Interaction of the WD40 Domain of a Myoinositol Polyphosphate 5-Phosphatase with SnRK1 Links Inositol, Sugar, and Stress Signaling

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In plants, myoinositol signaling pathways have been associated with several stress, developmental, and physiological processes, but the regulation of these pathways is largely unknown. In our efforts to better understand myoinositol signaling pathways in plants, we have found that the WD40 repeat region of a myoinositol polyphosphate 5-phosphatase (5PTase13; At1g05630) interacts with the sucrose nonfermenting-1-related kinase (SnRK1.1) in the yeast two-hybrid system and in vitro. Plant SnRK1 proteins (also known as AKIN10/11) have been described as central integrators of sugar, metabolic, stress, and developmental signals. Using mutants defective in 5PTase13, we show that 5PTase13 can act as a regulator of SnRK1 activity and that regulation differs with different nutrient availability. Specifically, we show that under low-nutrient or -sugar conditions, 5PTase13 acts as a positive regulator of SnRK1 activity. In contrast, under severe starvation conditions, 5PTase13 acts as a negative regulator of SnRK1 activity. To delineate the regulatory interaction that occurs between 5PTase13 and SnRK1.1, we used a cell-free degradation assay and found that 5PTase13 is required to reduce the amount of SnRK1.1 targeted for proteosomal destruction under low-nutrient conditions. This regulation most likely involves a 5PTase13-SnRK1.1 interaction within the nucleus, as a 5PTase13:green fluorescent protein was localized to the nucleus. We also show that a loss of function in 5PTase13 leads to nutrient level-dependent reduction of root growth, along with abscisic acid (ABA) and sugar insensitivity. 5ptase13 mutants accumulate less inositol 1,4,5-trisphosphate in response to sugar stress and have alterations in ABA-regulated gene expression, both of which are consistent with the known role of inositol 1,4,5-trisphosphate in ABA-mediated signaling. We propose that by forming a protein complex with SnRK1.1 protein, 5PTase13 plays a regulatory role linking inositol, sugar, and stress signaling.

Myoinositol (inositol) signaling pathways are important for many different developmental and physiological processes in eukaryotes (Boss et al., 2006; Berridge, 2007). In plants, inositol signaling is used in the response to abscisic acid (ABA); Sanchez and Chua, 2001; Xiong et al., 2001; Burnette et al., 2003; Gunesekeera et al., 2007; Lee et al., 2007), salt stress (DeWald et al., 2001; Takahashi et al., 2001), gravity (Perera et al., 2001, 2006), and pathogens (Ortega and Perez, 2001; Andersson et al., 2006). The inositol signaling pathway makes use of an inositol 1,4,5-trisphosphate (InsP$_3$) second messenger that triggers intracellular Ca$^{2+}$ release from various sources within the cell (Berridge, 1993; Tang et al., 2007), although the precise mechanism in plants is not yet known (Krinke et al., 2007). To terminate signaling events driven by InsP$_3$, cells utilize the myoinositol polyphosphate 5-phosphatases (5PTases; EC 3.1.3.56) to remove the 5-phosphate, thus initiating second messenger breakdown (Astle et al., 2007). Eukaryotes have a family of diverse 5PTases, with the Arabidopsis (Arabidopsis thaliana) genome containing 15 genes predicted to encode conserved 5PTases (Berdy et al., 2001). Of the 15 Arabidopsis 5PTase genes, four encode 5PTase enzymes with a large, N-terminal extension containing five to six WD40 repeat regions (Zhong and Ye, 2004). WD40 repeats are found in a number of eukaryotic regulatory proteins (Li and Roberts, 2001), where they are speculated to serve as a stable propeller-like platform to which proteins can bind either stably or reversibly. In some cases, sequences outside of the WD40 repeats may confer a new functional specificity to the protein even as conserved protein interactions are maintained through the WD40 repeats, potentially adapting basic cellular mechanisms to organism-specific processes. WD40-containing 5PTases are unique to plants and certain fungi, as no other genomes contain candidate genes for these proteins (Zhong and Ye, 2004).

By examining plants containing either a gain or loss of function in specific 5PTases, investigators have established that 5PTases are critical in plant develop-
ment and in ABA signaling. In particular, mutations in the CVP2 (5PTase6; At1g05470) and 5PTase13 (At1g05630) genes have been linked to altered cotyledon vascular patterning and/or blue light responses and phototropin 1 signaling (Carland and Nelson, 2004; Lin et al., 2005; Chen et al., 2008). In contrast, fra3 (At1g65580) and mhr3 (5PTase5; At5g65090) mutants are altered in fiber cell development (Zhong et al., 2004) and root hair initiation (Jones et al., 2006), respectively. With regard to stress signaling, ectopic expression of either 5PTase1 or 5PTase2 has been shown to decrease ABA signaling by increasing hydrolysis of InsP_3 (Sanchez and Chua, 2001; Burnette et al., 2003). Loss-of-function mutants in these same genes resulted in ABA hypersensitivity in germinating seeds and increased seedling hypocotyl elongation in the dark, which is accompanied by an increase in InsP_3 levels (Gunesecka et al., 2007).

5PTases have been reported to play a role in Glc sensing/metabolism in animal cells (Wada et al., 2001; Sasaoka et al., 2004; Kagawa et al., 2008). Overexpression of an SH2-containing inositol phosphatase 2 (SHIP2) inhibited insulin-induced signaling leading to Glc uptake and glycogen synthesis via hydrolysis of phosphatidylinositol 3,4,5-trisphosphate (PtdInsP_3) in 3T3-L1 adipocytes and L6 myotubes (Sasaoka et al., 2001; Wada et al., 2001). On the other hand, loss of SHIP2 in mice resulted in increased sensitivity to insulin, which is characterized by deregulated expression of genes involved in gluconeogenesis and perinatal death (Clement et al., 2001). Although there are recent reports delineating some relationship between inositol signaling and sugar sensing/metabolism in plant cells (Im et al., 2007; Lou et al., 2007), a specific role of 5PTases in sugar signaling has not yet been determined. The mechanisms by which plants sense sugars and regulate carbohydrate metabolism are complex and often facilitated by protein complexes. Therefore, we sought to identify 5PTase-interacting proteins involved in signaling and metabolic events.

We report here that the WD40 repeat region of the 5PTase13 gene interacts specifically with a Suc nonfermenting-1-related kinase (SnRK1.1, AIN10; At3g01090), which functions as a sensor of energy and stress in plants (Baena-Gonzalez et al., 2007; Hardie, 2007; Hue and Rider, 2007). Using T-DNA insertion mutants in the 5PTase13 gene, we present evidence that 5PTase13 is a nuclear protein that can act as a regulator of SnRK1.1 under low-nutrient conditions by decreasing the amount of SnRK1.1 degraded by the proteasome.

RESULTS

5PTase13 Is a Member of a Unique Group of WD40-Containing Proteins

Besides containing a conserved inositol polyphosphate 5-phosphatase catalytic domain, 5PTase13 also contains five WD40 repeat regions in the N terminus that may allow for unique protein interactions. Three other genes in the Arabidopsis genome, 5PTase12 (At2g43900), 5PTase14 (At2g31830), and FRA3 (At1g65580), encode similar proteins (Berdy et al., 2001; Zhong and Ye, 2004), and the amino acid identity of the WD40 repeat region varies from 45.5% to 75.3% among these proteins (Supplemental Table S1). We used the amino acid sequences corresponding to these WD40 repeat regions in BLASTp searches and obtained 12 related sequences with e-values less than 1. ClustalW and PAUP4.0 were used to generate phylogenetic trees using parsimony (Fig. 1), which showed that the 5PTase WD40 repeat regions are contained on a separate branch, thus indicating that they are more similar to one another than to other WD40 repeat regions.

The similarity of the 5PTases (Zhong and Ye, 2004) suggests that these proteins may function in a redundant manner. Microarray data from GENEVESTIGATOR (Zimmermann et al., 2004) indicates that FRA3 is broadly and abundantly expressed compared with 5PTase12, 5PTase13, and 5PTase14 (Supplemental Fig. S1A). We conclude that the WD40-containing 5PTases are a small group of proteins with the potential to form protein complexes and that functional redundancy may be present.

The WD40 Repeat Region of 5PTase13 Interacts with SnRK1.1

To investigate the ability of the WD40 regions of 5PTase13 to participate in protein complexes, we used the yeast two-hybrid system. The 533 N-terminal amino acids from 5PTase13 containing the WD40 repeats were used as bait in a yeast two-hybrid screen of an Arabidopsis 3-d-old etiolated seedling cDNA library. We screened over 1 million yeast transformants and obtained a positive clone containing the C-terminal domain of the SnRK1.1 gene (At3g01090; also known as AIN10). We retransformed this positive clone into yeast and verified the interaction (Fig. 2). Negative controls, including the empty DNA binding domain and activation domain vectors, established that SnRK1.1 binds to the WD40 repeat region of the 5PTase13 protein in yeast.

To examine the interaction between 5PTase13 and SnRK1.1 in vitro, we fused the sequence encoding the Xpress epitope tag to the C terminus of the WD40 repeat region of 5PTase13 (13WDX) and the V5 epitope tag sequence to the C terminus of SnRK1.1 (SnRKV5) and expressed both proteins in Escherichia coli. The SnRKV5 construct directs the expression of the 61.7-kD protein detected by an anti-V5 monoclonal antibody, and in most cases we detected two SnRKV5 bands, perhaps as a result of phosphorylation or proteolytic cleavage (Fig. 3, lane 1). The 13WDX construct directs the expression of a 65.9-kD protein detected by an anti-Xpress monoclonal antibody (Fig. 3, lane 2). These interactions are specific, as SnRKV5 is not detected by
the anti-Xpress antibody and 13WDX is not detected by the anti-V5 antibody (data not shown). To determine whether the SnRK1.1 recombinant protein can “pull down” 13WDX, immunoprecipitations using anti-V5:protein A-Sepharose beads were performed, and the resulting complex was then analyzed by western blotting with an anti-Xpress antibody (Fig. 3, IP lanes). As shown in Figure 3, the WD40 repeat region from 5PTase13 (13WDX) is only detected in this pull-down assay when it has been incubated in the presence of SnRKV5. We conclude that the WD40 repeat region from 5PTase13 and SnRK1.1 can form a protein complex in vitro.

SnRK1.1, along with its closely related gene family member SnRK1.2, encodes a Suc nonfermenting-1-related kinase implicated as a central integrator of energy signaling and metabolic regulation in yeast, plants, and animals. The interaction of 5PTase13 and SnRK1.1 is novel and is perhaps unique to plants in that yeast and animal 5PTases do not contain WD40 regions. This interaction may indicate that InsP$_3$ signal termination via 5PTase13 function affects the SnRK1 “energy sensor” in plants. Data obtained from the GENEVESTIGATOR database indicate that SnRK1.1, SnRK1.2, and 5PTase13 are detected in most plant tissues examined, although 5PTase13 expression levels are very low compared with SnRK1 gene expression (Supplemental Fig. S1B).

**5PTase13 Mutant Identification**

To further explore the link between InsP$_3$ signaling and the energy sensor, SnRK1.1, we isolated two independent T-DNA insertion mutants in the 5PTase13 gene. Two potential mutants were identified in the SALK T-DNA database and named 5ptase13-1 (SAIL_350_F01) and 5ptase13-2 (SALK_081991) and were compared with their corresponding wild-type accessions (Fig. 4A). The presence of the T-DNA insertion was verified by diagnostic PCR in each mutant using genomic DNA and primers specific for the T-DNA left border (LB) and gene-specific primers that flank the T-DNA insertion (Fig. 4, A and B). The resulting LB gene-specific fragments were sequenced, indicating that in 5ptase13-1 mutants, a second T-DNA insertion is found in tandem in the fourth exon (Fig. 4, A and B, LB-R band). This is in contrast to the previously reported analysis by Lin et al. (2005) showing that the 5ptase13-1 mutant contains a single T-DNA insertion.

Using primers specific for the 3’ end of 5PTase13, we detected a PCR product in both wild-type lines used...
(CS60000 and CS908) but not in 5ptase13-1 and 5ptase13-2 mutants (5PTase13 in Fig. 4C). Using primers that amplify the 5’ end, we detected a 1.68-kb product in both wild-type lines and in the 5ptase13-1 mutant, but not in 5ptase13-2 (Fig. 4C). We conclude that the 5ptase13-2 mutant is totally lacking 5PTase13 expression and that both 5ptase13-1 and 5ptase13-2 mutant lines do not express transcripts capable of encoding a full-length 5PTase13 protein.

To examine how the loss of 5PTase13 affects the expression of its binding partner SnRK1.1, we examined the expression of SnRK1.1 and its closely related isoform, SnRK1.2, in 7-d-old dark-grown 5ptase13 and wild-type seedlings. The results reveal that there are no large changes in SnRK1.1 and SnRK1.2 in 5ptase13 mutants (Fig. 4D). As shown in Figure 4D, we found that FRA3 expression remains unchanged in 5ptase13 mutants, and 5PTase14 expression is barely detectable but also unchanged. In contrast, the expression of 5PTase12 is increased in both 5ptase13 mutant lines, revealing a possible means of compensation for the loss of 5PTase13 function (Fig. 4D).

Under standard laboratory conditions, 5ptase13 mutants did not exhibit any abnormalities in plant growth or development. Since Lin et al. (2005) previously reported that 5ptase13-1 mutants are altered in cotyledon vein development, we carefully examined both 5ptase13-1 and 5ptase13-2 mutants and their corresponding wild-type accessions for cotyledon vein development, as described by Carland and Nelson (2004). Using 7-d-old light-grown seedlings of wild-type and 5ptase13 mutant soil-grown plants, we found no evidence for a cotyledon vein phenotype (Supplemental Tables S2 and S3). Our analysis revealed that 33.3% of total WT1 cotyledons and 26.18% of total 5ptase13-1 cotyledons have abnormal patterns and that only 15.9% of total WT2 cotyledons and 19.4% of total 5ptase13-2 cotyledons can be classified as abnormal (Supplemental Table S3). Thus, there is variation in cotyledon vein development within different wild-type accessions, but not between 5ptase13 mutants and their matched wild-type accessions.

**SnRK1 Activity Is Altered in 5ptase13 Mutants and Varies with Nutrient Conditions**

To determine whether 5PTase13 affects SnRK1 function, we measured the activity of SnRK1 in 5ptase13 mutants and wild-type seedlings grown under various nutrient conditions. It is well documented that SnRK1 regulates multiple transcription cascades in response to sugar or energy deprivation (Baena-Gonzalez et al., 2007); however, whether in planta SnRK1 activity changes under various nutrient conditions is not known. To address this, we measured SnRK1 activity from 7-d-old wild-type seedlings grown under low light (40 μE) with different added nutrients to establish various “low-energy” conditions: no nutrients (agar and water alone), low nutrients (0.5× Murashige and Skoog [MS] salts and agar), and optimal nutrients (0.5× MS salts, agar, and 3% Suc). In addition, we examined a stressful level of added sugar (0.5× MS salts, agar, and 6% Glc). For this work, we used a well-
established SnRK1 assay (Radchuk et al., 2006) and incubated a sucrose phosphate synthase (SPS) substrate peptide (Huang and Huber, 2001), radiolabeled $\gamma^{-32P}$ATP, and 5 $\mu$g of plant protein extract for 30 min. Validation that these conditions are in the linear range of product accumulation is shown in Supplemental Figure S2.

As expected from its role as a low-energy sensor, SnRK1 activity is higher in seedlings grown on low nutrients compared with extracts prepared from seedlings grown on optimal nutrients or 6% Glc (Fig. 5). Figure 6A shows that the activity of SnRK1 significantly increases in 5ptase13-1 and 5ptase13-2 mutants when seedlings are grown with no nutrients (8.1- and
When either low nutrients or 6% Glc is present, the absence of nutrients and as a positive regulator 5PTase13 as a negative regulator of SnRK1 activity in 5ptase13 mutants grown with no nutrients, we find less SnRKV5 accumulation in 5ptase13 extracts compared with wild-type extracts (Fig. 7D). We conclude that under the low-nutrient conditions, 5PTase13 is not required to stabilize SnRKV5 when seedlings are grown with no nutrients, and this is consistent with the switch in SnRK1 activity levels we found in 5ptase13 mutants grown with no nutrients (Fig. 6A). However, since we did not observe an increase in SnRKV5 stability in 5ptase13 extracts prepared from the no-nutrient conditions, we speculate that there is an additional mechanism that influences the elevated SnRK1 activity in 5ptase13 mutants under these conditions.

Figure 5. SnRK1 activity varies with different nutrient conditions. Seeds from wild-type (WT2-CS60000) plants were grown on 0.8% agar, no nutrients, low nutrients (0.5× MS salts), optimal nutrients (0.5× MS salts, 3% Suc), or 6% Glc (0.5× MS salts, 6% Glc) under low light for 7 d. The seeds grown in 6% Glc were allowed to germinate in 0.5× MS medium for 4 d and then transferred to 6% Glc medium for 3 d. SnRK1 activity with an SPS peptide was measured from crude plant extracts precipitated with ammonium sulfate according to the method described (Radchuk et al., 2006). Bars represent means ± se of three replicates. The experiment was independently repeated two times. *, P < 0.05 compared with no nutrients.

To test whether 5PTase13 is required for sugar and ABA responses, we analyzed age-matched seeds for germination in the presence of 0%, 1%, 3%, or 6% Glc.
or 0%, 1%, 2%, 3%, 6%, or 11% Suc. At low concentrations of sugar, there were no differences in the germination of wild-type and 5ptase13 seeds in the dark or light (data not shown). However, at a high exogenous sugar concentration (6% Glc and 11% Suc) in the dark, we found that 5ptase13 mutants were significantly less sensitive to sugar (Fig. 8D; Supplemental Figs. S3 and S4). We saw the same trend in sugar insensitivity when 5ptase13 mutant seeds were germinated in the presence of 6% Glc in the light, although the sugar insensitivity was less apparent (Supplemental Fig. S3). This sugar insensitivity was also noted for 5ptase13-1 and 5ptase13-2 mutant seeds germinated in the presence of 11% Suc, in which maximal increases of 2.5- and 2.2-fold for 5ptase13-1 and 5ptase13-2, respectively, were noted compared with wild-type seeds (Supplemental Fig. S4). Germination in the presence of mannitol, a nonmetabolizable sugar, was not altered, indicating that the sugar insensitivity of 5ptase13 mutants is not due to a general osmotic stress tolerance (Fig. 8D).

We also germinated 5ptase13 mutant seeds in the presence of 0, 1, 2, and 3 μM ABA and measured the impact on germination during a 6-d period. As expected, germination of wild-type seeds was delayed by ABA in a concentration-dependent manner during the 6-d period under light conditions (Fig. 8, E and F). In contrast, 5ptase13-1 and 5ptase13-2 mutant seeds were ABA insensitive, reaching 100% germination on day 2 in the presence of 1 μM ABA and 78% to 90% on day 6 in the presence of 3 μM ABA (Fig. 8, E and F). Since we did not find reduced seed dormancy in our mutants (data not shown), this ABA insensitivity of 5ptase13 mutants most likely does not correlate with changes in de novo synthesis of ABA (Gubler et al., 2005) and is in accordance with the previously reported results for ABA insensitivity of the 5ptase13-1 mutant (Lin et al., 2005).

Expression of Glc- and ABA-Regulated Genes Is Altered in 5ptase13 Mutants

To determine whether there are differences in the expression patterns of Glc- and/or ABA-regulated genes (RD29A, KIN1, and CAB1 genes; Price et al., 2004) in 5ptase13 mutants, we examined expression in 4-d-old dark-grown wild-type and 5ptase13 seedlings exposed to 6% Glc for 3 d (Fig. 9A). Treatment with 6% Glc strongly induced the expression of RD29A and KIN1 in wild-type seedlings. In contrast, 5ptase13-1

Figure 6. SnRK1 activity in 5ptase13 mutants. SnRK1 activity was measured in crude plant extracts from 7-d-old seedlings precipitated with ammonium sulfate according to the method described (Radchuk et al., 2006) using an SPS peptide. Values for SnRK1 activity (nmol inorganic phosphate min⁻¹ mg⁻¹ protein) are normalized to WT1. Extracts are from seedlings grown in no nutrients (A), low nutrients (0.5× MS salts; B), or 6% Glc (0.5× MS salts, 6% Glc; C). Bars represent means ± se of three replicates. The experiment was independently repeated two times. *, P < 0.05 compared with the wild type.
and 5ptase13-2 mutants grown in 6% Glc contained much smaller increases in RD29A and KIN1 transcript levels (Fig. 9A), indicating that induction of these genes is diminished, but not abolished, in 5ptase13 mutants. The regulation of CAB1, on the other hand, was not significantly altered in 5ptase13 mutants.

Complementation of the 5ptase13-1 Mutant

To ensure that the alterations noted in 5ptase13 mutants are due to loss of 5PTase13 expression, we expressed a 5PTase13:GFP fusion under the control of the 35S cauliflower mosaic virus promoter in wild-type and 5ptase13-1 plants (13:GFP and 13-1/13:GFP plants, respectively). We examined two lines of 13-1/13:GFP plants with good expression of the transgene (Fig. 9B) along with wild-type and 5ptase13-1 plants in ABA-sensitivity assays. Both 13-1/13:GFP lines exhibited more ABA sensitivity in germination assays compared with 5ptase13 mutants (Fig. 9C). Since 5ptase13-1 mutants contain decreased SnRK1 activity under low-nutrient conditions compared with wild-type plants (47%; Fig. 6B), we also measured SnRK1 activity in the 13-1/13:GFP-1 and 13-1/13:GFP-2 lines and found, as expected, similar or increased SnRK1 activity compared with wild-type plants (140% ± 10% for 13-1/13:GFP-1 and 102% ± 5% for 13-1/13:GFP-2). We conclude that expression of the 5PTase13:GFP transgene complements the 5ptase13-1 mutant.

Glc-Stimulated InsP3 Levels Are Altered in 5ptase13 Mutant Seedlings

We examined whether the Glc insensitivity of 5ptase13 mutants is accompanied by alterations in mass InsP3 levels by measuring mass InsP3 levels. The results in Figure 10 indicate that neither 5ptase13-1 nor 5ptase13-2 mutant seedlings differ significantly from wild-type seedlings in their InsP3 mass levels under control conditions. When wild-type seedlings are exposed to 6% Glc for 3 d, mass InsP3 levels increase 2.8- to 3.7-fold, which is a statistically significant elevation. However, when 5ptase13 mutants are grown for 3 d in the presence of 6% Glc, mass InsP3 level changes are smaller, with an increase of 1.6- to 2-fold over basal levels, and statistically significant only in the 5ptase13-2 mutant. More importantly, the...
Glc-stimulated InsP_3 levels in both 5ptase13 mutants differ significantly from the Glc-stimulated InsP_3 levels in wild-type seedlings. We conclude that 5ptase13 mutants are impaired in their ability to accumulate InsP_3 in response to Glc and that this correlates with the sugar and ABA insensitivity noted in the germination assays.

Subcellular Localization of the SPTase13:GFP Fusion Protein

To investigate the subcellular location of the SPTase13 protein, we performed imaging experiments with 5ptase13-1 mutants complemented with the SPTase13:GFP construct (13-1/13:GFP plants) and wild-type plants containing the same SPTase13:GFP construct. As the SPTase13:GFP construct we used allowed for complementation of the ABA and sugar insensitivity phenotype in 5ptase13-1 mutants (Fig. 9, B and C), it is likely that this fusion protein undergoes the same posttranslational modifications and subcellular localization as the native SPTase13 protein. We analyzed T2 progeny from two independent 13-1/13:GFP lines with fluorescence deconvolution microscopy and found a similar pattern in both lines. GFP fluorescence was associated with the nucleus in many, but not all, cells in cotyledon epidermis (Fig. 11, B and C), hypocotyls (data not shown), and roots (Fig. 11, D–I). Nuclei from some but not all guard cells contained the 13:GFP protein (Fig. 11C). To confirm the nuclear localization, we stained 13-1/13:GFP seedlings with the nuclear dye 4′,6-diamidino-2-phenylindole (DAPI) and imaged GFP and DAPI fluorescence simultaneously (Fig. 11, D–I). Once again, we found 13:GFP fluorescence in only a portion of root nuclei, while DAPI fluorescence was present in all nuclei. We conclude that SPTase13 protein is located in the nucleus of seedlings and that its presence in the nucleus is restricted in some cells.

DISCUSSION

Since control of second messengers is critical for signaling, there is interest in determining how the plant cell regulates levels of second messengers such as InsP_3. We identified a potential regulator of InsP_3 signaling by isolating protein interactors of 5PTase13. We focused on 5PTase13 because it contains several conserved WD40 repeats (for review, see van Nocker and Ludwig, 2003) and the presence of WD40 repeats within 5PTase genes is unique to plants and certain fungi (Zhong and Ye, 2004). To date, the WD40 repeats of 5PTases have not been functionally characterized, and no potential protein partners have been identified. We show here an important and novel interaction between the WD40 repeat region of 5PTase13 and the central metabolic regulator, SnRK1.1. SnRK1.1 and its orthologues in yeast and mammals is one subunit of a complex that modulates cellular metabolism in response to nutrient and environmental conditions (Baena-Gonzalez et al., 2007; Hardie, 2007; Hue and Rider, 2007). Recent work in Arabidopsis has elegantly delineated the role of SnRK 1.1 in the transcriptional activation of genes from catabolic pathways that provide routes for alternative energy sources and coordinate repression of a large number of anabolic genes (Baena-Gonzalez et al., 2007). Our discovery of a
The experiment was independently repeated two times. *, compared with the score for each group of seeds at day 3. Values are means ± SE of three to five replicates. The experiment was repeated three times. **, \( P < 0.05 \) compared with the control, untreated sample; ***, \( P < 0.05 \) compared with wild-type Glc sample; ****, \( P < 0.05 \) compared with the control, untreated sample.

Figure 9. A, Expression of Glc- and ABA-inducible genes from dark-grown 4-d-old wild-type and 5ptase13 seedlings that were either untreated (control; 0.5× MS salts, 0.8% agar) or treated with 6% Glc (0.5× MS salts, 0.8% agar) in the dark for 3 d. The experiment was independently repeated two times. B, Verification of overexpression of 5PTase13 in complemented lines and a 5PTase13:GFP line. Total RNA (1–2 µg) was isolated, and semiquantitative RT-PCR was performed with the indicated primers (Supplemental Table S4). C, The 5PTase13:GFP gene complements the ABA-insensitive phenotype of 5ptase13-1 mutants. Seeds from WT1, 5ptase13-1, and 5ptase13-1 containing a 5PTase13:GFP transgene were germinated and grown in light for 6 d on 0.5× MS salts, 0.8% agar, and 3 µM ABA. The germination rate was scored for each group of seeds at day 3. Values are means ± SE (n = 50). The experiment was independently repeated two times. *, \( P < 0.05 \) compared with the 5ptase13-1 mutant.

5PTase13:SnRK1.1 complex points to the novel interaction of this metabolic modulator and InsP3 signaling. It has been shown previously that a loss of function in the 5PTase13 gene leads to defects in auxin-regulated development (Lin et al., 2005) and in blue light responses (Chen et al., 2008). Because of the 5PTase13 interaction with SnRK1.1, we investigated the role of 5PTase13 in nutrient sensing and stress responses. Recent studies have shown that Arabidopsis plants overexpressing SnRK1.1 have greater Suc sensitivity and that mutants lacking SnRK1.1 use exogenous sugars more efficiently (Baena-Gonzalez et al., 2007). When virus-induced gene silencing is used to silence both SnRK1.1 and SnRK1.2, growth is severely affected, indicating the importance of the SnRK1 modulator to overall growth (Baena-Gonzalez et al., 2007). Repressing a pea (Pisum sativum) homolog of SnRK1 with antisense RNA results in seed maturation effects resembling ABA insensitivity (Radchuk et al., 2006), while a loss of function in SnRK1 in rice (Oryza sativa) delays seed germination and seedling growth (Lu et al., 2007). We report here that 5ptase13 mutants exhibit reduced root growth under limited nutrient conditions, but not when 3% Suc is supplied. The variability of the root growth phenotype under different nutrient conditions is thus similar, although not identical, to growth responses of SnRK1.1 mutants. In addition, we documented that 5ptase13 mutants are altered in their responses to ABA and sugar stress (Figs. 8–10). When 5ptase13 mutants are challenged with a sugar stress, alterations in germination, gene expression, and InsP3 accumulation occur, indicating that 5ptase13 mutants are sugar insensitive. This sugar insensitivity is similar to that noted for the regulator of G-protein signaling1 mutant (Chen et al., 2006b) but differs from the Glc-insensitive (Zhou et al., 1998; Arenas-Huertero et al., 2000; Moore et al., 2003; Lin et al., 2007) and sugar-insensitive (Laby et al., 2000; Gibson et al., 2001) mutants that are also insensitive to the developmental arrest induced by sugar that occurs over a period of 21 d. Since there are other WD40-containing 5PTases, we speculate that the sensitivity to developmental arrest noted in 5ptase13 mutants grown on high sugar may be due to this redundancy. Together, the data on nutrient and sugar stress responses of 5ptase13 mutants are consistent with the existence of a 5PTase13:SnRK1.1 complex that modulates seedling responses to nutrient conditions and stress.

To understand how a 5PTase13:SnRK1.1 complex might regulate nutrient and stress signaling, we measured SnRK1 activity in 5ptase13 mutants and wild-type seedlings grown under different nutrient conditions. Our data show that the presence of 5PTase13 affects SnRK1 activity and that nutrient availability is an important switch in 5PTase13 regulation of SnRK1.1. Specifically, 5PTase13 is required to maintain wild-type levels of SnRK1 activity when low nutrient or...
high sugar is present (Fig. 6, B and C). In contrast, under the greatest starvation conditions of no added nutrients, 5PTase13 appears to be a negative regulator of SnRK1, as 5ptase13 mutants contain significantly elevated SnRK1 activity (Fig. 6A). To further delineate the mechanism of 5PTase13 regulation of SnRK1.1, we used a cell-free degradation assay and showed that the decrease in SnRK1 activity in 5ptase13 mutants correlates with increased degradation of SnRK1.1 by the proteasome (Fig. 7D).

Interestingly, another WD40 repeat-containing protein called PRL1 (for PLEITROPIC REGULATOR LOCUS1) also interacts with SnRK1.1 and is able to inhibit the activity of SnRK1.1 and SnRK1.2 (Bhalerao et al., 1999; Farras et al., 2001). Recent evidence shows that there is a decrease in SnRK1 degradation in prl1 mutants, and additional evidence suggests that PRL1 acts in delivering SnRK1 to the CUL4-DDB1 complex for proteasomal degradation (Lee et al., 2008). Thus, we predict that 5PTase13 and PRL1 have opposing functions regarding SnRK1 stability under low-nutrient and sugar-stress conditions. The fact that prl1 mutants are ABA and sugar hypersensitive (Bhalerao et al., 1999), which is the opposite of 5ptase13 mutants, also supports opposing roles for 5PTase13 and PRL1 in regulating SnRK1 degradation.

Our data reported here, along with data from other groups, support the model of SnRK1 as a sensor of low nutrient status and cellular stress. Under low-nutrient or sugar-stress conditions, 5PTase13 acts as a positive regulator of SnRK1.1 activity by reducing the amount of SnRK1 targeted for proteasomal destruction. This regulation most likely involves a 5PTase13-SnRK1.1 interaction within the nucleus, as that is where we find 5PTase13:GFP (Fig. 11), and SnRK1 is most likely nuclear as well (Pierre et al., 2007; Thelander et al., 2007). A loss of 5Tase function, therefore, results in a lack of response, or insensitivity, to stress conditions. In contrast, nuclear PRL1 acts as a negative regulator of SnRK1.1 and facilitates its proteasomal destruction. Thus, the increased SnRK1 levels found in prl1 mutants could potentiate low-nutrient and/or stress signaling, resulting in the hypersensitivity to sugar and ABA found in this mutant. Nemeth et al. (1998) showed that PRL1 is located in the nucleus, as are CUL4 and other SnRK1-interacting proteasomal components such as SKP/ASK1 and a4/PAD1 (Farras et al., 2001; Chen et al., 2006a).

While data support this model, we currently do not know how second messenger InsP3 affects the SnRK1 pathway. We have shown that InsP3 levels increase when seedlings are given a sugar stress, implicating InsP3 as a second messenger under high-sugar conditions (Fig. 10). Given that the 5PTase13 enzyme has been shown to hydrolyze InsP3 in vitro (Zhong and Ye, 2004), we expected that a loss of 5PTase13 function would elevate InsP3. However, 5ptase13 mutants have, instead, lower InsP3 levels upon sugar stress (Fig. 10). The lack of increase in Glc-stimulated InsP3 accumulation in 5ptase13 mutants may indicate a more complex role for InsP3 in 5PTase13 function and/or could be the result of compensatory actions of the other 5Tase enzymes. Elevated InsP3 levels have been found in other 5ptase mutants, and this increase in...
InsP$_3$ is sometimes associated with ABA hypersensitivity (Carland and Nelson, 2004; Gunesekera et al., 2007). It should be noted that 5ptase13 mutant seeds are also ABA insensitive, a trait they share with other sugar-insensitive mutants. It is currently thought that the sugar-induced delay of germination involves the action of ABA, as ABA synthesis mutants are also resistant to the effects of sugars (Price et al., 2003; Rognoni et al., 2007).

There have been previous reports that inositol signaling and sugar sensing/metabolism are linked. Im et al. (2007) found that expressing a human phosphatidylinositol 4-phosphate 5-kinase (PIPK) gene in tissue culture cells dramatically elevated phosphatidylinositol 4-phosphate 5-kinase (PIP5K) gene and resulted in increased sugar use and oxygen uptake. This work importantly defined the PIP5K enzyme as the flux-limiting enzyme for PtdInsP$_2$ production and InsP$_3$ signaling in plant cells. In separate studies, Lou et al. (2007) found that one member of the PIP5K gene family, At3g09920, negatively regulates a cytosolic invertase by direct protein interactions. These investigators found that a gain of function in this same PIP5K gene results in less invertase activity and a resulting sugar insensitivity in transgenic seedlings. Additional evidence supporting the importance of PtdInsP$_2$ in sugar sensing/metabolism comes from transcriptional studies. A rigorous correlative analysis of gene expression showed that different PIP5K genes (At4g17080) of 278 genes coactivated by SnRK1.1 and sugar starvation and down-regulated by sugar treatment (Baena-Gonzalez et al., 2007). Together, these studies indicate that PIP5K-catalyzed production of PtdInsP$_2$ may play a critical role in mediating the response of plant cells to sugar. In addition to transcriptional changes in PIP5K, protoplasts overexpressing SnRK1.1 repress two genes required for inositol synthesis (At4g39800 and At3g02870) and activate three genes containing second messengers such as InsP$_3$.

Together, the data presented here support a unique role for 5PTase13 functioning as a binding partner and regulator of the SnRK1.1 modulator of energy and stress signaling. Given the previously established role of 5PTase13 in blue light signaling (Chen et al., 2008), this work identifies a new connection between inositol signaling and the sensing of cellular metabolism, energy, light, and stress.

**Yeast Two-Hybrid Screen**

The Matchmaker Two-Hybrid System 3 was used (BD Biosciences Clontech). The cDNA corresponding to amino acids 1 to 533 of 5PTase13 was amplified by PCR and ligated into pGBKKT bait vector (BD Biosciences Clontech), verified by DNA sequencing, and transformed into yeast strain AH109. The yeast strain AH109 containing the 5PTase13 WD40 repeat domain was transformed with an Arabidopsis (Arabidopsis thaliana) 3-dold etiolated seedling cDNA library (Thelosis). Screening for interactors was performed with SD/-Ade-His-Leu-Trp, 10 mg 3-amino-1,2,4-triazole, and 20 mg mL$^{-1}$ 5-bromo-4-chloro-3-indolyl-$b$-D-galactopyranoside plates. Candidate clones that grew were rescued from yeast and retested in the original bait and control strains. Prey plasmids that passed all tests were sequenced to identify the Arabidopsis gene insert.

**Mutant Isolation**

Arabidopsis ecotype Columbia was used for all experiments. Growth conditions of soil-grown plants have been described (Berdy et al., 2001). Regular-light conditions were 100 mE for 16-h days, while low-light conditions were 40 mE for 16-h days. Sptase13-1 (At1g05630, SAIL_350_F1) and Sptase13-2 (SALK_081991) mutants were identified from the SGnAR database (Alonso et al., 2003). Methods for PCR screening of mutants has been described (Ercetin et al., 2008) and utilized SALK and SAIL LB primers and 5PTase13 gene-specific primers (13-16or, 13-1rev, 13-2for, and 13-2rev; Supplemental Table S4). The resulting PCR fragments were sequenced to map the T-DNA insertions.

**Reverse Transcription-PCR**

Conditions for reverse transcription (RT)-PCR have been described previously (Ercetin et al., 2008) and used the primers listed in Supplemental Table S4. Amplification of the WD40 region of 5PTase13 was performed using primers 13WD40for and 13WD40rev. Conditions for acting (Berdy et al., 2001), KN1 (Knight et al., 1998), and RD29A (Sanchez and Chua, 2001) amplification have been described and generate 428-, 342-, and 715-bp products, respectively. Each RT-PCR experiment was independently repeated at least two times to verify the observed changes in expression.

**Seedling Growth and Seed Germination Assays**

Age-matched seeds used for assays were harvested from plants grown in parallel on the same shelf in a growth room, and seeds were harvested on the same day and ripened for 6 weeks at room temperature. Seeds were surface sterilized and plated on no salts or 0.5% MS salts solution (pH 5.7) containing 0.8% agar. Seeds were stratified at 4°C for 3 d and germinated at 23°C in the light or dark. For seed germination and root growth assays, seeds were plated on medium containing 0%, 1%, 3%, or 6% Glc or mannitol (Sigma-Aldrich) or 0%, 1%, 2%, 3%, or 6% Suc (Sigma-Aldrich) in the light or dark at 23°C. Germination was scored as positive when the radicle protruded through the seed coat. For ABA sensitivity experiments, ABA (Sigma-Aldrich) was dissolved in 100% ethanol and added to cold, sterile medium at a final concentration of 1, 2, or 3 µM ABA. The germination assay was performed as before. Hormone/sugar treatment experiments were repeated two or three times.

**Immunoprecipitations**

The entire open reading frame (encoding 335 amino acids) of Afl3g01090 was amplified with gene-specific primers SnRK1.1pFor and SnRK1.1pRev (Supplemental Table S4) and ligated to pCRT7/CT-TOPO vector. This resulted in p5nkRV5, which directed expression of a 61.7-kD SnRK1.1 protein with a C-terminal 6×His tag and V5 epitope tag (Invitrogen). The WD40 repeat region of 5PTase13 was amplified with gene-specific primers WD40For and WD40Rev (Supplemental Table S4) and ligated to pCRT7/NT-TOPO vector. This resulted in p31WDX, which directed expression of a 65.9-kD protein that corresponds to the WD40 region of 5PTase13 and an N-terminal 6×His tag and Xpress epitope (Invitrogen). For expression, p5nkRV5 and p31WDX were transformed in Escherichia coli BL21(DE3) pLysS cells (Invitrogen), and the resulting recombinant proteins 13WDX and SnRKV5 were purified using nickel-nitrilotriacetic acid agarose columns (Invitrogen). SnRKV5 protein...
Ponceau S staining was performed to ensure that equivalent amounts of mouse horseradish peroxidase-conjugated anti-V5 antibody (Amersham) were used. SDS-PAGE, followed by western blotting with a 1:10,000 dilution of goat anti-mouse monoclonal antibody, followed by a 1:20,000 dilution of goat anti-mouse horseradish peroxidase-conjugated antibody (Amersham), was used. Purified postipe protein (175 ng; Invitrogen) was loaded on the same gel to estimate the amount of recombinant V5- or Xpress-tagged proteins.

**SnRK1 Activity Assay**

Whole 7-d-old seedlings were ground in liquid nitrogen and resuspended in extraction buffer (50 mM Tris-HCl, pH 7.8, 1 mM EDTA, 2 mM dithiothreitol [DTT], 0.05% Nonidet P-40, 0.5 mM phenylmethylsulfonyl fluoride, 1 mM benzamidine, 3 mM glycerophosphate, and plant protease cocktail; Sigma). After centrifugation at 13.2 rpm for 15 min at 4 °C, ammonium sulfate was slowly added to the supernatant to 40% saturation while stirring for 10 min at 4 °C and centrifuged for 15 min at 13.2 rpm and 4 °C. Precipitated protein was resuspended in 50 mM Tris-HCl, pH 7.8, 1 mM EDTA, 1 mM DTT, 0.05% Nonidet P-40, 0.5 mM phenylmethylsulfonyl fluoride, 1 mM benzamidine, 3 mM glycerophosphate, 10% glycerol, and plant protease cocktail; Sigma). Protein concentration was determined according to the Bradford method (Bradford, 1976), and equal amounts of protein were added to each assay. SnRK1 activity assay was performed as described by Raczuk et al. (2006) with slight modifications. Protein extract (5 μg in 5 μL) was mixed with 5 μL of kinase buffer (50 mM Tris-HCl and 1 mM DTT, pH 7.0), 5 μL of sterile water, 5 μL of SPS peptide stock solution (peptide sequence RDHMPRIRSEMQIWSED; 200 μM), and 5 μL of labeled ATP stock solution (5 μM [γ-32P]ATP; 5 μM unlabeled ATP, and 125 mM magnesium chloride). The samples were incubated for 30 min at 30 °C, and 10-μL aliquots were spotted twice onto P81 paper. The pieces were washed three times in 125 mM phosphoric acid for 20 min each and transferred to scintillation vials for counting. Two different control reactions were examined along with each reaction assay. The first reaction control contained no SPS peptide, and the second control contained no protein extract. Activity was expressed as nanomoles of phosphate incorporated into peptide per minute per milligram of protein. The assay was performed using two independently prepared extracts and three replicates of each extract.

**Cell-Free Degradation Assay**

Conditions described by Lee et al. (2008) were followed. Briefly, 7-d-old light-grown seedlings were ground in liquid nitrogen, resuspended in buffer (25 mM Tris, pH 7.5, 10 mM MgCl₂, 5 mM DTT, and 10 mM NaCl), and centrifuged at 13.2 rpm for 10 min at 4°C. Purified recombinant SnRK1.1-V5 (SnRKV; 500 ng) and total cell extracts (30 μg) were mixed in a reaction buffer (25 mM Tris, pH 7.5, 10 mM MgCl₂, 5 mM DTT, 10 mM NaCl, and 10 μM ATP) and incubated at 30°C for the indicated times. Reactions were analyzed by SDS-PAGE, followed by western blotting with a 1:10,000 dilution of goat anti-mouse horseradish peroxidase-conjugated anti-V5 antibody (Amersham). One panel staining was performed to ensure that equivalent amounts of extracts were analyzed.

**Mutant Complementation and GFP Imaging**

The 3,408-bp coding region of 5PTase13 minus the stop codon was amplified by high-fidelity PCR, confirmed by sequencing, cloned into the pENTR/D-TOPO vector (Invitrogen), and recombined via the Gateway system (Invitrogen) using the manufacturer’s instructions into pK7FWG2. The resulting 35S cauliflower mosaic virus promoter:5PTase13:GFP construct was transformed into Agrobacterium tumefaciens by cold shock and was used in the transformation of 5ptase13-1 and wild-type plants as described (Bechtold et al., 1993). 13-1/5PTase13:GFP seedlings were identified on kanamycin plates and by screening for GFP production using a Zeiss Axioimager microscope equipped with fluorescence optics. Two independent complemented lines with detectable GFP expression (13-1/13:GFP-1 and 13-1/13:GFP-2) were used in growth assays and for subcellular localization. Seven-day-old seedlings were used for imaging utilizing Axiovision software (Zeiss). Z-stack series of 15 to 20 μm sections were collected, and deconvolution with an iterative algorithm was applied. The resulting deconvolved images were reconstructed into a single image using the maximal intensity projection function of Axiovision. To visualize nuclei, seedlings were stained with 1 μg mL⁻¹ DAPI (Molecular Probes) solution for 5 min, excess liquid was removed, and the seedlings were mounted in water. Photographs were taken with a Zeiss MC100 camera. GFP was imaged with a filter set consisting of an excitation filter of 540 to 580 nm, a dichroic mirror of 595 nm, and a barrier filter of 600 to 660 nm. DAPI staining was visualized with a standard UV fluorescence filter set.

**Extraction and Measurement of Mass InsP₃**

Filters containing whole 4-d-old dark-grown seedlings were floated on a 6% Glc solution (0.5× MS salts, pH 5.7) or on a control solution (0.5× MS salts, pH 5.7, only) in the dark for 3 d and then frozen in liquid nitrogen at the end of the treatment. Tissues were harvested and mass InsP₃ measurements were made as described previously (Guneseker et al., 2007; Ercetin et al., 2008). The assays were performed in triplicate, and the experiment was repeated two times. Raw values for InsP₃ in Glc-treated samples are as follows: WT1, 1,313 ± 203 pmol g⁻¹; WT2, 1,313 ± 372 pmol g⁻¹.

**Histological Characterization**

The histological analysis was performed as described (Carland et al., 1999) with slight modifications. Seven-day-old wild-type and mutant cotyledons of soil-grown seedlings in a growth room were fixed in ethanol:chloroform:acetic acid solution (6:3:1) overnight at 4°C and cleared sequentially in 80% ethanol, 95% ethanol overnight at 4°C, and 10% NaOH for 1 h at 42°C. Cotyledons then were stained briefly (30 s to 2 min) in 0.002% safranin-O (Sigma). Specimens were mounted on slides in 50% glycerol and visualized with differential interference contrast on a Zeiss Axioimager microscope and Spot digital camera (Zeiss). A total of 50 cotyledons per variant were used.

Sequence data from this article can be found in the GenBank/EMBL data libraries under the following accession numbers: 5PTase13, At1g5630, NP_172054; 5PTase11, At1g0190, NP_001115846; 5PTase12, At2g4300, NP_189118; 5PTase14, At2g31830, NP_180742; FRA3, At1g65580, NP_001118546; 5PTase12, At5g5880, NP_176736.

**Supplemental Data**

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

**Supplemental Figure S1.** GENEVISITATOR expression analysis.

**Supplemental Figure S2.** Validation of the SnRK1 activity assay.

**Supplemental Figure S3.** 5ptase13 mutants are less sensitive to 6% Glc under light.

**Supplemental Figure S4.** 5ptase13 mutants are less sensitive to 11% Suc under light.

**Supplemental Table S1.** Identity of the WD40 regions of 5PTases in Arabidopsis.

**Supplemental Table S2.** Normal cotyledon vein development patterns in wild-type and 5ptase13 plants.

**Supplemental Table S3.** Abnormal cotyledon vein development patterns in wild-type and 5ptase13 plants.

**Supplemental Table S4.** Primer sequences.

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LITERATURE CITED

tional mutagenesis of Arabidopsis thaliana. Science 301: 653–657
Andersson MX, Kourchtenko O, Dangl JL, Mackey D, Ellerstrom M (2006) Phospholipase-dependent signalling during the AvrRpm1- and
Arabidopsis glucos insensitive mutants, gin5 and gin6, reveals a central role of the plant hormone ABA in the regulation of plant
vegetative development by sugar. Genes Dev 14: 2085–2096
Astle MV, Horan KA, Ooms LM, Mitchell CA (2007) The inositol
polyphosphate 5-phosphatases: traffic controllers, waistsline
integrator of transcription networks in plant stress and energy signal-
ing, Nature 448: 938–942
Bechtold N, Ellis J, Pelletier G (2001) Molecular characteri-
azation of AtSPTase1, an inositol phosphatase capable of terminat-
ing IP3 signaling, Plant Physiol 126: 801–810
chem Soc Symp 74: 1–7
Bhalerao RP, Salchert K, Bako L, Okresz L, Szabados L, Muranaka T,
inositol polyphosphate 5-phosphatase: towards an understanding of subcellular signaling. Subcell Biochem 39: 181–205
Bradford MM (1976) A rapid and sensitive method for the quantitation of
microgram quantities of protein utilizing the principle of protein-dye
binding. Anal Biochem 72: 248–254
Burnette RN, Gineseker BM, Gillaspy GE (2003) An Arabidopsis ino-
sitol 5-phosphatase gain-of-function alters abscisic acid signaling. Plant
Physiol 132: 1011–1019
Carland FM, Berg BL, FitzGerald JN, Jinamorphongs S, Nelson T, Keith
Plant Cell 11: 2123–2137
inositol (1,4,5) triphosphate signal transduction is essential for closed
titin ligase with RBX1 and the CDD complex in mediating light control of
phate 5-phosphatase functions in PHOTOTROPIN signaling in Arabi-
signaling proteins involved in sugar and abscisic acid signaling in
Clement S, Krause U, Desmedt F, Tanti JF, Behrends J, Pesesse X, Sasaki
phosphatidylinositol(4,5)bisphosphate biosynthesis increases phospho-
DeWald DB, Torabinejad J, Jones CA, Shope JC, Cangelosi AR, Thompson
Clement S, Krause U, Desmedt F, Tanti JF, Behrends J, Pesesse X, Sasaki
phosphatidylinositol(4,5)bisphosphate biosynthesis increases phospho-
morphogenesis transcriptome reveals the molecular identity of six
genes with roles in root-hair development in Arabidopsis. Plant J 45:
83–100
Kawaga S, Soeda Y, Ishihara H, Oya T, Sasahara M, Yaguchi S, Oshita R,
Wada T, Tsuneki H, Sasaoka T (2008) Impact of transgenic overexpres-
sion of SH2-containing inositol 5-phosphatase 2 on glucose metabolism and
insulin signaling in mice. Endocrinology 149: 642–650
calcium signalling pathways in Arabidopsis. Plant J 16: 681–687
phate receptor in higher plants: is it real? J Exp Bot 58: 361–376
Laby RJ, Kincaid MS, Kim D, Gibson SI (2000) The Arabidopsis sugar-
insensitive mutants sis4 and sis5 are defective in abscisic acid synthesis
Lee JH, Terzaghi W, Gusmaroli G, Charrn JR, Yoon HJ, Chen H, He YJ,
proteins and their roles as substrate receptors for CUL4RING E3
Lee Y, Kim HY, Jeon BW, Park KY, Suh SJ, Jeo J, Kwak JM, Martinoia E,
Hwang I (2007) Phosphatidylinositol 4,5-bisphosphate is important for
stomatal opening. Plant J 48: 583–598
Li D, Roberts R (2001) WRD-repeat proteins: structure characteristics,
biological function, and their involvement in human diseases. Cell
Mol Life Sci 58: 2085–2097
Etoposic expression of ABSCISIC ACID2/GLUCOSE INSENSITIVE1 in
Arabidopsis promotes seed dormancy and stress tolerance. Plant
Physiol 143: 745–757
AtSPTase1 modulates cotyledon vein development through regulating
auxin homeostasis. Plant Physiol 139: 1677–1691
Lou Y, Gou JY, Xue HW (2007) PIP5K9, an Arabidopsis phosphatidylinositol
monophosphate kinase, interacts with a cytosolic invertebrate to nega-
Lu CA, Lin CC, Lee KW, Chen JL, Huang LE, Ho SI, Liu HJ, Hsing Y, Yu
SM (2007) The SnRK1A protein kinase plays a key role in sugar
signaling during germination and seedling growth of rice. Plant Cell
19: 2484–2499
Moore B, Zhou L, Rolland F, Hall Q, Cheng WH, Liu YX, Hwang I, Jones
T, Sheen J (2003) Role of the Arabidopsis glucose sensor HXK1 in
nutrient, light, and hormonal signaling. Science 300: 332–336
Nemeth K, Salchert K, Putnoky P, Bhalerao R, Koncz-Kalman Z, Stankovic-
Pleiotropic control of glucose and hormone responses by PRL1, a
Ortega X, Perez LM (2001) Participation of the phosphoinositide metabo-
ism in the hypersensitive response of Citrus limon against Alternaria
alternata. Plant Mol Biol 41: 43–50
inositol 1,4,5-trisphosphate in graviotropic signaling and the retention of
cold-perceived gravistimulation of oat shoot pulvinii. Plant Physiol 125:
1499–1507


