Endodermal ABA Signaling Promotes Lateral Root Quiescence during Salt Stress in Arabidopsis Seedlings

Lina Duan,a,b,c Daniela Dietrich,d Chong Han Ng,b Penny Mei Yeen Chan,b,c Rishikesh Bhalerao,e Malcolm J. Bennett,d and José R. Dinneny,a,b,c,1

The endodermal tissue layer is found in the roots of vascular plants and functions as a semipermeable barrier, regulating the transport of solutes from the soil into the vascular stream. As a gateway for solutes, the endodermis may also serve as an important site for sensing and responding to useful or toxic substances in the environment. Here, we show that high salinity, an environmental stress widely impacting agricultural land, regulates growth of the seedling root system through a signaling network operating primarily in the endodermis. We report that salt stress induces an extended quiescent phase in postemergent lateral root (LRs) whereby the rate of growth is suppressed for several days before recovery begins. Quiescence is correlated with sustained abscisic acid (ABA) response in LRs and is dependent upon genes necessary for ABA biosynthesis, signaling, and transcriptional regulation. We use a tissue-specific strategy to identify the key cell layers where ABA signaling acts to regulate growth. In the endodermis, misexpression of the ABA insensitive1-1 mutant protein, which dominantly inhibits ABA signaling, leads to a substantial recovery in LR growth under salt stress conditions. Gibberellic acid signaling, which antagonizes the ABA pathway, also acts primarily in the endodermis, and we define the crosstalk between these two hormones. Our results identify the endodermis as a gateway with an ABA-dependent guard, which prevents root growth into saline environments.

INTRODUCTION

Root system architecture is a complex emergent property of the root that arises due to the growth attributes of the primary root (PR), individual lateral roots (LRs), and adventitious roots. In many Eudicot species, the PR is specified during embryogenesis and establishes the majority of the root system through the production of LRs along its length. In Arabidopsis thaliana, the patterning of LR founder cell populations occurs at regular temporal intervals and leads to the specification of founder cell populations that later develop into LR primordia (De Smet et al., 2007; Moreno-Risueno et al., 2010). Subsequent to primordia formation, the process of LR emergence requires that the primordium communicate with the outer tissue layers of the root to coordinate changes in tissue integrity with the destructive process of outgrowth (Swarup et al., 2008). Much attention has focused on these initial stages of LR development, and auxin signaling and transport are critical components at every stage of this process (Benková et al., 2003; De Smet et al., 2007; Dubrovsky et al., 2008; Laskowski et al., 2008; Moreno-Risueno et al., 2010; Overvoorde et al., 2010).

Postemergence LR development is tightly controlled by environmental stimuli, such as nutrient and water availability (López-Bucio et al., 2003; Malamy, 2005). Nitrate, for example, can inhibit LR elongation through competition with auxin transportation and by affecting mitotic activity (Little et al., 2005; Okushima et al., 2011). Localized sources of iron, by contrast, promote LR growth by triggering auxin accumulation in root apices through the regulated expression of the auxin influx carrier AUXIN 1 (AUX1) (Giehl et al., 2012). Intrinsicherdifferences in the environmental response programs of PRs and LRs have also been observed. Phosphate promotes PR growth and inhibits LR growth (Zhang and Forde, 1998; Linkohr et al., 2002), though the mechanistic basis for these differences is unknown. The water stress-associated hormone abscisic acid (ABA) is a known inhibitor of LR development and has been shown to act at the postemergence stage (Signora et al., 2001; De Smet et al, 2003). ABA has a much stronger effect on LR than PR growth, suggesting that differences in environmental responsiveness between these two root types may be due to divergent hormone signaling networks.

Recent work using cell type-specific profiling methods has demonstrated that tissue identity is critical for providing a context...
in which environmental responses are interpreted at the cellular level (Dinneny, 2010; Wee and Dinneny, 2010). Changes in salinity, nutrient content, pH, and oxygen concentration all elicit a large number of transcriptional and posttranscriptional changes that substantially differ between each cell type (Dinneny et al., 2008; Gifford et al., 2008; Mustroph et al., 2009; Iyer-Pascuzzi et al., 2011). These data have led to a sophisticated understanding of the biological pathways targeted by environmental regulation, but it is still unclear what the tissue-specific regulatory mechanisms that control these changes are.

Plant hormones are important secondary signaling molecules that mediate responses to many environmental stimuli, and recent work has shown that several of these hormones promote growth primarily by acting in specific tissue layers. Brassinosteroid signaling acts in the epidermis to promote shoot growth (Savaldi-Goldstein et al., 2007) and regulate root meristem size (González-García et al., 2011; Hacham et al., 2011), while gibberellic acid (GA) signaling leads to the degradation of DELLA proteins, which negatively regulate growth primarily in the endodermal tissue layer (Ubeda-Tomás et al., 2008, 2009). These results suggest a possible mechanism by which environmental stimuli elicit localized transcriptional changes through the action of hormones that trigger tissue-specific signaling cascades. Hormones associated with stress responses also regulate growth of the plant (Spollen et al., 2000; Sharp and LeNoble, 2002; Achard et al., 2006), but it is not known whether the growth-suppressing activities of these hormones also act in specific cell layers.

Here, we dissect the tissue-specific function of ABA signaling in regulating root growth to reveal the mechanism by which salt controls root system architecture in the seedling. High salinity is an important and prevalent agricultural contaminant that affects yield (Flowers et al., 1997), and previous studies have shown that elevated salt levels can inhibit both PR and LR growth (Burssens et al., 2000; He et al., 2005). We reveal that PR and postemergence-stage LR growth show divergent temporal dynamics during salt stress, which is due to LR-specific ABA signaling. We identify the endodermis as the key tissue layer required for ABA-mediated growth repression of the LR during salt stress and dissect points of crosstalk with the GA pathway in this process. These data provide insight into a previously uncharacterized function of the endodermis as a sentinel cell layer, which guards against growth into saline environments.

RESULTS

The Growth of Primary and LRs Is Differentially Affected by Salt Stress

Root system architecture is determined by three fundamental parameters: (1) the number of LR branch points specified along the PR, (2) the initiation and emergence rates for these LR primordia, and (3) the growth rate of each root type, the PR, and postemergence LRs. Adventitious roots, which develop at the root-hypocotyl junction, are considered a special type of LR and are not studied here. To quantify the effect of salt stress on these three parameters, Arabidopsis seedlings were grown on standard media for 6 d postgermination (dpg) and then transferred to standard conditions or media supplemented with sodium chloride (NaCl) and grown for an additional 4 d post-transfer (dpt). This treatment established two regions along the PR: region A (for above transfer point), in which the PR had grown before the transfer to salt conditions; and region B (for below transfer point), in which the PR had grown after the transfer (Figure 1A). Based on recent work, the patterning of LRs is determined close to the root tip, and initiation of LR primordia from founder cells occurs later in development (De Smet et al., 2007; Moreno-Risueno et al., 2010; Overvoorde et al., 2010). Thus, LR patterning in region A is expected to have occurred before treatment of roots with high salinity. Consistent with this hypothesis, we found that high salinity had little effect on the total number of LRs initiated in region A (Figure 1B) (Malamy and Benfey, 1997). Furthermore, we did not observe any strong effects on the distribution of developmental stages for preemergent LR primordia (Figure 1B). In region B, the number of emerged LRs was clearly suppressed and the total number of LRs was lower, suggesting that LR initiation or founder cell specification is inhibited by salt treatment (Figure 1C). Further investigation of the mechanisms affecting the patterning of LRs in region B is the subject of another study and is not discussed further here.

While the total number of LRs initiated and the developmental progression of preemergent LR primordia was not affected in region A, measurement of the total or average postemergence LR length revealed that salt had a strong inhibitory effect on growth at this stage of development (Figure 1D; see Supplemental Figure 1A online). Interestingly, PR growth was much less sensitive to salt treatment compared with LR growth (49.6% compared with 84.3% reduction, respectively, at 100 mM NaCl) (Figure 1D). These differences were observed in multiple accessions of Arabidopsis, despite differences in overall salt sensitivity (see Supplemental Figures 1B and 1C online).

Developmental Age Affects the Sensitivity of the LR, but Not the PR, to Salt Stress

Our observed differences in the response of LRs and PRs to salt stress could have two potential causes, which are not mutually exclusive. First, in our experiments, we measured the response of PRs 6 dpd, but LR growth was measured after emergence from the PR. Thus, differences in salt response could be a consequence of the developmental age of the specific root type. A second possibility is that LRs and PRs differ in their salt responses due to an inherent difference in the signaling properties of the organ. Testing the first possibility is complicated by the fact that PRs and LRs are defined by fundamentally different times of origin (embryonic versus postembryonic) and positions of formation (de novo formation versus endogenous development within the PR). Thus, there is no stage of PR development that is equivalent to a postemergent LR. Nevertheless, we can ask the question of whether the salt response profile changes with developmental age for the PR or LR.

We transferred seedlings grown for between 2 to 8 dpd to media supplemented with 100 mM NaCl and found that the sensitivity of the PR did not significantly differ between these
Figure 1. The Growth of PRs and LRs Is Differentially Affected by Salt Stress.

(A) Morphology of the Col-0 root system grown on standard media for 6 dpg then transferred to standard, 100 mM NaCl, or 140 mM NaCl media conditions for 4 d. Arrowheads mark the position of the root tip at the time of transfer from standard to treatment conditions.

(B) and (C) Quantitation of preemergent and late stages of LR development in root region A (B) and region B (C) after transfer to standard or salt stress conditions (n > 10 seedlings).

(D) Average PR and LR length for seedlings transferred to 100 or 140 mM NaCl. Measurements are expressed as a percentage of the length under standard conditions (n > 20 seedlings).

(E) and (F) Suppression of LR (E) and PR (F) growth under 100 mM NaCl conditions measured as a percentage of reduction relative to standard conditions. Seedlings were transferred to salt at different days postgermination (dpg) and grown for 4 dpt (n > 8 seedlings).

(G) Number of emerged LRs present prior to salt treatment.

Error bars indicate SE. Asterisks mark significant changes in salt response based on a Student’s t test, P value < 0.05. [See online article for color version of this figure.]
seedlings (Figure 1F). Measurement of growth rates in roots that had just emerged from the seed coat also did not reveal any obvious hypersensitivity to salt stress at this earliest developmental stage (see Supplemental Figures 1E and 1F online). Thus, PRs consistently show relative resistance to salt treatment throughout initial seedling development. LRs, by contrast, showed a decrease in sensitivity to salt when seedlings were transferred at 8 dpg compared with 6 or 7 dpg (Figure 1E). This difference is correlated with the prevalence of postemergence stage LRs at the time of transfer (Figure 1G) and indicates that only newly emerged LRs are hypersensitive to salt treatment.

**Postemergence Growth Quiescence Induced in LRs by Salt Stress**

To understand how salt stress severely inhibits LR growth and why a specific LR developmental stage may be required, live imaging was used to monitor the temporal dynamics of growth regulation. Growth rates of postemergence LRs were quantified for 7 dpt on standard or saline media. Interestingly, growth was not stably suppressed throughout the salt response, but instead, LRs were maintained in a quiescent stage for several days (Figures 2A and 2B; see Supplemental Movie 1 online). Similar quiescent LRs were observed under standard conditions, however, much less frequently. After several days of quiescence, LRs resumed growth at rates similar to roots grown under standard conditions (Figure 2B). This response dynamic differed substantially for the PR, which showed an immediate recovery of growth after being transferred to salt stress conditions (see Supplemental Figure 1D online). We examined the expression of the cell cycle marker CYCB1;1::GUS (for β-glucuronidase) and found that expression was greatly reduced in quiescent-stage LR primordia (Figures 2C and 2D) (de Almeida Engler et al., 1999). No changes in the structure of the stem cell niche or radial organization, which could affect growth rates, were observed in quiescent stage LRs (see Supplemental Figure 2 online). Together, these results indicate that salt induces a period of quiescence in LRs, which may be, in part, a consequence of suppressed cell cycle activity.

**ABA Signaling Is Necessary for LR Growth Suppression during Salt Stress**

ABA acts as an important secondary signaling molecule during abiotic stress (Sharp and LeNoble, 2002; Zhu, 2002; De Smet et al., 2003; Hubbard et al., 2010). Under high salinity, the production of ABA is induced and its signaling pathways are necessary for salt tolerance (Achard et al., 2006). We first treated roots with varying concentrations of ABA to determine if this hormone treatment had an effect similar to high salinity. As previously shown (De Smet et al., 2003), ABA treatment affected LR growth more severely than PR growth (63.0% compared with 15.6% reduction, respectively, after treatment with 1 μM ABA) (Figure 3A). To determine whether LRs are generally more sensitive to the effects of growth-suppressing hormones, we treated seedlings with 1-aminocyclopropane-1-carboxylic acid (ACC), a biosynthetic precursor to ethylene, which is also induced under salt stress (Achard et al., 2006). Interestingly, ACC treatment caused the opposite trend in growth, having a much stronger effect on PR than LR growth (72.3% compared with 46.8% reduction, respectively, after treatment with 1 μM ACC) (Figure 3B). Other ecotypes also showed similar responses to NaCl, ABA, and ACC (see Supplemental Figure 3 online). Together, these results indicate that LRs and PRs are distinguished based on both their hormone and stress response profiles. The similarity in the response of the root system to NaCl and ABA suggests that this hormone may be most critical for eliciting the suppression of growth in LRs.

To test whether endogenous hormone signaling is responsible for root system architecture changes in the seedling during salt stress, we measured the effect of salt on average LR length for various mutants disrupted in the ABA pathway. This analysis identified mutations affecting ABA biosynthesis (ABA DEFICIENT2 alleles [aba2-1 or aba2-sail]), signal transduction (aba insensitive1-1 [abi1-1]), and transcriptional regulation (aba insensitive4-1 fusca3-3) that mediate salt-dependent LR growth suppression (Figure 3C; see Supplemental Figure 4A online). Importantly, none of these mutations significantly affected PR growth under these conditions, which suggests that the ABA response may be restricted to the LR during salt stress (see Supplemental Figure 4B online).

We also examined the role of ethylene signaling in LR growth suppression by salt stress. The ethylene insensitive 2-44 (ein2-44) mutant, which strongly disrupts ethylene signaling (Ghassemian et al., 2000), showed moderately higher sensitivity to salt stress, indicating that ethylene signaling may promote LR growth under these conditions (see Supplemental Figures 4C and 4D online). The like aux1 3 (jax3) mutant, which affects auxin transport and is necessary for LR emergence (Swarup et al., 2008), did not significantly impact the salt response of postemergent LR.

**The Endodermis Is the Target Cell Layer for ABA-Dependent Salt Stress Regulation of LR Growth**

Several plant hormones have been shown to regulate growth through tissue-specific signaling (Swarup et al., 2005; Savaldi-Goldstein et al., 2007; Ubeda-Tomás et al., 2008, 2009; González-García et al., 2011; Hacham et al., 2011). To determine if ABA operates by a similar principle, we used the GAL4-VP16/UAS enhancer trap system (Kiegle et al., 2000; Ubeda-Tomás et al., 2008) to drive tissue-specific expression of abi1-1, which dominantly suppresses ABA signaling. ABI1 encodes a PP2C-type protein phosphatase and ABA coreceptor, which dephosphorylates SnRK-type kinases to inhibit ABA-dependent signal transduction (Meyer et al., 1994; Fuji et al., 2009; Ma et al., 2009; Park et al., 2009). The abi1-1 mutation renders the protein constitutively active, and overexpression of the mutant coding sequence can be used to dominantly suppress ABA signaling (Leung et al., 1997). We verified that the enhancer-trap lines used in our study drive strong expression in the various cell layers of PRs and postemergence LRs using the associated UAS:ERGFP reporter (Figures 4A to 4C; see Supplemental Figure 5 online).

We first determined whether tissue-specific expression of abi1-1 could induce resistance to exogenous application of 10 μM ABA, which affects PR and LR growth to a similar extent as
100 mM NaCl. F1 seedlings generated by a cross between a UAS:abi1-1 transgenic plant and the various enhancer trap lines conferred reduced sensitivity of LR growth to ABA treatment when expression was driven in the ground tissue (J0571), endodermis and pericycle (Q2500), or stele (Q0990) (Figures 4D and 4F). This pattern of activity partially overlaps with the cell layers observed to be important for ABA signaling in the PR, where enhancer traps driving abi1-1 expression in the epidermis and cortex (J2812), ground tissue layers (J0571), and endodermis and pericycle (Q2500) had a significant effect, but stele expression (Q0990) did not (Figures 4D and 4G). These results indicate that abi1-1-dependent ABA signaling acts to regulate growth in a spatially restricted fashion and that the ground tissue layers are common sites for such regulation in the PR and LR.
We next asked whether salt stress–dependent ABA signaling acts through similar tissue layers as exogenous ABA treatment. Here, abi1-1 was most effective at rescuing LR growth when expressed in the endodermal and pericycle tissue layers of LRs (Q2500) or in the ground tissue as a whole (J0571) and less so when expressed in the epidermis and cortex (J2812) (Figures 4E and 4F). Use of a pericycle-specific enhancer trap (J0121) to drive abi1-1 expression had a very mild effect on LR length during ABA or salt treatment (see Supplemental Figure 6 online), which suggested that endodermal expression of abi1-1 in the Q2500 line is the primary cause of the reduced sensitivity to ABA or salt. Expression of abi1-1 in the various tissue layers had little effect on PR growth under salt stress conditions (Figures 4E and 4G), consistent with the LR specificity of the ABA response we inferred from our genetic analysis. Our media contains 1% Suc, but we found that this did not significantly affect the ABA-dependent regulation of LR growth during salt stress treatment (see Supplemental Figure 7 online). To confirm the function of ABA signaling in the ground tissue during salt stress, an endodermis-specific gene promoter, ProSCARECROW (ProSCR) (Figure 4H) (Di Laurenzio et al., 1996), and a cortex-specific promoter, ProCORTEX (ProCOR) (Lee et al., 2006), were used to drive the expression of an abi1-1:RFP (for red fluorescent protein) translational fusion. Two independent ProSCR:abi1-1:RFP transgenic lines showed rescue of LR growth under both ABA and salt treatment (see Supplemental Figure 8A online). Fluorescence imaging of these lines revealed weak but endodermal-specific localization (see Supplemental Figure 8C online). These results were further confirmed by transforming plants with a ProSCR:abi1-1:RFP transgene lacking any fluorescent tag (Figure 4I). The ProCOR:abi1-1:RFP lines also showed a rescue in LR growth with ABA or salt treatment (see Supplemental Figure 8B online). However, we found that the abi1-1:RFP fusion protein showed cortex and endodermal localization, suggesting that the promoter used to

**Figure 3.** LR Growth Is Hypersensitive to ABA Treatment and Mutants Disrupting the ABA Signaling Pathway Affect Salt-Mediated Suppression of LR Growth. 

(A) and (B) Relative PR and LR length quantified 4 dpt to media supplemented with 1 or 10 μM ABA (A) or 1 or 10 μM (B) ACC (n > 20 seedlings). The average length of all LRs per seedling was measured and the percentage of difference between salt stress and standard conditions was calculated and shown in the graph. Mutations in the Ler background are shown as white bars, Ws background as light-gray bars, and Col-0 background as dark-gray bars (n > 15 seedlings). TF, transcription factor.

All seedlings grown for 6 dpq and average LR length per seedling measured 4 dpt. Error bars indicate SE. Asterisks mark significant changes in salt response based on a Student’s t test, P value < 0.05, with Bonferroni correction.

[See online article for color version of this figure.]
Figure 4. The Endodermis Is an Important Site for ABA Signaling and Growth Regulation during Salt Stress.
drive transgene expression may not have cortex-specific activity or that the abi1-1 protein has limited ability to move between tissue layers (see Supplemental Figure 8D online). Together, our data indicate that ABA signaling acts primarily in the endodermis to limit LR growth during salt stress conditions, although we cannot exclude the possibility that the cortex may also have some importance in this process. These data also show that the pattern of activity for the ABA pathway is environmentally dependent, with salt acting through a more restricted domain of the root than with exogenous ABA treatment.

Our results suggest a model in which the endodermis is the source of a growth-promoting activity in the LR that is suppressed by salt stress through the execution of an ABA-dependent response in this cell type. To explore this model further, we next asked how loss of the endodermis might affect the salt responsiveness of the LR. SHORTROOT (SHR) is a GRAS family transcription factor necessary to promote periclinal division of the cortex/endodermal stem cell and the subsequent specification of endodermal identity in one of the two daughter cells (Helariutta et al., 2000; Cui et al., 2007). The shr mutant lacks an endodermis in both PRs and LRs and affects the morphogenesis of LRs (Lucas et al., 2011). To avoid some of the nonspecific effects of the shr mutant, we used a dexamethasone (DEX)-inducible SHRpro: SHR-GR, shr-2, ProSCR:erGFP line, which is able to fully rescue the mutant phenotype (Levesque et al., 2006; Sozzani et al., 2010). The ProSCR:erGFP reporter is directly regulated by SHR and is a marker for endodermal identity. Seedlings were initially grown on media supplemented with 1 μM DEX for 6 dpg and then transferred to media with or without 100 mM NaCl and/or 1 μM DEX (Figure 4J). We analyzed the expression of the SCR reporter and observed that an endodermis only formed in LRs developing on DEX-supplemented media (Figure 4K). Interestingly, we observed that LRs lacking DEX treatment grew more slowly under standard conditions and showed little response to salt, whereas DEX-treated LRs responded more similarly to wild-type plants (Figure 4L). These data are consistent with our model and support the hypothesis that endodermal signaling in the LR itself is key to the regulation of postemergence growth.

**Sustained ABA Signaling Is Associated with LR Quiescence during Salt Stress**

To determine if the quiescence of LRs is correlated with the timing of ABA signaling, we examined the activity of an ABA-responsive reporter driven by the RESPONsive TO ABA18 (RAB18) promoter, ProRAB18:GFP (Kim et al., 2011), in the root system after salt treatment. Under standard conditions, ProRAB18:GFP was expressed at very low levels in both the PR and LRs (Figures 5A, 5B, and 5D). During salt stress, ProRAB18:GFP was weakly expressed in the LR (Figure 5A). During LR development, reporter expression was also weak just after emergence but became highly induced ~2 d later (Figures 5A, 5C, and 5D). Reporter expression in postemergence LRs was maintained at very high levels for several days until the time at which growth resumed, when it decreased (Figures 5C and 5D). The substantially higher level of ABA signaling in the LR, compared with the PR, explains why the suppression of ABA signaling by our various perturbations had an LR-specific effect on growth. In response to ABA treatment, ProRAB18:GFP was also induced after LR emergence as it entered into a quiescent phase; however, expression was maintained for a longer time period (see Supplemental Figures 9B and 9C online). We analyzed the induction of ABA signaling in older root systems and found that LRs, which escaped growth suppression by salt stress, also lacked induction of the ProRAB18:GFP reporter.

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**Figure 4.** (continued).

(A) A false-colored drawing of a LR illustrating the different cell types present. COL, columella root cap; COR, cortex; END, endodermis; EPI, epidermis; LRC, lateral root cap; PER, pericycle; STE, stele without pericycle.

(B) Confocal images of the various GAL4-VP16/UAS enhancer trap lines used in this study showing the activity of the associated UAS:erGFP reporter in postemergent LRs.

(C) Heat map summarizing the relative expression level of the various enhancer trap lines.

(D) and (E) Images of root systems for the control genotype, Q25000>>abi1-1, and J0571>>abi1-1 transactivation lines 4 dpt to standard conditions, 10 μM ABA treatment (D), or 100 mM NaCl conditions (E). Red arrowheads mark the position of the root tip at the time of transfer.

(F) and (G) The average length of all LRs (F) or PRs (G) per seedling was measured and the percentage of difference between the treatment and standard conditions was calculated and shown in the graph. Growth suppression by 4 dpt to 100 mM NaCl conditions or 10 μM ABA conditions is shown (n > 20 seedlings).

(H) Confocal image of ProSCR:erGFP expression in a postemergent LR is localized to the endodermis.

(I) Percentage of difference in LR length of the ProSCR:abi1-1 transgenic line 4 dpt to 100 mM NaCl conditions (n > 18 seedlings).

(J) Diagram illustrating our experimental design for studying the effects of genetically ablating the endodermal tissue layer specifically in LRs. The ProSHR:SHR-GR, shr-2, ProSCR:erGFP line was grown on media supplemented with 1 μM DEX for 6 dpg then transferred to media with or without DEX. In the diagram, green portions of the root have an endodermis and ProSCR:erGFP reporter activity.

(K) Confocal images of the ProSHR:SHR-GR, shr-2, ProSCR:erGFP line in LRs, which developed after transfer to media with (left panel) or without (right panel) DEX. With DEX treatment, GFP expression driven by the ProSCR:erGFP marks the endodermis, while only a mutant cell layer (M) exists without it. c, cortex; e, endodermis.

(L) Quantitation of LR length in Col-0 and the ProSHR:SHR-GR, shr-2, ProSCR:erGFP transgenic line 4 dpt to standard conditions or 100 mM NaCl-supplemented media with or without 1 μM DEX (n > 17 seedlings). Error bars indicate se. Red asterisks mark significant differences based on a two-way analysis of variance and Student’s t test, P value < 0.05. Bars = 50 μm.
while LRs entering quiescence showed strong expression (see Supplemental Figure 9D online). Together, these data show that salt-induced growth quiescence is tightly correlated with the activation of ABA signaling.

To confirm and extend our results, we used high-throughput quantitative RT-PCR (qRT-PCR) to analyze salt-regulated changes in expression for a collection of ABA-responsive genes. Seedlings were grown on standard media for 6 dpg and then transferred to standard conditions or media supplemented with NaCl and grown for an additional 3 dpt. RNA was isolated from either region A or B of the root system from Columbia-0 (Col-0), aba2-sail, the Q2500>>abi1-1 transactivation line, or the associated control. Known ABA-inducible genes (RESPONSIVE TO DESICCATION 29A [RD29A] and ABI2) and others showed strong transcriptional activation by salt stress predominantly in region A, which contains postemergent LRs, while expression was much reduced in region B (Figure 5E). Induction of RAB18 expression by salt was alleviated in the aba2-sail mutant and also strongly reduced in the Q2500>>abi1-1 transactivation line (Figure 5E). The tissue-specific expression pattern of the ProRAB18:GFP reporter is not informative since it uses a non-localized GFP reporter protein, which may translocate between cell layers. However, based on cell type–specific transcriptional profiling data (Winter et al., 2007; Dinneny et al., 2008), RAB18 is most strongly activated in the endodermis and moderately induced in the cortex (see Supplemental Figure 9A online). However, an important caveat applies to the analysis of such data, since it was generated from PRs treated with 140 mM NaCl.

Figure 5. ABA Signaling Is Selectively Induced in LRs and Correlates with Growth Quiescence.

(A) Seedlings with the ProRAB18:GFP reporter were transferred to standard or 100 mM NaCl media and PRs or LRs imaged by confocal microscopy. Bars = 50 μm.

(B) and (C) Quantification of LR length and ProRAB18:GFP fluorescence intensity in seedlings transferred to standard conditions for 1 to 3 d post-transfer (dpt) (B) or 100 mM NaCl for 1 to 5 dpt (C) (n = 15 seedlings). Error bars indicate se. a.u., arbitrary units.

(D) Time-lapse images of ProRAB18:GFP expression in LRs after transfer to standard or 100 mM NaCl conditions.

(E) Gene expression analysis of ABA-dependent salt responsive genes using high-throughput qRT-PCR. Expression was measured 3 dpt to standard or 100 mM NaCl media from root region A and region B, separately. Expression measured in Col-0, aba2-SAIL, enhancer trap control, and the Q2500>>abi1-1 transactivation line. Heat map shows normalized gene expression values averaged from two biological replicates and three technical replicates. Gene expression profiles were organized by hierarchical clustering. Bold highlighted genes showed dependency on endodermal ABA signaling.
Analysis of other salt-responsive genes identified several that were also strongly dependent upon ABA signaling in the endodermis, including HOMEOBOX12 (HB12), an ABA-induced gene necessary for growth suppression during water stress (Figure 5E) (Olsson et al., 2004; Son et al., 2010). These data show that ABA signaling is enriched in the region of the root-containing LRs and that endodermal ABA signaling is necessary for many of these responses.

Live-imaging analysis of the Q2500>>abi1-1 transactivation line revealed that postemergent LRs showed an immediate activation of growth under salt conditions (Figure 6; see Supplemental Movie 2 online), which confirms that ABA signaling in this sub-domain of the LR is critical for maintaining quiescence. Interestingly, LR growth was not sustained in Q2500>>abi1-1, and after a few days, growth became arrested and the roots senesced (Figures 6B and 6C; 66.7% of all LRs). In control plants, LR growth was sustainable after recovery from quiescence throughout the duration of the imaging session (Figure 6A).

**Figure 6.** Suppression of Endodermal ABA Signaling Leads to a Loss of Salt-Induced Quiescence and Unsustainable Growth Recovery.  
(A) and (B) Time-lapse images of enhancer trap control (A) or Q2500>>abi1-1 transactivation seedlings (B) after transfer to 100 mM NaCl conditions. Red arrowheads mark quiescent LRs. Red asterisks mark LRs that showed unsustainable growth recovery.  
(C) Heat map showing the growth profiles of individual LRs quantified from time-lapse imaging data. Each pixel in the heat map represents the total growth over 2 h. Data quantified from enhancer trap control seedlings (n = 12 LRs) or Q2500>>abi1-1 transactivation seedlings (n = 15 LRs) transferred to 100 mM NaCl conditions and imaged for 7 d. Pie charts to the right of the heat maps quantify the proportion of LRs that are quiescent for different lengths of time. Red asterisks mark LRs that show unsustainable growth recovery, 10 out of 15.

**Nutrients Enhance the ABA-Dependent Effect of NaCl on LR Growth**

Differences in the response of PRs and LRs to changes in nutrient levels have previously been described, and responses to high levels of nitrate are known to be ABA dependent (Signora et al., 2001). Thus, we asked whether nutrient concentration in our media might also modulate the response to salt stress. In the PR, we found that increasing concentrations of Murashige and Skoog (MS) nutrient salts reduced the effect of NaCl on growth (see Supplemental Figure 10A online). This effect is consistent with previous studies showing that increasing levels of certain nutrients, such as potassium (Zhu et al., 1998), can provide protection against the ionic stress caused by sodium. Interestingly, we saw the opposite trend for LRs, which became more sensitive to NaCl as the concentration of MS nutrients increased (see Supplemental Figure 10B online). The suppression of LR growth at full-strength MS is strongly affected in the Q2500>>abi1-1 transactivation line, but this transgene had little
impact in media containing quarter-strength MS. These data provide evidence that nutrients can alter the decision-making process of the root, as it relates to salt response, and that these effects are dependent on root type— and tissue-specific ABA signaling.

**GA Signaling Acts in Parallel and Antagonistically with ABA in LRs**

Our data show that postemergence LRs undergo a period of quiescence during salt stress, which is induced by ABA. Another phase of plant development characterized by an ABA-dependent period of growth arrest is seed dormancy (Finkelstein et al., 2008). Here, GA acts antagonistically to promote germination. To test whether GA may also suppress LR quiescence, we treated roots with varying concentrations of GA during salt stress. With GA treatment, the suppression of LR growth by salt was significantly reduced (Figure 7A). Time-lapse imaging of LRs under GA-supplemented salt stress conditions showed that the timing of growth recovery was hastened by hormone treatment (Figure 7B). PR growth was less sensitive to GA treatment, again highlighting important differences in hormone signaling between the two root types (see Supplemental Figure 11A online).

To understand what function endogenous GA levels might have in regulating growth during salt stress, we studied the effects of the GA biosynthesis inhibitor paclobutrazol (PAC). Under standard conditions, PAC treatment suppressed LR growth, indicating that GA biosynthesis is important under this condition (Figure 7C). The abi1-1 mutation partially suppressed this effect, indicating that PAC treatment affects LR growth, in part, by elevating ABA signaling. PAC was not able to fully suppress the abi1-1 phenotype under salt stress conditions, which suggests that abi1-1 also acts to regulate growth independent of changes...
DISCUSSION

Our study reveals an important interplay between developmental and environmental pathways that ultimately transforms the root system architecture of the seedling to suit a new environmental condition. We show that LRs and PRs have intrinsically different response programs to salinity and the associated hormones elicited downstream with ABA being the critical effector of LR growth. The site of action for this hormone is primarily localized to the endodermis, thus highlighting how cell identity provides an important signaling context for salt stress responses. Finally, the ultimate outcome of the stress signaling network outlined here is to dynamically control the growth of LRs, which will cause a transformation in the emergent properties of the seedling root system due to changes in the ratio between LR and PR growth. We hypothesize that these differences are likely related to the unique functions of each root type in a typical Eudicot taproot system; the PR functions to extend the root system into deeper terrain, while LRs exploit the resources identified at specific depths. We speculate that LR-specific activation of ABA signaling will shift the balance of root growth toward soil exploration, away from resource acquisition.

The Root System Is Composed of Root Types with Distinct Salt Stress and Hormone Signaling Properties

The Arabidopsis root system is composed of primary, lateral, and adventitious roots. LRs are differentiated from the PR due to their postembryonic origin from the pericycle cell layer of the PR (Malamy and Benfey, 1997). Other differences in growth have been described; however, the mechanistic basis controlling such distinctions is unclear. The response to gravity, for example, differs between the PR and LR (Kiss et al., 2002; Guyomarc’h et al., 2012). The PR is able to immediately respond to the gravity vector and, in flax, is responsive even before germination (Ma and Hasenstein, 2006). On the other hand, newly emerged LRs in Arabidopsis show little sign of gravitropism initially and become more gravitropic over time. These differences have important effects on root system architecture and enable larger domains of the soil environment to be explored. Work by Guyomarc’h et al. revealed that the expression pattern of PIN transporters changes extensively through postemergence LR development, which suggests a potential mechanism to control the observed differences in gravitropism (Guyomarc’h et al., 2012).

In our study, we show that hormone and salt stress signaling differ significantly between LRs and PRs. Salt stress activates high levels of ABA signaling exclusively in LRs, as demonstrated by the LR-specific expression of the ProRAB18:GFP reporter. These data suggest that ABA signaling may have little activity in the PR at the concentrations of salt used here (100 mM NaCl). Indeed, all mutants affecting ABA signaling primarily rescue growth of the LR, while the PR is unaffected or grows slower. These data suggest that salt may induce LR-specific ABA biosynthesis or that the LR may be hypersensitive to ABA. Exogenous application of ABA clearly shows that the LR exhibits significantly higher sensitivity to ABA than the PR. Thus, we propose that LRs and PRs diverge in their salt response due to differences downstream of ABA biosynthesis. Root type-specific
In the shoot, well-characterized sets of transcription factors determine the identities of different meristem types (Bowman et al., 1993; Wellmer and Riechmann, 2010). The vegetative meristem of Arabidopsis transitions to an inflorescence meristem, and lateral branch meristems are specified as floral meristems. While currently uncharacterized, the various meristem types in the shoot may also have unique hormone and stress signaling properties that allow the plant to regulate context appropriate environmental responses. We speculate that post-emergent LRs may have a unique identity, in the developmental sense, that specifies the suite of genes controlling hormone, stress signaling, and gravity response. In species such as maize (Zea mays), where additional root types like seminal, crown, and brace roots occur, mechanisms to determine the identity of each root type may exist (Hochholdinger and Tuberosa, 2009). To understand how plants tailor stress signaling for particular environmental and field conditions, it will be important to understand the mechanisms that regulate the identity of each root type and the associated stress signaling networks.

**Tissue-Specific ABA Signaling Regulates Root Growth**

In our study, we examined the function of tissue-specific ABA signaling in the context of exogenous ABA treatment as well as an environmental stress that acts through the ABA pathway to regulate growth. These studies focused on LR growth but also examined the PR. This comparative approach enabled us to distinguish where ABA is able to regulate growth from where ABA actually acts during a physiologically relevant execution of the pathway. Interestingly, we find that the endodermis is not the only cell type where ABA signaling can regulate growth. With exogenous ABA treatment, the stele is also important. In other cell types, ABA signaling may control the same downstream pathways to affect growth. Alternatively, ABA signaling may regulate growth by distinct developmental mechanisms in each cell type. We find that the salt-induced expression of some ABA-regulated genes (e.g., RD29A and ABI2) is not strictly dependent on endodermal ABA signaling (Figure 5E). Consistent with these results, cell type–specific analysis shows that such genes are induced by salt stress in all root tissue layers (see Supplemental Figure 9A online). Thus, we propose that ABA signaling may be active in multiple tissue layers of the root during salt stress but that certain functions, such as growth regulation, are controlled by tissue-specific ABA signaling pathways.

Several recent studies have also highlighted the role of developmental pathways in the control of stress response. Iyer-Pascuzzi et al. (2011) have shown that the scr mutant, which disrupts the asymmetric division of the cortex/endodermal stem cell, has defects in germination upon ABA treatment (Cui et al., 2012). This work and another study by Cui et al. (2012) have shown that SCR can directly regulate the expression of several transcription factors controlling ABA response, including HB12, which we find here to be regulated by ABA signaling in the endodermis. Work presented here, along with these other studies, have now clearly demonstrated that developmental regulation and tissue-specific signaling are important for generating a stress response program and for salt tolerance (Møller et al., 2009), a model we proposed in our original work studying tissue-specific responses to abiotic stress (Dinneny et al., 2006).

**The Endodermis as a Gateway with an ABA-Dependent Guard**

The endodermis is an evolutionarily conserved cell type found in all flowering plants and acts as a semipermeable barrier due to the presence of a specialized cell wall modification termed the Casparian strip. Recent work has shown that lignin deposition in the Casparian strip primarily contributes to its function as a barrier to the apoplastic diffusion of solutes into the vascular stream (Roppolo et al., 2011; Naseer et al., 2012). Consequently, transport of solutes must occur by entering the symplast of the outer tissue layers or the endodermis itself. Thus, this cell type can be considered a gateway for entry into the vascular stream. Indeed, several transporters are expressed in this tissue layer and are localized to a specific side of the cell, consistent with their function in nutrient uptake (Ma et al., 2006, 2007; Alasimone et al., 2010; Takano et al., 2010; Sasaki et al., 2012).

For salinity stress, in particular, the extent to which a plant absorbs sodium into the vascular stream and transports it to the shoot is negatively correlated with overall stress tolerance (Munns, 2002; Møller and Tester, 2007). The endodermis may cause a bottleneck in the diffusion of sodium ions through the root, and as a consequence, ABA signaling may be most strongly activated there and growth arrested. This hypothesis is consistent with data from Møller et al. (2009) showing that, after salt treatment, sodium ions accumulate in the outer tissue layers of the root while the inner stele has lower levels. Use of a Casparian strip marker, CASPARIAN STRIP MEMBRANE DOMAIN PROTEIN 1:GFP (Roppolo et al., 2011), confirms that quiescent stage LRs form a well-developed Casparian strip under salt stress, while under standard conditions, LRs show less marker expression (see Supplemental Figure 13 online). These data are consistent with previous findings showing that salt can induce thickening of the Casparian strip in maize (Kararaha et al., 2004). Endodermal cells may also be particularly sensitive to the presence of sodium ions and activate ABA signaling at a lower threshold than other cell types. As a consequence of either mechanism, ABA signaling would act as a guard at the endodermal gate, ensuring that growth of the LR does not proceed into high saline environments (Figure 8). The quiescent phase of growth induced by ABA appears to be necessary for acclimation and ultimate growth recovery, as plants suppressed in endodermal ABA signaling exhibit unsustainable growth. Thus, other processes besides growth regulation, such as the control of ion homeostasis, may be downstream of endodermal ABA signaling and enable long-term growth and acclimation.

**METHODS**

**Plant Materials**

Arabidopsis thaliana ecotypes Col-0, Landsberg erecta (Ler), and Wassilewskija (Ws) were used in this study. The abi1-1 (Armstrong et al., 1996), abi2-1 (Luong et al., 1997), abi3-1 (Nambara et al., 1995), and det3 quadruple...
plants (Tyler et al., 2004), and the ProRGA:GFP.RGA (Achard et al., 2006) and ProSCR.gai-1::GFP (Ubeda-Tomás et al., 2008) transgenes are in the Ler background. The abid1-1 (Finkelstein et al., 1998), aba2-1 (Léon-Kloosterziel et al., 1996), aba2-sail (SAIL_407_E12), nced2 (Toh et al., 2008), nced3 (Ruggiero et al., 2004), nced5 (Toh et al., 2008), nced9 (Toh et al., 2008), fis3-3 (Keith et al., 1994), era1-1 (Cutler et al., 1996), ein2-44 (Ghassemian et al., 2000), ein3 ein4 (An et al., 2010), iax3 (Swarup et al., 2008), UAS:abi1-1, ProRAB18:GFP (Kim et al., 2011), CYCB1:1:GUS (Colón-Carmona et al., 1999), ProSHR:SHR:GR, shr-2, ProSCR:GFP (Levesque et al., 2006), and CASP1:GFP (Roppolo et al., 2011) transgenes are in the Col-0 background. The abid5-1 (Finkelstein, 1994) mutation is in the Ws background. GAL4-VP16:UAS enhancer trap lines J3411, J0951, J3812, J0571, Q2500, Q6990, and J0121 (Kiegle et al., 2000; Ubeda-Tomás et al., 2008) are in the C24 background.

### Plant Growth Conditions

Seeds were surface sterilized by washing in a 95% ethanol solution for 5 min followed by a 5-min wash in a 20% bleach/0.1% Tween 20 solution. Seeds were then rinsed in sterile deionized water four times and stored in water for 2 d at 4°C. Sterilized seeds were grown on sterile 1% agar media containing 1× MS nutrients (MS01-50LT; Caisson), 1% Suc, and 0.5× MES, adjusted to pH 5.7 with KOH (termed “standard media”). Seedlings were grown for 6 d before transfer to standard media supplemented with NaCl or other chemicals for 3 to 7 d. Supplements include NaCl (Sigma-Aldrich), ABA (Sigma-Aldrich), ACC (Sigma-Aldrich), GA (Sigma-Aldrich), PAC (Sigma-Aldrich), and DEX (Sigma-Aldrich). The position of the root tip was marked at the time of transfer to distinguish the two regions of the root, A and B. Growth of seedlings was performed in a Percival CU41L4 incubator at a constant temperature of 22°C with long-day lighting conditions (16 h light and 8 h dark). Plates were partly sealed with Parafil (Alcan Packaging) on three sides, while the top of the plate was sealed with micropore tape (3M) to allow for gas exchange. This was important for allowing a sufficient number of LRs to grow for observation.

### Transgenic Construction

The UAS:abi1-1 construct was generated by mobilizing the abi1-1 cDNA (with N-terminal triple myc tag) in plasmid pDONR207 (gift from J. Leung) into the plasmid pUAS-KWG (Karimi et al., 2002) between the UAS and NOS terminator sequences using a Gateway (Invitrogen) cloning approach. Transgenic plants were generated by a standard floral dip method (Clough and Bent, 1998).

To generate the ProSCR:abi1-1::RFP and ProCOR:abi1-1::RFP constructs, the primers 5′-caccATGAGGAAATATCTCGGCCG-3′ and 5′-GCCAGGTTGGCTCTGTGAG-3′ were used to PCR amplify the mutated abi1-1 coding sequence without its stop codon from cDNA synthesized using abi1-1 inflorescence RNA and cloned into the Gateway-compatible D-TOPO vector. Multisite Gateway (Invitrogen) recombination was used to introduce ProSCR:abi1-1:1::RFP and ProCOR:abi1-1:1::RFP into a dpGreen-based binary vector, which contains a kanamycin-resistant gene for plant selection.

To generate the ProSCR:abi1-1 construct, the primers 5′-caccATGAGGAAATATCTCGGCCG-3′ and 5′-TACGTTCAAGGTTGCTCTTGT-3′ were used to PCR amplify the mutated abi1-1 coding sequence. Multisite Gateway (Invitrogen) recombination (Karimi et al., 2005) was used to introduce a ProSCR:abi1-1::GUS fusion into a modified dpGreen-based binary vector, which contained a 35S:PM-mCherry selection marker in place of the kanamycin resistance gene (pCherry-picker7).

Transgenic plants were generated using a modified floral dip method as follows. GV3101 cells were cultured in large Petri dishes (150-mm diameter) on selective Luria-Bertani agar media for 2 d at 28°C. Fifty milliliters of infiltration media (half-strength MS salts, 0.03% Silwet L77, and 5% Suc adjusted to pH 5.7 with KOH) was poured on top of the Luria-Bertani agar media and cells were scraped off into the solution. The cell suspension was homogenized by gentle shaking for 5 s in a 50-mL conical tube. The cell suspension was then diluted with an additional 150 mL of infiltration media and the combined solution used for floral dip–mediated plant transformation. Seeds were harvested from treated plants and selected visually based on mCherry fluorescence using an M165 FC fluorescence microscope (Leica).

### Microscopy Analysis

For quantitation of LR developmental stages, roots were mounted in a modified Hoyer’s solution (chloral hydrate:water:glycerol in proportions 8:2:1, g/mL/mL) and then imaged using a Leica DMi6000 inverted compound microscope. The GUS staining protocol was performed as previously described (Swarup et al., 2008). For confocal microscope imaging, roots were mounted in an FM4-64 solution (Invitrogen; Levesque et al., 2008) or propidium iodide (Invitrogen) and imaged using a Leica SP5 point-scanning confocal microscope. The imaging settings were 488-nm excitation and 505- to 550-nm emission for GFP and 488-nm excitation and >585-nm emission for FM4-64 or propidium iodide.

### Phenotypic Analysis

For end-point analysis of root length, seedlings were grown on standard media for 6 d and then transferred to various conditions for 4 d. Images of 10 dpq seedlings were captured using a CanonScan 9000F flattened scanner (Canon). The length of all visible LRs in root region A and PR length of region B was quantified using ImageJ (Abramoff et al., 2004). The average LR length and standard error of the mean per treatment was calculated from the average LR length of each seedling. Sample size of the various experiments is indicated in the corresponding figure legends.

For live-imaging analysis of LR growth, a custom live-imaging system was developed and consisted of the following: Samples were manipulated using a circular platform with six tissue-culture plate holders and controlled by an automated Theta/360 degree rotary stage and MFC-2000 controller (Applied Scientific Instrumentation), samples were backlit using an infrared light-emitting diode panel, and images were captured using a digital monochrome camera (CoolSnap) fitted with an NF Micro-Nikon 60-mm lens (Nikon) and infrared filter. Micro-Manager Software (Vale Lab, University of California, San Francisco) was used to control the stage and automate image acquisition (Edelstein et al., 2010). Images were taken every 15 min (wild-type growth under standard or salt stress conditions) or 20 min (transactivation of abi1-1 studies) for 7 d. Sequential images were collated as a stack for further analysis using ImageJ. The StackReg plug-in was used to align the stack of image slices before root growth quantitation. Quantification of data from the time-lapse movies was conducted using the Manual Tracking plug-in for ImageJ. The growth of the root tip was tracked manually from frame to frame. Growth rate data were imported into TM4 microarray software suite (www.tm4.org/mev/) to generate heat maps (Saeed et al., 2006). Hierarchical clustering using Pearson correlation as the distance metric was performed to organize the growth data.

To determine the effect of salt stress on cell cycle activity in LRs, the expression of the CYCLINB1;1::GUS reporter was quantified in seedlings transferred to standard or 100 mM NaCl conditions for 3 d. The total number of GUS-positive and GUS-negative LRs was counted for LRs where obvious elongation and maturation zones had developed (long LRs) or where no such zonation had developed (short LRs).

To monitor dynamic changes in ABA signaling in LRs, fluorescence of the ProRAB18::GFP reporter was imaged on a Leica fluorescence dissecting microscope once per day of treatment. LR length and GFP fluorescence intensity at each time point was quantified using ImageJ (Abramoff et al., 2004). GFP was quantified by calculating the average
pixel intensity over the area of the entire LR. To monitor GFP:RGA accumulation, we quantified the average fluorescence intensity of nuclei in the endodermis using ImageJ.

Two-way analysis of variance and Student's t test were used to test for statistical significance in root length measurements using a P value threshold of <0.05 and a Bonferroni correction based on the number of tests performed.

Gene Expression

For qRT-PCR analysis of gene expression, RNA was extracted from root region A or region B separately using the RNaseasy plant mini kit (Qiagen) according to manufacturer's instructions. cDNA was prepared using the iScript advanced cDNA synthesis kit (Bio-Rad) from 300 ng of total RNA. qRT-PCR was performed on a Fluidigm BioMark 96.96 Expression Chip using EvaGreen (Bio-Rad) as the fluorescence probe according to the Fluidigm Advanced Development Protocol #37. AT3G07480 was used as a control gene because of its stable expression under various stresses (see Supplemental Figure 14 online). Two biological replicates and three technical replicates were used in this analysis. Technical replicates were averaged first, then average expression values were calculated from the biological replicates and imported into TM4 microarray software suite (www.tm4.org/index.html) to generate expression heat maps (Saeed et al., 2006). Expression values were unlog-transformed assuming two-fold amplification efficiency with each PCR cycle and normalized. Hierarchical clustering using Pearson correlation as the distance metric was performed to organize the expression data. Primers used in these experiments are listed in Supplemental Table 1 online.

Genetic Analysis

To selectively express abi1-1 in different tissue types, various enhancer trap lines were crossed to plants harboring the UAS:abi1-1 transgene. Wild-type plants of the C24 ecotype were crossed with UAS:abi1-1 plants to generate the control genotype. Phenotypic and gene expression analysis were performed using the F1 seeds. To evaluate whether salt-dependent changes in RGA protein stability were ABA signaling dependent, ProRGA:GFP:RGA transgenic plants were crossed with the abi1-1 mutant or to the Ler ecotype to generate the control genotype. F1 seeds from these crosses were used for confocal image analysis of GFP:RGA fluorescence under standard or salt stress conditions.

Accession Numbers

Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under accession number At4g26080 (ABI1). Gene IDs for genes used for real-time quantitative PCR can be found in Supplemental Table 1 online.

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure 1. LR Growth Is More Strongly Suppressed Than PR Growth by Salt among Different Ecotypes.

Supplemental Figure 2. Quiescent LRs Have Normal Tissue Organization.

Supplemental Figure 3. ABA and ACC Treatments Show Different Trends in Their Effects on LR and PR Growth and Are Ecotype Independent.

Supplemental Figure 4. ABA Signaling Primarily Affects LR Growth under Salt Stress Conditions.

Supplemental Figure 5. Six Enhancer-Trap Lines Show Distinct Expression Patterns in Arabidopsis PRs.

Supplemental Figure 6. ABA Signaling in the Pericycle Does Not Significantly Regulate LR Growth during Salt Stress.

Supplemental Figure 7. Sucrose in the Media Does Not Affect the Salt Response of LRs in Control or Q2500>>abi1-1 Transactivation Genotypes.

Supplemental Figure 8. Endodermis-Specific Expression of abi1-1 Partially Rescues LR Growth during ABA and Salt Treatment.

Supplemental Figure 9. NaCl Regulates the Expression of ABA-Responsive Genes and Activates Prolonged Expression of the ProRAB18:GFP Reporter.

Supplemental Figure 10. Nutrient Conditions Modulate the Response of LRs and PRs to Salt Stress.

Supplemental Figure 11. The Salt Response of PRs Is Moderately Affected by Exogenous GA Treatment and by Endogenous GA Signaling.

Supplemental Figure 12. Interactions between ABA and GA Signaling and the Effect of DELLA Activity on LR Growth and Gene Expression.

Supplemental Figure 13. ProCASP1:GFP Expression Is Induced at an Earlier LR Stage and Is More Intensely Expressed under Salt Stress Conditions.

Supplemental Figure 14. AT3G07480 Serves as the Control Gene for qRT-PCR Expression Assays in This Work Due to Its Stable Expression under Various Abiotic Stress Conditions.

Supplemental Table 1. Primer Sequences Used for Real-Time qPCR Experiments.

Supplemental Movie 1. Time-Lapse Movie of Col-0 Lateral Root Growth.

Supplemental Movie 2. Time-Lapse Movie of Lateral Root Growth in Control and Q2500>>abi1-1 Transactivation Line under 100 mM NaCl.

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AUTHOR CONTRIBUTIONS

L.D. and J.R.D. designed the research plan. L.D. performed most of the experiments and data analysis. M.J.B., D.D., and R.B. originated the concept for inhibiting ABA signaling using abi1-1 misexpression and generated the UAS:abi1-1 transgenic line. C.H.N. generated the pCherry-pickerT vector. P.M.Y.C. established protocols for live imaging, image analysis, and high-throughput qRT-PCR. L.D. and J.R.D. wrote this article, which was read by all the authors.

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Lina Duan, Daniela Dietrich, Chong Han Ng, Penny Mei Yeen Chan, Rishikesh Bhalerao, Malcolm J. Bennett and José R. Dinneny

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