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PHT4;2 Influences Starch Accumulation and Leaf Size

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The sink-specific plastidic phosphate transporter PHT4;2 influences starch accumulation and leaf size in Arabidopsis

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ABSTRACT
Nonphotosynthetic plastids are important sites for the biosynthesis of starch, fatty acids and amino acids. The uptake and subsequent use of cytosolic ATP to fuel these and other anabolic processes would lead to accumulation of inorganic phosphate (Pi) if not balanced by a Pi export activity. However, the identity of the transporter(s) responsible for Pi export is unclear. The plastid-localized Pi transporter PHT4;2 of Arabidopsis thaliana is expressed in multiple sink organs but is nearly restricted to roots during vegetative growth. We identified and used pht4;2 null mutants to confirm that PHT4;2 contributes to Pi transport in isolated root plastids. Starch accumulation was limited in pht4;2 roots, which is consistent with inhibition of starch synthesis by excess Pi as a result of a defect in Pi export. Reduced starch accumulation in leaves and altered expression patterns for starch synthesis genes and other plastid transporter genes suggest metabolic adaptation to the defect in roots. Moreover, pht4;2 rosettes, but not roots, were significantly larger than those of wild type with 40% greater leaf area and twice the biomass when plants were grown with a short (8 hr) photoperiod. Increased cell proliferation accounted for the larger leaf size and biomass as no changes were detected in mature cell size, specific leaf area or relative photosynthetic electron transport activity. These data suggest novel signaling between roots and leaves that contributes to the regulation of leaf size.
INTRODUCTION

Unlike chloroplasts, nonphotosynthetic plastids cannot synthesize the ATP needed to fuel anabolic processes within these organelles, which include biosynthesis of starch, fatty acids and amino acids (Neuhaus and Emes, 2000). Metabolism in such plastids, therefore, is dependent on the import of ATP from the cytosol. Nucleotide transporters (NTTs) located in the plastid inner envelope fulfill this energy requirement by catalyzing the stoichiometric exchange of cytosolic ATP for stromal ADP (Reiser et al., 2004; Reinhold et al., 2007). A consequence of the unbalanced phosphate moieties associated with this exchange is that inorganic phosphate (Pi) would accumulate to deleterious levels within the stroma if not countered by Pi export. Despite the fundamental importance of Pi homeostasis to plastid functions and, in turn, to plant growth and development, the identity of the transporter(s) responsible for this activity has been elusive.

In addition to ATP/ADP exchange, NTTs can accept Pi as a third substrate to transport Pi-Pi homo-exchange or to co-transport Pi plus ADP in exchange for ATP (Trentmann et al., 2008). Although the latter activity would preclude the necessity for a distinct Pi transporter, studies with cauliflower bud amyloplasts revealed Pi export activity in the absence of potential counter-exchange substrates, i.e., unidirectional transport, indicating that transporters other than NTTs contribute to net Pi export (Neuhaus and Maass, 1996).

Plastid-localized Pi transport systems identified to date include members of the plastidic Pi translocator (pPT), PHT2 and PHT4 families. pPTs catalyze exchange of Pi with phosphorylated compounds, and therefore would not support net Pi export (Flügge, 1999; Eicks et al., 2002). In contrast, members of the PHT2 and PHT4 families mediate H+ and/or Na+-dependent Pi transport (Daram et al., 1999; Versaw and Harrison, 2002; Zhao et al., 2003; Guo et al., 2008; Ruiz Pavón et al., 2008). The role of these transporters in nonphotosynthetic plastids is unclear. Gene expression studies suggest that all of these transporters are present in limited types of heterotrophic tissues, but most are expressed predominantly in autotrophic tissues (Rausch et al., 2004; Guo et al., 2008). PHT4;2 (At2g38060) is the sole exception to this expression pattern as its transcripts are nearly restricted to roots (Guo et al., 2008).

In this study, we used Arabidopsis thaliana mutants that lack a functional copy of the PHT4;2 gene to gain insight into the roles of the encoded transporter in plastid and whole plant physiology. We confirmed that PHT4;2 contributes to Pi transport in isolated root plastids, and found that starch accumulation is reduced in both roots and leaves of the mutant plants. We also
discovered a surprising conditional growth phenotype in which rosette leaves of *pht4;2* mutants were larger than those of wild type when plants were grown with a short (8 hr) photoperiod, and increased cell proliferation accounted for the greater leaf area and biomass.

Mature leaf size is an intrinsic trait that is a function of both the size and number of cells in the organ. The mechanisms that control final leaf size are poorly understood. In some cases, a reduction in cell number through mutation or transgenes is offset by a compensatory increase in cell size (Hemerly et al., 1995; De Veylder et al., 2001; Horiguchi et al., 2006). In contrast, a number of modifiers of leaf size have been identified that primarily affect cell number, but these represent diverse functional classes, including transcriptional regulation, auxin signaling, protein synthesis and DNA replication (for review, see Gonzalez et al., 2009; Krizek, 2009). A recent comparative analysis of a subset of these modifiers confirmed that multiple independent pathways contribute to the control of cell proliferation in leaves (Gonzalez et al., 2010). Because none of the previously described modifiers of leaf size appear to be involved in plastid functions, and because *PHT4;2* expression during vegetative growth is restricted to roots, the effect of *pht4;2* mutations on cell proliferation suggests novel signaling between roots and developing leaves that also contributes to the regulation of leaf size.

**RESULTS**

**Localization of *PHT4;2* Expression in Arabidopsis**

We reported previously that *PHT4;2* expression was restricted to roots and exhibited a diurnal but not circadian pattern in which transcript abundance was greater in the dark than the light (Guo et al., 2008). Expression data in the Genevestigator microarray database (Hruz et al., 2008) support these patterns and suggest that *PHT4;2* is also expressed in floral tissues. To investigate spatial expression in greater detail, a *PHT4;2* promoter-GUS transcriptional fusion construct was introduced into Arabidopsis and GUS activity was assessed in progeny of the transgenic plants. To maximize expression, plants were grown with a short (8 hr) photoperiod.

As expected, GUS activity was readily detected in roots but not in leaves (Fig. 1A). GUS activity was also detected in sepals, stamens and carpels, as well as silique valves and septum (Fig. 1A), but not in the mature seeds. Quantitative RT-PCR analysis of *PHT4;2* transcripts in wild-type plants corroborated these findings and indicated that expression was greatest in roots, followed by intermediate levels in flowers and siliques, and barely detectable levels in rosette
and cauline leaves (Fig. 1B). These results extend our previous studies to suggest that in addition to roots, PHT4;2 is broadly expressed in carbon import-dependent sink organs. However, even in roots, PHT4;2 transcript levels were low suggesting that abundance of the encoded protein would also be limited. Indeed, PHT4;2 was detected by western blot analysis in small amounts in total protein extracts from roots, but not in extracts from other organs, and the detection was further complicated by the presence of several nonspecific bands (data not shown). Next, we isolated microsomes from different organs to enrich the cellular membrane protein fractions and detected PHT4;2 as a 35 kDa band in samples prepared from roots, but the abundance of this protein in flowers and siliques still remained below our limit of detection (Supplemental Fig. S1). The identity of the 35 kDa band as PHT4;2 was confirmed in blots prepared from mutant and wild-type root plastids, and this enrichment also eliminated nonspecific signals (see below). The significance of expression in other carbon import-dependent sink organs such as flowers and siliques is unclear because the encoded protein could be detected by western blot analysis only in roots (Supplemental Fig. S1).

Identification of pht4;2 T-DNA Insertion Mutants

To obtain insight into the role of PHT4;2 in plastid function and overall plant metabolism, we required mutants that lacked a functional PHT4;2 allele. We obtained two independent T-DNA insertion lines from the SALK collection (Alonso et al., 2003), SALK_019289 (pht4;2-1) and SALK_070992 (pht4;2-2). Insertions were confirmed by PCR and sequencing of amplicons that spanned the insertion junctions. The pht4;2-1 line has a tandem T-DNA insertion 2170 bp downstream of the translation start site, and a 7 bp deletion at the insertion site. The pht4;2-2 line has a single T-DNA inserted within the first intron, 738 bp downstream of the translation start site, and also has a deletion of 4 bp at the insertion site. The structures of the mutant loci are represented in Fig. 2A.

Homozygous mutant lines were identified by PCR then screened by DNA gel blot for the presence of additional insertions. At least one backcross to wild type was required to segregate ectopic insertions from the genetic backgrounds. Only homozygous mutants that lacked ectopic insertions were used in subsequent studies.
RT-PCR analysis of root RNA using primers that anneal downstream of the insertion sites revealed no PHT4;2 transcripts in the homozygous mutants suggesting that both mutant alleles are null (Fig. 2B).

**Localization of PHT4;2 to Root Plastids**

To confirm localization of the endogenous PHT4;2 protein to root plastids, intact plastids were isolated from roots of hydroponically-grown wild-type Arabidopsis, and the proteins were subjected to western blot analysis. As shown in Fig. 3A, a single 35 kDa band reacted with the antibody. The signal intensity varied in proportion to the amount of total protein loaded in the well and no reaction was detected when the corresponding pre-immune serum was used, indicating that the antibody is highly specific. No reacting proteins were detected in leaf chloroplasts (data not shown), in accord with transcript localization results (Fig. 1).

Western blot analysis was also carried out with plastids isolated from roots of pht4;2 mutants. Equal amounts of total plastid protein (15 µg protein per lane) from wild type and each mutant were evaluated. The 35 kDa band was detected in the wild-type preparation but in neither of the mutants (Fig. 3B), which further verified that the mutations are null.

PHT4;2 encodes a 512 aa protein (UniProt Q7XJR2) that is predicted by TargetP (Emanuelsson et al., 2007) to include an N-terminal 44 aa transit peptide. The theoretical molecular mass of the processed protein is 50.5 kDa, which differs significantly from the 35 kDa estimated from SDS-PAGE. Two other members of the PHT4 family, PHT4;1 and PHT4;4, also exhibit anomalous migration in SDS-PAGE (Roth et al., 2004; Ruiz Pavón et al., 2008). The faster gel migrations of PHT4;2 and these other PHT4 proteins may be a function of over-proportional binding of SDS, which is common for highly hydrophobic integral membrane proteins (Rath et al., 2009).

**Pi Transport in Isolated Root Plastids**

Previous studies indicated that PHT4;2, as well as the other five members of the Arabidopsis PHT4 family, mediate Pi transport that is dependent on the presence of a H+ electrochemical gradient when expressed in yeast (Guo et al., 2008). Interestingly, one of these same proteins, PHT4;1, was found to catalyze Na⁺-dependent Pi transport when expressed in *Escherichia coli* (Ruiz Pavón et al., 2008). Thus, different heterologous systems, perhaps due to
differences in membrane lipid composition and cell physiology, impart or reveal unique qualities of the Pi transport process. Here we have taken advantage of the availability of \textit{pht4;2} knockout lines to study PHT4;2 in its native context of root plastids.

Plastids isolated from roots of hydroponically-grown wild-type and \textit{pht4;2} plants had equivalent intactness (70-75\%) as measured by phase contrast microscopy. To assess purity and yield of the plastid preparations, the activities of organelle-specific marker enzymes were measured in crude extracts (filtered root homogenates) and in the enriched plastid preparations (Table I). Specific activities of the marker enzymes could be determined only for the enriched plastid preparations due to the presence of bovine serum albumin (BSA) in the isolation buffer. Approximately 7\% of the total activity of the plastid marker nitrite reductase (NiR) was recovered in each plastid preparation, indicating that plastid enrichment was equivalent for each plant genotype. Contamination of these preparations by other organelles was also equivalent for each plant genotype with 0.33-0.44\% of the total activity of NADH-malate dehydrogenase (NADH-MDH), a marker for combined mitochondrial, cytosolic and peroxisomal contamination, and 0.55-0.86\% of the total activity of the vacuolar marker \(\alpha\)-mannosidase. Microbial contamination of plastid preparations, which was assessed by growth on LB agar medium, was also similar with 200-250 colony forming units mg\(^{-1}\) protein, which corresponds to a low number of microbial cells (6 to 8) in samples used for Pi transport measurements. Pi uptake by root plastids was linear for 60 s regardless of genotype but the rate of transport measured with wild-type plastids was twice that of plastids isolated from \textit{pht4;2} mutants (Supplemental Fig. S2). Because the purity and enrichment of root plastids were equivalent, we attribute the reduced transport rates with mutant plastids solely to the absence of PHT4;2.

Pi uptake by root plastids isolated from wild-type plants was significantly greater than those isolated from \textit{pht4;2} mutants when 5 mM NaCl was included in the assay, but not when this salt was omitted (Fig. 4A) or when all sources of Na\(^+\) were substituted with K\(^+\) (data not shown). Pi transport activities for all plastids were greater at pH 7.5 than at pH 6.5, but significant differences between mutants and wild-type were observed only in the presence of NaCl. These results suggest that PHT4;2 catalyzes Na\(^+\)-dependent Pi transport in root plastids.

To better understand the mechanism of the observed Pi uptake activity we investigated the effects of select pharmacological agents. Transport assays were conducted in the presence of NaCl and at pH 6.5, and Pi uptake was expressed as a percentage of the amount of Pi
accumulated in untreated wild-type plastids. All three of the compounds we tested, mersalyl, pyridoxal 5’-phosphate (PLP) and carbonyl cyanide p-(trifluoromethoxy)-phenyl-hydrazone (FCCP) had a pronounced inhibitory effect on Pi transport in wild-type plastids and a lower but consistent effect in mutant plastids (Fig. 4B), suggesting that PHT4;2 is the predominant but not sole source of Pi transport activity under these assay conditions. Addition of mersalyl, a thiol-reactive agent (Amores et al., 1994), reduced wild-type transport to 35%, indicating the importance of sulfhydryl groups for PHT4;2 activity. Similarly, treatment with PLP, a lysine-reactive agent known to inhibit pPTs (Gross et al., 1990; Flügge, 1992), reduced wild-type activity to 23%. Addition of the protonophore FCCP reduced transport in wild-type plastids to 40%, indicating that the concentration and/or membrane potential components of the H⁺ electrochemical gradient also affect PHT4;2 activity. Dissipation of membrane potential by FCCP would also diminish the magnitude of a Na⁺ electrochemical gradient.

The Pi uptake assays described above verified that root plastids could be preloaded with Pi to levels sufficient to assess its export. Plastids were preloaded in the presence of 5 mM NaCl at pH 6.5 or 7.5 then washed and resuspended in buffer with or without NaCl. After incubation, the amounts of Pi exported and retained in the plastid were both measured, and export was calculated as a percentage of total Pi to account for differences in preloading. As shown in Fig. 4C, Pi export was significantly reduced in root plastids isolated from pht4;2 mutants under most of the conditions tested, confirming that PHT4;2 contributes to this activity in wild-type plastids. Export activities for wild-type plastids measured at pH 7.5 were equivalent when pH, NaCl or both were varied after preloading (Fig. 4C), suggesting an apparent lack of specificity for either H⁺ or Na⁺. In contrast, similarly low export activities were detected in mutant and wild-type plastids when the composition of the preloading and export solutions were identical (data not shown).

**Starch Accumulation**

Pi is an allosteric inhibitor of ADP-glucose pyrophosphorylase (AGPase), which catalyzes the first committed step in starch biosynthesis within plastids (Preiss, 1982; Ballicora et al., 2004). Therefore, we reasoned that starch accumulation in roots could serve as an in vivo indicator for defects in plastidic Pi homeostasis associated with the absence of PHT4;2. To test this hypothesis, we measured starch contents of roots that were harvested from hydroponically-
grown, 6-wk-old wild-type and *pht4;2* plants at multiple times throughout an 8 hr light/16 hr dark photocycle. As reported previously for hydroponically-grown Arabidopsis (Malinova et al., 2011), starch was distributed throughout the entire root rather than restricted to the root tip (data not shown). Starch levels in wild-type roots doubled during the first half of the photoperiod then returned to near-starting levels by the end of the photoperiod and remained relatively constant throughout the dark portion of the photocycle (Fig. 5). In contrast, starch levels in *pht4;2* roots were nearly constant throughout the entire photocycle and were significantly lower than those in wild-type roots at nearly all time points measured. This reduced capacity for starch accumulation is consistent with Pi inhibition of starch synthesis due to a defect in Pi export.

Starch contents of leaves were also lower in the mutants than wild type but only during the light portion of the photocycle (Fig. 5). Because *PHT4;2* transcripts and protein in wild-type plants were restricted to roots during vegetative growth, the diminished accumulation of starch in mutant leaves presumably reflects a secondary consequence of metabolic changes in the roots. Consistent with this idea, we detected root- and leaf-specific changes in the expression of a number of genes that are either directly or indirectly involved in starch synthesis (Fig. 6). Specifically, transcript levels for *GPT1* and *GPT2*, which encode Glc6P/Pi translocators (GPTs), and *NTT1* and *NTT2*, which encode ATP/ADP nucleotide translocators (NTTs), were induced in roots but not in leaves of *pht4;2* plants (Fig. 6A). The proposed functions of GPT and NTT are to provide nonphotosynthetic plastids with the carbon and energy, respectively, needed for starch synthesis and the oxidative pentose phosphate pathway, although mutant analyses indicate that these proteins also have roles in photosynthetic tissues (Niewiadomski et al., 2005; Reinhold et al., 2007). In addition, expression of the AGPase small subunit gene, *APS1*, was induced only in roots of *pht4;2* plants, whereas AGPase large subunit genes, *APL1* and *APL3*, were induced only in leaves (Fig. 6B). Because transcript levels for these genes were inversely correlated with starch levels, the altered expression patterns may reflect compensatory responses to the effects of *pht4;2* mutations. Similarly, the cytosolic fructose-1,6-bisphosphatase gene (*cFBP*), which encodes a key enzyme in sucrose biosynthesis (Stitt et al., 2010), was induced only in *pht4;2* leaves (Fig. 6B) but we detected no significant changes in the accumulation of sucrose in leaves or roots of *pht4;2* plants (Fig. 5).

**Plant Size and Biomass**
The *pht4;2* mutants have no obvious morphological or developmental phenotype when grown with a 14 hr photoperiod. However, when plants were grown with a shorter photoperiod (8 hr light), fully-expanded *pht4;2* rosettes were notably larger than those of wild type (Fig. 7A). Both mutant and wild-type plants had 43 ± 1 rosette leaves when flowering initiated, indicating that the increase in rosette size was not due to additional leaves but rather an increase in leaf size (Fig. 7B). Because individual leaves emerge and complete the expansion phase of organ growth at different chronological times, differences in rosette sizes were not detected until plants had grown for at least 5 wk but then increased to a maximum when flowering initiated at 8 to 9 wk (Fig. 7C).

The large-leaf phenotype was also observed when plants were grown hydroponically, and the magnitude of the size difference was typically greater than that seen with plants grown in soil. Rosette area of fully-expanded *pht4;2* mutants was 45 to 65% greater than wild type, and biomass was at least twice that of wild type (Table II). Roots at the time of harvest (8 wk) were tangled, so it was not possible to accurately assess growth of individual root systems. Therefore, we used the combined weight of root systems to derive an average fresh weight per plant. Results from four independent growth experiments consisting of 74 to 113 plants indicated no significant differences in root growth (0.86 ± 0.17, 0.88 ± 0.18 and 0.73 ± 0.10 g plant^{-1} for wild type, *pht4;2-1* and *pht4;2-2*, respectively). In addition, when plants were allowed to complete their life cycle, no differences between *pht4;2* mutants and wild type were detected in flowering time, flower size or seed size (data not shown).

It was possible that, in addition to increased area, leaf thickness also contributed to the greater biomass of *pht4;2* leaves. To estimate thickness of leaves throughout a rosette we took advantage of the relationship between leaf thickness and the product of specific leaf area (leaf area per unit dry mass) and leaf dry matter content (ratio of dry mass to fresh mass) (Vile et al., 2005). Leaf punches from the centers of ten fully expanded rosette leaves were collected from mutant and wild-type plants. Both specific leaf area, 0.51 ± 0.02 cm\(^2\) mg\(^{-1}\), and leaf dry matter content, 0.10 ± 0.01 mg mg\(^{-1}\) were identical for *pht4;2* and wild-type leaves. From these values and a leaf density of 1 g cm\(^{-3}\) (Vile et al., 2005), we calculated an average thickness of 196 ± 2 μm for both mutant and wild-type leaves, which is similar to values obtained from direct measurements of leaf sections and three-dimensional imaging of mature leaves (Wuyts et al., 2010). Consequently, the increased area of *pht4;2* leaves fully accounts for increased biomass.
In the course of our investigation of leaf size we noted that \textit{PHT4;2} is located adjacent to \textit{DET2} on chromosome 2 (loci At2g38060 and At2g38050, respectively) with 550 bp between the \textit{PHT4;2} stop codon and the \textit{DET2} translation start site. \textit{DET2} encodes a steroid 5\(\alpha\)-reductase that catalyzes a key step in BR biosynthesis (Noguchi et al., 1999). This close proximity raised the possibility that the large-leaf phenotype we attributed to disruption of \textit{PHT4;2} was instead caused by over-expression of \textit{DET2} via T-DNA activation (Weigel et al., 2000; Ren et al., 2004). However, this possibility is inconsistent with our finding that the large-leaf phenotype (assessed by rosette fresh weight) segregated as a recessive trait in F2 progeny of a cross between wild type and the homozygous mutant (14 wild type \([1.69 \pm 0.06 \text{ g}]\); 45 hemizygous mutant \([1.56 \pm 0.10 \text{ g}]\); 15 homozygous mutant \([2.17 \pm 0.12 \text{ g}]\); \(\chi^2=3.48, p=0.17\). Furthermore, quantitative RT-PCR indicated that \textit{DET2} transcript level was unaffected by \textit{pht4;2} mutation (data not shown).

\textbf{Leaf Cell Size, Number and Ploidy}

Leaf size is dependent on both the number and the size of cells in the organ. To determine whether enlargement of \textit{pht4;2} leaves was associated with increased cell number, cell size or a combination of these parameters, we compared the ninth rosette leaves of \textit{pht4;2} and wild-type plants at different stages of leaf expansion (Table III). When plants were grown for 7 wk the ninth rosette leaves were fully expanded and the leaf-size phenotype was apparent with 23\% greater leaf area for \textit{pht4;2} mutants. The size of epidermal cells, however, was not significantly different. Consequently, \textit{pht4;2} leaves had 30-35\% more cells than wild type. Expansion of the ninth rosette leaf was incomplete when plants were grown for 5 wk, and the sizes of \textit{pht4;2} and wild-type leaves were indistinguishable. Nevertheless, the mutant leaves had 34-44\% more epidermal cells than wild type, but the cells were also 26\% smaller. Together, these results indicate that the large-leaf phenotype of \textit{pht4;2} plants is the result of increased cell proliferation with subsequent expansion of individual cells to normal size at maturity. This phenotype appears to be conditional for growth with a short photoperiod because when the same analysis was conducted with plants grown with a longer (14 hr) photoperiod, all genotypes exhibited equivalent leaf areas, epidermal cell sizes and epidermal cell numbers (2.64 ± 0.16 cm\(^2\), 2120 ± 120 \(\mu\text{m}\(^2\), 124,530 ± 10,330 cells, respectively).

Repeated cycles of endoreduplication gives rise to somatic polyploidy, which is positively correlated with cell and leaf size (Melaragno et al., 1993; Vlieghe et al., 2005).
However, endoreduplication appears to be unaffected by loss of PHT4;2 because the distribution of ploidy levels in cells of mature pht4;2 and wild-type leaves were nearly identical (Supplemental Fig. S3). Endoreduplication occurs after cell division ceases and continues until leaf expansion is complete (Beemster et al., 2005). Consequently, the similar distribution of ploidy levels also suggests that additional cell proliferation in pht4;2 leaves occurs predominantly, if not exclusively, at early stages in leaf development.

**Photosynthetic Activity**

Leaf enlargement and altered starch accumulation in pht4;2 mutants suggested the possibility of a change in photosynthetic activity. The two mutant lines presented similar fluorescence characteristics to wild-type plants, as reflected in the $F_0$ and $F_m$ values (data not shown). Therefore, the maximum quantum yield for photosystem II photochemistry ($F_v/F_m$) was not significantly different in the wild-type, pht4;2-1 and pht4;2-2 plants, 0.812 ± 0.011, 0.809 ± 0.005 and 0.801 ± 0.003, respectively.

To assess and compare the light saturation of photosynthetic performance in mutant and wild-type plants we recorded a light response curve of chlorophyll fluorescence and calculated the relative electron transport rate (ETR). This analysis revealed that all plants display similar ETR values and reach saturation at approximately 500 µmol photons m$^{-2}$ s$^{-1}$ (Supplemental Fig. S4). Together, these data indicate that the lack of PHT4;2 does not modify photosynthetic electron transport in leaves. As expected from these data, total chlorophyll contents were not significantly different in wild-type, pht4;2-1 and pht4;2-2 leaves (1.14 ± 0.1, 1.15 ± 0.1 and 1.15 ± 0.1 mg g$^{-1}$ FW, respectively).

**DISCUSSION**

Although ATP/ADP exchange fulfills the energy requirement of plastids with limited or no photosynthetic activity, the unbalanced phosphate moieties would lead to detrimental accumulation of Pi and negative charge within the stroma if not countered by Pi export. Earlier biochemical data suggested that an unknown transporter in cauliflower bud amyloplasts catalyzes Pi export in the absence of exogenous counter-exchange substrates (Neuhaus and Maass, 1996), but the effect of Na$^+$ and H$^+$ levels on this activity was not investigated. Our data presented here suggest that PHT4;2 mediates Na$^+$-dependent Pi transport in Arabidopsis root
plastids and, given its spatial expression pattern, is likely to carry out the same function in other sink organs. Our analysis of *pht4;2* null mutants also revealed pleiotropic effects in starch accumulation throughout the plant and cell proliferation in leaves, which underscore the importance of plastidic Pi homeostasis to plant growth and physiology.

We evaluated Pi transport in root plastids isolated from *pht4;2* and wild-type plants and found that export and import activities were both reduced in mutant plastids, indicating that PHT4;2 is capable of catalyzing reversible Pi transport under our assay conditions (Fig. 4). Given that ATP consumption would generate considerable amounts of Pi within the stroma, it is possible that Pi import does not occur under physiological conditions. Moreover, because root plastids and chloroplasts are developmentally related and share many of the same biosynthetic and transport processes (Bräutigam and Weber, 2009; Daher et al., 2010), stromal pH and inner envelope membrane potential of root plastids are likely to be similar to those of dark-adapted chloroplasts (pH ~7; 20-100 mV, interior negative) (Demmig and Gimmler, 1983; Wu et al., 1991), and these conditions would favor Pi export.

Pi uniport, such as the activity described by Neuhaus and Maass (1996), is an attractive possible mechanism for PHT4;2, as this would rectify both the Pi concentration and charge imbalances derived from ATP/ADP exchange. Nevertheless, coupled transport, e.g., symport, as indicated from the effects of Na⁺ and H⁺ we observed with isolated plastids, is also possible provided that the associated charge imbalance is rectified by an independent activity. Indeed, the addition of Na⁺, but not K⁺, to plastids clearly stimulated the Pi import activity of PHT4;2. Pi import was not stimulated by external acidic conditions (pH 6.5) in the absence of Na⁺ (Fig. 4A), indicating that the presence of a H⁺ gradient alone is insufficient to activate Pi transport. However, Na⁺-dependent Pi import was largely inhibited when the imposed H⁺ gradient was dissipated by a protonophore (FCCP, Fig. 4B), suggesting that other aspects of the H⁺ gradient are important for Pi transport. For example, a change in stromal pH would affect the fraction of Pi ionic forms, H₂PO₄⁻ and HPO₄²⁻. Thus, if PHT4;2 preferentially transports monoanionic H₂PO₄⁻, then its inward electrochemical gradient would be considerably less when the stroma and external solution have the same pH than when the stroma is more alkaline than the external solution. Alkalinization of the stroma may occur via Na⁺/H⁺ exchange, as this transport process has been reported to play an important role in homeostasis of cytosolic and stromal pH (Song et al., 2004).
Pi export mediated by PHT4;2 was also found to be Na\(^+\) dependent (Fig. 4C). We propose that this export activity is the *in vivo* function of PHT4;2 in nonphotosynthetic plastids, as this would enable homeostasis with respect to Pi concentration. Although Pi export under conditions where Na\(^+\) was present both during loading and export would suggest that transport can occur in the absence of a Na\(^+\) gradient, a Na\(^+\) gradient may in fact exist under these conditions if Na\(^+\)/H\(^+\) exchangers use the imposed H\(^+\) gradient to import Na\(^+\) into the stroma. Additional studies using mutants with limited plastidic Na\(^+\)/H\(^+\) exchange activities are needed to resolve this possibility. However, it is important to note that the lack of significant Pi export when the composition of the import and export solutions were the same (no pH or Na\(^+\) gradient) strongly supports the idea that PHT4;2 requires a Na\(^+\) gradient. Furthermore, this requirement may be common to PHT4 transporters because mutational analysis identified a Ser that is critical for the Na\(^+\)-dependency of Pi transport by PHT4;1, and this amino acid is fully conserved in the PHT4 family (Ruiz Pavon et al., 2010).

*pht4;2* mutants exhibited abnormal starch accumulation as well as increased leaf cell proliferation with a concomitant increase in biomass. These pleiotropic effects could initiate with altered homeostasis of Pi levels in the stroma of root plastids. Specifically, we expected a defect in Pi export to result in elevated stromal Pi levels. Confirmation of this effect through direct measurement in isolated plastids was compromised by ongoing metabolism during the isolation procedure. However, reduced starch accumulation in *pht4;2* roots was consistent with an increase in stromal Pi concentrations, as Pi is an inhibitor of starch biosynthesis (Preiss, 1982; Ballicora et al., 2004).

Starch levels were also reduced in *pht4;2* leaves. The lack of *PHT4;2* transcripts and protein in wild-type leaves suggests that the starch phenotype in mutant leaves is a secondary consequence of the metabolic defect in roots. For example, although the altered expression of other plastid transporter genes and *APS1* in roots but not leaves (Fig. 6A) suggests a local response to the defect in root plastids, altered expression of a subset of starch and sucrose metabolism genes in leaves but not roots (Fig. 6B) implicates a defect in long-distance signalling of metabolic status. The strong induction of *GPT2* in *pht4;2* roots further substantiates the relationship between PHT4;2 and starch accumulation because *GPT2* is also induced in a number of starch-deficient mutants (Kunz et al., 2010).
The partitioning of carbon between starch and sucrose biosynthetic pathways is tightly regulated to coordinate carbon demands throughout the day and night. Thus, it might be expected that reduced starch accumulation in *pht4;2* plants would be accompanied by an increase in sucrose levels. However, we detected no significant changes in sucrose levels in roots or leaves of *pht4;2* mutants. This finding does not preclude the possibility that the partitioning of carbon to other sugars, amino acids or lipids is altered, and that one or more of these metabolites are directly involved in stimulating cell proliferation in *pht4;2* leaves. Additional studies that include defining the full metabolite profiles of roots and leaves of *pht4;2* and wild-type plants are needed to resolve these possibilities.

An increase in the number of cells in *pht4;2* leaves fully accounted for the greater rosette area and biomass, as epidermal cell sizes and leaf thickness were equivalent to wild type when leaves were fully expanded. When leaves were examined prior to full expansion the increase in cell number was already evident, indicating that cell proliferation was augmented early in leaf development, which is in accord with the timing of the proliferative phase of leaf growth (Beemster et al., 2005). However, there was a corresponding reduction in cell size at this time point such that leaf sizes were equivalent to wild type. Thus, the leaf-size phenotype was apparent only after cells expanded to normal sizes. Although relative photosynthetic electron transport rate (normalized to leaf area) was unaffected by *pht4;2* mutations, the greater total leaf area of the mutants would provide a proportional increase in the photosynthetic activity per plant and thereby support a concomitant increase in biomass.

The mechanisms controlling the size of leaves and other organs are poorly understood, largely because this trait is regulated through multiple pathways (Gonzalez et al., 2009; Krizek, 2009). Moreover, Gonzalez et al., (2010) recently demonstrated that several of these pathways, including those associated with auxin, jasmonate, brassinosteroid and GA synthesis or signaling, as well as transcriptional regulation by GROWTH-REGULATING FACTOR5 (GRF5), which is not associated with changes in phytohormones, independently control cell proliferation and leaf size. Indeed, the phenotypes of lines evaluated in this study differ from each other and from *pht4;2*, primarily with respect to leaf positions. For example, overexpression of AVP1, a vacuolar pyrophosphate-dependent H⁺ pump that is involved in auxin transport (Li et al., 2005), yields rosettes with a greater number of leaves, all of which are enlarged. Overexpression of JAW1, which is involved in the regulation of jasmonate biosynthesis (Palatnik et al., 2003;
Schommer et al., 2008), leads to enlargement of only the first rosette leaves and these have a characteristic uneven shape. Leaf enlargement is also restricted to the first few rosette leaves in lines overexpressing GRF5 and BRI1, a brassinosteroid receptor (Wang et al., 2001; Horiguchi et al., 2005), whereas overexpression of GA20OX1, an enzyme involved in GA synthesis (Huang et al., 1998), yields enlarged young but not older leaves.

Cell cycle progression is responsive to carbon status (Riou-Khamlichi et al., 2000), and it is possible that the influence of pht4;2 mutations on starch accumulation constitutes one of the multiple pathways that converge on control of cell proliferation and organ size. Starch-deficient mutants do not have abnormally large leaves (Ventriglia et al., 2008), suggesting that reduced starch accumulation is unlikely to be causal for the increase in cell proliferation underlying the pht4;2 leaf-size phenotype. However, starch accumulation in such mutants is affected throughout the entire plant so comparisons may be misleading if the leaf-size phenotype is related to carbon balance between source and sink organs. In addition, our results indicate that the mechanism responsible for increased cell proliferation is non-autonomous, suggesting that PHT4;2 plays a key role in coordinating metabolic signals throughout the entire plant.
MATERIALS AND METHODS

Plant Material and Growth Conditions

*Arabidopsis thaliana* T-DNA insertion lines SALK_019289 (*pht4;2-1*), and SALK_070992 (*pht4;2-2*) were obtained from ABRC. Insertion sites and zygosity were confirmed by PCR using combinations of *PHT4;2*-specific primers and primers that anneal to the T-DNA right and left borders. The sequence of amplicons that spanned insertion junctions was confirmed. Homozygous *pht4;2-1* and *pht4;2-2* plants were screened by DNA gel blot analysis using the *nptII* segment of the binary plasmid pROK2 as probe to identify individuals with insertions at a single locus.

Plants were grown in chambers at 21°C with 70% relative humidity and an 8 hr photoperiod (150 μmol m\(^{-2}\) s\(^{-1}\)) in soil (SunGro Redi-earth, Bellevue, WA) or hydroponically as previously described (Noren et al., 2004; Guo et al., 2008).

Analysis of *PHT4;2* promoter-GUS Expression

A *PHT4;2* promoter-GUS transcriptional fusion (Guo et al., 2008) was introduced into *Agrobacterium tumefaciens* strain GV3101 and used to transform *Arabidopsis* (Clough and Bent, 1998). Transgenic seedlings were selected on half-strength Murashige and Skoog medium (Murashige and Skoog, 1962) containing 25 μg mL\(^{-1}\) kanamycin. At least 12 independent lines were examined for GUS activity by histochemical detection. Seedlings, flowers and siliques were fixed in 50 mM sodium phosphate pH 7.0, 4% (v/v) formaldehyde then infiltrated with 50 mM sodium phosphate pH 7.0, 0.1% (v/v) Triton X-100, 0.5 mM potassium ferrocyanide, 0.5 mM potassium ferricyanide, 10 mM EDTA, 0.05% (w/v) 5-bromo-4-chloro-3-indolyl glucuronide (X-gluc), and then incubated in the dark at 37°C overnight. Green tissues were de-stained with 70% (v/v) ethanol prior to observation. Images were captured with a Nikon Coolpix 4300 camera mounted on an Olympus LMS 225R dissection microscope.

Quantitative RT-PCR Analysis

Total RNA was isolated from roots or leaves using TRI reagent (Sigma Aldrich, St. Louis, MO), and traces of contaminating DNA were eliminated with TURBO DNA-free (Ambion, Austin, TX). Three biological replicates were used for each analysis. First-strand cDNA was synthesized from 1 μg total RNA using SuperScript first-strand cDNA synthesis kit.
(Invitrogen, Carlsbad, CA). Real-time PCR was conducted with Power SYBR Green Master Mix and the ABI Prism 7500 sequence detection system (Applied Biosystems, Foster City, CA). Expression levels were normalized to *EIF4A-2* (Guo et al., 2008).

**Root Plastid Isolation and Pi Transport Assays**

Plastids were isolated from the entire root system of 6-wk-old plants as described (Emes and England, 1986). Briefly, roots (10-15 g) of hydroponically-grown plants were washed in distilled water, chilled at 4°C for 30 min then ground in four 5 s bursts in a blender with 40 mL of cold isolation buffer: 50 mM K-Tricine (pH 8.0), 0.3 M sorbitol, 1 mM EDTA, 2 mM MgCl₂ and 0.1 % (w/v) freshly added BSA. The homogenate was filtered through two layers of Miracloth to yield a crude extract fraction. The crude extract was centrifuged at 1500 g for 3 min at 4°C. The pellet was suspended in 0.5 mL isolation buffer then transferred to a tube containing 40 mL 10% (v/v) Percoll in isolation buffer. The mixture was centrifuged at 3000 g for 10 min at 4°C. Intact plastids were pelleted while broken plastids and cell debris remained at the top of the Percoll solution. The intact plastids were suspended in 20 mL isolation buffer without BSA then pelleted again by centrifugation at 1500 g for 3 min. The pellet was finally suspended in 0.5 mL isolation buffer without BSA and maintained at 4°C. Root plastids were typically 70-75% intact as estimated by phase contrast microscopy. The protein content of the plastid suspension was determined using the Bradford assay with BSA as a standard (Bradford, 1976). For plastid enrichment, the activity of nitrite reductase (NiR) was assayed as previously described (Takahashi et al., 2001). For plastid purity, vacuolar contamination was determined by α-mannosidase activity (Stitt et al., 1989), and combined mitochondrial, cytosolic and peroxisomal contamination was determined by NADH-malate dehydrogenase (NADH-MDH) activity (Schneider and Keller, 2009). Yields of marker enzyme activities in plastid preparations were calculated as a percent of the activities in crude extracts.

Pi uptake by freshly isolated root plastids was assayed in 30 µL assay buffer at a final protein concentration of 1 mg mL⁻¹ for up to 60 sec at 22°C. The assay buffer contained 50 mM Mg-HEPES adjusted to either pH 6.5 or 7.5 as indicated, 0.3 M sorbitol, 1 mM MgCl₂ and 0 or 5 mM NaCl as indicated. The final concentration of [³²P]orthophosphate (60 mCi/mmol; 1 mCi = 37 MBq; Perkin Elmer, USA) was 50 µM. Transport inhibition studies were carried out with 150 µM mersalyl, 6 mM pyridoxal-5'-phosphate (PLP) or 50 µM FCCP, added 1 min before addition.
of Pi. Transport was terminated by the addition of 170 µL of cold assay buffer followed by centrifugation at 16000 g for 1 min. The pellet was washed with 200 µL cold assay buffer then suspended in 30 µL of 2 mM dodecyl maltoside and incubated for 5 min on ice. Accumulated Pi was measured by liquid scintillation spectrometry in 1.5 mL distilled water. Control experiments performed in the presence of 10 mM potassium phosphate buffer (pH 6.5 and 7.5) indicated 10-15% nonspecific binding, which was subtracted from the corresponding measured activity. Root plastids isolated from wild type and mutants were assayed in parallel for each experiment. Results are the average of three independent experiments performed in triplicates ± SD.

Pi export studies were conducted with the same buffers as for uptake. Plastids were first preloaded with Pi for 60 sec in the indicated buffer then washed twice with the same buffer to remove excess Pi. Preloaded plastids were then incubated for 60 sec in the indicated assay buffer at the same temperature. The assay was terminated by centrifugation, and the amounts of Pi remaining in the plastid pellet and exported to the assay buffer were determined by scintillation counting as described above. Results are the average of two independent experiments performed in triplicates ± SD.

**SDS-PAGE and Western Blotting**

Microsomes were prepared as described (Bush, 1989) from wild-type organs to enrich for cellular membrane proteins. The resulting membrane protein fractions and root plastid proteins were separated by electrophoresis using 14 % (w/v) acrylamide SDS-gels. Following electroblotting to PVDF membranes (Millipore, Bedford, MA), the PHT4;2 protein was immunodetected using a specific antibody against the N-terminal peptide RYSSESDGKRRNA produced in rabbit by Innovagen (Lund, Sweden). The blots were further reacted with secondary donkey anti-rabbit antibody conjugated with horseradish peroxidase and chemiluminescent substrate kit (GE Healthcare, UK).

**Extraction and Measurement of Sucrose and Starch**

Leaf and root samples were harvested at the indicated times in the photoperiod then immediately frozen in liquid nitrogen. Sucrose was extracted with 80% (v/v) ethanol, 5% (v/v) formic acid at 80°C for 20 min then the extraction was repeated with 80% ethanol. The supernatants were pooled, lyophilized, suspended in 10 mM Na-acetate (pH 5.5) then assayed for
sucrose. Pellets were dried and starch was gelatinized and hydrolyzed as described (Smith and Zeeman, 2006). Sucrose and starch concentrations were quantified using coupled assays in which the formation of NADH was monitored by absorbance at 340 nm using a Synergy HT microplate reader. All assays were performed in duplicate and values were determined from standard curves. At least four independent samples were collected for each time point and results are reported as mean ± SE.

**Leaf and Cell Size Analysis**

The area of rosettes and individual rosette leaves was determined from digital images that included reference objects using the histogram function of Adobe Photoshop. To determine epidermal cell area, nail polish impressions of the abaxial epidermis were prepared from the ninth rosette leaf of at least three wild-type and *pht4;2* plants. Images were captured with constant magnification (400x) for three different locations in each leaf between 25% and 75% of the distance between the tip and the base of the blade, halfway between the mid-rib and leaf margin. Cells in each field of view were counted, and cell areas (mean ± SE) were determined from a total of at least 500 cells.

**Ploidy Analysis**

Nuclei were isolated from fully expanded leaves of 8-wk-old plants as described previously (Galbraith et al., 1983). Briefly, leaves were finely chopped with a razor blade in 2 ml of ice-cold buffer containing 45 mM MgCl₂, 30 mM sodium citrate, 20 mM MOPS, pH 7 and 1 mg ml⁻¹ Triton X-100. Tissue fragments were removed by filtration through 36 μm nylon mesh, and DNase-free RNase (10 μg ml⁻¹ final) and propidium iodide (100 μg ml⁻¹ final) were added. Nuclei were incubated for 30 min in the dark at room temperature before analyzing the distribution of DNA content using a FACScan flow cytometer. Results are from three independent plants of each genotype and at least 4000 nuclei for each analysis.

**Photosynthetic Activity in Detached Leaves**

Chlorophyll fluorescence was measured using a pulse-amplitude fluorometer model PAM-210 (Walz, Effeltrich, Germany) in leaves detached from 16-h dark-adapted plants. The dark-adapted state fluorescence yield, \( F_0 \), was recorded by using a weak measuring light whereas
the maximum fluorescence yield, $F_m$, was measured after application of a 1-sec pulse of saturating visible light (Schreiber et al., 1986). The maximum quantum yield of photosystem II photochemistry ($F_v/F_m$) was calculated using the equation $F_v/F_m = (F_m-F_0)/F_m$. The photosynthetic electron transport rate (ETR) was determined in detached leaves using PAM-210 by measuring the quantum yield of PSII photochemistry ($Y'$) after every 20 sec of illumination with photosynthetically active radiation (PAR) of 0 to 1850 µmol photons m$^{-2}$ s$^{-1}$, increased stepwise. The relative ETR was calculated by the equation $ETR = 0.84 \times 0.5 \times PAR \times Y$ (Genty et al., 1989). It was assumed that 84% of the incident light was absorbed (factor 0.84), and that an equal fraction of the absorbed quanta is distributed to photosystem II and I (factor 0.5). Results are the mean of three independent measurements ± SD.

Sequence data from this article can be found in the GenBank/EMBL data libraries under the following accession numbers: PHT4;2 (NM_129362), EIF4A-2 (NM_104305), GPT1 (NM_124861), GPT2 (NM_104862), NTT1 (NM_106679), NTT2 (NM_101419), APS1 (NM_124205), APL1 (NM_121927), APL3 (NM_120081), cFBP (NM_103492) and DET2 (NM_129361).

**Supplemental Data**

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

**Supplemental Figure S1.** Localization of PHT4;2 protein in Arabidopsis organs by western blot analysis.

**Supplemental Figure S2.** Rate of Pi uptake by isolated root plastids.

**Supplemental Figure S3.** Distribution of ploidy in rosette leaves.

**Supplemental Figure S4.** Plot of relative electron transport rate as a function of intensity of photosynthetically active radiation.

**ACKNOWLEDGMENTS**

We thank David Galbraith, Georgina Lambert and Katy Kao for assistance with flow cytometry, and Pallavi Mukherjee and Tom McKnight for critical reading of the manuscript.
LITERATURE CITED


Figure legends

**Figure 1.** Localization of *PHT4;2* expression in Arabidopsis. A, Representative PHT4;2-GUS staining patterns for a 5-wk-old plant (left), flower (center) and mature silique (right). B, Quantitative RT-PCR analysis of *PHT4;2* expression in wild-type plants (RT, root; RL, rosette leaf; CL, cauline leaf; F, flower; S, silique). Expression levels were normalized to *EIF-4A2*. Values shown are mean ± SE for three independent analyses.

**Figure 2.** Molecular characterization of *pht4;2* T-DNA insertion mutants. A, Scheme indicating the positions of T-DNA insertions in *pht4;2-1* and *pht4;2-2* lines. Insertions were confirmed by PCR and DNA sequence. Arrows represent primers used for RT-PCR. B, RT-PCR analysis of total RNA isolated from roots of wild-type and *pht4;2* plants.

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**Figure 5.** Starch and sucrose contents. Roots and rosette leaves of 6-wk-old wild-type (WT) and *pht4;2* plants were collected throughout an 8 hr light/16 hr dark photocycle. Values shown are
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**Figure 6.** PHT4;2 modifies the expression of multiple genes involved in plastid transport and carbon metabolism. Expression levels were determined by quantitative RT-PCR of total RNA from plants grown in hydroponics and expressed relative to wild-type levels. A, Genes involved in plastid transport, including Glc6P/Pi exchange ($GPT1$ and $GPT2$) and ATP/ADP exchange ($NTT1$ and $NTT2$). B, Genes involved in starch synthesis ($APS1$, $APL1$ and $APL3$) and sucrose synthesis ($cFBP$). Values shown are mean ± SE for three independent analyses. * Significantly different from wild type ($P<0.05$, Student’s $t$ test).

**Figure 7.** Increased size of $pht4;2$ rosette leaves. A, Rosettes at the end of vegetative growth (8 wk). B, Rosette leaf number 3, 6, 8 and 14 of representative 8 wk wild-type (WT) and $pht4;2$ plants. C, Increase in rosette area during vegetative growth. Values shown are mean ± SE, n=8. * Significantly different from wild type ($P<0.05$, Student’s $t$ test). As shown in panel A, both of the allelic loss of function mutants exhibited the same growth pattern so for clarity only the results for $pht4;2-1$ are shown in panels B and C.
Table I. Distribution of organelle marker enzyme activities in isolated root plastid preparations

Yields are defined as the percentage of total activity in crude extracts (filtered root homogenates). Specific activities are expressed as nkat mg^{-1} protein. Values are mean ± SE of three independent determinations. No significant differences from wild type were found in the *pht4;2-1* and *pht4;2-2* mutants (*P*<0.05, Student’s *t* test).

<table>
<thead>
<tr>
<th></th>
<th>Wild type</th>
<th><em>pht4;2-1</em></th>
<th><em>pht4;2-2</em></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Yield (%)</td>
<td>Specific activity</td>
<td>Yield (%)</td>
</tr>
<tr>
<td>NiR (plastid)</td>
<td>6.91 ± 0.52</td>
<td>42.51 ± 14.34</td>
<td>7.73 ± 1.01</td>
</tr>
<tr>
<td>NADH-MDH (mitochondrion, cytosol, peroxisome)</td>
<td>0.33 ± 0.21</td>
<td>3.83 ± 1.17</td>
<td>0.39 ± 0.32</td>
</tr>
<tr>
<td>α-Mannosidase (vacuole)</td>
<td>0.56 ± 0.03</td>
<td>3.00 ± 0.67</td>
<td>0.86 ± 0.61</td>
</tr>
</tbody>
</table>
Table II. Rosette area and biomass at the end of the vegetative growth stage

Plants were grown in hydroponics for 8 wk. Values are mean ± SE, n=10. * Significantly different from wild type (P<0.05, Student’s t test).

<table>
<thead>
<tr>
<th></th>
<th>Area (cm²)</th>
<th>FW (g)</th>
<th>DW (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wild type</td>
<td>37.06 ± 2.82</td>
<td>0.78 ± 0.08</td>
<td>0.06 ± 0.01</td>
</tr>
<tr>
<td>pht4;2-1</td>
<td>53.76 ± 3.52 *</td>
<td>1.47 ± 0.13 *</td>
<td>0.12 ± 0.01 *</td>
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<tr>
<td>pht4;2-2</td>
<td>61.20 ± 3.87 *</td>
<td>1.78 ± 0.13 *</td>
<td>0.16 ± 0.01 *</td>
</tr>
</tbody>
</table>
Table III. *Leaf cell sizes and numbers*

The ninth rosette leaf was harvested from plants at growth times corresponding to incomplete and complete leaf expansion. Values are mean ± SE for three leaves. Cell areas are calculated from three regions within each leaf. * Significantly different from wild type (P<0.05, Student’s t test).

<table>
<thead>
<tr>
<th></th>
<th>Leaf 9 area (cm²)</th>
<th>Epidermal cell area (μm²)</th>
<th>Epidermal cell number</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Incomplete expansion (5 wk)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wild type</td>
<td>1.6 ± 0.1</td>
<td>2,540 ± 140</td>
<td>63,000 ± 3,940</td>
</tr>
<tr>
<td>pht4;2-1</td>
<td>1.6 ± 0.1</td>
<td>1,900 ± 60*</td>
<td>84,410 ± 5,280*</td>
</tr>
<tr>
<td>pht4;2-2</td>
<td>1.7 ± 0.1</td>
<td>1,870 ± 250*</td>
<td>90,980 ± 5,350*</td>
</tr>
<tr>
<td><strong>Complete expansion (7 wk)</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Wild type</td>
<td>2.2 ± 0.1</td>
<td>2,790 ± 450</td>
<td>78,870 ± 3,590</td>
</tr>
<tr>
<td>pht4;2-1</td>
<td>2.7 ± 0.2*</td>
<td>2,640 ± 200</td>
<td>102,250 ± 7,570*</td>
</tr>
<tr>
<td>pht4;2-2</td>
<td>2.7 ± 0.1*</td>
<td>2,520 ± 210</td>
<td>107,030 ± 3,960*</td>
</tr>
</tbody>
</table>
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