaling response to an increase in VPD.

Previous studies showed that ABA and hyperosmolarity activate the OST1/SnRK2.6/SRK2E protein kinase in Arabidopsis culture cells, roots, and seedlings (32, 50, 91, 92). Moreover, the OST1/SnRK2.6/SRK2E-like protein kinases are also activated by ABA in guard cells (31, 47, 93–96). Although low humidity/high VPD activates the OST1/SnRK2.6/SRK2E protein kinase in leaves (50), it remains to be determined whether, and if so how, low humidity may enhance OST1/SnRK2s activity in guard cells. In-gel kinase assays demonstrated that Raf-like kinases are required for OST1/SnRK2s kinase activation during ABA and hyperosmolarity stimulation in seedlings (52–54, 78). In the present study, we have found that the guard-cell-expressed M3Kδ5-protein kinase is activated in parallel to the OST1/SnRK2.6 in plants (Fig. 5a). The higher basal stomatal conductance in m3kδ5 single-mutant alleles and m3kδ1s5/δ6/δ7 quadruple mutants (Fig. 5 and 6 and SI Appendix, Fig. S7) may be due to a lower basal OST1 activity in mutant guard cells. The slower high VPD–induced stomatal closure in m3kδ1s5/δ6/δ7 quadruple-mutant leaves (Fig. 6 and SI Appendix, Fig. S7) may be caused by a reduction in Raf-like kinase-mediated OST1/SnRK2s activity in guard cells. Time-resolved analyses of OST1/SnRK2 protein kinase activity (47) in response to VPD changes could allow testing of this model. Unlike the m3kδ5 single-mutant and m3kδ1s5/δ6/δ7 quadruple-mutant alleles, no obvious high VPD–triggered stomatal response phenotype was observed in m3kδ1s5/δ6/δ7 triple, m3kδ1s5/δ6/δ7, and an ok-quatudec quattuordecuple mutant (Fig. 4D). There are 80 M3K genes encoded in the Arabidopsis genome (97–99). The OK-quatudec line includes mutations in 14 Raf-like kinase genes, including several osmotic stress-activated B4 family members (53). The present findings indicate a less important role of these B4-family Raf-like kinase members in the VPD response, compared to the osmotic stress response in whole seedlings. The present results further suggest that M3Kδ5/RAF6 plays an important role in VPD-induced stomatal regulation.

A previous study suggested that the receptor-like pseudokinase GHR1 mediates stomatal closure in response to ABA, CO2, ozone, and darkness, but may not be required for high VPD–triggered stomatal closure (80). In the present study, the severe ghr1-3 mutant allele showed a clear impairment in high VPD–induced stomatal closure (Fig. 7 and SI Appendix, Fig. S9). GHR1 can directly activate the SLAC1 anion channel in Xenopus oocytes when these two proteins are artificially linked via BiFC constructs (79, 80). GHR1 is required for stomatal closure in response to ABA, hydrogen peroxide, ozone, high CO2, and low light intensity (79, 80), and GHR1-mediated activation of the SLAC1 anion channel relies on a different signaling pathway that makes use of GHR1. It has been suggested that the receptor-like pseudokinase GHR1 serves as a scaffolding protein to recruit the calcium-dependent protein kinase CPK3 for regulating SLAC1 anion-channel activation (80). Higher-order cpk mutants analyzed here included cpk3 mutation, but did not affect the high VPD response (SI Appendix, Fig. S6 A and B). Therefore, GHR1 may recruit additional, yet to be discovered signaling components to regulate the stomatal high VPD response.

Classical research showed that high VPD/low humidity triggers a transient stomatal opening before stomatal closing (56, 100–102). Notably, this transient increase in stomatal conductance in response to an increase in VPD (wrong-way response) may in part result from a simplified assumption of saturated vapor pressure inside leaves during stomatal conductance recordings at high stomatal conductances (103). The wrong-way response was found in WT and most of the studied mutants, including ost1-3, slac1-3, slac1-3slat12-1, and mpk12/4GC mutants, with high steady-state stomatal conductances. These data indicate that the larger stomatal conductance of ghr1 leaves is not solely due to the above-simplified assumption. Interestingly, the lack of a clear wrong-way response observed in both ghr1 mutant alleles points to an additional role of GHR1. It has been proposed that the initial hydropassive loss of turgor in epidermal cells adjacent to guard cells during high VPD/low-humidity stimulation drives the transient stomatal opening (2, 104). In a previous study, a transient stomatal opening could only be observed in stomata with living subsidiary cells in Tradescantia pallida or when guard cells were surrounded by living epidermal cells in Vicia faba (3). Therefore, the lack of high VPD–triggered transient stomatal opening in ghr1 mutants suggests a function of GHR1 in the mechanical interaction between guard cells and neighboring pavement cells. It remains to be elucidated whether mechanical and hyperosmolarity-sensing mechanisms are involved in high VPD–induced stomatal closure. OSCA channels have been reported to be mechanical- and osmotic-sensing calcium channels in higher plants, with structural analyses resolving possible mechanisms (59–64). In the present study, we generated higher-order osca mutants, based on guard-cell transcriptome analyses from independent laboratories (SI Appendix, Fig. S4) (58, 65). However, the stomatal responses to high VPD were not altered in osca1-2/1.3/2.2/3.3/3.1 quintuple-mutant leaves (Fig. 2D), and hyperosmotic stress was insufficient to activate the rice ortholog OsOSCA2;1 when expressed alone in Xenopus oocytes (64). It has been reported that the activation of Raf-like kinases or SnRK2s by hyperosmolarity was not altered in an osca septuple mutant (53). These results suggest that high VPD–induced stomatal closure may be independent of OSCAs and, in particular, the five more guard-cell-expressed OSCAs analyzed here. A recent study has identified an additional OSCA that is not highly expressed in guard cells, OSCA1.7, but is together with OSCA1.3 required for PAMP-induced stomatal closing during pathogen signaling (66). However, we found WT-like high VPD responses in osca1.3/1.7 double-mutant
leaves. We show that guard-cell cytosolic Ca^{2+} concentrations transiently increase in response to both hyperosmolality and hypo-osmolality (Fig. 2 A–C), but whether Ca^{2+} signaling is required for the stomatal high-VPD response remains to be investigated. Although no obvious stomatal closure phenotype in response to VPD elevation was found in the investigated calcium-permeable channel mutants, cpk quintuple, and higher-order cbl mutants (Fig. 2 D–G and SI Appendix, Fig. S6), we do not exclude a possible role of calcium in stomatal high-VPD signal transduction. Additional higher-order mutants in Ca^{2+} signaling components may be needed to impair the stomatal high-VPD response.

In summary, the identification of B3-family Raf-like kinases and the transmembrane receptor-like GHR1 in the stomatal VPD response advances the understanding of molecular and genetic mechanisms that mediate the hydroactive stomatal VPD response. Interestingly, the initial transient stomatal opening wrong-way response to a VPD increase that precedes stomatal closing (56, 100) was disrupted in ghr1 mutant alleles, indicating a function of GHR1 in guard-cell–pavement-cell communication during VPD signaling. Furthermore, our data exclude the involvement of the early CO_{2} signal-transduction pathway in rapid high VPD–induced stomatal closure. Further research will be required to elucidate the upstream high VPD–sensing mechanisms, which are shown here to require B3-family Raf-like kinases and the receptor-like (pseudo) protein kinase GHR1. Understanding the detailed molecular and genetic mechanisms underlying the VPD response is relevant for models that predict plant transpiration in response to climate change and could become relevant for the future development of plants with enhanced resilience to climate change.

Materials and Methods

Plant Materials and Growth Conditions. Arabidopsis thaliana Columbia-0 (Col-0) ectype plants were used as WT. Arabidopsis mutant lines investigated in this study: ost-1 (32), slact-3 (17, 18), almt12-1 (16), slact-3almt12-1, osca1-1 (76), ghr1-2, ghr1-3, ghr1-2/3, and ghr1-2/3/4 (105), cngc5-11/12/13, cpg3/7/8/9 (105), cagg2 (Salk_129133), cpgk3-1 (58). Plants were isolated from 3- to 4-wk-old Arabidopsis leaves were transfected with 20 μg of pUC18 plasmids carrying 35S:Raf-M3K-GFP:nosT using the polyethylene glycol-mediated method (107). After overnight incubation in incubation buffer (10 mM 2-[N-morpholino]ethanesulfonic acid [MES]-KOH, pH 6.0, 0.4 M mannitol, 20 μM AVG, and 1 mM CaCl_{2}), protoplasts were harvested by centrifugation and lysed in sodium dodecyl sulfate–polyacrylamide gel electrophoresis sample buffer.

Analyses of Guard-Cell Cytosolic Calcium Dynamics. For analyses of cytosolic Ca^{2+} dynamics in guard cells, we used A. thaliana Col-0 plants expressing Yellow Cameleon YC3.60 under control of the GGH5 promoter (58). Plants were grown on soil under long-day conditions (16:8 h at a photon fluence rate of 75 μmol m^{-2} s^{-1} and a temperature of 20°C). Small pieces of mature leaves of 4-wk-old plants were glued to the cover slide by using a medical adhesive (Hollister Inc.); upper cell layers were gently removed with a razor blade, thus isolating epidermal strips. The leaf epidermal strips were preincubated in a buffer solution (5 mM KCl, 1 mM MES, and 50 μM Ca^{2+}, pH 6.5, adjusted to 300 μM NO_{3}^{-} for 2 to 3 h with blue light (450 μmol m^{-2} s^{-1}) to induce the stomatal opening to get turgid guard cells. The epidermal strips were then mounted on an open-top imaging chamber and perfused with the same solution (5 mM KCl, 10 mM MES, and 50 μM Ca^{2+}, pH 6.5) as described (58, 108). To induce the hyperosmotic stress, after 5 min, the strips were perfused with the standard solution supplemented with 500 mM sorbitol. Epidermal strips were kept in this solution for 10 min, and then the hypertonic solution was replaced again with the standard buffer solution, without sorbitol, to induce the hypo-osmotic stress.

To perform Ca^{2+} imaging experiments, images of epidermises with stomatal guard cells were acquired with a Nikon TE300 inverted fluorescence microscope, with a FLUORESCEIN Module ILLUMINATOR (TE-FM) epi-fluorescence attachment (Nikon Inc.) using a Nikon 60× Plan Apo oil objective with a numerical aperture (n.a.) of 1.4. Excitation light was produced by a 75-W xenon fluorescent lamp (Osram) that was attenuated by 97% (3% light transmission) by using both 4x and 8x neutral density filters to reduce exposure of the fluorescent reporters and cells to epifluorescence excitation. Cameleon YC3.60 specific excitation and detection was achieved by using a yellow fluorescent protein/CFP filter set: 440/20 nm excitation, 485/40 nm emission for CFP, and 535/30 nm emission for cpVenus with a 455 nm dichroic long-pass filter (Chroma Technology). The filter wheel, shutter, and CoolSNAP charge-coupled device camera from Photometrics (Roper Scientific) were controlled by Fluor Systeem Software (MDS Inc.). Images were acquired every 3 s, on a time range of >30 min.

Fluorescence intensity was determined over regions of interest (ROIs) that corresponded to single guard cells. cpVenus and CFP emissions of the analyzed
ROIs were used for the ratio (R) calculation (cpVenus/CFP). cpVenus/CFP ratios were normalized to the initial ratio (R0) and plotted vs. time (ΔR/Δt). Background subtraction was performed independently for both channels before calculating the ratio. Data are representative of n = 6 experiments. P values were calculated with an unpaired Student's t-test. Data are illustrated as scatter dot plots using GraphPad.

**Data Availability.** All study data are included in the article and/or Dataset 51. Sequences of genes described in this article can be found in the Arabidopsis Genome Initiative database under the following accession numbers: OST1 (AT4G33950), SLAC1 (AT1G12480), ALMT12/QUA1 (AT4G17970), OSCA1 (AT4G04340), OSCA1.3 (AT1G11960), OSCA1.7 (AT4G02900), OSCA2.2 (AT1G00900), OSCA2.3 (AT3G10000), OSCA3.1 (AT3G0360), GLR3.2 (AT3G35290), GLR3.3 (AT3G25400), CNGC5 (AT3G57940), CNGC8 (AT3G23980), CNGC19 (AT3G17690), CNGC20 (AT3G17700), CPK3 (AT2G34360), CPK5 (AT4G02800), CPK7 (AT3G53310), CPK9 (AT2G17290), CPK11 (AT1G35670), CPK23 (AT4G04740), CBL1 (AT1G17615), CBL2 (AT4G55990), CBL3 (AT4G26570), CBL4 (AT5G24270), CBL5 (AT4G10420), CBL8 (AT1G64480), CBL9 (AT4G57100), MPK4 (AT1G3070), MPK12 (AT2G64070), MKII3 (AT3G11850), MKII3 (AT2G27510).

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