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ABA homeostasis and long-distance translocation is redundantly regulated by ABCG ABA importers

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Abstract:

The effects of abscisic acid (ABA) on plant growth, development and response to the environment depend on local ABA concentrations. Here, we exploited a genome-scale amiRNA screen, targeting the *Arabidopsis* transportome, to show that ABA homeostasis is regulated by two previously unknown ABA transporters. ABCG17 and ABCG18 are localized to the plasma membranes of leaf mesophyll and stem cortex cells to redundantly promote ABA import, leading to conjugated inactive ABA sinks, thus restricting stomatal closure. *ABCG17* and *ABCG18* double knockdown revealed that the transporters encoded by these genes not only limit stomatal aperture size, conductance and transpiration while increasing water-use efficiency but also control ABA translocation from the shoot to the root to regulate lateral root emergence. Under abiotic stress conditions, *ABCG17* and *ABCG18* are transport mechanism mediated by ABCG17 and ABCG18 allows plants to maintain ABA homeostasis under normal growth conditions.

Introduction:

Abscisic acid (ABA) is a plant hormone that regulates growth and responses to the changing environment. For example, seed dormancy, germination, drought tolerance, stomatal closure and lateral root emergence are modulated by this important hormone under normal conditions and in response to stimuli (1-7). The ABA response is regulated at multiple steps: biosynthesis, transport, catabolism, perception, and signal transduction. ABA travels long distances throughout the plant, a characteristic that was shown about 50 years ago to affect stomatal conductance and responses to drought (8-11). For decades, the concept stated in textbooks was that in response to drought, root-derived ABA travels to shoot guard cells via the xylem sap to prevent water loss (12, 13). Indeed, several studies support the idea that root-derived ABA synthesis is required for ABA-induced stomatal closure to elicit drought tolerance (14-16). Recent studies, however, provide evidence that stomatal closure is triggered by shoot-specific ABA synthesis (17-20). While the complete map of ABA biosynthesis is not entirely clear yet, studies showed that ABA could be synthesized directly in guard cells during abiotic stresses and in the leaf vasculature (21).

Multiple lines of evidence support the hypothesis that ABA transport is necessary for proper ABA responses (22-24). First, ABA, being a weak acid, exists primarily in its charged and membraneimpermeable form (depending on the pH), suggesting the need for transporter-facilitated movement across membranes (25-29). Second, using deuterium-labeled ABA and reciprocal grafting between wild-type and ABA-biosynthetic mutant tomatoes, it was found that foliage-derived ABA promotes primary root growth but inhibits the development of lateral roots (30). A recent study demonstrated that leaf-derived ABA participates in rice seed development in response to environmental temperature variations (31). These highlights the importance of long-distance ABA transport in governing plant physiological and morphological aspects, beyond stomata control. Third, phloem-specific ABA synthesis complements ABA activity in stomatal aperture, indicating that ABA movement within the leaf is important (32-34). Fourth, ABA transport is required for processes such as germination (23, 35, 36).

In recent years, several ABA transporters have been identified and characterized (15, 23, 37, 38). For example, in seeds, ABCG25 and ABCG31 export ABA out of endosperm while ABCG40 and ABCG30 import ABA into the embryo to regulate seed germination. In leaves, *ABCG25* is mainly expressed in the vasculature, promoting ABA transport to guard cells, while ABCG40 imports

ABA into guard cells to control the stomatal aperture (23, 37, 39-41). MtABCG20 was recently characterized as an ABA importer that is involved in germination and lateral-root formation in *Medicago truncatula* (42). Additional transporters include DTX50, a transporter of the multidrug and toxic compound extrusion (MATE) transporter family, and NITRATE TRANSPORT1/PEPTIDE TRANSPORTER FAMILY (NPF) 4.6 (also known as NRT1.2 and AIT1) (15, 23, 25, 34, 43, 44).

In addition to transport, ABA concentrations are tightly controlled by ABA metabolism (36, 45). ABA is conjugated with glucose by UDP-glucosyltransferases (ABA-glucosyltransferase) to catalyze the formation of the stored form of ABA, ABA-glucosyl ester (ABA-GE) (46). Previous studies showed that there are 26 subfamilies of UDP-glucosyltransferases (UGTs) in *Arabidopsis*. Seven UGTs, which belong to different groups of UDPs subfamily 1 (encoded by *UGT84B1*, *UGT75B1*, *UGT84B2*, *UGT71B6*, *UGT75B2*, *UGT73B1* and *UGT71C5*) have ABA to ABA-GE catalysis activity (47, 48). However, our understanding on the spatiotemporal aspects of ABA homeostasis and the mechanisms underlying ABA translocation to control ABA responses under normal and stress conditions are limited.

Here we exploited a genome-scale amiRNA screen targeting the *Arabidopsis* transportome to identify novel ABCG ABA transporters. ABCG17 and ABCG18 are localized to the plasma membrane and import ABA. *ABCG17-* and *ABCG18-*knockdown lines displayed enhanced ABA accumulation, induced ABA response, and reduced stomatal aperture size, conductance, and transpiration with increased water-use efficiency compared to WT. In addition, seedlings with double-loss-of-function of *ABCG17* and *ABCG18* showed enhanced long-distance ABA translocation from the shoot to the root, which led to lateral root outgrowth inhibition. *ABCG17* and *ABCG18* are primarily expressed in the shoot mesophyll and stem cortex cells, where they drive ABA import, allowing ABA-GE formation. In turn, these ABA-GE sinks prevent active ABA accumulation in guard cells and limit ABA long-distance travel to lateral root formation sites. Under abiotic stress conditions, *ABCG17* and *ABCG18* are transcriptionally repressed, promoting active ABA movement, accumulation, and response.

Results

Transportome-scale amiRNA screen indicates involvement of class A ABCGs in ABA mediated activity.

Over 75% of the genes in the *Arabidopsis* genome belong to gene families, complicating functional analyses (49). In order to partially overcome functional redundancy and reveal genetic factors involved in ABA homeostasis and translocation, we utilized a transportome-specific amiRNAs collection of 3,000 lines targeting multiple transporters from the same family (50). The transporter amiRNA sub-library is part of the broad PHANTOM amiRNA tool (49). The seedlings in the amiRNA collection were screened for defects in shoot growth and photosynthesis-related parameters. The screen identified *amiRNA-1228* line as an interesting candidate; this line had slight but significant shoot growth inhibition (Fig. 1A, Fig. S1A). In addition, the *amiRNA-1228* line showed a significant reduction in photosystem II quantum yield of light-adapted samples at steady state (Fig. 1B), indicating limited activity of photosystem II (51). We therefore hypothesized that the genes targeted by *amiRNA-1228* might be related to ABA activity and homeostasis (52).

amiRNA-1228 putatively targets four class A ABCG subfamily members with different extents of complementarity in the potential recognition sites (Fig. S1*B*). ABCG17 and ABCG18 are grouped in one branch, ABCG1 and ABCG20 are classified into another, of the *Arabidopsis* ABCG family phylogenetic tree (Fig. 1*C*). To test if *amiRNA-1228* indeed targets these genes, we examined their expression levels by qRT-PCR in *amiRNA-1228* plants compared to that in wild-type (WT) plants (Fig. 1*D*). *ABCG1, ABCG17,* and *ABCG18* were significantly down-regulated in *amiRNA-1228,* but there was no difference in levels of *ABCG20* (Fig. 1*D*). Next, we tested the transcriptional responses of these genes to ABA treatment. *ABCG1 and ABCG20* were induced by 5 µM ABA treatment for 3 hours, whereas *ABCG17* and *ABCG18* were repressed (Fig. 1*E*). ABCG1 and ABCG16 were previously shown to regulate pollen and reproductive development (53, 54) and suberin formation in roots (55). ABCG20, ABCG2, and ABCG18, located on separate phylogenetic branch (Fig. 1*C*), have not been characterized. Altogether, our results indicate that *ABCG17* and *ABCG18* genes are interesting candidates in the context of ABA-mediated activity.

ABCG17 and ABCG18 are redundantly required for ABA response and stomatal closure.

To determine whether *ABCG17* and *ABCG18* contribute to the *amiRNA-1228* phenotype and to dissect their activity, we obtained T-DNA insertion lines for both genes (Fig. 2A). Since *ABCG17* and *ABCG18* are genetically linked on chromosome 3, we could not generate double mutants by crossing T-DNA-insertion lines. Therefore, two independent amiRNA double-knockdown lines

for *ABCG17* and *ABCG18* were generated. The first was an amiRNA targeting both *ABCG17* and *ABCG18* (*mir17*, *18*). The second was an amiRNA targeting *ABCG17* (*mir17*) transformed into the *abcg18-1* T-DNA-insertion background (*mir17*, *g18-1*) (Fig. 2A). We examined the expression levels of *ABCG17* and *ABCG18* in the single- and double-knockdown lines compared to WT. Both *ABCG17* and *ABCG18* were significantly downregulated in the respective backgrounds (Fig. 2*B-C*). Next, we monitored the shoot phenotypes of *abcg17* and *abcg18* single-mutant and double-knockdown lines grown in soil under normal growth conditions. None of the single mutants showed shoot-growth impairment, but both double-knockdown lines, *mir17*, *18* and *mir17*, *g18-1*, displayed significant reductions in shoot surface area compared to WT and the single mutants (Fig. 2*D-E*). Similarly, 50-day-old double-knockdown plants had shorter inflorescence stems compared to WT and single mutants (Fig. 2*F-G*), but showed no differences in bolting time (except *abcg18-2*) (Fig. S2). The results are in line with the *amiRNA-1228* phenotype and indicate that *ABCG17* and *ABCG18* are redundantly required for plant shoot growth.

Since *amiRNA-1228* showed photosynthesis-related phenotypes, we tested whether it resulted from *ABCG17/ABCG18*-dependent alteration of stomatal aperture size. Single *abcg17* and *abcg18* mutants did not differ from WT plants in stomatal aperture size. However, both *mir17,18* and *mir17,g18-1* had significantly smaller stomatal apertures than WT and single-mutant plants (Fig. 3*A*).

To further validate if the phenotypes observed in mir17,18 and mir17,g18-1 are indeed an ontarget effect driven by ABCG17 and ABCG18, we generated ABCG17 and ABCG18complementation lines, pABCG17:ABCG17 in the background of mir18,g17 double knockdown (g17 is a T-DNA line and thus can be complemented by ABCG17) and pABCG18:ABCG18 in the background of mir17,g18-1 (g18-1 is a T-DNA line and thus can be complemented by ABCG18). The complementation lines showed complete rescue of stomatal apertures phenotype (Fig. 3B), together validating the activity of ABCG17 and ABCG18 in regulating stomatal aperture.

To further study ABCG17 and ABCG18 activity in guard cell regulation, we tested stomatal conductance and gas exchange (i.e., transpiration and photosynthesis) in the mutant lines. The double-knockdown line had significantly reduced stomatal conductance compared to WT and single mutants (Fig. 3*C*). In line with stomatal conductance and aperture defects, the double-

knockdown line had significantly lower transpiration rates than WT and single mutants (Fig. 3*D*). However, photosynthesis rates were not changed within experimental error in both single mutants and the double-mutant *mir17,18* line compared to WT plants (Fig. 3*E*). In agreement with this finding, the double-mutant line, *mir17,18* showed an increase in instantaneous water use efficiency under well-watered conditions (Fig. 3*F*). Analysis of water-loss rates from excised leaves of single mutants and *mir17,18* showed delayed water-loss rates compared to WT when exposed to air (Fig. 3*G*, Fig. S3). Evaluation of leaf temperature utilizing thermal imaging showed higher leaf temperature in both the double-knockdown lines compared to WT and single mutants (Fig. 3*H-I*). These data support the hypothesis that ABCG17 and ABCG18 are required for stomatal activity under normal conditions.

To further characterize ABCG17 and ABCG18 activity, we generated transgenic plants overexpressing (*35S* promoter) each of the two genes. Two independent lines for each gene showed a 50-100-fold change in transcriptional activation (Fig. S4A-B). *ABCG17-* or *ABCG18-* overexpressing lines showed no differences in shoot surface area from that of WT plants, but *ABCG18* overexpression resulted in a slight plant height increase (Fig. S4C-F). Neither *ABCG17-* nor *ABCG18-* overexpressing lines showed differences in bolting time (Fig. S4G). *ABCG17 or ABCG18* overexpression resulted in reduced stomatal aperture (Fig. S5A-B), elevated leaf temperature (Fig. S5*C-D*), and induced ABA response, reflected by the induction of *pRAB18:GFP* (56) and *pMAPKKK18:LUC (57)* ABA reporters (Fig. S5*E-F)* compared to WT plants. Overall, we observed that some overexpression lines presented opposite phenotypes to the loss-of-function lines while others were similar. This may result from responses that depend on local ABA biosynthesis sites. Altogether, these results suggest that ABCG17 and ABCG18 redundantly regulate stomatal conductance and aperture.

ABCG17 and ABCG18 are plasma membrane localized ABA importers.

To determine the subcellular localizations of ABCG17 and ABCG18, we cloned the coding sequences of *ABCG17* and *ABCG18* and constructed vectors for expression of these proteins with N-terminal YFP tags under the control of the *35S* promoter. Confocal microscopy of *p35S:YFP-ABCG17* and *p35S:YFP-ABCG18* transgenic lines in the root meristem, revealed that ABCG17 and ABCG18 are localized to the plasma membrane (Fig. 4A).

To examine whether ABCG17 and ABCG18 facilitate ABA transport, we synthesized fluorescently labeled ABA molecules by adding a fluorescein-tag to the carboxylic group (Fig. 4*B*). Unlike the fluorescently labeled gibberellin molecules we previously reported (58), the fluorescent ABA (ABA-FL) was only slightly bioactive in root growth assays, suggesting low binding affinity for the PYR/PYL/RCAR ABA receptor (Fig. S6). We used the new ABA-FL molecule to test ABCG17 and ABCG18 transport activity. Isolated protoplasts from leaf mesophylls of transgenic *Arabidopsis* plants overexpressing *ABCG17* or *ABCG18*, and WT as a control were treated with 5 μ M ABA-FL for 12 hours and imaged. Protoplasts overexpressing *ABCG17* or *ABCG18* had enhanced ABA-FL fluorescent signal compared to WT protoplasts (Fig. 4*C-D*), suggesting that both *ABCG17* and *ABCG18* promote ABA import.

To validate ABCG17 and ABCG18 ABA transport activity, a biochemical transport assay with radiolabeled ABA ([¹⁴C]ABA) in tobacco plants that overexpress each of these two transporters was carried out. ABCG17 or ABCG18 overexpression in tobacco leaves showed plasma membrane localization (Fig. 4E) and resulted in a significant reduction in measured protoplast export activity compared to the control (Fig. 4F). Reduced net export is in line with an import activity for both transporters that would function here as re-importers of $[^{14}C]ABA$. Additionally, we examined the transport activity of ABCG17 and ABCG18 in Xenopus oocytes. Both genes were expressed individually and also in combination. No transport activity was observed for ABCG17 and ACBG18 when expressed individually or together (Fig. S7, Table S1), suggesting that additional in planta cofactors might be required for ABCG17 and ABCG18 activity. In order to determine the specificity of ABCG17 and ABCG18 in ABA transport activity, we tested the effects of overexpressing ABCG19, the closest gene in the cluster (Fig. 1C). Notably, overexpression of ABCG19 did not result in significant ABA transport activity (Fig. 4F). We also evaluated transport of a radioactively labeled auxin ([³H]IAA), like ABA, an organic acid and a known substrate of members of the ABCB subclass. ABCG17 or ABCG18 overexpressing lines did not reveal altered transport of auxin compared to the control (Fig. 4G). In summary, these experiments indicate that ABCG17 and ABCG18 are specific plasma membrane-localized ABA importers.

ABCG17 and ABCG18 limit long-distance ABA transport to regulate lateral root development.

In addition to the stomatal conductance and aperture size phenotypes, double knockdown of *ABCG17* and *ABCG18* resulted in reduced lateral root numbers compared to WT and single mutants (Fig. 5A). More specifically, *mir17,18* and *mir17,g18-1* showed normal lateral root initiation (Fig. S9*B-C*, Tables S2-3) but reduced lateral root primordia outgrowth compared to WT under normal conditions (Fig. 5*B*, Fig. S9*A*). This result is intriguing since the expression of *ABCG17* and *ABCG18* is largely restricted to the shoot: luciferase reporter lines (*pABCG17:LUC* and *pABCG18:LUC*), GUS reporter lines (*pABCG17:GUS* and *pABCG18:GUS*), YFP reporter lines (*pABCG17:NLS-YFP* and *pABCG18:NLS-YFP*) and qRT-PCR measurements showed strong signals in shoots (Fig. 5*C-E*, Fig. S10). A very weak signal was found in lateral root emerging primordia using the *pABCG17:NLS-YFP pABCG18:NLS-YFP* lines, only when applying maximum laser power and using a high sensitivity gallium arsenide phosphide (GaAsP) detector (Fig. 5*E*, Fig. S11*A-B*). ABA regulates key stages of lateral root post-emergence development (59). Specifically, ABA signaling plays crucial role in salt stress-induced lateral root suppression (5, 60) and lateral root growth recovery (61), associated with an increase in *miR165a* and a reduction in *PHB* (*PHABULOSA*) levels (62).

In order to test if shoot-born ABA can affect lateral root development, we applied ABA specifically to shoots and quantified lateral root length and lateral root number. The results indicated that ABA application to WT shoots inhibited lateral root elongation and lateral root development (Fig. S12*A*-*B*). Moreover, ABA application to the shoots induced expression of the *pMAPKKK18:GUS* ABA reporter (57) in the root (Fig. S12*C*). Shoot-specific ABA application inhibited lateral root emergence in the single mutants similarly to WT, but the *mir17,18* double knockdown mutant was insensitive to ABA application to shoots (Fig. 5F). We speculated that *mir17,18* accumulates high ABA levels in the root compared to WT under normal conditions.

To test this hypothesis, we quantified ABA levels in roots. Indeed, the double-knockdown *mir17,18* plants accumulated significantly higher ABA amounts in the root compared to WT (Fig. 5G). The data showed more than 85% accuracy, among different replicates and experiments, suggesting good recovery of the analytes after sample preparation steps, with minimal losses (Table S4). In addition, we analyzed the ABA reporter *pRAB18:GFP* in the background of *ABCG17* and *ABCG18* loss-of-function lines. We found enhanced *pRAB18:GFP* signal in

mir17,18 roots compared to control roots (Fig. 5*H*), supporting the hypothesis that *ABCG17* and *ABCG18* knockdown causes accumulation of abnormally high levels of ABA in the root.

In order to further understand an apparent ABCG17 and ABCG18 activity in limiting ABA shootto-root translocation, we treated shoots with radioactively labeled ABA (0.1 μ Ci/ml [³H]ABA), and quantified counts in roots after 24 hours. The *mir17,18* roots accumulated higher levels of [³H]ABA than the WT roots (Fig. 5*I*), supporting the hypothesis that ABCG17 and ABCG18 activities restrict long-distance transport of ABA to the root. Since auxin is a key mobile molecule required for lateral root formation, we carried similar experiments with shoot-applied radioactively labeled IAA. Root extractions did not reveal significant differences between *mir17,18* and WT (Fig. S13). To further confirm ABCG17 and ABCG18 activity in long-distance ABA translocation, reciprocal WT and *mir17,18* grafting assays were carried out. The results showed that, within experimental error, WT scion/*mir17,18* rootstock grafts developed the same number of lateral roots as WT self-grafted plants. However, *mir17,18* scion/WT rootstock grafts developed significantly fewer lateral roots, with lateral root numbers similar to *mir17,18* self-grafted plants (Fig. 5*J*). These results indicate that ABCG17 and ABCG18 redundantly restrict shoot-to-root ABA translocation, mediating lateral root development under normal conditions.

ABCG17 and ABCG18 are expressed in shoot mesophyll and cortex cells under non-abiotic stress conditions.

The obtained results suggest that ABCG17 and ABCG18 limit ABA levels in guard cells and roots, through a process that takes place in the shoot. Therefore, we speculated that ABCG17 and ABCG18 redundantly regulate ABA homeostasis, possibly by allowing ABA storage. To test this hypothesis, we evaluated ABA and ABA-GE levels in shoots. The double-knockdown *mir17,18* plants accumulated higher ABA levels in the shoot than WT or single mutant plants (Fig. 6A), consistent with results showing that double-knockdown lines have reduced stomatal aperture size, conductance, reduced transpiration, and elevated leaf temperature (Fig. 3). Interestingly, the double-knockdown *mir17,18* shoots accumulated less ABA-GE than WT shoots (Fig. 6B), suggesting that ABCG17 and ABCG18 might promote ABA-GE production catalyzed by UDP-glucosyltransferases. In support of higher ABA content in the *mir17,18* shoots, the ABA response gene *ABI1* was significantly induced in *miR17,18* compared to WT plants (Fig. 6C). To better characterize *ABCG17* and *ABCG18* function in ABA metabolism, we quantified additional ABA

metabolites. Several ABA inactive catabolites and conjugated forms showed profound effects in *ABCG17* and *ABCG18* mutants (Fig. 6D, Fig. S14). High accuracy and repeatability were measured for shoot ABA and ABA metabolites recovery method (Table S4).

To identify the exact cell types that accumulate ABA due to transport facilitated by ABCG17 and ABCG18, we characterized the expression patterns of *ABCG17* and *ABCG18* in leaf and stem sections. *pABCG17:NLS-YFP* and *pABCG18:NLS-YFP* lines showed that *ABCG17* and *ABCG18* are primarily expressed in parenchyma shoot cells, which are the leaf mesophyll cells (Fig. 6*E*-*F*, Fig. S15*A*-*B*). We did not detect expression in the leaf or stem veins or bundle sheath cells (Fig. 6*E*-*F*). These results imply that under normal conditions, ABCG17 and ABCG18 promote ABA uptake into shoot parenchyma cells, leading to increased ABA-GE levels. Thus, under non-stress conditions, low levels of free shoot-born ABA are available to guard cells and for long-distance movement to roots.

Next, we wanted to elucidate how ABCG17 and ABCG18 function in ABA-mediated homeostasis under abiotic stress conditions, when high levels of ABA are needed. Both of the two double knockdown lines exhibited stronger resistance by showing more lateral roots and longer roots with 100 mM NaCl treatment compared to WT and single mutants (Fig. S9A, *D-E*). Characterizing lateral root number, lateral root length and lateral root primordium of *ABCG17* overexpressing lines with and without 100 mM NaCl treatment did not show significant difference with NaCl treatment compared to WT (Fig. S16, Tables S5-6). However, one of the *ABCG18* overexpressing lines presented more lateral root compared to WT with 100 mM NaCl treatment (Fig. S17, Tables S7-8).

To study how ABCG17 and ABCG18 function in ABA-mediated homeostasis under stress conditions, we monitored the expression levels of ABCG17 and ABCG18 in response to abiotic stresses using pABCG17:NLS-YFP and pABCG18:NLS-YFP plants as well as pABCG17:LUC and pABCG18:LUC plants. ABCG17 and ABCG18 were transcriptionally repressed following ABA or NaCl treatments (Fig. 6G, Fig. S18A-B), consistent with qRT-PCR experiments following ABA Similarly, long-period water-withhold experiments treatment (Fig. 1*E*). showed that ABCG17 and ABCG18 are repressed as luminescence intensity of promoter-driven luciferase reporters was reduced in plants not irrigated for several days (14 and 24 days) (Fig. 6H-I, Fig. S18C).

In summary, our experiments imply that ABCG17 and ABCG18 are necessary for ABA homeostasis through creation of ABA-GE sinks in mesophyll and cortex cells. These ABA-GE sinks limit ABA accessibility to guard cells and ABA availability for long-distance translocation to regulate lateral root formation. The ABA homeostasis shoot sink is released once the plant senses abiotic stress conditions.

Discussion

Here we utilized a transportome-scale amiRNA screen to overcome functional redundancy to identify two previously unstudied ABCG transporters, *ABCG17* and *ABCG18*. Compared to WT and single *abcg17* and *abcg18* mutants, the double-knockdown lines had higher ABA content and reduced ABA-GE content in shoots. Since ABCG17 and ABCG18 are primarily expressed in the shoot cortex and mesophyll cells, localized to the plasma membrane, and promote ABA import, we hypothesized that the two proteins are redundantly required for ABA accumulation and storage in these cells (Fig. 7). The ABCG17- and ABCG18-dependent ABA sink in mesophyll cells limits the hormone translocation and activity to guard cells and the lateral-root emergence sites, two distinct ABA-mediated processes. The two redundant transporters are required for ABA homeostasis under normal conditions, whereas the suggested conjugated-ABA-sink mechanism is restricted in abiotic stress environments, enabling rapid ABA responses at distinct target sites (Fig. 7).

Our model raises several questions. It is unknown how free ABA, which is not taken into the shoot cortex or mesophyll cells, finds its way to the guard cells or lateral-root formation sites, which requires short or long-distance transport, respectively. ABA might move through the apoplast, passively diffuse, or actively translocate over the plasma membrane. Several ABA transporters have been identified so far. In *Arabidopsis*, ABCG40 imports ABA into guard cells (15, 45), and in tomato AIT1.1 was recently shown to play a role in ABA import into guard cells in a DELLA-dependent manner (63). In addition, NPF4.6 regulates vasculature ABA import and ABCG25 and DTX50 regulate vasculature ABA export (24). We speculate that it is unlikely that free ABA can travel from shoot to root by simple diffusion and that there are transporters that remain to be identified that mediate shoot-to-root ABA transport. These transporters are expected to load ABA into the shoot phloem and unload ABA in the root to regulate lateral root development. Our

proposed model suggests that ABCG17 and ABCG18 mediate the rate-limiting step for ABA accumulation in shoots, thus allowing ABA translocation to roots while ABA-GE accumulates within the shoot cortex and mesophyll cells (Fig. 7). However, it is currently unclear whether ABA-GE itself can travel from root to shoot and *vice versa (29)* and further tools are required to characterize such mechanisms.

Previous studies have shown that ABA translocation from root to shoot via the xylem is required for stomatal closure, which restricts transpiration and water loss under drought stress (*12-16*). Little is known, however, about shoot-to-root transport via the phloem. McAdam *et al.* presented evidence that most of the ABA found in the tomato root is synthesized in the shoot (30). It is therefore possible that ABCG17 and ABCG18 might serve a role in the mechanism regulating this process in *Arabidopsis*. Further studies in tomato and other species are needed to confirm whether the ABCG17- and ABCG18-mediated shoot-to-root ABA translocation mechanism is conserved. Ramzi *et al.* showed that the accumulation of ABA in roots after long-term water stress largely relies on shoot-to-root ABA transport (64), a process that is likely conserved in *Arabidopsis* (17), citrus (65) and other species. We show that *ABCG17* and *ABCG18* are downregulated under abiotic stress conditions, supporting the idea that their reduced activity promotes shoot-to-root ABA translocation under stress conditions.

The model proposed here suggests that ABCG17 and ABCG18 drive an inactive ABA-GE sink that restricts stomatal activity and lateral root formation. However, we currently have little information on the cell-type or subcellular distributions of ABA and ABA-GE or of the associated ABA-glucosyltransferase localization. A detailed map of ABA-glucosyltransferase activity is required to plot ABA and ABA-GE movement in high resolution. Such a task remains challenging as this large family of enzymes is likely regulated at the protein level and profiling the expression pattern will not be sufficient. High-resolution fluorescent probes specific for ABA-glucosyltransferase that can be used in live plants would be an extremely useful tool in the field. There are some indications for ABA-GE accumulation in intracellular storage organelles (such as vacuoles) (66), but it remains unclear exactly how ABA and ABA-GE distribute within the mesophyll cells. We speculate that ABCG17 and ABCG18 activity leads to high levels of free ABA within the mesophyll and shoot cortex cells, which is then converted to ABA-GE by UDP-glycosyltransferases. Current models suggest that the ABA-GE is translocated to the endoplasmic

reticulum and vacuole for storage (45, 67). Burla *et al.* suggested that two distinct membrane transport mechanisms are active in *Arabidopsis* mesophyll cells: a proton gradient-driven and an ATP-binding cassette transporter-mediated. They demonstrated that ABCC1 and ABCC2 are localized in tonoplasts and exhibit ABA-GE transport activity in a yeast heterologous expression system (68, 69). Therefore, it is possible that ABCC and ABCC transporters work in concert: ABCG17 and ABCG18 import free ABA into mesophyll cells, and ABCC1 and ABCC2 mediate storage of ABA-GE in vacuoles. In addition to the need to identify subcellular ABA-GE transporters, there is no direct evidence of ABA and ABA-GE distribution to different organelles. New biochemical and genetic tools are required to generate a comprehensive ABA profile at subcellular resolution.

Methods

Plant material and growth conditions.

All *Arabidopsis thaliana* lines are in Col-0 background. For assays on plates, sterilized seeds were plated on 16×16 cm square petri dishes or 8.5 cm round petri dishes with growth media containing 0.5 × Murashige-Skoog (MS) medium (pH 5.95 - pH 6.1, 1% sucrose, 0.8% plant agar). The sterilized seeds were transferred to growth chambers (Percival, CU41L5) at 21 °C, 100 μ Em⁻² S⁻¹ light intensity under long-day conditions (16 h light/8 h dark) after 3-day-stratification at 4 °C in the dark. For soil experiments, seedlings were transferred onto soil and grown in dedicated growth rooms under long-day conditions (16 h light/8 h dark) at 21 °C. The following ABA reporters were previously described: *pMAPKKK18:GUS*, *pMAPKKK18:LUC* (*57*) and *pRAB18:GFP* (*56*).

Agrobacterium transformation.

GV3101 electro-competent *Agrobacterium tumefaciens* strain was incubated on ice with ~100 ng plasmid for 5 min and electroporated using a MicroPulser (Bio-Rad Laboratories; 2.2 kV, 5.9 ms). Immediately after electroporation, 600 μ l LB medium was added, and sample was shaken for 2 h at 28 °C. The *Agrobacterium* was then plated on selective LB agar plates containing the relevant antibiotics for 2 days at 28 °C.

Arabidopsis transformation.

Agrobacterium vectors were validated by colony PCR and sequencing before growing in 100 ml LB medium containing 25 µg/ml gentamycin, 50 µg/ml rifampicin, and vector-specific antibiotic

for ~36 hours at 28 °C. *Agrobacteria* was harvested by centrifugation for 10 min at 4000 rpm, supernatant was discarded, and the bacteria pellet was resuspended in 40 ml 5% sucrose + $0.05 \times$ MS + 0.02% Silwet L-77. *Arabidopsis* flowers were dipped into the *agrobacterial* solution for ~5 minutes. After dipping, plants were kept in the dark for ~18 hours and grown until seeds were harvested. T1 seeds were sown on 0.5 × MS media containing the appropriate antibiotics for selecting transgenic plants or on soil with 0.1% basta spraying on ~10-day-old plants.

Genotyping.

T-DNA insertion lines for single mutants ordered from Gabi Kat (<u>https://www.gabi-kat.de</u>) and The *Arabidopsis* Information Resource (<u>https://www.arabidopsis.org/</u>) are listed in Table S9. Primers for the T-DNA insertion mutants genotyping were designed using the T-DNA Primer Design Tool powered by Genome Express Browser Server (GEBD) (<u>http://signal.salk.edu/</u> <u>tdnaprimers.2.html</u>). Homozygous mutants were characterized by PCR carried out with primers listed in Table S10.

Cloning.

ABCG17, ABCG18, and ABCG19 coding regions were amplified using Physican High-fidelity Polymerase (New England Biolabs) from Col-0 cDNA using primers listed in Table S11. Promoters of ABCG17 and ABCG18 were amplified from Col-0 DNA using Phusion High-fidelity Polymerase (New England Biolabs) using primers listed in Table S11. Promoters of ABCG17 and ABCG18 are 1276 and 1580 bp long respectively, including 5' UTR. ABCG17, ABCG18, and ABCG19 coding regions and as well as ABCG17 and ABCG18 promoter fragments were cloned into pENTR/D-TOPO (Invitrogen K2400), verified by sequencing, and subsequently cloned into binary destination vectors using LR Gateway reaction (Invitrogen 11791). p35S:YFP-ABCG17 and p35S:YFP-ABCG18 were generated using the pH7WGY2 vector and were selected using spectinomycin in E. coli and hygromycin in plants. p35S:ABCG17 and p35S:ABCG18 were generated using pH2GW7 vector and selected using spectinomycin in E. coli and hygromycin in plants. p35S:XVE:ABCG17 and p35S:XVE:ABCG18 were generated using pMDC7 vector and selected using spectinomycin in E. coli and hygromycin in plants. pABCG17:LUC and pABCG18:LUC were generated using the pFlash vector and selected using spectinomycin in E. coli and gentamycin in plants. pABCG17:NLS-YFP and pABCG18:NLS-YFP were generated using R1-R2:NLS-YFP in pART27 vector and selected using spectinomycin in E. coli and kanamycin in plants. *pABCG17:GUS* and *pABCG18:GUS* were generated using pWGB3 vector, kanamycin and hygromycin in *E. coli* and hygromycin in plants. *p35S:amiRNA* lines were generated using the pH2GW7 vector, spectinomycin in *E. coli* and hygromycin in plants. Using primers with restriction enzyme AscI, *pABCG17* and *pABCG18* were ligated into *ABCG17 CDS* and *ABCG18 CDS* using AscI site. LR gateway reaction was carried with the binary vector pGWB1. *pABCG17:ABCG17* and *pABCG18:ABCG18* were transformed into *mir18,g17* and *mir17,g18-1* respectively for phenotype complementation assays.

WMD3 website was used to design amiRNAs (<u>http://wmd3.weigelworld.org/cgi-bin/webapp.cgi</u>) (Table S12). The amiRNA sequences were synthesized by Syntezza Bioscience Ltd. and were cloned into the pUC57 vector with Gateway system borders, and cloned the amiRNAs into the pH2GW7 destination vector using the Gateway system.

Plant genetics.

All transgenic lines were generated from the destination vectors transformed into Col-0. T1 seeds were collected and selected on 1/2 MS plates containing appropriate antibiotics or on soil with basta spraying. At least 10 independent lines for each construct were generated. Two representative homozygous lines were obtained for further characterization. To generate overexpression lines in the background of the ABA reporters, the homozygous *ABCG17* or *ABCG18* overexpression lines were crossed with the ABA reporter lines *pRAB18:GFP*, *pMAPKKK18:GUS* or *pMAPKKK18:LUC*. F1 heterozygous seeds for both constructs were used for analysis. As a control, the respective ABA reporters were crossed with Col-0 to generate F1 seeds. To generate double-knockdown line *mir17,18* in the background of the ABA reporters, the homozygous of the ABA reporters, the homozygous of the ABA reporters, the homozygous seeds were used for analysis. As a control, the respective ABA reporters were crossed with Col-0 to generate F1 seeds. To generate double-knockdown line *mir17,18* in the background of the ABA reporters, the homozygous seeds were used for further analysis. As a control, the *pMAPKKK18:GUS* to obtain F1 seeds. Heterozygous seeds were used for further analysis. As a control, the *pMAPKKK18:GUS* ABA reporter was crossed with Col-0 to generate F1 seeds.

Phylogenetic tree.

A phylogenetic tree of *Arabidopsis* ABCG family members, based on protein sequences, was constructed using sequences alignment by Clustal W (<u>http://www.clustal.org/clustal2/e</u>). The one neighbor-joining phylogenetic tree was constructed using MEGA7.0 (Molecular Evolutionary Genetics Analysis) software with 1000 bootstrap replications.

Gas-exchange measurements.

Gas exchange was measured using a LI-COR LI-6800 portable gas-exchange system. Plants were grown in short-day conditions (8 h light/16 h dark). Photosynthesis was induced under 150 μ mol m⁻² s⁻¹ light with 400 μ mol mol⁻¹ CO₂ surrounding the leaves. The amount of blue light was set to 10% of the photosynthetically active photon flux density. The flow rate was set to 200 μ mol air s⁻¹, and the leaf-to-air vapor pressure deficit was kept around 1kPa during the measurement. Leaf temperature was ~22 °C. Measurements were performed between 10:00 AM and 1:00 PM (three hours after the light was switched on.).

Fluorescently labeled ABA synthesis.

The synthesis and characterization of ABA-FL are described in the supplemental information (Figs. S19-20).

Fluorescently labeled ABA protoplast experiments.

ABA-FL was dissolved in DMSO to 5 mM stock solution. Protoplasts isolation was carried as previously described (70) with the following adaptations: Protoplasts were treated with ABA-FL for 12 hours at a final concentration of 5 μ M ABA-FL, washed with 500 μ l W5 solution (2 mM solution (pH 5.7) containing 154 mM NaCl, 125 mM CaCl₂ and 5 mM KCl) three times, resuspended in 100 μ l W5 solution, and mounted on slides for confocal microscopy.

Stomatal aperture measurements.

Stomatal aperture was determined using a rapid and almost permanent imprinting technique previously described (71). Leaves of similar size were cut from 25-day-old plants. The abaxial side of the leaves was attached to 0.5 ml silicone impression material (elite HD+, Zhermack Clinical). The leaves were removed after the silicone impression material dried. Subsequently, transparent nail varnish was placed on the epidermal side. The samples were placed on slides with cover-slips and imaged with a light microscope (71). The stomatal aperture size was quantified using Fiji software.

RNA extraction, cDNA synthesis, and quantitative PCR.

RNA was isolated using SV Total RNA Isolation System (Promega). For cDNA synthesis, a High-Capacity cDNA Reverse Transcription Kit (Applied Biosystems) was used according to manufacturer information. Fast SYBR Green Master Mix (Applied Biosystems) was used for quantitative PCR as described before (72). The primers used for quantitative PCR are listed in Table S13.

Thermal imaging

Thermal images were obtained using a FLIR T6xx series instrument. The camera was placed vertically ~50 cm above the plants. Leaf temperature was quantified using a customized region-of-interest tool implemented according to the manufacturer's instructions.

Histology, microscopy, and phenotypic characterization.

Sections were made using the Vibratome technology as previously described (73). Sections were stained with 0.01% (g/ml) Fluorescent Brightener 28 in 1 x PBS for 20 minutes, washed with 1 x PBS three times (10 min each wash), and subsequently imaged with a Zeiss LSM780 confocal microscope. Plant shoot images were taken using a Nikon camera (WEM ED IF Aspherical Micro 1:1 Φ 62). For root growth assays, seedlings were grown on plates and scanned using HP Scanjet G3110. Leaf size and root length were measured using Fiji software (https://fiji.sc/).

GUS staining and histology.

Histochemical detection of GUS activity was carried out using 5-bromo-4-chloro-3-indolyl- β -D-glucuronide as a substrate as previously described (50). Samples were placed on slides with glass cover-slips and imaged using Zeiss binocular microscope. Histology of GUS stained samples was performed using protocol as described previously (74).

Leaf water-loss assay.

Leaves of 30-day-old plants were cut and weighed immediately to determine the fresh weight. Leaves were then exposed to room temperature air and weighed at 30 min, 60 min, 90 min, 120 min, 180 min, and 480 min.

Bioluminescence LUC assays.

ABCG17 and *ABCG18* promoters were cloned into the pFlash vector using the Gateway system to generate *promoter:LUC* reporter constructs. Constructs were then transformed into Col-0 plants, and homozygous transgenic plants were treated with 10 mg/ml luciferin (Promega) and placed in the dark for 5 min. The LUC signal intensity was subsequently detected using the Biospace system (camera: IS1643N7056).

Lateral root primordium and bolting time quantifications.

Five-day-old seedlings were treated with 100 mM NaCl for 5 more days (control plants were grown on regular MS media), lateral root primordium number of 10-day-old seedlings at different developmental stages were monitored. The seedlings were fixed in fresh 4% PFA (4 grams of PFA

powder dissolved into 100 ml 1 X PBS solution), with 0.01% Triton in vacuum for 1 h. Samples were washed 3 times with 1 X PBS, each time 3 min. The solution was removed and ClearSee solution was added (10% Xylitol, 25% Urea, 15% Sodium Deoxycholate in water, Sodium Deoxycholate was added at last) for at least 3 days at room temperature. Samples were placed on slides and lateral root primordium at different developmental stages were monitored using a Zeiss fluorescent binocular Axio Zoom V16.

For plant bolting time quantification, a standard 5 mm stem inflorescence was set as a bolting point. Plants were monitored every day to mark the bolting point.

Grafting assays.

Six-day-old plants were used for grafting assay according to procedures described previously (75). Both of the two cotyledons were cut from seedlings, media used in the grafting assay contained 0.5% sucrose, and grafted plants grew vertically in short-day conditions.

Radioactive ABA translocation and transport assays.

ABA long-distance shoot-to-root translocation was measured using [³H]ABA. Shoots were treated with 2 μ l of 0.1 μ Ci/ml [³H]ABA. After 24 h, roots were excised and placed into 3 ml of scintillation liquid (Opti-Fluor) for at least 24 h in darkness before monitoring in a liquid scintillation counter.

Simultaneous [¹⁴C]ABA and [³H]IAA from tobacco (*N. benthamiana*) mesophyll protoplasts were analyzed as described (76). Tobacco mesophyll protoplasts were prepared 4 days after agrobacterium-mediated transfection of *p35S:YFP-ABCG17*, *p35S:YFP-ABCG18*, *p35S:YFP-ABCG19* or empty vector control. Relative export from protoplasts is calculated from exported radioactivity into the supernatant as follows: (radioactivity in the supernatant at time t = x min.) -(radioactivity in the supernatant at time t = 0)) * (100%)/ (radioactivity in the supernatant at t = 0 min.); presented are mean values from 8 ([³H]ABA) and 4 ([¹⁴C]IAA) independent transfections.

Xenopus oocytes transport assay

For cRNA synthesis Coding DNA sequences of the ABCG genes were cloned into *Xenopus laevis* oocyte expression vector pNB1u using USER cloning as described previously (77). DNA template for in vitro transcription was generated by PCR using Phusion High-Fidelity DNA Polymerase (NEB) using forward primer (5'– AATTAACCCTCACTAAAGGGTTGTAATACGACTCACTATAGGG – 3') and reverse primer

X. laevis oocytes, stage V or VI, were purchased from Ecocyte Bioscience (Germany). Oocytes were injected with 25 ng cRNA for each transporter expressed, i.e. regardless of whether the transporters were expressed individually or in combination. cRNA was injected using a Drummond NANOJECT II (Drummond Scientific Company, Broomall Pennsylvania). The injected oocytes were incubated for three days at 16 °C in HEPES-based kulori buffer (90 mM NaCl, 1 mM KCl, 1 mM MgCl₂, 1 mM CaCl₂, 5 mM HEPES pH 7.4) supplemented with gentamycin (100 μ g/mL).

Three days after cRNA injection, transport assay was performed as described previously (78), with some modifications. Oocytes were preincubated in Kulori pH 5.8 for 5 min and then the oocytes were incubated in 10 uM ABA containing Kulori buffer (pH 5.8) for 30 min. Subsequently, oocytes were washed 4 times in Kulori buffer (pH 5.8) and then homogenized with 50 % methanol and stored at -20 °C overnight. Oocyte extracts were spun down at 15000 X g for 10 min at 4 °C, supernatant was spun down at 10000 X g for 10 min at 4 °C. The supernatant was then diluted with water and filtered through a 0.22 μ m PVDF-based filter plate (MSGVN2250, Merck Millipore) and analyzed by analytical liquid chromatography coupled to mass spectrometry (LC-MS/MS) detailed in the Supplemental data.

Hormone quantification.

Plant samples frozen in liquid nitrogen were grounded using motor and pestle. Around 200 mg of samples (shoot or root) was measured from ground powder and extracted with ice cold methanol/water/formic acid (15/4/1 v/v/v) added with isotope labelled internal standards (IS). Similar concentrations of internal standards (IS) of ABA and its metabolites were added into samples and calibration standards. The samples were purified using Oasis MCX SPE cartridges (Waters) according to manufacturer's protocol. The samples were injected on Acquity UPLC BEH C18 column (1.7 μ m, 2.1x100 mm, Waters; with gradients of 0.1% acetic acid in water or acetonitrile), connected to Acquity UPLC H class system (Waters) coupled with UPLC-ESI-MS/MS triple quadrupole mass spectrometer for identification and quantification of hormones. The hormones were measured using MS detector, both in positive and negative mode, with two

MRM transitions for each compound. External calibration curves that were constructed with serial dilutions of hormone standards, added with internal standards, were used for quantification, and calculated through Target Lynx (v4.1; Waters) software by comparing the ratios of MRM peak areas of analyte to that of internal standard (endogenous levels). To evaluate reproducibility and linearity of the method, parameters such as accuracy and precision were measured from five replicates for each of *Arabidopsis thaliana* (Col-0) shoot and root samples, and were spiked with three different concentrations of non-labelled hormone standards (low/ medium/ high), along with added internal standards. Accuracy was measured after comparing the known concentration of standard with that of endogenous hormone levels and spiked samples. LODs and LOQs were given as signal to noise ratios of 3:1 and 10:1 respectively. The repeatability of the method, known as precision was calculated in terms of percent relative standard deviation (RSD %; Table S4). Heat map was generated using the BAR HeatMapper Plus tool (http://bar.utoronto.ca/ntools/cgi-bin/ntools_heatmapper_plus.cgi).

Statistical analysis.

Multiple comparisons were performed by using one-way ANOVA with the least significant difference (LSD) Post Hoc test in SPSS 19.0. Two-tailed Student's t-test was used for graphs with only two groups. Statistical significance was determined at P < 0.05 unless otherwise claimed.

Data availability

All the data supporting the findings of this study are available within the article and Supplementary Information files or from the corresponding author upon reasonable request.

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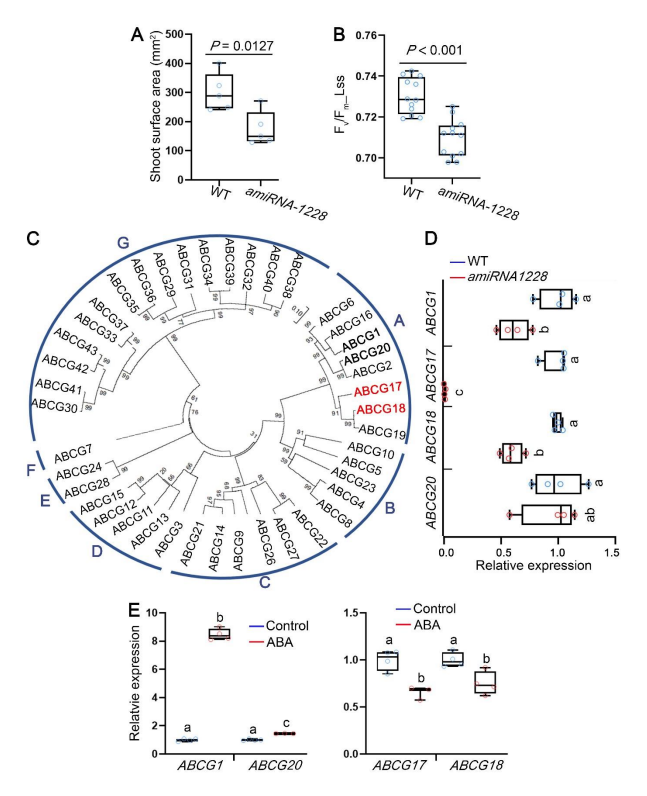


Fig. 1. Reduced expression of several class A ABCG members results in moderate shoot growth and decreased photosynthesis rate. (*A*) Shoot surface area of 18-day-old WT and *amiRNA-1228*. Shown are averages (\pm SD), n = 5, P value indicates significant differences, student's t-test. (*B*) Fv/Fm_Lss, indicating for photosystem II quantum yield of light-adapted samples at steady state, measured for 18-day-old WT

and *amiRNA-1228*. Shown are averages (\pm SD), n = 12, P < 0.001 (P = 1.15208e-05) indicates significant differences, student's t-test. (*C*) Phylogenetic tree of *Arabidopsis* ABCG family based on amino acid sequences. Red and bold fonts indicate proteins coded by putative *amiRNA-1228* target genes. (*D*) Relative expression of the indicated *ABCG amiRNA-1228* targeted genes in 12-day-old WT and *amiRNA-1228* seedlings, quantified by qRT-PCR. Shown are averages (\pm SD), n = 4, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*E*) Relative expression, quantified by qRT-PCR, of the indicated *ABCG amiRNA-1228* targeted genes in response to ABA treatment (5 µM ABA for 3 hours) in 7-day-old seedlings. Shown are averages (\pm SD), n = 4, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05).

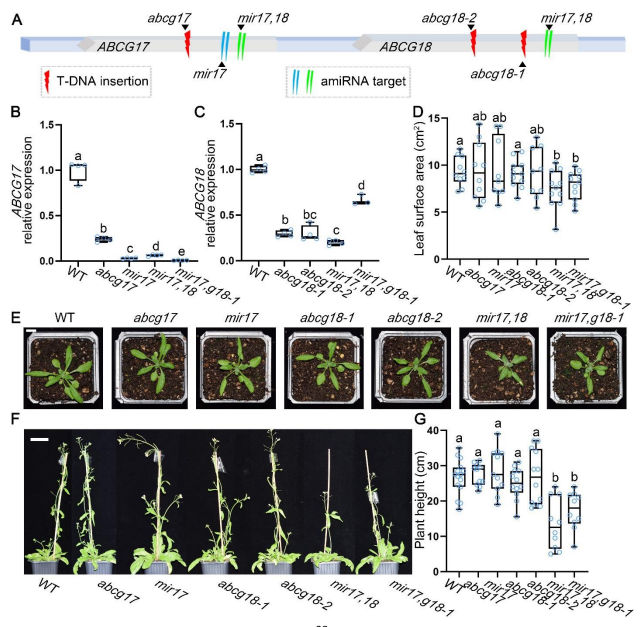


Fig. 2. *ABCG17* and *ABCG18* redundantly regulate plant growth. (*A*) Illustration of the *ABCG17* and *ABCG18* genome region. T-DNA insertions and amiRNA knockdown sites are indicated. (*B-C*) Relative expression levels of *ABCG17* (*B*) and *ABCG18* (*C*) in the indicated genotypes. *mir17,18* is the *abcg17*, *abcg18* double-knockdown amiRNA line; *mir17,g18-1* is *mir17* (*amiRNA-ABCBG17*) transformed into the background of *abcg18-1* T-DNA insertion line. Shown are averages (±SD), n = 4, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*D*) Leaf surface area of the indicated lines. Shown are averages (±SD), n ≥ 9 plants, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*E*) Shoot phenotypes of 25-day-old plants grown in soil under normal conditions. Scale bar = 1 cm. (*F*) Phenotypes of 50-day-old plants grown in soil under normal conditions. Scale bar = 2 cm. (*G*) Quantification of inflorescence stem height of the indicated genotypes. Shown are averages (±SD), n ≥ 10 plants, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05).

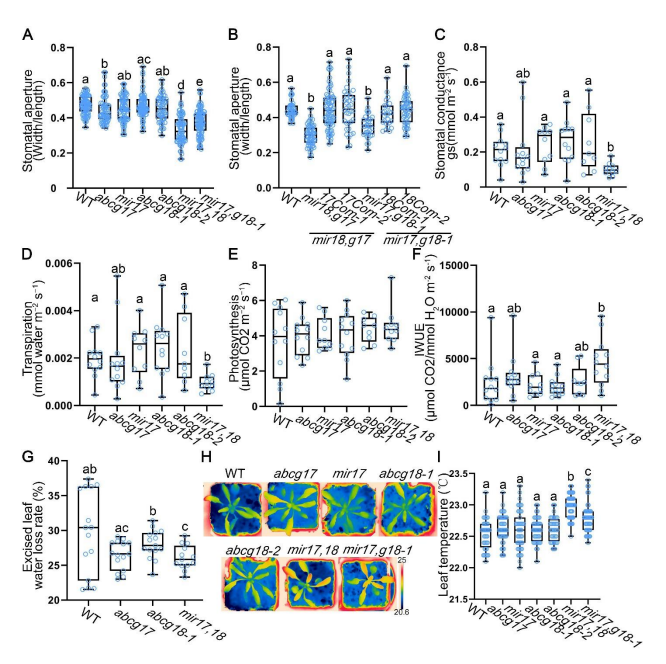


Fig. 3. Stomatal aperture, conductance, and transpiration is regulated by *ABCG17* and *ABCG18*. (*A*) Stomatal aperture measurements of 25-day-old plants of the indicated genotypes. *mir17,18* is *abcg17*, *abcg18* double-knockdown amiRNA line, *mir17,g18-1* is *mir17* (*amiRNA-ABCBG17*) transformed into the background of *abcg18-1* T-DNA line. Shown are averages (±SD), $n \ge 45$, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*B*) Thirty-day-old plants stomatal aperture complementation lines (*pABCG17:ABCG17* or *pABCG18:ABCG18*). Shown are averages (±SD), $n \ge 26$, different letters represent significant differences, one-way ANOVA with student differences, one-way ANOVA with student's t-test (P < 0.05). (*C*) Plant stomatal conductance measurements of 25-day-old plants of the indicated genotypes. Shown are averages (±SD), n = 5 plants, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*D*) Leaf transpiration rates of the 25-day-old plants of the

indicated genotypes. Shown are averages (\pm SD), n = 5 plants, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*E*) Leaf photosynthetic properties of 25-day-old plants of the indicated genotypes. n = 5 plants. Results were not significant at P > 0.05 by one-way ANOVA with student's t-test (P < 0.05). (*F*) Instantaneous water use efficiency (IWUE) of plants of the indicated genotypes under normal conditions. Shown are averages (\pm SD), n = 5 plants, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*G*) Water-loss rates of leaves excised from 30-day-old plants of the indicated genotypes (90 minutes, room temperature air). Shown are averages (\pm SD), n \geq 13, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*H*) Thermal images of 25-day-old plants of the indicated genotypes grown on soil under normal conditions. Color-coded scale bar indicates for temperature. Black scale bar = 1 cm. (*I*) Leaf temperatures of plants of the indicated genotypes measured at sites indicated in panel (H). Shown are averages (\pm SD), n \geq 40, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05).

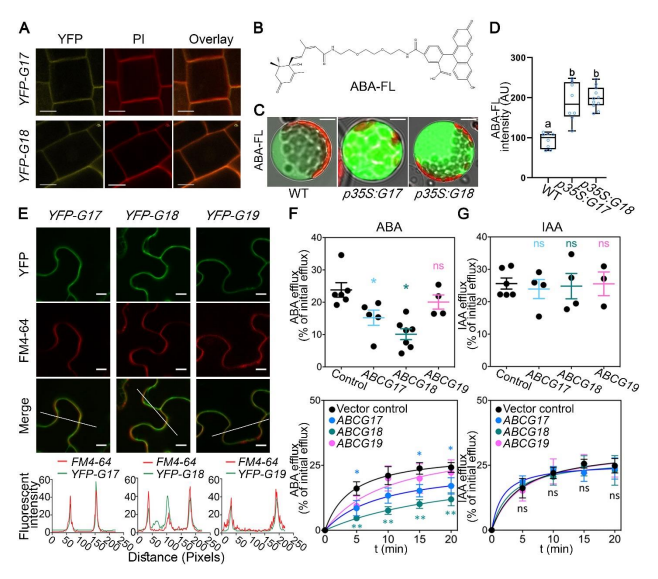


Fig. 4. ABCG17 and ABCG18 are ABA importers localized to the plasma membrane. (A) ABCG17 and ABCG18 subcellular localization shown by p35S:YFP-ABCG17 (YFP-G17) and p35S:YFP-ABCG18 (YFP-G18) stable transgenic Arabidopsis lines. Fluorescence was imaged in the root meristem epidermis layer. Yellow indicates YFP-ABCG17 or YFP-ABCG18 fluorescence, red indicates propidium iodide (PI). Scale bar = $5 \mu m$. (B) Molecular structure of ABA-FL, a fluorescein moiety is linked to the ABA carboxylic group. (C) ABA-FL fluorescent signal in Arabidopsis protoplasts overexpressing ABCG17 or ABCG18. p35S:G17 and p35S:G18 indicate p35S:ABCG17 and p35S:ABCG18, respectively. Protoplasts were treated with 5 μ M ABA-FL for 15 min. Scale bar = 10 μ m. (D) Quantification of ABA-FL fluorescence intensity in protoplasts overexpressing ABCG17 or ABCG18. Shown are averages (\pm SD), n \geq 8 protoplasts, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (E) Confocal imaging of tobacco leaves transfected with p35S:YFP-ABCG17 (YFP-G17), p35S:YFP-ABCG18 (YFP-G18) or p35S:YFP-ABCG19 (YFP-G19), stained with FM4-64 plasma membrane marker. Scale bar = 10 μm. Bottom graphs are quantified fluorescence intensities of pixels along the white line within the images. X-axis represents distance along the line. (F) $[^{14}C]ABA$ efflux from tobacco protoplasts as percentage of initial efflux. Top graph is 15 min [¹⁴C]ABA, $n \ge 4$, *P < 0.05, Welch's t-test. (G) [³H]IAA efflux from tobacco protoplasts as percentage of initial efflux. Top graph is 15 min [³H]IAA, $n \ge 4$, ns indicates not significant.

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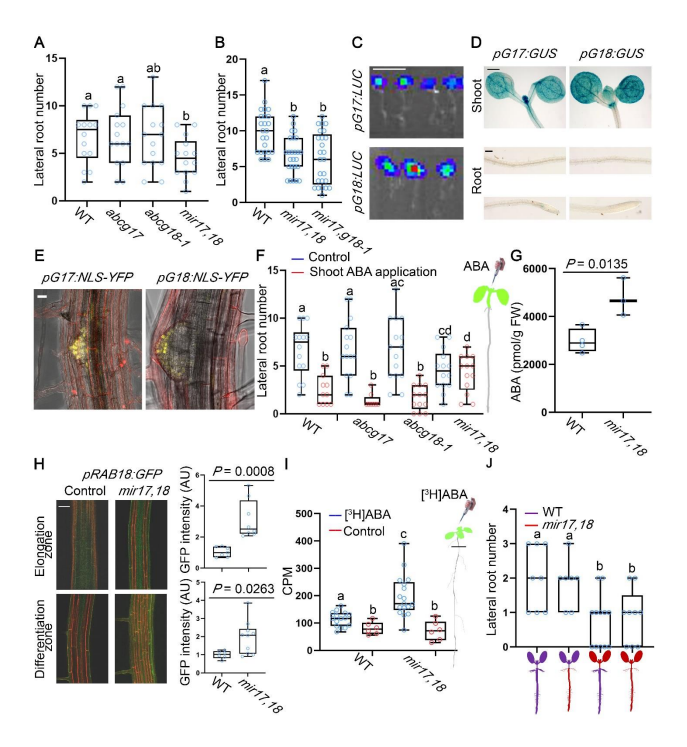


Fig. 5. ABCG17 and ABCG18 are required for long-distance ABA transport and regulation of lateral root formation. (*A*) Lateral root numbers of 10-day-old seedlings for indicated genotypes. *mir17,18* is the *abcg17, abcg18* double-knockdown amiRNA line. Shown are averages (±SD), $n \ge 14$, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). ns indicates for not significant. (*B*) Lateral root numbers of 10-day-old plants for indicated genotypes. *mir17,18* is the *abcg17, abcg18* double-knockdown amiRNA line. *mir17,g18-1* is *mir17 (amiRNA-ABCBG17)* transformed into the background of *abcg18-1* T-DNA insertion line. Shown are averages (±SD), $n \ge 18$, different letters

represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (C) Bioluminescence signal driven by pABCG17:LUC (pG17:LUC) and pABCG18:LUC lines (pG18:LUC). Scale bar = 1 cm. (D) GUS staining for pABCG17:GUS (pG17:GUS) and pABCG18:GUS (pG18:GUS) in shoot and root of 5-day-old seedlings. Scale bar = 0.5 mm for shoots; scale bar = 0.1 mm for roots. (E) Representative images of pABCG17:NLS-YFP (pG17:NLS-YFP) and pABCG18:NLS-YFP (pG18:NLS-YFP) lateral root emergence sites (YFP in yellow). Scale bar = $20 \,\mu m$. (F) Lateral root numbers for the indicated genotypes. Shoots of 4-day-old plants were treated with 5 μ M ABA for 4 days. Shown are averages (±SD), n \geq 13, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). To the right is an illustration of shoot-specific ABA application. (G) Quantification of ABA amounts in roots of indicated genotypes (12-day-old plants). FW: fresh weight. Shown are averages (\pm SD), n \geq 3, P value indicates significant differences, student's t-test. (H) pRAB18:GFP signal in the background of miR17,18 and control roots (left) and the quantification of respective GFP signal intensity (right). Scale bar = $20 \mu m$. Shown are averages (\pm SD), n \geq 7, P value indicates significant differences, student's t-test. (1) [³H]ABA counts per minute (CPM) in roots only (roots were isolated as indicated by a black line in the illustration on the right). Controls were samples not treated with 0.1 µCi/ml [³H]ABA. Shown are averages (±SD), n \geq 10, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (J) Lateral root number quantification of the indicated grafted lines, grafting performed on 6-day-old seedlings, lateral roots were scored 8 days after grafting. mir17,18 is abcg17, abcg18 double-knockdown amiRNA line. Shown are averages (\pm SD), n \geq 7, different letters represent significant differences, oneway ANOVA with student's t-test (P < 0.05).

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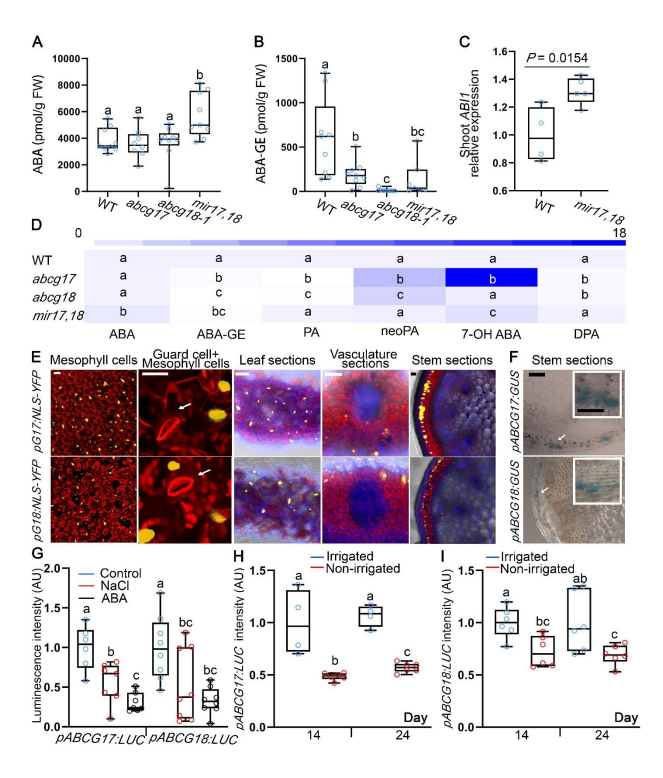


Fig. 6. *ABCG17* and *ABCG18* are primarily expressed in mesophyll and cortex cells, allowing ABA-GE formation under normal conditions. (*A-B*) Quantification of ABA (*A*) and ABA-GE (*B*) in shoots of 12-day-old plants of the indicated genotypes. FW: fresh weight. Shown are averages (±SD), $n \ge 7$, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*C*) Expression of *ABI1* in 12-day-old WT and *mir17,18* seedlings, quantified by qRT-PCR. Shown are averages (±SD), n

= 4, P value indicates significant differences, student's t-test. (*D*) ABA metabolites profile heat-map in shoots of 12-day-old plants of the indicated genotypes. Absolute values and abbreviations are presented in Fig. S14. Shown are averages (±SD), $n \ge 7$, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*E*) *pABCG17:NLS-YFP* (*pG17:NLS-YFP*) and *pABCG18:NLS-YFP* (*pG18:NLS-YFP*) signal (yellow) in leaves and stem. YFP signal is detected in mesophyll and cortex cells. Chlorophyll in red. Fluorescent brightener in blue. White arrows point to the guard cells. Scale bars = 20 µm. (*F*) GUS staining for *pABCG17:GUS* (*pG17:GUS*) and *pABCG18:GUS* (*pG18:GUS*) in inflorescence stem of 30-day-old plants. Scale bar = 100 µm. Insets are magnifications of areas indicated by white boxes. Scale bar = 50 µm. (*G*) Luminescence intensity of *pABCG17:LUC* (*pG17:LUC*) and *pABCG18:LUC* (*pG18:LUC*) with and without 5 µM ABA or 100 mM NaCl treatment. Shown are averages (±SD), $n \ge 6$, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05). (*H-1*) Luminescence intensity of *pABCG17:LUC* (*pG17:LUC*) (*H*) and *pABCG18:LUC* (*pG18:LUC*) (*I*). Plants were irrigated for 15 days following by water withhold for 3 weeks. Control plants (irrigated) were watered throughout the experiment. Shown are averages (±SD), $n \ge 4$, different letters represent significant differences (±SD), n ≥ 4, different letters represent significant differences (±SD), n ≥ 4, different letters represent significant differences (±SD), n ≥ 4, different letters represent significant differences, one-way ANOVA with student's t-test (P < 0.05).

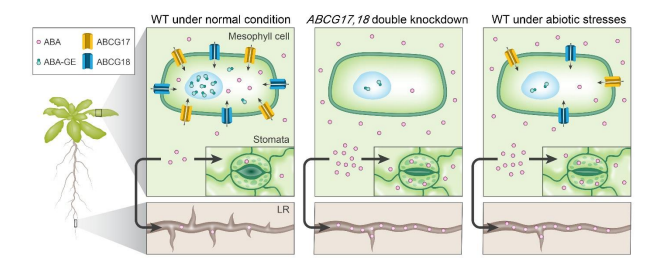


Fig. 7. Proposed model illustrating ABCG17 and ABCG18 function in regulating ABA homeostasis under normal and abiotic stress conditions. LR, lateral root.