ATP-Dependent Binding Cassette Transporter G Family Member 16 Increases Plant Tolerance to Abscisic Acid and Assists in Basal Resistance against Pseudomonas syringae DC30001[W][OPEN]

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Plants have been shown previously to perceive bacteria on the leaf surface and respond by closing their stomata. The virulent bacterial pathogen Pseudomonas syringae pv tomato DC3000 (PstDC3000) responds by secreting a virulence factor, coronatine, which blocks the functioning of guard cells and forces stomata to reopen. After it is inside the leaf, PstDC3000 has been shown to up-regulate abscisic acid (ABA) signaling and thereby suppress salicylic acid-dependent resistance. Some wild plants exhibit resistance to PstDC3000, but the mechanisms by which they achieve this resistance remain unknown. Here, we used genome-wide association mapping to identify an ATP-dependent binding cassette transporter gene (ATP-dependent binding cassette transporter G family member16) in Arabidopsis (Arabidopsis thaliana) that contributes to wild plant resistance to PstDC3000. Through microarray analysis and β-glucuronidase reporter lines, we showed that the gene is up-regulated by ABA, bacterial infection, and coronatine. We also used a green fluorescent protein fusion protein and found that transporter is more likely to localize on plasma membranes than in cell walls. Transferred DNA insertion lines exhibited consistent defective tolerance of exogenous ABA and reduced resistance to infection by PstDC3000. Our conclusion is that ATP-dependent binding cassette transporter G family member16 is involved in ABA tolerance and contributes to plant resistance against PstDC3000. This is one of the first examples, to our knowledge, of ATP-dependent binding cassette transporter involvement in plant resistance to infection by a bacterial pathogen. It also suggests a possible mechanism by which plants reduce the deleterious effects of ABA hijacking during pathogen attack. Collectively, these results improve our understanding of basal resistance in Arabidopsis and offer unique ABA-related targets for improving the innate resistance of plants to bacterial infection.

Bacterial pathogens of plants disperse through the global hydrological cycle (Morris et al., 2008) and are likely to exert selection favoring defenses of wild plants at large spatial scales. In agriculture, these pathogens cause significant losses and necessitate extensive spraying of pesticides (Oerke, 2006). Although there is great potential to learn the mechanisms that underlie this resistance (Todesco et al., 2010), the genetic underpinnings of resistance in wild plants remain essentially unknown. Identification of natural mechanisms by which resistant wild plants prevent the growth of bacterial pathogens has the potential to improve crop yields and reduce exogenous pesticide use in agriculture.

The immune system of plants is sophisticated, involves the coordinated action of hundreds of genes (Jones and Dangl, 2006), and requires the presence of a single major hormone, salicylic acid (SA; Vlot et al., 2009). This pathway is cross regulated by other signaling pathways in the plant, including those controlled by jasmonic acid (Vlot et al., 2009) and abscisic acid (ABA; de Torres-Zabala et al., 2007). However, how and to what extent plants can regulate these hormonal-dependent pathways and cross talk is still not clear. SA-dependent defenses are effective against many bacterial pathogens, including the causal agents of bacterial spot disease, Pseudomonas syringae (Vlot et al., 2009). In some cases, plants perceive bacteria on the surface of the leaf and close the stomata, thereby preventing entry (de Torres-Zabala et al., 2007). However, at least one particularly virulent strain, P. syringae pv tomato (PstDC3000), has evolved a novel virulence mechanism, in which it secretes coronatine to force stomata to reopen (de Torres-Zabala et al., 2007; Montillet et al., 2013). After bacteria have entered the interior of a leaf, these virulent bacteria also use coronatine to hijack plant ABA biosynthesis, which elevates ABA levels and suppresses Isochorismate synthase1 expression, leading to down-regulation of SA-dependent defenses in the infected tissue (de Torres-Zabala et al., 2009; Zheng et al., 2012). What has remained unknown is whether plants can counteract this suppression and if so, how.

ATP-dependent binding cassette (ABC) transporters have received particular attention recently for their...
importance in moving hormones, metal ions, and other compounds across membranes (Rea, 2007; Kang et al., 2011). The largest subfamily of ABC transporters in Arabidopsis (Arabidopsis thaliana), the Arabidopsis ATP-dependent binding cassette transporter G Family (AtABCG) group, contains 43 genes that share a common nucleotide-binding domain and transmembrane domain orientation (Verrier et al., 2008). To date, three members of this AtABCG family (AtABCG22, AtABCG25, and AtABCG40) have been associated with ABA transport directly. AtABCG22 and AtABCG40 are found on the guard cells specifically, and AtABCG25 is found on the vascular bundles (Kang et al., 2010; Kuromori et al., 2010, 2011). AtABCG36 Penetration Resistance3 (PEN3)/Pleiotropic Drug Resistance8 (PDR8) is the only gene in AtABCG subfamily that has been reported for involvement in plant resistance to bacterial or fungal pathogens (Kobae et al., 2006; Stein et al., 2006; Xin et al., 2013). The functions of other AtABCGs and particularly, their possible roles in ABA transport or plant-pathogen interaction remain largely unknown. Here, we present evidence that one of the AtABCG transporters, AtABCG16, identified by genome-wide association (GWA) mapping is involved in ABA responses and plant resistance against PstDC3000.

RESULTS
GWA Mapping and Microarray Analysis Suggest the Involvement of AtABCG16 in Plant Resistance to Bacterial Pathogen PstDC3000

To identify candidate genes that contribute to differences in defense against bacterial infection in wild plants, we assessed a GWA map (Fig. 1A) constructed from 96 genotypes of the RegMap panel of Arabidopsis (Supplemental Table S1) consisting of a worldwide collection of plants for which over 200,000 single nucleotide polymorphisms (SNPs) have been identified (Atwell et al., 2010). GWA mapping has emerged as a particularly powerful method for identifying novel genes, because it uses genetic polymorphisms that are already present in wild populations and therefore, is likely to pinpoint genetic variation that is important in nature (Weigel, 2012). To assess resistance to infection, those plants were grown on soil and challenged by blunt syringe with a 10⁴ colony-forming unit (cfu) mL⁻¹ solution of PstDC3000 as described previously (Atwell et al., 2010). Our assessment of the GWA map revealed 13 major loci (Fig. 1A) and 54 candidate genes (Supplemental Table S2), of which one of the top candidates was an ABCG transporter (AtABCG16) at locus 10 on chromosome 3 of Arabidopsis.
3; Fig. 1B). This locus also included two additional gene candidates, At3g55070 and At3g55080, that contained equally significant SNPs (Supplemental Tables S3 and S4) but about which little is known.

To better understand possible defense-related functions at this locus, we first mined published microarray data for the widely used ecotype Columbia-0 of Arabidopsis (Col-0) accession and found that AtABCG16 had a 2.76-fold up-regulation 2 h after exposure to PstDC3000 that diminished to 0.72-fold at 24 h after exposure (Fig. 1C). The early response to bacteria is consistent with functional role in the initial stages of plant responses to infection. Given the key roles of SA, ABA, indole-3-acetic acid (IAA), and methyl jasmonate (MeJA) in plant signaling and known transport roles of ABCG transporters in moving these compounds (Dean and Mills, 2004; Vlot et al., 2009; Kuromori et al., 2010), we then asked whether AtABCG16 expression was altered after exogenous application of these hormones to plants. We found that AtABCG16 was induced 3.8-fold and 3.5-fold by 10 μM exogenous SA and IAA, respectively, and not induced by MeJA (Fig. 1D). In contrast, AtABCG16 was up-regulated nearly 40-fold by ABA. Indeed, this was the 34th highest response of 22,591 Arabidopsis genes, and this gene was the only one within the 250-kb neighborhood to exhibit a response (Fig. 1D; Supplemental Table S5). Furthermore, tissue-specific microarrays showed that AtABCG16 was up-regulated 65-fold in mesophyll cells and nearly 20-fold in guard cells treated with exogenous ABA (Fig. 1E). Collectively, these data showed that this candidate gene responds to PstDC3000 and responds even more strongly to exogenous ABA.

Transferred DNA Insertion Mutants of AtABCG16 Show a Loss of ABA Tolerance in Plate Assays

To assess experimentally whether AtABCG16 is required for plant tolerance of exogenous ABA, SA, or IAA, we used a standard hormone assay on Murashige and Skoog medium (MS) plates (Kuromori et al., 2010) to compare the tolerance of two transferred DNA (T-DNA) insertion knockout lines relative to the background (Col-0) in experiments where all three lines were grown simultaneously. To characterize the knockouts, we first ran quantitative reverse transcription (qRT)-PCR (Fig. 2A; Supplemental Tables S6 and S7) and found that they reduced the AtABCG16 gene expression by 92% (SALK_087501; hereafter abcg16-1) and 84% (SALK_119868C; hereafter abcg16-2), respectively. Because SA and IAA likely operate at different concentrations from ABA in planta, we included both low-range (1 and 3 μM) and high-range (100 and 300 μM) tests of ABA, SA, and IAA. Through the plate assay, we found
that both abc16-1 and abc16-2 had significantly reduced germination in the presence of exogenous ABA relative to the Col-0 background (\(F_{4,45} = 18.49, P < 0.0001;\) Fig. 2B; Supplemental Table S8), whereas no such response was detected in the presence of IAA (Fig. 2C; Supplemental Table S8) or SA (Fig. 2D; Supplemental Table S8). The loss of ABA tolerance was dramatic. Specifically, in the presence of 3 \(\mu\)M exogenous ABA, the abc16-1 line had a 94.8% reduction in germination (Dunnett \(T = -8.7, P < 0.001\)) and the abc16-2 line had a 43.5% reduction in germination (Dunnett \(T = -4.0, P = 0.002\)) relative to the background. Our conclusion from these assays was that \(AtABCG16\) was necessary for successful germination in the presence of exogenous ABA but not SA or IAA at the concentrations tested.

To further assess the tolerance of abc16-1 and abc16-2 to exogenous ABA, we also tested these lines across a smaller range of 0 to 1.5 \(\mu\)M exogenous ABA using an established protocol (Kuromori et al., 2010), and again, we found highly significant defects in the tolerance of both knockout lines relative to the Col-0 background in terms of reduced germination (\(F_{6,108} = 3.71, P = 0.002;\) Fig. 2E; Supplemental Table S9), reduced root growth (\(F_{6,108} = 7.03, P < 0.001;\) Fig. 2, F and H; Supplemental Figs. S1 and S2; Supplemental Table S9), and elevated mortality of the knockouts (\(F_{6,103} = 3.92, P = 0.001;\) Fig. 2G; Supplemental Table S9). To assess the possibility of off-target effects of the T-DNA insertions in the abc16-1 and abc16-2 lines, we measured the expression of \(AtABCG17\) and \(AtABCG18\), which are directly adjacent to \(AtABCG16\). We found no evidence of off-target effects of the T-DNA insertions (Supplemental Fig. S3), which was indicated by consistent expression of \(AtABCG17\) and \(AtABCG18\) in abc16-1 and abc16-2 relative to the Col-0 background line (CS60000). Collectively, these experiments showed that knockouts of \(AtABCG16\) were defective in ABA tolerance, that the response was present at multiple stages of plant development, including germination and subsequent growth and survival of germinated seeds, and that the defects were likely to be caused specifically by the reduced expression of \(AtABCG16\) (Fig. 2A) in these T-DNA insertion knockout lines.

**AtABCG16 Potentially Localizes to the Plasma Membrane**

To localize \(AtABCG16\), we use a construct consisting of the cauliflower mosaic virus 35S promoter driving the GFP fused to \(AtABCG16\) (2X35S::GFP-\(AtABCG16\)). Subcellular localization of the fusion protein was observed using the filtered epifluorescence microscopy imaging of the green fluorescence signals in cells of the 2X35S::GFP-\(AtABCG16\)-transformed plants (Fig. 3). In root cells, fluorescence was highest on the cell surface, which was consistent with GFP-\(AtABCG16\) localization in the plasma membrane or cell wall (Fig. 3A; Supplemental Fig. S4). To assess the hypothesis that the protein was in the cell wall, we treated the root tip cells with 20% sucrose. A to H show root cells. A, B, E, F, I, and J show plants transformed with 35S::GFP-\(AtABCG16\). C, D, G, H, K, and L show untransformed plants with at least 2 times exposure time. E to H show plasmolyzed root cells after treatment with 20% Suc. I to L show leaf epidermal cells. Red arrows indicate stomata. Positive controls are shown in Supplemental Figure S4. Bars = 10 \(\mu\)m.

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**Figure 3.** Plasma membrane localization of 2X35S::GFP-\(AtABCG16\) fusion protein. A, C, E, G, I, and K show fluorescence levels under GFP expression conditions. B, D, F, H, J, and L were taken under white light. A to H show root cells. A, B, E, F, I, and J show plants transformed with 35S::GFP-\(AtABCG16\). C, D, G, H, K, and L show untransformed plants with at least 2 times exposure time. E to H show plasmolyzed root cells after treatment with 20% Suc. I to L show leaf epidermal cells. Red arrows indicate stomata. Positive controls are shown in Supplemental Figure S4. Bars = 10 \(\mu\)m.
with 20% Suc to create a high osmotic environment following a published protocol (Kuromori and Shinozaki, 2010). On the osmotic treatment cells, the fluorescence signal was observed to associate more with the cell interior and plasma membrane than on the cell wall (Fig. 3, E and F). In leaf cells, we observed fluorescence signals in both pavement cells and guard cells (Fig. 3, I and J). Collectively, these results are consistent with localization of AtABCG16 on the plasma membrane and seem to rule out localization on the cell wall but cannot rule out localization on vacuolar membranes or elsewhere in the cell interior.

**AtABCG16 Gene Expression Is Induced by Hormones and Bacteria**

To further investigate how this gene is regulated, we created transgenic lines containing the GUS reporter gene driven by a native *AtABCG16* promoter (Fig. 4A). This native promoter of *AtABCG16* (Fig. 4B) contains important binding motifs for the ABA response factor, ABA Response Element, at −370, −250, and −80 as well as a binding site at −470 for WRKY38, a transcription factor directly linked to infection by *PstDC3000* (Kim et al., 2008). Our *AtABCG16 pro:GUS* transgenic lines exhibited inducible GUS signals in control seeds during the germination process (Fig. 4C), which were significantly increased in seeds that were treated with 6 μM ABA (Fig. 4D). In *AtABCG16 pro:GUS* transgenic plants, in the absence of exogenous ABA, the GUS signals were observed in the center of the roots and hypocotyl (Fig. 4, E and F) and the veins of leaves (Fig. 4, G and H), consistent with association with vascular bundles. After treatment with 3 μM ABA for 24 h, GUS signals were observed ubiquitously in both roots and leaves (Fig. 4, E–H), indicating significant up-regulation in all types of cells. To see if *AtABCG16* was up-regulated by bacterial pathogens, we tested the transgenic plants with the coronatine secreting strain (COR+) and *PstDC3000* and separately tested them with a solution containing only the virulence factor coronatine. We found that *PstDC3000* induced expression clearly in leaves and that coronatine up-regulated *AtABCG16* even more strongly than ABA itself (Fig. 4I). Also, we observed elevated GUS levels in both guard cells and pavement cells under ABA induction (Fig. 4J), which confirmed our earlier microarray

**Figure 4.** *AtABCG16* expression in seeds, roots, and leaves. A, Diagram of the *AtABCG16 pro:GUS* construct including Hygromycin Resistance (HygR) selectable marker. B, The *AtABCG16* promoter region includes three conserved motifs for the important ABA transcription factors ABA Insensitive3 (ABI3) and ABA Insensitive5 (ABI5) (Brady et al., 2003), one for WRKY38 (a known *PstDC3000*-induced transcription factor; Kim et al., 2008), and one for NAC-domain containing protein69 (NAC69) (a drought stress-related transcription factor; Xue et al., 2011) and positions of Homeobox protein 2a (Hox2a), myelocytomatosis oncogene_myeloblastosis oncogene (MYC_Myb) binding motifs as determined in Athamap. Expression of *AtABCG16* on days 1, 2, and 3 in control seeds (C) or seeds treated with 6 μM ABA (D). E to H, GUS staining without (left) and with (right) ABA in root (E), hypocotyl (F), cotyledon (G), and rosette leaf (H) at 21 d. I, GUS staining of leaves in response to control solution (10 μM MgCl₂) or 10 μM MgCl₂ containing 10⁸ cfu ml⁻¹ of DC3000 (COR+), 3 μM ABA, and 1 μM COR (left to right). J, GUS staining at stomata: control (left), 3 μM ABA (center), and 1 μM coronatine (right) at 24 h after treatment. Guard and pavement cells are identified by white and yellow arrows, respectively.
evidence that AtABCG16 was responding in multiple cell types within the leaf (Fig. 1E). Collectively, our GUS assay results showed that AtABCG16 responded to bacterial pathogen infection as well as to exogenous ABA and coronatine.

**AtABCG16 Reduces Bacterial Growth in Planta and Increases Plant Performance**

To assess the role of AtABCG16 in resistance to infection, we challenged plants with *PstDC3000* using soil-grown plants of the Col-0 background and the T-DNA insertion knockout lines *abcg16-1* and *abcg16-2*. In the first experiment, we compared growth of *PstDC3000* in leaves of the Col-0 background and *abcg16-1* after inoculation at $5 \times 10^4$ cfu mL$^{-1}$ by blunt syringe. The *PstDC3000* strain grew significantly more quickly and reached higher concentrations in the *abcg16-1* mutant relative to the background line ($F_{1,49} = 5.1, P = 0.028$; Fig. 5A). To assess the possible effects on plant performance under more realistic conditions, we then performed a second experiment, where we spray inoculated soil-grown plants with bacteria and measured both bacterial and plant growth. To assess the possible role of AtABCG16, we sprayed both T-DNA insertion knockouts *abcg16-1* and *abcg16-2* with the COR+ and COR− bacterial strains and compared their responses relative to the Col-0 background. We found that the COR+ strain *PstDC3000* was faster to colonize the interior of leaves and achieved higher titers ($F_{1,31} = 15.4, P < 0.001$; Fig. 5B; Supplemental Table S10), and it caused a greater reduction in shoot biomass than the COR− bacterial strain *PstDC3118* but only in two T-DNA *abcg16* knockout lines ($F_{1,67} = 4.6, P = 0.034$; Fig. 5, C and D; Supplemental Table S11). Together, these bacterial-challenge experiments suggested that AtABCG16 has an important role in the resistance of Arabidopsis against bacterial infection.

**DISCUSSION**

Although ABC transporters have been shown previously to transport ABA (Kang et al., 2010; Kuromori et al., 2010, 2011), this function has not been previously linked to resistance to bacterial infection. Here, we have shown that an ABCG transporter, AtABCG16, is important for ABA tolerance and also assists in plant basal resistance against *PstDC3000*. Both Atabcg16 T-DNA knockouts exhibited greater susceptibility to bacterial growth and reduced performance relative to transformed plants. Together, these results suggest an important role of the gene in resistance to these bacteria. As such, the study provides one of the first demonstrations, to our knowledge, of a link between an ABC transporter and plant resistance to bacterial pathogens. Previously, only one other ABCG transporter, AtABCG36 (PEN3/PDR8), has been found to be involved in plant defense against pathogens (Kobae et al., 2006), but in that case, auxin transport was implicated (Strader and Bartel, 2009; Xin et al., 2013).

Although it seems clear that AtABCG16 is involved in both plant basal resistance to *PstDC3000* and tolerance to exogenous ABA, we do not yet know whether these two effects are linked together. We have two
reasons, however, for suspecting that a link does exist. First, our GUS assays indicated strong AtABCG16 expression in response to both the bacterial pathogen and hormone treatments. Indeed, AtABCG16 expression is induced by flagellin 22 amino acid fragment alone (C. Danna, personal communication). Interestingly, when we applied ABA to AtABCG16pro::GUS plants, the signal was uniformly detected across the leaf surface, including guard cells and nonguard cells. However, when we applied coronatine, we observed a dramatic response in nonguard cells but only moderate up-regulation in guard cells (shown by the arrows in Fig. 3I). This difference may reflect ABA movement into guard cells after synthesis in other cell types (Bauer et al., 2013; Boursiac et al., 2013). Second, the T-DNA insertion knockouts showed that AtABCG16 suppresses bacterial growth and increases tolerance to the exogenous ABA.

The bacterial virulence factor, coronatine, strongly induced expression of AtABCG16, but the exact nature of the relationship remains unclear. Most notably, we found that a bacterial strain that lacks coronatine, PstDC3118, had significantly higher growth in the T-DNA knockout lines, abc16-1 and abc16-2, than in the untransformed background plants. In other words, even the avirulent strain lacking COR secretion was able to achieve higher titers in the plants that lacked expression of this particular gene. These data suggest, therefore, that AtABCG16 has a role in resistance that is independent of mitigation of the effects of coronatine.

How the gene functions in ABA tolerance remains to be further clarified. AtABCG16 and two other ABC transporters implicated in ABA transport (AtABCG22 and AtABCG25) are half transporters, meaning that their products must form either homodimers or heterodimers to become functional transporters. Heterodimerization has been shown previously for ABC transporters (McFarlane et al., 2010) and provides a mechanism by which a large number of transporter structures could be assembled from a relatively small number of genes. It is possible that the transport function of AtABCG16 will involve the proteins from these transporters or other half ABCG transporters. Interestingly, the expression pattern of AtABCG16 is different from AtABCG25, AtABCG22, and AtABCG40 (Kang et al., 2010; Kuromori et al., 2010, 2011). In leaves, AtABCG25 is located on xylem, whereas AtABCG22 and AtABCG40 are expressed mostly on the membrane of guard cells. This would seem to suggest that AtABCG16 may not form heterodimers with these particular proteins and is consistent with the finding that AtABCG22 and AtABCG40 do not seem to interact in planta (Kuromori et al., 2011). Our analysis of ABCG family genes (Supplemental Fig. S5) indicates that AtABCG16 is most similar to AtABC1, and our data suggest that the gene is most likely to have generalized expression across cell types, which may only be turned on under stress, whereas AtABCG25, AtABCG22, and AtABCG40 may have evolved more specialized functions with restricted expression that is cell-type specific (Ukitsu et al., 2007; Kuromori et al., 2011).

AtABCG16 exists on the third chromosome as a part of a tandem repeat of three ABC transporters that includes AtABCG17 and AtABCG18. There is no indication, however, that these three genes have overlapping function, and indeed, the differential responses of these genes to exogenous ABA, SA, and IAA are striking (Fig. 1D). However, because of the proximity of these three genes, it was particularly important to assess possible off-target effects of the T-DNA insertions in the abc16-1 and abc16-2 lines. Although both T-DNA insertion lines exhibited defectively low expression of AtABCG16 (Fig. 2A), they possessed normal expression of AtABCG17 and AtABCG18 relative to the Col-0 background (Supplemental Fig. S3). The strong defects that we observed in tolerance of abc16-1 and abc16-2 to ABA (Fig. 2, E–H) were, therefore, not caused by off-target effects on expression of AtABCG17 or AtABCG18. These data are consistent with a role of AtABCG16 specifically in tolerance of exogenous ABA.

To understand whether particular coding and promoter region differences among wild lines affect the function of AtABCG16, it will be important test the relative activities of alleles from susceptible and resistant wild lines in a common null background in the future. To assess more clearly whether AtABCG16 transports ABA, membrane transport assays (Kuromori et al., 2010) should also be performed. GWA mapping has attracted a great deal of interest in the literature (Atwell et al., 2010), but examples of successful use of this methodology remain very rare in plants. Our study is one of the first, to our knowledge, to identify a candidate gene directly from a GWA map and find supporting experimental evidence of relevant gene function.

In summary, we have provided evidence that AtABCG16 is involved in ABA tolerance and contributes to plant resistance against PstDC3000. We also confirm here the evidence from published microarrays that indicates very low levels of constitutive expression but dramatic induction of this gene by ABA and coronatine as well as its expression in both mesophyll and guard cells. Collectively, these results improve our understanding of basal resistance in Arabidopsis and offer unique ABA-related targets for improving the innate resistance of plants to bacterial infection.

MATERIALS AND METHODS

GWA Mapping

To identify candidates in plant resistance to PstDC3000, we created a GWA map for 96 accessions from the RepMap panel using data published previously (Atwell et al., 2010) of Arabidopsis (Arabidopsis thaliana) collected across the worldwide range. The dataset contained 205,803 SNPs derived from the 250K SNP data, version 3.06 (https://cynin.gmi.oeaw.ac.at/home/resources/atpolydb). All SNPs used in the analysis were diallelic and had the minor nucleotide represented in more than 5% of the accessions. For the GWA map, we used accession averages of leaf bacterial titer from three independent experiments where plants were infected by blunt syringes with a 1 × 10^6 cfu mL^-1 solution of DC3000, and bacterial titers were measured at 0, 1, 4, and 7 d postinfection (Atwell et al., 2010; Supplemental Table S1). Wilcoxon rank sum tests were used to calculate the significance of association between each SNP and the phenotypic values using standard methods (Atwell et al., 2010). To handle confounding caused by
populcell, we used the standard approach of mixed-model analysis known as Efficient Mixed-Model Association (Kang et al., 2008). We present log-transformed $P$ values using the standard approach (Atwell et al., 2010) and refer to them as the GWA score. To control the bias of rare single extreme GWA score values, we smoothed the map by averaging the GWA score of every 10 adjacent SNPs. The 13 loci identified represented the subset common to both the Wilcoxon and Efficient Mixed-Model Association maps (Supplemental Table S2).

Microarray Data

Expression data for all genes in the 250-kb neighborhood of AtABCG16 (Supplemental Table S3) were extracted from the following published microarray datasets: PtdDC3000 (NASCARRAYS-120; Nurnberger Laboratory), ABA (NASCARRAYS-176; Shimada Laboratory), MeJ/A (NASCARRAYS-174; Shimada Laboratory), IAA (NASCARRAYS-175; Shimada Laboratory), SA (NASCARRAYS-192; Shimada Laboratory), ABA leaf (Pandey et al., 2010), ABA seed (NASCARRAYS-183; Kamiya Laboratory), ABA guard cells (Pandey et al., 2010), and ABA mesophyll cells (Yang et al., 2008). All genes within the prescribed interval were included. Each of these published datasets included at least two biological replicates per treatment.

Plant Materials and Seed Sterilization

The wild-type Col-0 background (CS60000) and the two T-DNA insertion AtABCG16 knockouts, abcg16-1 (SALK_087501) and abcg16-2 (SALK_119868C), were acquired from the Arabidopsis Biological Resource Center at Ohio State. Seeds were sterilized before experiments using a standard bleach protocol consisting of 70% (v/v) ethanol for 1 min and 50% (v/v) bleach with 0.05% (v/v) Tween-20 (BP337-100; FisherBiotech) for 10 min followed by four washes with sterilized distilled water before use.

Plate Assays with Exogenous ABA, IAA, and SA

To assess plant tolerances of exogenous ABA, IAA, and SA, we followed a standard plate assay (Kurumori et al., 2010) using 15 cm petri dishes containing 0.8% agar with one-half-strength MS (M5524; Sigma-Aldrich). ABA, IAA, and SA were added to cooling agar (45°C or less) to the specified concentrations just before pouring the plates. Seeds were sterilized and washed as described above. Each MS plate received 25 seeds followed by cold treatment at 4°C in the dark for 1 week to stratify the seeds. These plates were then moved to a plant growth cabinet at 22°C under a 12-h-light/12-h-dark photoperiod with 55% humidity, which was considered day 1 of the experiment. Plates were measured for percentage of germination on day 4, root length on day 10, and mortality on day 30. For measuring root length on day 10, all roots were traced on acetate paper, digitally scanned, and measured for pixel number by ImageJ. Root length was calculated as root pixel number divided by pixel number of a line of 1 cm in length. All experiments included 5, 6, or 10 independent replicate MS plates as indicated (Supplemental Tables S8 and S9).

Expression Analyses Using qRT-PCR

Total RNA was isolated from leaves, stems, flowers, and root tissues of 6-week-old plants using TRIzol Reagent according to the manufacturer’s protocol (MRC). Residual genomic DNA was removed first by treatment with RNase-free DNase 1 (Invitrogen). First-strand complementary DNA was synthesized using approximately 2 µg of total RNA, 0.5 µg of oligo(dT)12-18, and SuperScript III reverse transcriptase according to the manufacturer’s instructions (Invitrogen). Reverse transcription (RT) -PCR was carried out in an Eppendorf mastercycler using GoTaq Green Master Mix (Promega), which was programmed as follows: 2 min at 95°C for predenaturation and 25 to 40 cycles of 15 s at 94°C, 15 s at 55°C, and 20 s at 72°C for each gene. qRT-PCR was carried out in an ABI-7900 thermal cycling system using a Real-Time PCR Power Mix Kit (ABI) and programmed at the same condition. For control reactions, either no sample was added or RNA alone was added without RT to test if the RNA sample was contaminated with genomic DNA. Arabidopsis thaliana Actin protein2 (Arabidopsis thaliana Actin protein2 was used as a reference gene. Oligonucleotide primer sets (Supplemental Table S6) were designed with the Primer Express 2.0 software (Applied Biosystems-Perkin-Elmer). To test the suitability of these primer sets, the specificity and identity of the RT-PCR products were monitored by melting curve analysis (65°C-99°C, 5°C s−1) of the reaction products, which can distinguish the gene-specific PCR products from nonspecific PCR products. All primers were synthesized by Integrated DNA Technologies. Data were analyzed by a standard method (Yuan et al., 2006).

Fluorescence Microscopy

To construct the GFP fusion lines, we first amplified the 2,211-bp AtABCG16 complementary DNA using PCR and cloned it into the PENTR/D/TOPO vector (Invitrogen). After sequence confirmation, we then integrated this construct (pPENTR/AtABCG16) into the GFP fusion protein vector pmD838 using LR clonase (Invitrogen). The construct of 2X35S::GFP-AtABCG16 was introduced into the recipient Col-0 plant through Agrobacterium sp-mediated transformation using the floral-dip method (Clough and Bent, 1998). Hypogymnecorrresistant lines (T1) were screened and PCR confirmed. We used tissue from 1-week-old seedlings of T2 plants for study of the subcellular localization of the GFP fusion protein. Microscope images were taken using fluorescein isothiocyanate (FITC) -filtered epifluorescence microscopy (Olympus BX51 model) with blue light at 400× magnification. For plasmolysis, root sections were treated with 20% Suc for 10 min.

GUS Expression Assays with the AtABCG Promoter

For AtABCG16 promoter-driven GUS expression lines, a 1,388-bp AtABCG16 promoter region was amplified by using K7 Taq (Takara) with the primers listed in Supplemental Table S6. Then, the PCR product was cloned into the pCR8/ GW/TOPO vector (Invitrogen), sequence confirmed by Sanger sequencing, and integrated into the promoter analysis vector pmDC162 through the LR reaction (Curtis and Grossniklaus, 2003). Transgenic plants were made as described above. T2 lines were used for the GUS expression analysis. To test the expression profile of ABCG16 and the response to ABA, 1-week-old seedlings were soaked in 10 μM MgCl2 (control) or 3 μM ABA for 24 h. For other treatments, leaves of 3-week-old plants were soaked in 10 μM MgCl2 (control), 108 cfu mL−1 of DC3000, 3 μM ABA, and 1 μM coronatine for 24 h. The expression pattern of ABCG16 during seeds germination and the response to ABA were tested as described above. Seed germination assays were performed by soaking seeds in 10 μM MgCl2 (control) or 6 μM ABA for 1, 2, or 3 d. GUS staining was performed following standard procedures (Jefferson et al., 1987).

Bacterial Growth Assays

To assess the possible role of AtABCG16 on plant resistance, we performed bacterial growth assays using PtdDC3000 alone and in conjunction with the COR – T-DNA insertion mutant bacterial strain PtdDC3118 obtained from D. Cuppels and J. Tambong at Agriculture and Agri-Food. To generate the inoculation solution, we first streaked bacteria on a King’s B media agar plate with 50 µg L−1 of rifampicin (R3501-5G; Sigma-Aldrich), and after 3 d, we transferred several colonies to 50 mL of liquid King’s B media also containing 50 µg L−1 of rifampicin and shook this solution in an incubator for 24 h at 28°C. We then transferred a aliquot of 5 mL to a fresh tube of 50 mL of liquid King’s B media, shook this solution for 4 h at 28°C to achieve midlogarithmic stage growth, spun the tube for 5 min at 3,000 rpm to obtain a pellet, poured off the supernatant, resuspended the pellet in 1 mL of 10 mM MgSO4, vortexed, measured bacterial concentration by spectrophotometry, and diluted 10-fold using 10 mM MgSO4 to 108 cfu mL−1. For each plant, we inoculated two leaves with approximately 0.1 mL of 5×106 cfu mL−1 bacterial solution by blunt syringe. For the T-DNA insertion knockout lines, we sampled six independent plants per time point. We measured the knockouts and wild-type background plants simultaneously in each experiment. To measure leaf bacterial titers, we used a hole punch to collect a leaf disc (0.28 cm2) and then ground the leaf disc in 200 µL of 10 mM MgSO4 buffer, 40 µL of the diluted mixture were then plated on the agar plates with 50 µg μL−1 of rifampicin and incubated at 28°C for 3 d before counting the number of colonies.

Plant Growth Assay

To compare the effects of bacterial infection by COR+ and COR− bacteria on the performance of plants with and without functional AtABCG16, we grew the wild-type background line (Col-0) and both T-DNA insertion knockouts (abcg16-1 and abcg16-2) and then exposed them to either the COR+ strain (PtdDC3000) or the COR− strain (PtdDC3118) by repeated spray inoculation. Seeds were sown on soil (Promix BX) in standard 36-cell flats, cold treated for 1 week, and then placed in a growth room at 20°C under a 12-h-light/12-h-dark photoperiod. Seeds germinated the next day, which was considered day 1 of the experiment. Flats were distributed to one at least one time per week to reduce positional effects. On day 11, we thinned seedlings to 5 plants per cell for a total of 120 plants per line. On day 35, we randomly assigned plants to receive either PtdDC3000 or PtdDC3118 by spray inoculation.
On day 36, we sprayed the plants to saturation with 1 × 10^6 cfu mL⁻¹ of PstDC3118 or PstDC3000 in 10 mM MgSO₄ buffer. We then covered the flats with clear plastic covers. Given that Pseudomonas sp. infection is likely to be amplified by cold stress and high humidity, we then placed all plants in a 4°C cabinet for 48 h and sprayed the bacterial solution every 12 h for 2 d. After 48 h, we moved the flats back to the growth room but left the clear plastic covers on to maintain high humidity. On day 54 of plant growth (the 18th day after infection), we removed discs from plants in one-half of the cells for measurement of bacterial titer. The leaf disc was sterilized in 70% ethanol for 2 s, ground in 200 μL of 10 mM MgSO₄ buffer, and then, diluted in 10 mM MgSO₄ buffer and plated on King’s B media with 50 μg mL⁻¹ of rifampicin. Plates were counted after 3 d at 28°C. On day 95, we harvested the remaining unsampled plants from each cell, photographed them, dried them at 60°C for 2 weeks, and weighed them to determine shoot dry biomass.

**Supplemental Data**

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

**Supplemental Figure S1.** Root tracings following ABA treatment.  
**Supplemental Figure S2.** Photograph of roots shown in Fig. 2H.  
**Supplemental Figure S3.** Expression of AtABC2C17 and AtABCG18.  
**Supplemental Figure S4.** 35S::GFP positive controls for root and leaf epidermis.  
**Supplemental Figure S5.** Gene tree of ABC transporter G family group.  
**Supplemental Table S1.** Titers of PstDC3000 in RegMap panel lines.  
**Supplemental Table S2.** Candidate genes from genome-wide association map.  
**Supplemental Table S3.** Base calls for SNP nos. 118594–118611.  
**Supplemental Table S4.** Base positions and map scores for SNP nos. 118594–118611.  
**Supplemental Table S5.** Expression response to bacterial infection, ABA, IAA, and SA.  
**Supplemental Table S6.** Primers used for RT-PCR, gene and promoter cloning.  
**Supplemental Table S7.** RT-PCR of AtABCG16.  
**Supplemental Table S8.** Effects of ABA, IAA, and SA on germination.  
**Supplemental Table S9.** Effects of ABA on germination, root growth, and mortality.  
**Supplemental Table S10.** Bacterial titer following surface spray inoculation.  
**Supplemental Table S11.** Shoot dry biomass following surface spray inoculation.

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