FERONIA receptor kinase pathway suppresses abscisic acid signaling in Arabidopsis by activating ABI2 phosphatase

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Plant growth and development are controlled by a delicate balance of hormonal cues. Growth-promoting hormones and growth-inhibiting counterparts often antagonize each other in their action, but the molecular mechanisms underlying these events remain largely unknown. Here, we report a cross-talk mechanism that enables a receptor-like kinase, FERONIA (FER), a positive regulator of auxin-promoted growth, to suppress the abscisic acid (ABA) response through activation of ABI2, a negative regulator of ABA signaling. The FER pathway consists of a FER kinase interacting with guanine exchange factors GEF1, GEF4, and GEF10 that, in turn, activate GTPase ROP11/ARAC10. Arabidopsis mutants disrupted in any step of the FER pathway, including fer, gelf1gelf4gef10, or rop11/arac10, all displayed an ABA-hypersensitive response, implicating the FER pathway in the suppression mechanism. In search of the target for the FER pathway, we found that the ROP11/ARAC10 protein physically interacted with the ABI2 phosphatase and enhanced its activity, thereby linking the FER pathway with the inhibition of ABA signaling.

A-type protein phosphatase 2C | signal transduction | small GTPase

Originally discovered as a hormone for senescence and abscission, abscisic acid (ABA) is generally considered an inhibitor of cell growth (1, 2). In addition, ABA plays a central role in plant stress responses. The content of ABA in plant cells significantly increases under drought or salinity stress conditions, inducing stomatal closure and gene expression to cope with stress conditions (3–5). Stress adaptation often involves the inhibition of vegetative growth, consistent with the known role of ABA. Recent work has identified several putative ABA receptors (3–5), in particular, the PYR/PYL/RCAR proteins linking ABA perception to a biochemical pathway consisting of A-type protein phosphatase 2Cs (PP2Cs) and SNF1-related protein kinase 2 (SnRK2)-type protein kinases (6–9). It is widely accepted that ABA binding to PYR/PYL/RCAR proteins initiates physical interaction and inhibition of PP2Cs, negative regulators in ABA signaling (3–5). Deactivation of PP2Cs releases SnRK2 kinases from the PP2C–SnRK2 complexes, activating those kinases that target proteins downstream.

While this primary “linear” pathway was being defined, parallel studies suggested that ABA signaling is regulated by a number of other pathways initiated by hormones and environmental factors (3, 4). However, little is understood about the molecular links that allow these alternate signaling pathways to intercept the primary ABA signaling pathway. We report that the FERONIA (FER) receptor kinase pathway, a signaling module that is required for auxin response (10), suppresses ABA signaling through activation of ABI2, a PP2C member located in the primary signaling pathway of ABA response.

Results and Discussion

fer Mutants Are Hypersensitive to ABA. Receptor-like protein kinases (RLKs), which contain an extracellular receptor domain and a cytosolic Ser/Thr kinase domain, function in a number of signaling processes in response to hormones and environmental factors (11). In previous work, we studied the transcriptional activity of a diverse superfamily of 604 RLKs associated with hormones and environmental stress (12). In particular, in an attempt to connect RLK-mediated signaling with hormonal responses, we focused on those RLKs that are highly regulated by hormones. We identified FER as one of the RLK members that displayed opposing expression patterns after treatment of Arabidopsis seedlings with auxin and ABA. FER mRNA was up-regulated by auxin and down-regulated by ABA. This finding was confirmed by available online databases and quantitative RT-PCR analysis (Fig. S1A). The FER gene was first identified as a regulator in the male–female interaction during pollen tube reception (13–15). FER was later shown to function in several other growth regulatory pathways, including root hair elongation regulated by auxin and cell growth induced by other hormones (10, 16–18).

We were curious as to whether the opposite patterns of regulation of FER gene expression by auxin and ABA, two antagonizing hormones in cell growth (2), may implicate a role of FER in both auxin and ABA signaling pathways. To follow up on this possibility, we examined ABA responses of fer mutants to determine whether disruption of FER function impacts ABA sensitivity. Three FER mutants were analyzed: two null mutants (fer-4 and sm) and a truncated allele called fer-5 that shows a partial defect in FER function (10). When germinated on Murashige and Skoog (MS) medium supplemented with ABA, fer-4 and sm seedlings showed dramatic growth arrest and turned brown in response to ABA (Fig. 1A a and b, B, and C), whereas the wild-type (WT) and fer-5 seedlings grew well and remained green. In the absence of exogenous ABA, cotyledons of all five genotypes were expanded and green. When supplemented with 0.25 μM ABA, both fer-4 and sm failed to produce true leaves and eventually died (Fig. 1Ab and C).

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We examined the \textit{fer-4} mutant in comparison with the WT in the ABA-inhibited growth assay. Without ABA, the primary root growth of WT and \textit{fer-4} was largely comparable. However, when supplemented with 3 \( \mu \text{M} \) ABA, root growth in \textit{fer-4} seedlings was significantly decreased compared with WT \((P < 0.0001)\) (Fig. 1D). This assay and the one described immediately above demonstrate that the inhibition of growth by ABA was dramatically enhanced in the \textit{fer} mutants.

We then determined whether ABA-induced environmental responses such as stomatal closure are altered in \textit{fer} mutants. We performed stomatal assays using isolated rosette leaves from \textit{fer-4} and WT plants. Maximum stomatal opening was observed when WT intact rosette leaves were floated on stomata-opening buffer and illuminated for 3 h. The same treatment induced maximal aperture of \textit{fer}-4 mutants in 6 h, indicative of slower stomatal opening. After maximal aperture was achieved in both the WT and \textit{fer-4} mutant, samples were exposed to 1 \( \mu \text{M} \) ABA for 1 h, and stomatal apertures were again measured. Fig. 1E shows that stomatal opening in the \textit{fer-4} mutant was more sensitive to ABA than the WT \((P < 0.0001)\).

**FER May Regulate Reactive Oxygen Species (ROS)-Mediated ABA-Signaling Pathway.** Previous work showed that FER regulates auxin-stimulated ROS accumulation in the root and root hair (10), as well as in the leaf under fungal invasion (19). Ample evidence supports a role for ROS as a second messenger in the ABA signaling pathway (20-22). It is possible that FER may regulate ROS accumulation in the ABA response. Because the \textit{FER} mutation caused ABA hypersensitivity in stomatal closing, we analyzed the accumulation of ROS in guard cells. We found higher levels of ROS in the \textit{fer-4} guard cells either in the absence or presence of exogenous ABA (Fig. 1F). Whereas ROS accumulation in the WT plants increased 87\% after 45-min treatment with 25 \( \mu \text{M} \) ABA (from 286.032 \( \pm \) 46.340 to 535.530 \( \pm \) 38.618), the same treatment resulted in a 500\% increase in ROS levels in the \textit{fer-4} mutant (from 525.041 \( \pm \) 119.109 to 3483.611 \( \pm \) 413.820) (Fig. 1F and Fig. S1G). From these data, we suggest that ABA hypersensitivity of \textit{fer-4} guard cells may be caused, at least in part, by increased ROS production compared with WT.

**ROP11/ARAC10 Interacted with GEF1, GEF4, GEF10, and AB2.** Duan et al. (10) reported that FER directly interacts with guanine nucleotide exchange factors (RopGEF) to activate GTPase (ROP/ARAC) in response to auxin. Furthermore, some ROPs, such as ROP10 and ROP6, have been shown to negatively regulate ABA responses (23, 24). It is possible that FER may regulate ABA responses through activation of some ROPs/ARACs that interact with components in the primary ABA signaling pathway involving the PYR/PYL/RCAR receptors, A-type PP2Cs, and SnRK2s. To test this hypothesis, we carried out yeast two-hybrid (Y2H) assays to identify physical interactions among FER, GEFs, ROPs, A-type PP2Cs (25, 26), and SnRK2s. It has been shown that FER interacts with the five GEFs that function as activators of ROPs (10). To test this hypothesis, we carried out yeast two-hybrid (Y2H) assays to identify physical interactions among FER, GEFs, ROPs, A-type PP2Cs, and SnRK2s. To test this hypothesis, we carried out yeast two-hybrid (Y2H) assays to identify physical interactions among FER, GEFs, ROPs, A-type PP2Cs, and SnRK2s. To test this hypothesis, we carried out yeast two-hybrid (Y2H) assays to identify physical interactions among FER, GEFs, ROPs, A-type PP2Cs, and SnRK2s. To test this hypothesis, we carried out yeast two-hybrid (Y2H) assays to identify physical interactions among FER, GEFs, ROPs, A-type PP2Cs, and SnRK2s.
data suggest that the GTP-binding form, but not the GDP-binding form, of ROP11/ARAC10 interacted with ABI2 (Fig. 2 A–D). These results suggest that the interaction of ROP11/ARAC10 with ABI2 occurs after activation by FER–GEFs and may serve as a cross-talk mechanism between the FER pathway and the primary ABA signaling pathway.

Interestingly, although ROP10 and ROP11/ARAC10 are similar in amino acid sequence, and both function in ABA responses, ROP10 did not interact with ABI2 in the Y2H and in BiFC assays, suggesting that ROP10 may regulate ABA responses through different partner proteins and pathways. Indeed, mutation in either ROP10 or ROP11 causes ABA hypersensitivity in mutant plants (23) (Fig. 3), consistent with the idea that they are not functionally redundant.

We next analyzed the expression patterns of ROP11/ARAC10 and found that ROP11/ARAC10 was highly expressed in stomatal guard cells and several other tissues (Fig. 2D a–d). This expression pattern coincided with those for FER and GEF1 (Fig. 2D e–h and ref. 10), making it possible for these genes to work together in responses to ABA.

ROP11/ARAC10 Can Activate the Phosphatase Activity of ABI2. We next examined the biochemical consequence of the ROP11/ARAC10–ABI2 interaction by measuring the phosphatase activity of ABI2 in the presence and absence of ROP11/ARAC10. As shown in Fig. 3 A, the WT ROP11/ARAC10 and CA-ROP11/ARAC10 loaded with GTP-γ-S activated ABI2 phosphatase (P < 0.0001), but the DN-ROP11/ARAC10 loaded with GDP did not (Fig. 3A). Another ROP/ARAC, ROP10, also failed to interact with ABI2 or activate its phosphatase activity in the presence of GTP-γ-S. In the ABA signaling pathway, ABI2, as one of the A-type PP2Cs, inhibits the SnRK2.6/OST1 kinase activity (3–5). If ROP11/ARAC10 enhances ABI2 activity, the presence of ROP11/ARAC10 should further reduce the kinase activity of OST1. In a kinase assay, the activity of OST1 was reduced by the addition of ABI2, consistent with earlier studies. Adding ABI2 together with ROP11/ARAC10 further reduced kinase activity (Fig. 3B). OST1 activity was not affected by ROP11/ARAC10 in the absence of ABI2, confirming that ROP11/ARAC10 inhibits ABA signaling through activation of the negative regulator ABI2.

rop11/arac10 Mutants Were Hypersensitive to ABA. If ROP11/ARAC10 interacts with ABI2, thereby activating its phosphatase activity, the rop11/arac10 mutant, like the fer mutant, should be hypersensitive to ABA. To pursue this possibility, we isolated three mutant alleles of ROP11/ARAC10 for ABA response analysis: rop11/arac10-1, Salk_039681; rop11/arac10-2, Salk_087260C; and rop11/arac10-3, Salk_063154 (Fig. S1, B and C). We found that all three mutants were hypersensitive to ABA in the seed germination assay (Fig. 3E and Fig. S1 D–F). In addition, rop11/arac10 10 mutants resembled fer-4 in being hypersensitive to ABA in the seedling growth assay (Fig. 3F and Fig. S1D). Finally, when the rop11/arac10 10 mutants were analyzed in the root growth and stomatal assays, they resembled fer-4 in displaying increased sensitivity to ABA (P < 0.0001) (Fig. 3E and Fig. S1E).

gef1/4 Double and gef1/4/10 Triple Mutants Were Hypersensitive to ABA. To confirm the role for the FER–GEF1/4/10–ROP11–ABI2 pathway in the ABA response, we analyzed this property in null mutants of gef1 (GK_586B11), gef4 (Salk_107520), and gef10 (Salk_009456) and found no difference in these single mutants compared with WT (Fig. 4A and B), suggesting functional redundancy of the three GEF genes. By contrast, when the gef1/4 double and gef1/4/10 triple mutants were isolated and examined in the same assays, they displayed ABA hypersensitivity in both the root growth and stomatal assays. Furthermore, the gef1/4/10 triple mutant showed a stronger phenotype compared with the gef1/4 double mutant (Fig. 4A and B). This result suggests that these three GEFs interact with FER and ROP11/ARAC10 and share functions in relaying the FER signal in responding to ABA.

Concluding Remarks

We have identified a pathway linked to the FER receptor kinase that negatively regulates ABA responses through activation of an A-type PP2C phosphatase, ABI2. In the absence of ABA, the phosphatase interacts and inhibits an SnRK2-type protein kinase that acts as a positive regulator of ABA responses. Increased levels of ABA induce an interaction of PP2C with PYR/PYL/RCAR receptors that triggers release of active SnRK2s, thereby activating downstream targets (25–27). These targets include...
transcription factors that activate ABA-responsive gene expression (8), ion channels that regulate guard cell turgor and stomatal aperture (25, 28), and enzymes that control the production of ROS (29). The A-type PP2Cs and SnRK2s are partitioned in multiple subcellular locations, including the nucleus, cytosol, and plasma membrane, thus enabling them to regulate targets present in different compartments of the cell. Although the FER receptor kinase positively regulates the auxin response in the growth of roots and root hairs (10), the present results suggest that FER–RopGEF–ROP/ARAC modules negatively regulate...
ABA responses, thus serving as a cross-talk node for antagonizing the ABA response by the auxin signaling pathway.

We propose that the FER–ROPGEF1/4/10–ARAC10/ROP11 pathway channels a signal to activate ABI2 that, in turn, inhibits the SnRK2-mediated ROS production characteristic of the ABA response (Fig. 4C). Disruption of the FER pathway lowers ABI2 activity and leads to hyperactivation of the SnRK2s with consequent elevated levels of ROS production. The increased ROS induces stomatal closure, cell death, and growth inhibition as reflected in ABA-induced seedling arrest and root inhibition. It is possible that ion channels such as SLAC1 located in the plasma membrane participate in this cross-talk and also function in the stomatal response.

Materials and Methods

Phenotype and ROS Analyses. Arabidopsis thaliana plants were grown in a room with constant temperature at 23 °C under 16-h-light/8-h-dark cycles. Seeds for the T-DNA insertion lines were purchased from the Arabidopsis Biological Resource Center and the GABI-Kat collections, and the double and triple mutants were obtained by crosses of single mutants. Arabidopsis seeds were surface-sterilized and germinated on 1/2 MS medium supplemented with 1% sucrose and solidified with 0.9% agar. ABA (mixes isomers; Sigma-Aldrich A1049) or an equal volume of ethanol as a control was added to the medium to the concentrations indicated in the figure legends.

For stomatal aperture assays, leaves were floated on stomatal opening buffer (5 mM Mes, 5 mM KCl, 50 μM CaCl2, pH 5.6) under light for the indicated time and then treated with ABA for 1 h. Thereafter, stomatal apertures were measured as described (30). ROS measurements were performed in guard cells essentially as described (30). Briefly, epidermal tissues were incubated for 3 h in a buffer containing 30 mM KCl and 10 mM Mes-KOH, pH 6.15. Then, 30 μM 2′,7′-dichlorodihydrofluorescein diacetate (Sigma) was added to the incubation buffer. After 20-min incubation, the dye was removed by three washes with distilled water. Epidermal tissues were incubated for 45 min with 25 μM ABA or equal volume of ethanol as a control. The fluorescence was detected by using a confocal microscope (OLYMPUS FV1000) with 488-nm excitation light.

Gene Expression Studies. Total RNA was prepared with TRIzol (Invitrogen) after DNase treatment (17); the primers used in these experiments are shown in Table S1. For β-glucuronidase (GUS) expression analysis, the promoter region containing the ATG start codon of ROP11/ARAC10 and GEF1 were amplified (primers are shown in Table S1) and cloned into the pGADT7 vector by using the primers described in Table S1, cotransformed into yeast cells AH109, and plated on synthetic dropout medium (SD-Trp-Leu-His) supplemented with 10 mM 3-aminotriazole for 1 wk (31).

Protein–Protein Interaction Analysis. Full-length cDNA sequences of the PpPP2Cas were subcloned into the pgADT7 vector by using the primers described in Table S1, cotransformed into yeast cells AH109, and plated on synthetic dropout medium (SD-Trp-Leu-His) supplemented with 10 mM 3-aminotriazole for 1 wk (31).

According to a ROP10 study, mutations Q66L and D123A in the ROP10 protein produce a CA and a DN form of ROP10, respectively (23). The same mutations in ROP11/ARAC10 were made to produce CA–ROP11/ARAC10 and DN–ROP11/ARAC10 by using PCR-based site-directed mutagenesis. Two primers (ROP11–CD-F and ROP11–CD-R) were used to amplify the full-length coding region of ROP11/ARAC10. Two mutagenesis primer pairs (ROP11–CA-TF paired with ROP11–CA-TR and ROP11–DN-TR paired with ROP11–DN-TF) were used to produce Q66L and D123A mutations, respectively. The coding

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Fig. 4. ABA response of gef1 mutants and a working model for the FER signaling pathway in the regulation of ABA responses. (A) ABA-induced growth inhibition in roots. The WT, single mutants, gef1/4 double mutant, and gef1/4/10 triple mutant were grown on MS medium for 5 d before transfer to MS containing 12 μM ABA. Root length was then measured 3 d after transfer. Each value is the average ± SE of 3 replicates of 30 plants. Three separate experiments were conducted. (B) ABA-induced stomatal closure in WT and gef mutants. Leaves were treated with 5 μM ABA for 1 h before stomatal apertures were measured. Each value is the average ± SE of 3 independent experiments with 10 apertures each. (C) Working model for FER pathway in regulating response to ABA. The FER–GEF–ROP/ARAC pathway is an essential module for the auxin response in cell growth. ABI2 is a major component of ABA signaling pathway. The ROP11/ARAC10–ABI2 interaction mediates cross-talk between the auxin and ABA signals. Arrows denote activation, and bars indicate repression. Details are described in the text.
regions of ROP11/ARAC10, CA-ROP11/ARAC10 and DN-ROP11/ARAC10 were subcloned to the pGBK7 separately. The primers are listed in Table S1. The DNA sequence of ROP11/ARAC10 and 2PCAs were amplified by PCR and cloned into plasmids pE3228 and pE3449, respectively, for BIFC assays in mesophyll protoplasts (32). Protoplasts were isolated from 5-wk-old Arabidopsis rosette leaves as essentially described (33). Leaf strips were incubated in the cell wall-degrading enzyme solution in the dark for 3.5 h. Protoplasts were purified (33) and transfected with 20 μg of plasmid DNA and an equal volume of PEG solution. The transfected protoplasts were incubated in the dark at 23 °C for 16 h to allow expression of the BIFC proteins. Fluorescence detection was performed using a confocal microscope (LSM 5 LIVE; Zeiss) with an excitation wavelength of 488 nm.

**ABI2 and OST1 Enzyme Assays.** The coding DNA sequences of ROP10, ROP11/ARAC10, and ABI2 were amplified by PCR and cloned into pGEX4T-1 (Amersham Biosciences) expression vector. All constructs were transformed into BL21(DE3) for protein expression according to a described procedure (25). Proteins were purified as GST fusions by using immobilized glutathione according to instructions of the manufacturer (Pierce). The glutathione solution containing ABI2 was concentrated and dialyzed vs. a buffer containing 50 mM imidazole (pH 7.2) and 5 mM MgCl2. The recombinant ROP10, ROP11/ARAC10, CA-ROP11/ARAC10, and DN-ROP11/ARAC10 were first incubated with 10 μM EDTA for 5 min at room temperature, and then the GTPases were loaded with 10 mM GTP-γ-S (Sigma; G8634) or 10 mM GDP (Sigma; G7127) for 20 min at room temperature, followed by the addition of 40 mM MgCl2. Loaded proteins were washed four times in 50 mM imidazole (pH 7.2) and 5 mM MgCl2.

Phosphatase activity was measured by using the Ser/Thr Phosphatase assay kit (Promega; V2460) as described (6). Reactions were performed in a 50-μL volume containing 9 μg of GST-ABI2, in the presence of other factors, including 10 μg of GST-ROP11/ARAC10, GST-CA-ROP11/ARAC10, GST-DN-ROP11/ARAC10, or GST-ROP10. After incubation with the RR(A)phosphoT(AY) peptide substrate in 50 mM imidazole (pH 7.2) and 5 mM MgCl2, 5 mM MgCl2, 0.5 μM/L BSA, and 0.1% β-mercaptoethanol for 40 min at room temperature, the reaction was stopped by adding 50 μL of molybdate dye. After incubation for an additional 20 min at room temperature, absorbance was measured at 620 nm.

For OST1 kinase assays, GST–OST1 was expressed and purified as described (25). The kinase buffer contained 20 mM Tris HCl (pH 7.5), 2.5 mM MnCl2, 2.5 mM MgCl2, 1 mM CaCl2, 1 mM DTT, and 1x Roche EDTA free proteinase inhibitors. The protein combinations indicated in the legend of Fig. 3 were included in a total volume of 20 μL. The reaction was started by adding 1μM [32P] ATP. After incubation for 45 min at 30 °C, the reaction was stopped by adding 4 μL of 6x loading buffer and then separated by SDS/PAGE using a 12% (wt/vol) SDS-acrylamide gel. 32P was detected by autoradiography using a Typhoon imager (Molecular Dynamics).

**Note.** When this paper was under preparation, another group reported that ROP11 interacted with ABI1 and regulated ABA responses (34, 35), although authors found that ABI1 interaction with ROP11 occurs when the regulatory domain of ABI1 is deleted (35). It seems likely that different cross-talk nodes are present in subcellular compartments, such as the nucleus and cytosol, where they link alternate signals to ABA. Further work should highlight the importance of these cross-talk nodes in hormonal interactions that control plant growth and development.

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Fig. S1. Hormone-regulated FER expression and phenotypic analyses of rop11/arac10. (A) Quantitative RT-PCR analysis of FER transcripts under auxin and ABA treatments (Materials and Methods). Three independent experiments yielded similar results. (B) Scheme of ROP11/ARAC10 gene organization and position of T-DNA insertions. The ATG start codon and TGA stop codon are indicated; boxes are exons, and lines are introns. The exact site for T-DNA insertions (indicated by triangles) was mapped by PCR and DNA sequencing of the PCR products. (C) RT-PCR analysis showing levels of ROP11/ARAC10 transcripts in rop11/arac10-1, rop11/arac10-2, and rop11/arac10-3 mutant lines. ACTIN2 were used as loading control. (D) ABA sensitivity of WT and mutant seedlings. Physiologically comparable seeds of WT and rop11/arac10-2 mutant were sterilized and plated on MS medium supplemented with 0 or 2 μM ABA at 23 °C for 12 d. (E) ABA inhibition of root elongation, rop11/arac10-3 seedlings are more sensitive to inhibition of root elongation by ABA compared with WT. Data represent average ± SE of three independent experiments with 10 seedlings each. (F) rop11/arac10-3 seeds are more responsive to ABA inhibition during germination. Data represent average ± SE of three independent experiments with ~100 seeds each. (G) ROS production in guard cells of WT and fer-4 mutant with or without ABA, indicated by fluorescent dye 2′, 7′-dichlorodihydro-fluorescein diacetate.
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